Essential Orthopaedics

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Essential Orthopaedics
(Including Clinical Methods)

FIFTH EDITION

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Dedicated to

My patients
for giving me an opportunity to make a difference in their lives

and

My family & friends
for continuous support at all times
This book written by Dr Maheshwari, is designed to introduce the trainee doctor and the young surgeon to orthopaedic surgery as he will meet it in the developing countries. There have been many good books on orthopaedic surgery and trauma written by experienced authors from Britain and America but their exposure to the real orthopaedic problems experienced in developing countries has often been limited, and today the difference in presentation of orthopaedic surgery in these countries and the presentation of conditions in developing countries like India and Africa is so different that this book, written by a surgeon with a good grounding of clinical experience in India, is most appropriate for the trainee from the developing countries.

I have read a number of chapters and have been impressed with the simple text and clarity with which the different conditions are explained.

Dr Maheshwari has visited my centre in Nottingham for a period of two months. I was impressed with his clarity of thought and his depth of understanding of orthopaedic conditions. I anticipate that this book will be one of many that he writes in future years and is likely to be a major contribution to orthopaedic training in developing countries.

WA Wallace
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The unique position of this book is, that it is meant for undergraduate medical students and those preparing for PG entrance exam. We have had temptation to expand various chapters every time we are called upon to revise the book. But, we keep reminding ourselves of a medical student who has to cover so much curriculum while he is in final year or preparing for PG entrance exam, and thus restrain ourselves from adding ‘too much’.

Though there are rapid advances in medicine, but only few qualify place in a basic book such as this. Every few years when we look at the book we find that some concepts have changed. What was being practiced in only big cities and was mentioned only for the sake of completion in the earlier edition, has become a routine. Also, since the book has remained a popular choice for those appearing for PG entrance, we have added a section on MCQ’s after each chapter.

Like as we would, to add a lot of new information, but all such thoughts go through the mental screening whether it is required for the target audience! We are cognisant of the fact that no change should be made for the heck of it. We hope the essential changes in this new edition would be appreciated by the readers. Feedback welcome.

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J. Maheshwari
Vikram A Mhaskar
What was the thought behind this book in 1993

As an undergraduate, though exposed to orthopaedics only for a short period, I was impressed by the ease with which I could understand the wonderful texts I studied. The problems were that their contents did not exactly meet the requirements of an undergraduate, and most of these books, written by authors from developed countries, did not provide adequate information about diseases peculiar to tropical and underdeveloped countries. Above all, I thought that the concepts could be presented with still more clarity, and improved by way of presentation.

This feeling continued to haunt me everytime I was called upon to teach undergraduates. A couple of years later, an experience at home helped me give a practical shape to this feeling. My wife, who was preparing for PG entrance examination, expected me to teach her orthopaedics. I tried out my ideas on her, and the result was extremely gratifying. Soon after, many more such occasions of teaching undergraduates gave me further opportunities for refining the material. It was on the request of the students that I decided to give it the shape of a book.

The book is primarily addressed to undergraduates and those preparing for the postgraduate entrance tests. General practitioners, particularly in the early stage of their practice would find it useful reference. It would enable nurses and physiotherapists to understand the basic concepts in orthopaedics. Junior postgraduates would find it an enjoyable reading.

Following are the salient features of the book:

1. Most chapters begin with a brief review of the relevant anatomy. This is because by the time a student comes to clinical departments, he has forgotten most of the anatomy he had learnt in the dissection hall.

2. While discussing treatment of a condition, a brief mention of principles is made first, followed by various methods and their indications. This is followed by treatment plan; a practical plan of treatment which is either being followed or can be developed in an average hospital. A brief mention of recent developments is also made.

3. The book has three additional chapters. These are "Approach to a Patient with Limb Injury", "Approach to a Patient with Back Pain", and "Recent Advances in Treatment of Fractures". The first two present a practical approach to handling these frequently encountered emergencies, and the third chapter updates the reader with the latest in this rapidly developing field. Due emphasis has been given to aspects of rehabilitation, considering the recent recommendations of Medical Council of India for including 'rehabilitation' in undergraduate curriculum.

4. Simple line diagrams have been used to supplement the text. Most of them have been developed by myself while teaching the undergraduates. Simplified line diagrams, rather than photographs, enable students understand the basic concepts better.

5. Self-explanatory flow charts are made use of wherever they would help to develop a concept in decision-making.

6. Tables have been used liberally. These serve two purposes: Firstly, they present the text matter in a concentrated form and allow review at a glance. Secondly, they permit quick and easily understandable comparison between related conditions.

7. Necessary information on instruments and implants commonly used in orthopaedics has been provided as an appendix, purely considering the requirement of such knowledge for final professional examination.

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An injury to the musculo-skeletal system can result in damage to bones, joints, muscles and tendons. In addition, the neurovascular bundle of the limb may be damaged. This section will outline the broad principles used in the diagnosis and management of these injuries. These principles can be applied, with suitable modifications, in the management of any musculo-skeletal injury.

**CLASSIFICATION OF FRACTURES**

A fracture is a break in the continuity of a bone. It can be classified on the basis of aetiology, the relationship of the fracture with the external environment, the displacement of the fracture, and the pattern of the fracture.

**ON THE BASIS OF AETIOLOGY**

**Traumatic fracture:** A fracture sustained due to trauma is called a traumatic fracture*. Normal bone can withstand considerable force, and breaks only when subjected to excessive force. Most fractures seen in day-to-day practice fall into this category e.g., fractures caused by a fall, road traffic accident, fight etc.

**Pathological fracture:** A fracture through a bone which has been made weak by some underlying disease is called a pathological fracture. A trivial or no force may be required to cause such a fracture e.g., a fracture through a bone weakened by metastasis.

Although, traumatic fractures have a predictable and generally successful outcome, pathological fractures often go into non-union.

**Stress Fracture:** This is a special type of fracture sustained due to chronic repetitive injury (stress) causing a break in bony trabeculae. These often present as only pain and may not be visible on X-rays.

**ON THE BASIS OF DISPLACEMENTS**

**Undisplaced fracture:** These fractures are easy to identify by the absence of significant displacement.

**Displaced fracture:** A fracture may be displaced. The factors responsible for displacement are: (i) the fracturing force; (ii) the muscle pull on the fracture fragments; and (iii) the gravity. While describing the displacements of a fracture, conventionally, it is the displacement of the distal fragment in relation to the proximal fragment which is mentioned. The displacement can be in the form of shift, angulation or rotation (Fig-1.1).

* An unqualified word ‘fracture’ usually means a traumatic fracture.

** Terms, simple fracture for closed fracture, and compound fracture for open fracture is being dropped, as it is confusing. In fact, new terminology of simple and complex fracture is being introduced (page 2) to mean whether the fracture is simple or complex to treat.
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is extensive devascularisation of fracture ends. Such fractures are often unstable, and slow to heal.

Low-velocity injury: These fractures are sustained as a result of mild trauma force, as in a fall. There is little associated soft tissue injury, and hence these fractures often heal predictably. Lately, there is a change in the pattern of fractures due to shift from low-velocity to high-velocity injuries. The latter gives rise to more complex fractures, which are difficult to treat.

ON THE BASIS OF PATTERN

Transverse fracture: In this fracture, the fracture line is perpendicular to the long axis of the bone. Such a fracture is caused by a tapping or bending force (Fig-1.2).

Oblique fracture: In this fracture, the fracture line is oblique. Such a fracture is caused by a bending force which, in addition, has a component along the long axis of the bone.

Spiral fracture: In this fracture, the fracture line runs spirally in more than one plane. Such a fracture is caused by a primarily twisting force.

Comminuted fracture: This is a fracture with multiple fragments. It is caused by a crushing or compression force along the long axis of the bone.

ON THE BASIS OF RELATIONSHIP WITH EXTERNAL ENVIRONMENT

Closed** fracture: A fracture not communicating with the external environment, i.e., the overlying skin and other soft tissues are intact, is called a closed fracture.

Open fracture: A fracture with break in the overlying skin and soft tissues, leading to the fracture communicating with the external environment, is called an open fracture. A fracture may be open from within or outside, the so called internally or externally open fracture respectively.

a) Internally open (from within): The sharp fracture end pierces the skin from within, resulting in an open fracture.

b) Externally open (open from outside): The object causing the fracture lacerates the skin and soft tissues over the bone, as it breaks the bone, resulting in an open fracture.

Exposure of an open fracture to the external environment makes it prone to infection. This risk is more in externally open fractures.

ON THE BASIS OF COMPLEXITY OF TREATMENT

Simple fracture: A fracture in two pieces, usually easy to treat, is called simple fracture, e.g. a transverse fracture of humerus.

Complex fracture: A fracture in multiple pieces, usually difficult to treat, is called complex fracture, e.g. a comminuted fracture of tibia.

ON THE BASIS OF QUANTUM OF FORCE CAUSING FRACTURE

High-velocity injury: These are fractures sustained as a result of severe trauma force, as in traffic accidents. In these fractures, there is severe soft tissue injury (periosteal and muscle injury). There

is extensive devascularisation of fracture ends. Such fractures are often unstable, and slow to heal.

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Comminuted fracture: This is a fracture with multiple fragments. It is caused by a crushing or compression force along the long axis of the bone.
Monteggia fracture-dislocation: Fracture of the proximal third of the ulna, with dislocation of the head of the radius (page 110).

Galeazzi fracture-dislocation: Fracture of the distal third of the radius with dislocation of the distal radio-ulnar joint (page 111).

Night-stick fracture: Isolated fracture of the shaft of the ulna, sustained while trying to ward off a stick blow.

Colles’ fracture: A fracture occurring in adults, at the cortico-cancellous junction of the distal end of the radius with dorsal tilt and other displacements (page 111).

Smith’s fracture: A fracture occurring in adults, at the cortico-cancellous junction of the distal end of the radius with ventral tilt and other displacements (reverse of Colles’).

Barton’s fracture (Marginal fracture): Intra-articular fractures through the distal articular surface of the radius, taking a margin, anterior or posterior, of the distal radius with the carpals, displaced anteriorly or posteriorly (page 114).

Chauffeur fracture: An intra-articular, oblique fracture of the styloid process of the radius.

Bennett’s fracture-dislocation: It is an oblique, intra-articular fracture of the base of the first metacarpal with subluxation of the trapezio-metacarpal joint (page 117).

Boxers’ fracture: It is a ventrally displaced fracture through the neck of the 5th metacarpal, usually occurs in boxers.

Side-swipe fracture: It is an elbow injury sustained when one’s elbow, projecting out of a car, is ‘side-swept’ by another vehicle. It has a combination of fractures of the distal end of the humerus and fractures of proximal ends of radius and/or ulna. It is also called baby car fracture.

Bumper fracture: It is a comminuted, depressed fracture of the lateral condyle of the tibia.

Pott’s fracture: Bimalleolar ankle fracture.

Cotton’s fracture: Trimalleolar ankle fracture.

Massonaise’s fracture: It is a type of ankle fracture in which fracture of the neck of the fibula occurs.

Pilon fracture: It is a comminuted intra-articular fracture of the distal end of the tibia.

Aviator’s fracture: Fracture of neck of the talus.

Chopart fracture-dislocation: A fracture-dislocation through inter-tarsal joints.

Jone’s fracture: Avulsion fracture of the base of the 5th metatarsal.

Rolando fracture: Fracture of the base of the first metacarpal (extra-articular).

Jefferson’s fracture: Fracture of the first cervical vertebra.

Whiplash injury: Cervical spine injury where sudden flexion followed by hyperextension takes place.

Chance fracture: Also called seat belt fracture, the fracture line runs horizontally through the body of the vertebra, through and through, to the posterior elements.

March fracture: Fatigue fracture of the shaft of 2nd or 3rd metatarsal.

Burst fracture: It is a comminuted fracture of the vertebral body where fragments “burst out” in different directions (page 268).

Clay-Shoveller fracture: It is an avulsion fracture of spinous process of one or more of the lower cervical or upper thoracic vertebrae.

Hangman’s fracture: It is a fracture through the pedicle and lamina of C2 vertebra, with subluxation of C2 over C3, sustained in hanging.

Dashboard fracture: A fracture of posterior lip of the acetabulum, often associated with posterior dislocation of the hip.

Straddle fracture: Bilateral superior and inferior pubic rami fractures.

Malgaigné’s fracture: A type of pelvis fracture in which there is a combination of fractures, pubic rami anteriorly and sacro-iliac joint or ilium posteriorly, on the same side.

Mallet finger: A finger flexed at the DIP joint due to avulsion or rupture of extensor tendon at the base of the distal phalanx.

**PATHOLOGICAL FRACTURES**

A fracture is termed pathological when it occurs in a bone made weak by some disease (Fig-1.3). Often, the bone breaks as a result of a trivial trauma, or even spontaneously.

**CAUSES**

A bone may be rendered weak by a disease localised to that particular bone, or by a generalised bone disorder. Table–1.1 gives some of the common
causes of pathological fractures. Osteoporosis is the commonest cause of pathological fracture. The bones most often affected are the vertebral bodies (thoracic and lumbar). Other common fractures associated with osteoporosis are fracture of the neck of the femur and Colles’ fracture.

A local or circumscribed lesion of the bone, responsible for a pathological fracture, may be due to varying causes in different age groups (Table 1.2). In children, it is commonly due to chronic osteomyelitis or a bone cyst. In adults, it is often due to a bone cyst or giant cell tumour. In elderly people, metastatic tumour is a frequent cause.

**Table 1.1: Causes of pathological fractures**

<table>
<thead>
<tr>
<th>Localised Diseases</th>
</tr>
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<tbody>
<tr>
<td>Inflammatory</td>
</tr>
<tr>
<td>• Pyogenic osteomyelitis</td>
</tr>
<tr>
<td>• Tubercular osteomyelitis</td>
</tr>
<tr>
<td>Neoplastic</td>
</tr>
<tr>
<td>• Benign tumours</td>
</tr>
<tr>
<td>• Giant cell tumour, Enchondroma</td>
</tr>
<tr>
<td>• Malignant tumours</td>
</tr>
<tr>
<td>• Primary</td>
</tr>
<tr>
<td>• Osteosarcoma, Ewing’s tumour</td>
</tr>
<tr>
<td>• Secondary</td>
</tr>
<tr>
<td>• In males: lung, prostate, kidney</td>
</tr>
<tr>
<td>• In females: breast, lung, genitals</td>
</tr>
<tr>
<td>Miscellaneous</td>
</tr>
<tr>
<td>• Simple bone cyst</td>
</tr>
<tr>
<td>• Aneurysmal bone cyst</td>
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<tr>
<td>• Monostotic fibrous dysplasia</td>
</tr>
<tr>
<td>• Eosinophilic granuloma</td>
</tr>
<tr>
<td>• Bone atrophy secondary to polio etc.</td>
</tr>
<tr>
<td>Generalised Diseases</td>
</tr>
<tr>
<td>Hereditary</td>
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<tr>
<td>• Osteogenesis imperfecta</td>
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<tr>
<td>• Dyschondroplasia (Ollier’s disease)</td>
</tr>
<tr>
<td>• Osteopetrosis</td>
</tr>
<tr>
<td>Acquired</td>
</tr>
<tr>
<td>• Osteoporosis</td>
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<td>• Osteomalacia</td>
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<tr>
<td>• Rickets</td>
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<tr>
<td>• Scurvy</td>
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<tr>
<td>• Disseminated malignancy in bones</td>
</tr>
<tr>
<td>• Multiple myeloma</td>
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<tr>
<td>• Diffuse metastatic carcinoma</td>
</tr>
<tr>
<td>• Miscellaneous</td>
</tr>
<tr>
<td>• Paget’s disease</td>
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<tr>
<td>• Polyostotic fibrous dysplasia</td>
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</tbody>
</table>

**DIAGNOSIS**

A fracture sustained without a significant trauma should arouse suspicion of a pathological fracture.

Often the patient, when directly questioned, admits to having suffered from some discomfort in the region of the affected bone for some time before the fracture. The patient may be a diagnosed case of a disease known to produce pathological fractures (e.g., a known case of malignancy), thus making the diagnosis of a pathological fracture simple. At times, the patient may present with a pathological fracture, the cause of which is determined only after a detailed work up.

**TREATMENT**

Treatment of a pathological fracture consists of: (i) detecting the underlying cause of the fracture; and (ii) making an assessment of the capacity of the fracture to unite, based on the nature of the underlying disease.

A fracture in a bone affected by a generalised disorder like Paget’s disease, osteogenesis imperfecta or

**Table 1.2: Causes of pathological fractures at different ages**

<table>
<thead>
<tr>
<th>Age</th>
<th>Causes</th>
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</thead>
<tbody>
<tr>
<td>At birth</td>
<td>Osteogenesis imperfecta</td>
</tr>
<tr>
<td>0-5 years</td>
<td>Osteogenesis imperfecta</td>
</tr>
<tr>
<td></td>
<td>Osteomyelitis</td>
</tr>
<tr>
<td>5-20 years</td>
<td>Osteomyelitis</td>
</tr>
<tr>
<td></td>
<td>Simple bone cyst</td>
</tr>
<tr>
<td></td>
<td>Primary bone malignancy</td>
</tr>
<tr>
<td>20-50 years</td>
<td>Cystic lesions of the bone</td>
</tr>
<tr>
<td></td>
<td>Malignancy</td>
</tr>
<tr>
<td></td>
<td>Osteomalacia</td>
</tr>
<tr>
<td></td>
<td>Giant cell tumour</td>
</tr>
<tr>
<td>After 50 years</td>
<td>Osteoporosis</td>
</tr>
<tr>
<td></td>
<td>Multiple myeloma</td>
</tr>
<tr>
<td></td>
<td>Secondaries in the bone</td>
</tr>
</tbody>
</table>
osteoporosis is expected to unite with conventional methods of treatment. A fracture at the site of a bone cyst or a benign tumour will also generally unite, but the union may be delayed. Fractures occurring in osteomyelitic bones often take a long time, and sometimes fail to unite despite best efforts. Fractures through metastatic bone lesions often do not unite at all, though the union may occur if the malignancy has been brought under control with chemotherapy or radiotherapy.

With the availability of facilities for internal fixation, more and more pathological fractures are now treated operatively with an aim to: (i) enhance the process of union by bone grafting (e.g. in bone cyst or benign tumour); or (ii) mobilise the patient by surgical stabilisation of the fracture. Achieving stable fixation in these fractures is difficult because of the bone defect caused by the underlying pathology. The defect may have to be filled using bone grafts or bone cement.

INJURIES TO JOINTS

Joint injuries may be either a subluxation or a dislocation. A joint is subluxated when its articular surfaces are partially displaced but retain some contact between them (Fig-1.4).

A joint is dislocated when its articular surfaces are so much displaced that all contact between them is lost. A dislocated joint is an emergency, and should be treated at the earliest.

INJURIES TO LIGAMENTS

An injury to a ligament is termed as a sprain. This is to be differentiated from the term ‘strain’ which means stretching of a muscle or its tendinous attachment.

CLASSIFICATION

Sprains are classified into three degrees (Fig-1.5):

First-degree sprain is a tear of only a few fibres of the ligament. It is characterised by minimal swelling, localised tenderness but little functional disability.

Second-degree sprain is the one where, anything from a third to almost all the fibres of a ligament are disrupted. The patient presents with pain, swelling and inability to use the limb. Joint movements are normal. The diagnosis can be made on performing a stress test as discussed subsequently.

Third-degree sprain is a complete tear of the ligament. There is swelling and pain over the torn ligament. Contrary to expectations, often the pain in such tears is minimal. Diagnosis can be made by performing a stress test, and by investigations such as MRI or arthroscopy.

PATHOLOGY

A ligament may get torn in its substance (mid-substance tear) or at either end. In the latter case, it often avulses with a small piece of bone from its attachment (Fig-1.6).

DIAGNOSIS

A detailed history, eliciting the exact mechanism of injury, often indicates the likely ligament injured. The examination helps in finding the precise location and severity of the sprain, which can then be confirmed by investigations.
Sometimes, following a complete tear of ligament, the blood seeps out of the joint through the rent in the ligament, and it appears as if there is no haemarthrosis.

**Clinical examination:** A localised swelling, tenderness, and ecchymosis over a ligament indicates injury to that ligament. Usually, a haemarthrosis is noticed in second and third-degree sprains within 2 hours. It may be absent* despite a complete tear, or if the torn ligament is covered by synovium (e.g., intra-synovial tear of anterior cruciate ligament).

**Stress test** (Fig-1.7): This is a very useful test in diagnosing a sprain and judging its severity. The ligament in question is put to stress by a manoeuvre. The manoeuvre used for testing of individual ligaments will be discussed in respective chapters. When a ligament is stressed, in first and second-degree sprains, there will be pain at the site of the tear. In third-degree sprain, the ligament will ‘open up’ as well.

**INVESTIGATIONS**

A plain X-ray of the joint is usually normal. Sometimes, a chip of bone may be seen in the region of the attachment of the ligament to the bone. An X-ray taken while the ligament is being stressed (stress X-ray) may document an abnormal opening up of the joint in a third-degree sprain.

Other investigations required in a few cases are MRI or arthroscopy.

**TREATMENT**

There has been a significant change in the treatment of sprains. All sprains are treated initially with rest, ice therapy, compression bandage, elevation (RICE). Suitable analgesics and anti-inflammatory medication is given. This is enough for first-degree sprains. Second and third-degree sprains are immobilised in a brace or a plaster cast for a period of 1-2 weeks, mainly for pain relief. No longer is plaster immobilisation advised for long periods. In fact, early mobilisation and walking with support enhances healing of ligaments. In some third-degree sprains, surgery may be required.

**INJURIES TO MUSCLES AND TENDONS**

Muscles are ruptured more often than tendons in young people, while the reverse is true in the elderly. The most frequent cause of partial or complete rupture of a muscle or a tendon is sudden vigorous contraction of a muscle. It may be by overstretching of a muscle at rest. Such an injury to muscle is termed strain (and not sprain, which is ligament injury). A muscle or tendon injury may also be produced by a sharp object such as a sword.

**PATHOLOGY**

A rupture occurs within a tendon only if it is abnormal and has become weak, either due to degeneration or wear and tear. Degenerative tendon ruptures commonly occur in rheumatoid arthritis, SLE, senile degeneration, etc. Tendon rupture related to wear and tear commonly occurs in the biceps (long head), and in extensor pollicis longus tendons. Some tendons known to rupture commonly are as given in Table–1.3. Diagnosis of a ruptured tendon is usually easy. The patient complains of pain and

**Table–1.3: Common sites of tendon rupture**

- Supraspinatus tendon
- Achilles tendon
- Biceps tendon – long head
- Extensor pollicis longus tendon
- Quadriceps tendon
- Patellar tendon
inability to perform the movement for which the tendon is meant.

**TREATMENT**
The best treatment of a fresh rupture is to regain continuity by end-to-end repair. When the gap is too much, it can be filled with the help of a tendon graft. In cases where the repair is not possible, a tendon transfer may be performed. In some old tendon ruptures, especially in the elderly, there may be only a minimal functional disability. These patients do well without treatment.

**Further Reading**

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### What have we learnt?

When we see a fracture, we must find out:

- Is it purely a traumatic fracture or is there an underlying pathology (weak bone)?
- What is the pattern of the fracture, so as to know the inherent stability of the fracture. Stable fractures can generally be treated non-operatively, unstable fractures often need surgery.
- What is the force causing the fracture? Is it a ‘high-velocity’ injury where fracture is likely to be unstable with lot of associated injury or a ‘low-velocity’ injury. The type of force producing the injury has bearing on healing of the fracture.
- It is a simple or complex fracture? The latter may be a badly comminuted fracture as a result of a bad trauma, and hence may be associated with lot of soft tissue damage. Such fractures often need surgical treatment.
- An open fracture has additional problem of getting infected. Hence, appropriate care in the early part of management is important.
- Sprain and strain are not interchangeable terms.
- A ‘no bony injury’ on X-ray does not mean no injury. Look for ligament or muscle injuries.

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### Additional information: From the entrance exams point of view

Best diagnostic test for unilateral stress fracture is MRI, and for bilateral is bone scan: The most common cause of pathological fracture is osteoporosis.
Bones may be classified into four types on the basis of their shape i.e., long, short, flat and irregular. For practical purposes, anatomy of a typical long bone only is being discussed here.

**Structure of a typical long bone**: In children, a typical long bone, such as the femur, has two ends or *epiphyses* (singular *epiphysis*), and an intermediate portion called the *shaft* or *diaphysis*. The part of the shaft which adjoins the epiphysis is called the *metaphysis* – one next to each epiphysis. There is a thin plate of growth cartilage, one at each end, separating the epiphysis from the metaphysis. This is called the *epiphyseal plate*. At maturity, the epiphysis fuses with the metaphysis and the epiphyseal plate is replaced by bone. The articular ends of the epiphyses are covered with articular cartilage. The rest of the bone is covered with periosteum which provides attachment to tendons, muscles, ligaments, etc. The strands of fibrous tissue connecting the bone to the periosteum are called *Sharpey’s fibres*.

Microscopically, bone can be classified as either woven or lamellar. *Woven bone or immature bone* is characterized by random arrangement of bone cells (osteocytes) and collagen fibres. Woven bone is formed at periods of rapid bone formation, as in the initial stages of fracture healing. *Lamellar bone or mature bone* has an orderly arrangement of bone cells and collagen fibres. Lamellar bone constitutes all bones, both cortical and cancellous. The difference is, that in cortical bone the lamellae are densely packed, and in cancellous bone loosely.

The basic structural unit of lamellar bone is the *osteon*. It consists of a series of concentric laminations or lamellae surrounding a central canal, the *Haversian canal*. These canals run longitudinally and connect freely with each other and with *Volkmann’s canals*. The latter run horizontally from endosteal to periosteal surfaces. The shaft of a bone

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**Fig-2.1 Parts of a child’s bone**
is made up of cortical bone, and the ends mainly of cancellous bone. The junction between the two, termed the **cortico-cancellous junction** is a common site of fractures (Fig-2.2).

**Structural composition of bone:** The bone is made up of bone cells and extra-cellular matrix. The matrix consists of two types of materials, organic and inorganic. The organic matrix is formed by the collagen, which forms 30-35 percent of dry weight of a bone. The inorganic matrix is primarily calcium and phosphorus salts, especially **hydroxyapatite** \(\text{Ca}_10(\text{PO}_4)_6(\text{OH})_2\). It constitutes about 65-70 percent of dry weight of a bone.

**Bone cells:** There are three main cell types in the bone. These are:

a) **Osteoblasts:** Concerned with ossification, these cells are rich in alkaline phosphatase, glycolytic enzymes and phosphorylases.

b) **Osteocytes:** These are mature bone cells which vary in activity, and may assume the form of an osteoclast or reticulocyte. These cells are rich in glycogen and PAS positive granules.

c) **Osteoclasts:** These are multi-nucleate mesenchymal cells concerned with bone resorption. These have glycolytic acid hydrolases, collagenases and acid phosphatase enzymes.

**Growth of a long bone**

All long bones, with the exception of the clavicle, develop from cartilaginous primordia (enchondral ossification). This type of ossification commences in the middle of the shaft (primary centre of ossification) before birth. The secondary ossification centres (the epiphyses) appear at the ends of the bone, mostly* after birth.

The bone grows in length by a continuous growth at the epiphyseal plate. The increase in the girth of the bone is by subperiosteal new bone deposition.

At the end of the growth period, the epiphysis fuses with the diaphysis, and the growth stops. The secondary centres of ossification, not contributing to the length of a bone, are termed **apophysis** (e.g., apophysis of the greater trochanter). The time and sequence of appearance and fusion of epiphysis has clinical relevance in deciding the true age (bone age) of a child. Sometimes, an epiphyseal plate may be wrongly interpreted as a fracture.

**Remodelling of bone:** Bone has the ability to alter its size, shape and structure in response to stress. This happens throughout life though not perceptible. According to Wolff’s law of bone remodelling, bone hypertrophy occurs in the plane of stress.

**Blood supply of bones**

There is a standard pattern of the blood supply of a typical long bone. Blood supply of individual bones will be discussed wherever considered relevant. The blood supply of a typical long bone is derived from the following sources (Fig-2.3):

a) **Nutrient artery:** This vessel enters the bone around its middle and divides into two branches, one running towards either end of the bone.

b) **Metaphyseal vessels:** These are numerous small vessels derived from the anastomosis around the joint. They pierce the metaphysis along the line of attachment of the joint capsule.

c) **Epiphyseal vessels:** These are vessels which enter directly into the epiphysis.
d) Periosteal vessels: The periosteum has a rich blood supply, from which many little vessels enter the bone to supply roughly the outer-third of the cortex of the adult bone. Blood supply to the inner two-thirds of the bone comes from the nutrient artery and the outer one-third from the periosteal vessels.

**FRACTURE HEALING**

The healing of fractures is in many ways similar to the healing of soft tissue wounds, except that soft tissue heals with fibrous tissue, and end result of bone healing is mineralised mesenchymal tissue, i.e. bone. A fracture begins to heal soon after it occurs, through a continuous series of stages described below (Table–2.1).

**STAGES IN FRACTURE HEALING OF CORTICAL BONE (FROST, 1989)**

- Stage of haematoma
- Stage of granulation tissue
- Stage of callus
- Stage of remodelling (formerly called consolidation)
- Stage of modelling (formerly called remodelling)

**Stage of haematoma:** This stage lasts up to 7 days. When a bone is fractured, blood leaks out through torn vessels in the bone and forms a haematoma between and around the fracture. The periosteum and local soft tissues are stripped from the fracture ends. This results in ischaemic necrosis of the fracture ends over a variable length, usually only a few millimetres. Deprived of their blood supply, some osteocytes die whereas others are sensitised to respond subsequently by differentiating into daughter cells. These cells later contribute to the healing process.

**Stage of granulation tissue:** This stage lasts for about 2-3 weeks. In this stage, the sensitised precursor cells (daughter cells) produce cells which differentiate and organise to provide blood vessels, fibroblasts, osteoblasts etc. Collectively they form a soft granulation tissue in the space between the fracture fragments. This cellular tissue eventually gives a soft tissue anchorage to the fracture, without any structural rigidity. The blood clot gives rise to a loose fibrous mesh which serves as a framework for the ingrowth of fibroblasts and new capillaries. The clot is eventually removed by

<table>
<thead>
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<th>Table–2.1: Stages of fracture healing (Frost, 1989)</th>
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<tr>
<td>Stage of healing</td>
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<tr>
<td>Stage of haematoma</td>
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<tr>
<td>Stage of granulation tissue</td>
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<tr>
<td>Stage of callus</td>
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<tr>
<td>Stage of remodelling</td>
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<tr>
<td>Stage of modelling</td>
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</table>
macrophages, giant cells and other cells arising in the granulation tissue.

From this stage, the healing of bone differs from that of soft tissue. In soft tissue healing the granulation tissue is replaced by fibrous tissue, whereas in bone healing the granulation tissue further differentiates to create osteoblasts which subsequently form bone.

**Stage of callus:** This stage lasts for about 4-12 weeks. In this stage, the granulation tissue differentiates further and creates osteoblasts. These cells lay down an intercellular matrix which soon becomes impregnated with calcium salts. This results in formulation of the callus, also called woven bone. The callus is the first sign of union visible on X-rays, usually 3 weeks after the fracture (Fig-2.4). The formation of this bridge of woven bone imparts good strength to the fracture. Callus formation is slower in adults than in children, and in cortical bones than in cancellous bones.

**Stage of remodelling:** Formerly called the stage of consolidation. In this stage, the woven bone is replaced by mature bone with a typical lamellar structure. This process of change is multicellular unit based, whereby a pocket of callus is replaced by a pocket of lamellar bone. It is a slow process and takes anything from one to four years.

**Stage of modelling:** Formerly called the stage of remodelling. In this stage the bone is gradually strengthened. The shaping of cortices occurs at the endosteal and periosteal surfaces. The major stimulus to this process comes from local bone strains i.e., weight bearing stresses and muscle forces when the person resumes activity. This stage is more conspicuous in children with angulated fractures. It occurs to a very limited extent in fractures in adults.

**HEALING OF CANCELLOUS BONES**

The healing of fractured cancellous bone follows a different pattern. The bone is of uniform spongy texture and has no medullary cavity so that there is a large area of contact between the trabeculae. Union can occur directly between the bony trabeculae. Subsequent to haematoma and granulation formation, mature osteoblasts lay down woven bone in the intercellular matrix, and the two fragments unite.

**PRIMARY AND SECONDARY BONE HEALING**

*Primary fracture healing* occurs where fracture haematoma has been disturbed, as in fractures treated operatively. The bone heals directly, without callus formation, and it is therefore difficult to evaluate union on X-rays. *Secondary fracture healing* occurs in fractures where fracture haematoma is not disturbed, as in cases treated non-operatively. There is healing, with callus formation, and can be evaluated on X-rays. It also occurs in fractures operated without disturbing the fracture haematoma, as in fractures fixed with relative stability (e.g. comminuted fractures).

**FACTORS AFFECTING FRACTURE HEALING**

a) **Age of the patient:** Fractures unite faster in children. In younger children, callus is often visible on X-rays as early as two weeks after the fracture. On an average, bones in children unite in half the time compared to that in adults. Failure of union is uncommon in fractures of children.

b) **Type of bone:** Flat and cancellous bones unite faster than tubular and cortical bones.

c) **Pattern of fracture:** Spiral fractures unite faster than oblique fractures, which in turn unite faster than transverse fractures. Comminuted fractures are usually result of a severe trauma or occur in osteoporotic bones, and thus heal slower.

d) **Disturbed pathoanatomy:** Following a fracture, changes may occur at the fracture site, and may hinder the normal healing process. These are: (i) soft tissue interposition; and (ii) ischaemic fracture ends. In the former, the fracture ends pierce through the surrounding soft tissues, and...
get stuck. This causes soft tissue interposition between the fragments, and prevents the callus from bridging the fragments. In the latter, due to anatomical peculiarities of blood supply of some bones (e.g. scaphoid), vascularity of one of the fragments is cut off. Since vascularised bone ends are important for optimal fracture union, these fractures unite slowly or do not unite at all.

e) Type of reduction: Good apposition of the fracture results in faster union. At least half the fracture surface should be in contact for optimal union in adults. In children, a fracture may unite even if bones are only side-to-side in contact (bayonet reduction).

f) Immobilisation: It is not necessary to immobilise all fractures (e.g., fracture ribs, scapula, etc). They heal anyway. Some fractures need strict immobilisation (e.g., fracture of the neck of the femur), and may still not heal.

g) Open fractures: Open fractures often go into delayed union and non-union (discussed subsequently on page 21).

h) Compression at fracture site: Compression enhances the rate of union in cancellous bone. In cortical bones, compression at the fracture site enhances rigidity of fixation, and possibly results in primary bone healing.

Further Reading


What have we learnt?

- There are different parts of a long bone such as diaphysis, metaphysis and epiphysis. There are diseases which typically affect only some parts of the bones.
- Growing skeleton is identified by presence of growth plate.
- The structure of the bone is complex, made of the basic structural unit called osteon.
- Different bone cells have different functions.
- Fracture healing follows a series of stages.
- Fracture healing depends upon a number of factors.

Additional information: From the entrance exams point of view

Pathognomonic sign of traumatic and fresh fracture is crepitus.
Most common cause of non-union is inadequate immobilisation.

Markers of bone formation:
- Serum bone specific alkaline phosphatase
- Serum osteocalcin
- Serum peptide of type 1 collagen

Markers of bone resorption:
- Urine and serum crosslinked ‘N’ telopeptide
- Urine and serum crosslinked ‘C’ telopeptide
- Urine total free deoxypyridinoline

Rate of mineralisation determined by labelled tetracycline.
Treatment of a fracture can be considered in three phases:
- Phase I - Emergency care
- Phase II - Definitive care
- Phase III - Rehabilitation

**PHASE I - EMERGENCY CARE**

*At the site of accident:* Emergency care of a fracture begins at the site of the accident. In principle, it consists of RICE, which means:
- Rest to the part, by splinting.
- Ice therapy, to reduce occurrence of swelling
- Compression, to reduce swelling
- Elevation, to reduce swelling

*Rest to the part* (splinting) is done by splinting. 'Splint them where they lie'. Before applying the splint, remove ring or bangles worn by the patient. Almost any available object at the site of the accident can be used for splinting. It may be a folded newspaper, a magazine, a rigid cardboard, a stick, an umbrella, a pillow, or a wooden plank. Any available long piece of cloth can be used for tying the splint to the fractured limb. Some of the examples of splinting a fractured extremity at the site of the accident are shown in Fig-3.1. One may correct any gross deformity by gentle traction. Feel for distal pulses, and do a quick assessment of nerve function.
supply before and after splinting. The advantages of splinting are:

- Relief of pain, by preventing movement at the fracture.
- Prevention of further damage to skin, soft tissues and neurovascular bundle of the injured extremity.
- Prevention of complications such as fat embolism and hypovolaemic shock.
- Transportation of the patient made easier.

**Ice therapy:** An immediate application of ice to injured part helps in reducing pain and swelling. This can be done by taking crushed ice in a polythene bag and covering it with a wet cloth. Commercially available ice packs can also be used. Any wound, if present, has to be covered with sterile clean cloth.

**Compression:** A crepe bandage is applied over the injured part, making sure that it is not too tight.

**Elevation:** The limb is elevated so that the injured part is above the level of the heart. For lower limb, this can be done using pillows. For upper limb, a sling and pillow can be used.

**In the emergency department:** Soon after a patient with a musculo-skeletal trauma is received in an emergency department, one has to act in a coordinated way. It is most important to provide, if required, basic life support (BLS). If in shock, the patient is stabilised before any definitive orthopaedic treatment is carried out. A quick evaluation of the extent of injury at this stage enables a doctor to understand the seriousness of the problem. Particular attention is paid to head injury, chest injury and abdominal injury. These can be cause of early fatality. Any bleeding is recognised and stopped by local pressure. The fractured limb is examined to exclude injury to nerves or vessels. As soon as the general condition of the patient is stabilised, the limb is splinted. It is important to check the bandaging done elsewhere, as it may be too tight. Some of the splints used in the emergency department are as shown in Fig-3.2.

In addition to splintage, the patient should be made comfortable by giving him intramuscular analgesics. In a case with suspected head injury, narcotic analgesics should be avoided. A broad spectrum antibiotic may be given to those with open fractures. It is only after the emergency care has been given, and it is ensured that the patient is stable. He should be sent for suitable radiological and other investigations, under supervision.

**PHASE II - DEFINITIVE CARE**

**Philosophy of fracture treatment:** Over the years, treatment of fractures has undergone change in philosophy. In the past, the aim of treatment was a mere fracture union. This could be achieved in most cases by immobilisation, which would cause joint stiffness, muscle wasting etc., and may result in less than optimal functional recovery. The aim now is to get the limb functions back to pre-injury level. For this, early mobilisation of the limb is desirable, as this helps in preserving joint movements and muscle functions.

Perfect anatomical reduction and stable fixation is preferred for intra-articular fractures, as only then early mobilisation can be done. In diaphyseal fractures, the aim is to achieve union in good alignment and length. This can be done by non-operative methods, if the fracture is stable. Operative methods are required for unstable fractures. With currently available techniques of surgery, the trend is towards treating more and more fractures operatively as this gives more predictable results, early recovery and better functions. The discussion that follows will give the reader a guideline.

**Fundamental principles of fracture treatment:**
The three fundamental principles of treatment of a fracture are: (i) reduction; (ii) immobilisation; and (iii) preservation of functions.
Reduction is the technique of ‘setting’ a displaced fracture to proper alignment. This may be done non-operatively or operatively, so-called closed and open reduction respectively.

Immobilisation is necessary to maintain the bones in reduced position. This may be done by external immobilisation such as plaster etc., or by internal fixation of the fracture using rods, plates, screw etc.

To preserve the functions of the limb, physiotherapy all throughout the treatment, even when the limb is immobilised, is necessary.

Methods of treatment: Not all the three fundamental treatment principles discussed above apply to all fractures. Treatment of a particular fracture can fall in one of the following categories:

a) Treatment by functional use of the limb: Some fractures (e.g., fractured ribs, scapula) need no reduction or immobilisation. These fractures unite despite functional use of the part. Simple analgesics and splinting are needed for the initial few days, basically for pain relief.

b) Treatment by immobilisation alone: In some fractures, mere immobilisation of the fracture in whatever position, is enough. Fractures without significant displacement or fractures where the displacement is of no consequence (e.g., some fractures of surgical neck of the humerus) are treated this way.

c) Treatment by closed reduction followed by immobilisation: This is required for most displaced fractures treated non-operatively. The reduction could be done under mild sedation or under anaesthesia. Immobilisation is usually in a plaster cast. There is trend towards use of image intensifier (page 32) to aid closed reduction.

d) Closed reduction and percutaneous fixation: This is done for fractures, which though can be reduced by closed manipulation, but are unstable, and are likely to displace subsequently. These fractures are reduced under image intensifier, and fixed with percutaneous devices such as K-wire, rush pins etc., which hold the fracture in position. External support of a plaster or splint is usually required, in addition.

e) Open reduction and internal fixation: There are some fractures, such as intra-articular fractures, where accurate reduction, stable fixation and early mobilisation are very important to regain joint functions. Such fractures are best treated by open reduction and internal fixation. Some unstable fractures are also treated by this method.

f) Minimally invasive surgery (MIS): There is trend towards treating fractures with minimally invasive techniques. In this, image intensifier is used to aid reduction without opening the fracture. The fracture is, then stabilised internally using special devices such as rods, plates etc. These devices are introduced through small incisions using special instrumentation. MIS has the advantage that the blood supply of the bone is preserved, and thus early union occurs. Less pain, early recovery and cosmesis are other advantages.

Which of the above method is used in a particular fracture depends upon a number of factors such as patient’s profession, whether the injured limb is dominant or not, surgeon’s experience, availability of facilities, patient’s affordability etc. It is therefore common to see differing opinions on the treatment of a particular fracture.

Discussed below are the three fundamental principles of fracture treatment: reduction, immobilisation and preservation of functions.

Reduction of Fractures

Indications: Not all fractures require reduction, either because there is no displacement or because the displacement is immaterial to the final outcome. For example, a child’s clavicle fracture does not need reduction because normal function and appearance will be restored without any intervention.

In general, imperfect apposition of fragments can be accepted more readily than imperfect angulatory alignment or rotational mal-alignment. Perfect anatomical reduction is desirable in some fractures, even if for this an operation is required (e.g. intra-articular fractures).

Methods: Reduction of a fracture can be carried out by one of the following methods:

a) Closed manipulation: This is the standard initial method of reducing most of the common fractures. It is usually carried out under general anaesthesia* and requires experience. It is an art of realigning a displaced bone by feeling
through the soft tissues. The availability of an image intensifier has greatly added to the skills of closed reduction. It is not necessary that perfect anatomical reduction be achieved in all cases. Displacements compatible with normal functions are considered 'acceptable'. Most fractures reduced by closed manipulation need some kind of immobilisation (PoP, brace, bandaging etc.) discussed subsequently.

b) Continuous traction: It is used to counter the forces which will not allow reduction to happen or would cause redisplacement. These are muscle forces and the force of gravity. A common example is that of an inter-trochanteric fracture, in which the muscles attached to different fragments cause displacements. A continuous traction can counter this force, and bring the bones in proper alignment.

Continuous traction has its own problem of keeping the patient in bed for long time with its complications such as bedsores etc. It is for this reason that once the fracture so treated becomes 'sticky', and has little possibility of redisplacement, the traction is discontinued and the fracture supported in a plaster cast till healing occurs. It is because of uncertainty of result and need for in-bed immobilisation that many of these fractures are now treated operatively. Different methods of applying traction are discussed in Chapter 4.

c) Open reduction: In this method, the fracture is surgically exposed, and the fragments are reduced under vision. Some form of internal fixation is used in order to maintain the position. This is commonly referred as 'open reduction and internal fixation' or ORIF. This is one area of fracture treatment which is continuously evolving. There used to be times when orthopaedic wards used to be full of patients in traction and huge plaster casts for months. Today, with advancement in surgical treatment, the paradigm has shifted to operative treatment. The big deciding factor for adopting ORIF as the treatment of choice is the facilities available and training of the surgeon. The potential risks of surgery are sometimes worse than the disadvantages of non-operative treatment.

One reason to do open reduction is when other methods of achieving reduction have failed. There are fractures which are so unstable that one knows that these fractures will redisplace in due course. In such fractures, open reduction and secure internal fixation is carried out in the first instance.

Some of the widely accepted indications of ORIF are given in Table–3.1.

### Immobilisation of fractures

<table>
<thead>
<tr>
<th>Table–3.1 Indications for open reduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Absolute</td>
</tr>
<tr>
<td>• Failure of closed reduction</td>
</tr>
<tr>
<td>• Displaced intra-articular fractures</td>
</tr>
<tr>
<td>• Some displaced epiphyseal injuries (types III and IV)</td>
</tr>
<tr>
<td>• Major avulsion fractures e.g., fracture of patella</td>
</tr>
<tr>
<td>• Non-union</td>
</tr>
<tr>
<td>Relative</td>
</tr>
<tr>
<td>• Delayed union and malunion</td>
</tr>
<tr>
<td>• Multiple fractures</td>
</tr>
<tr>
<td>• Pathological fractures</td>
</tr>
<tr>
<td>• Where closed reduction is known to be ineffective e.g., fracture of the neck of the femur</td>
</tr>
<tr>
<td>• Fractures with vascular or neural injuries</td>
</tr>
</tbody>
</table>

**Indications:** Not all fractures require immobilisation. The reasons for immobilising a fracture may be:

a) To prevent displacement or angulation: In general, if reduction has been necessary, immobilisation will be required.

b) To prevent movement that might interfere with the union: Persistent movement might tear the delicate early capillaries bridging the fracture. More strict immobilisation is necessary for some fractures (e.g., scaphoid fracture).

c) To relieve pain: This is the most important reason for the immobilisation of most fractures. As the fracture become pain free and feels stable, guarded mobilisation can be started.

**Methods:** Immobilisation of a fracture can be done by non-operative or operative methods.

**Non-operative methods**

Most fractures can be immobilised by one of the following non-operative methods:

**Strapping:** The fractured part is strapped to an adjacent part of the body e.g., a phalanx fracture,
where one finger is strapped to the adjacent normal finger (see Fig-16.3 on page 118).

**Sling:** A fracture of the upper extremity is immobilised in a sling. This is mostly to relieve pain in cases where strict immobilisation is not necessary e.g., triangular sling used for a fracture of the clavicle.

**Cast immobilisation:** This is the most common method of immobilisation. Plaster-of-Paris casts have been in use for a long time. Lately, fibreglass casting tapes have become popular. The latter provide durable, light-weight, radiolucent casts.

Plaster of Paris (Gypsum salt) is CaSO$_4$·½H$_2$O in dry form, which becomes CaSO$_4$·2H$_2$O on wetting. This conversion is an *exothermic* reaction and is *irreversible*. The plaster sets in the given shape on drying. The setting time of a plaster varies with its quality, and temperature of the water. Names of some of the plaster casts commonly used are given in Table–3.2.

Table–3.2 Plaster casts and their uses

<table>
<thead>
<tr>
<th>Name of the cast</th>
<th>Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minerva cast</td>
<td>Cervical spine disease</td>
</tr>
<tr>
<td>Risser’s cast</td>
<td>Scoliosis</td>
</tr>
<tr>
<td>Turn-buckle cast</td>
<td>Scoliosis</td>
</tr>
<tr>
<td>Shoulder spica*</td>
<td>Shoulder immobilisation</td>
</tr>
<tr>
<td>U-slab</td>
<td>Fracture of the humerus</td>
</tr>
<tr>
<td>Hanging cast</td>
<td>Fracture of the humerus</td>
</tr>
<tr>
<td>Colles’ cast</td>
<td>Colles’ fracture</td>
</tr>
<tr>
<td>Hip spica</td>
<td>Fracture of the femur</td>
</tr>
<tr>
<td>Cylinder cast</td>
<td>Fracture of the patella</td>
</tr>
<tr>
<td>PTB cast</td>
<td>Fracture of the tibia</td>
</tr>
</tbody>
</table>

* A spica is a cast where a limb and a part of the trunk are included, e.g., hip spica, shoulder spica.

**Types of plaster bandages:** There are two types of plaster bandages in use—one prepared by impregnating rolls of starched cotton bandages with plaster powder (home-made bandages); the other are ready-made bandages available as a proprietary bandage.

**Use of Plaster of Paris:** It can be applied in two forms i.e., slab or a cast.

A plaster slab covers only a part of the circumference of a limb. It is made by unrolling a plaster bandage to and fro on a table. An average slab is about twelve such thicknesses. The slab is used for the immobilisation of soft tissue injuries and for reinforcing plaster casts. A plaster cast covers the whole of the circumference of a limb. Its thickness varies with the type of fracture and the part of the body on which it is applied. Some of the fundamental principles to be remembered while applying a plaster cast are as follows:

- Immobilise the joints above and below the fracture.
- Immobilise joints in a functional position*.
- Pad the limb adequately, especially on bony prominences.

**After care of a plaster:** This involves noticing any cracks in the plaster, avoiding wetting the plaster, and graduated weight bearing for lower limb fractures. Exercising the muscles within the plaster and moving the joints not in the plaster, is necessary to ensure early recovery.

**Complications of plaster treatment:** The following are some of the common complications of plaster treatment:

- Impairment of circulation (tight cast) (Fig-3.3): A plaster cast is a closed compartment. Haematoma and tissue oedema following a fracture can result in increased pressure inside the cast, leading to impaired circulation of the extremity. Early diagnosis, by a high index of suspicion, can prevent disastrous complications like gangrene. Unrelenting pain, especially stretch pain (see page 47), swelling over the fingers, inability to move the fingers, hypoesthesia and bluish discoloration of

![Fig-3.3 Disastrous complication of plaster cast. (a) X-ray of a child with rather simple fracture (b) Ischaemic and deformed foot due to tight plaster](https://kat.cr/user/Blink99/)
the digits are signs of a tight cast. A tight cast can be prevented by adequately padding the cast and elevating the extremity for the first 2-3 days following a cast application.

- Plaster sores: These are caused by inadequate padding, irregularity of the inner surface of the cast, or foreign bodies in the plaster. A sore formation within a plaster cast can be suspected by the following:
  - Pain, out of proportion to fracture
  - Fretfulness
  - Disturbed sleep
  - Recurrence of swelling over toes or fingers
  - Low grade fever
  - Patch of blood/soakage over the cast.

A plaster sore can be prevented by examination of the suspected area through a window in the cast. It is possible to dress a small sore through this window. Occasionally, the plaster has to be removed and reapplied.

Functional bracing (Fig-3.4): A brace is a type of cast where the joints are not included, so that while the fracture is kept in position, the joints can also be mobilised. This method is commonly used for stable fractures of the tibia and humerus. It is based on the principle that continuous use of the affected limb while the fracture is kept adequately supported, encourages union and prevents joint stiffness. The brace is usually applied after the fracture becomes 'sticky'. In experienced hands, the rate of fracture healing by this method is comparable to other methods (see details on page 33). It is a useful option at places where facilities for surgical treatment are not available.

Splints and traction: Splints of various designs are used for the definitive treatment of fractures. Thomas splint is still very popular for the treatment of fractures of the lower limb. Disadvantages of this method of treatment are prolonged hospitalisation and confinement to the bed. This can be hazardous, especially in elderly people who develop complications secondary to recumbency (e.g., bed sores, chest infection etc.). For details about splints, see Chapter 4.

OPERATIVE METHODS

Wherever open reduction is performed, fixation (internal or external) should also be used. External fixation is usually indicated in situations where for some reason, internal fixation cannot be done.

Internal fixation: In this method, the fracture, once reduced, is held internally with the help of some metallic or non-metallic device (implant), such as steel wire, screw, plate, Kirschner wire (K-wire), intra-medullary nail etc. These implants are made of high quality stainless steel to which the body is inert.

Indications: Internal fixation of fractures may be indicated under the following circumstances:

a) When a fracture is so unstable that it is difficult to maintain it in an acceptable position by non-operative means. This is the most frequent indication for internal fixation.

b) As a treatment of choice in some fractures, in order to secure rigid immobilisation and to allow early mobility of the patient.

c) When it has been necessary to perform open reduction for any other reason such as an associated neurovascular injury.

Methods: A fracture can be fixed internally by any one or combination of implants given in Table-3.3.

a) Steel wire: A gauge 18 or 20 steel wire is used for internal fixation of small fractures (e.g., fracture of the patella, comminuted fragments of large bones etc.).

b) Kirschner wire: It is a straight stainless steel wire, 1-3 mm in diameter. It is used for the fixation of small bones of the hands and feet.

c) Intra-medullary nail: It is erroneously called 'nail', but in fact is a hollow rod made...
Advantages of internal fixation: With the use of modern techniques and implants, there is minimal need for external immobilisation. It allows early mobility of the patient out of bed and hospital. Joints do not get stiff and the muscle functions remain good. The complications associated with confinement of a patient to bed are also avoided.

Disadvantages: The disadvantages of internal fixation are infection and non-union. It needs a trained orthopaedic surgeon, free availability of implants and a good operation theatre; failing which, the results of internal fixation may not only be poor but disastrous.

External fixator: It is a device (Fig-3.5) by which the fracture is held in a steel frame outside the limb. For this, pins are passed percutaneously to hold the bone, and are connected outside to a bar with the help of clamps. This method is useful in the treatment of open fractures where internal fixation cannot be carried out due to risk of infection.

These are of the following type:

i. Pin fixators: In these, 3–4 mm sized pins are passed through the bone. The same are held outside the bone with the help of a variety of tubular rods and clamps [Fig-3.5 (a)]

ii. Ring fixators: In these, thin ‘K’ wires (1–2 mm) are passed through the bone. The same are held outside the bone with rings [Fig-3.5 (b)]. (For details, page 32).

PHASE III - REHABILITATION OF A FRACTURED LIMB

Rehabilitation of a fractured limb begins at the time of injury, and goes on till maximum possible functions have been regained. It consists of joint

Table 3.3 Some implants used in treatment of fractures

<table>
<thead>
<tr>
<th>Intra-medullary nails</th>
</tr>
</thead>
<tbody>
<tr>
<td>PFN, DFN, PHN, Recon nail supracondylar nail</td>
</tr>
<tr>
<td>- Kuntscher’s nail</td>
</tr>
<tr>
<td>- Smith-Petersen nail</td>
</tr>
<tr>
<td>- Talwalkar’s nail</td>
</tr>
<tr>
<td>- V-nail</td>
</tr>
<tr>
<td>- Ender’s nail</td>
</tr>
<tr>
<td>- Rush nail</td>
</tr>
<tr>
<td>- Hartshill rectangle</td>
</tr>
<tr>
<td>- GK nail</td>
</tr>
<tr>
<td>- Gamma nail</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Plates and screws</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Compression plate</td>
</tr>
<tr>
<td>- Neutralisation plate</td>
</tr>
<tr>
<td>- Buttrress plate</td>
</tr>
<tr>
<td>- Locking compression plate</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Special implants</th>
</tr>
</thead>
<tbody>
<tr>
<td>- SP nail-plate</td>
</tr>
<tr>
<td>- Dynamic hip screw (DHS)</td>
</tr>
<tr>
<td>- Condyilar blade-plate</td>
</tr>
<tr>
<td>- T-plate</td>
</tr>
<tr>
<td>- Spoon plate</td>
</tr>
<tr>
<td>- Cobra plate</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Steel wire</td>
</tr>
<tr>
<td>- K-wire</td>
</tr>
</tbody>
</table>
mobilisation, muscle re-education exercises and instructions regarding gait training.

**Joint mobilisation:** The joint adjacent to an injured bone tends to get stiff due to: (i) immobilisation; (ii) inability to move the joints due to pain; and (iii) associated injury to the joint as well. To prevent stiffness, the joint should be mobilised as soon as possible. This is done initially by *passive mobilisation* (some one else does it for the patient). Once the pain reduces, patient is encouraged to move the joint himself with assistance (active assisted), or move the joint by himself (active mobilisation). Motorized devices which slowly move the joint through a predetermined range of motion can be used. These are called **continuous passive motion (CPM) machines**. Techniques such as hot fomentation, gentle massage and manipulation aid in joint mobilisation.

**Muscle re-education exercises:** Because of lack of use, the muscles get wasted quickly. Hence, it is desirable that muscle activity be maintained all through the treatment. This can be done even during immobilisation (static contractions) or after removal of external immobilisation (dynamic contractions), as discussed below:

a) **During immobilisation:** Even while a fracture is immobilised, the joints which are out of the plaster, should be moved to prevent stiffness and wasting of muscles. Such movements do not cause any deleterious effect on the position of the fracture. The muscles working on the joints inside the plaster can be contracted without moving the joint (static contractions). This maintains some functions of the immobilised muscles.

b) **After removal of immobilisation:** After a limb is immobilised for some period, it gets stiff. As the plaster is removed, the following care is required:
   - The skin is cleaned, scales removed, and some oil applied.
   - The joints are moved to regain the range of motion. Hot fomentation, active and active-assisted joint mobilising exercises are required for this (page 81).
   - The muscles wasted due to prolonged immobilisation are exercised.

**Functional use of the limb:** Once a fracture is on way to union, at a suitable opportunity, the limb is put to use in a guarded way. For example, in lower limb injuries, gradual weight bearing is started – partial followed by full. One may need to support the limb in a brace, caliper, cast etc. Walking aids such as a walker, a pair of crutches, stick etc. may be necessary.

A general plan of management for a usual fracture is shown in Flow chart-3.1.

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**Flow chart-3.1 General plan for treatment of fractures**

---

* Open reduction straightaway is required for some fractures
MANAGEMENT OF OPEN FRACTURES

A fracture is called open (compound) when there is a break in the overlying skin and soft tissues, establishing communication between the fracture and the external environment. Three specific consequences may result from this.

a) **Infection of bone:** Contamination of the wound with bacteria from the outside environment may lead to infection of the bone (osteomyelitis).

b) **Inability to use traditional methods:** A small wound can be managed through a window in a plaster cast. But, it may not be possible to manage a big wound through a window. The presence of a wound may also be a deterrent to operative fixation of the fracture.

c) **Problems related to union:** Non-union and mal-union occur commonly in open fractures. This may be because of one or more of the following reasons: (i) a piece of bone may be lost from the wound at the time of the fracture, the gap thus created predisposes to non-union; (ii) the fracture haematoma, which is supposed to have osteogenic potential, is lost from the wound; (iii) the 'vascular' cover by the overlying soft tissues, so important for fracture union, may be missing; and (iv) the bone may get secondarily infected, and thus affect union.

It is because of these possible consequences that open fractures deserve utmost care throughout their management. Open fractures have been classified into three types, depending upon the extent of soft tissue injury (Table–3.4).

### TREATMENT

The principle of treatment is to convert an open fracture into a closed fracture by meticulous wound care. Thereafter, the treatment of open fracture is essentially on the lines of closed fractures. The following discussion emphasises the points pertinent to the treatment of open fractures.

#### Phase I - Emergency Care

**At the site of accident:** The following measures are taken at the site of the accident:

a) The bleeding from the wound is stopped by applying firm pressure using a clean piece of cloth. At times it may be necessary to use a tight circular bandage proximal to the wound in order to stop bleeding.

b) The wound is washed with clean tap water or saline, and covered with a clean cloth.

c) The fracture is splinted.

At times, a piece of bone devoid of all soft tissue attachments may be lying out of the wound. It should be washed and taken to the hospital in a clean cloth. It may be useful in reconstruction of the fracture.

**In the emergency department:** Open fractures are known to be associated with neurovascular injuries more often than the simple fractures. Hence, one should carefully look for these associated injuries. The following treatment is performed in the emergency department:

a) **Wound care:** Care in the emergency room consists of washing the wound under strict aseptic conditions and covering it with sterile dressing. Sometimes, the bone may be jetting out of the skin, causing stretching of the skin around the wound. Replacing the projecting bone is necessary in order to prevent devascularisation of the skin. A piece of bone with intact soft tissue attachments hanging out of the wound, should...
be washed and put back in the wound. All this is done under proper aseptic conditions.

b) **Splintage** as described on page 13.

c) **Prophylactic antibiotics** should be given to all patients. Cephalexin is a good broad spectrum antibiotic for this purpose. In serious compound fractures, a combination of third generation cephalosporins and an amino-glycoside is preferred.

d) **Tetanus prophylaxis** is given after evaluating the tetanus immunisation status of the patient.

e) **Analgesics** to be given parenterally to make the patient comfortable.

f) **X-rays** are done to evaluate the fracture in order to plan further treatment.

**Phase II - Definitive Care**

Definitive care of an open fracture is possible at a place equipped with a high class aseptic operation theatre, plenty of orthopaedic instruments and implants, and a competent orthopaedic surgeon. In some compound fractures, the damage to soft tissues is so much that it is wise to consult a plastic surgeon right at the beginning. The patient may need plastic surgery techniques, such as flap reconstruction, at the time of the first operation itself. Longer a bone is exposed to outside environment, more it gets desicated, resulting in subsequent non-union.

In principle, in the treatment of open fractures, care of the wound goes hand in hand with that of the fracture.

**Wound care:** This consists of early wound debridement and subsequent care.

a) **Wound debridement:** Wound debridement is needed in all cases. There may be only a puncture wound, needing minimal debridement, irrigation and wound closure; or the limb may be so badly crushed that repeated debridement may be required. While debriding the wound, the skin should be excised as little as necessary. The muscles and fascia can be excised liberally. The most reliable indicator of the viability of a muscle is its contractility, on pinching it with a forceps. Only badly lacerated tendons are excised. The ends of a cut tendon are approximated with non-absorbable sutures so that they can be identified at a later date, and a definitive repair performed. Bone ends are cleaned thoroughly with normal saline. The margins of the fractured ends may be nibbled. A bone fragment with attached soft tissues is replaced at the fracture site. Small fragments without soft tissue attachments can be discarded. Sometimes, the limb is so badly injured that the prospects of salvaging the limb to a reasonable function is poor. In such cases, amputation, straight away, may be a better option. It is recommended that opinion of at least one more surgeon be taken before taking such a drastic decision.

b) **Definitive wound management:** Once the wound is debrided, decision regarding its closure is to be made. **Primary** closure by suturing the skin edges or by raising a flap, can be okay for clean wounds. In all wounds debrided after 6-8 hours, immediate closure should not be done. The wound, in such cases should be covered with sterile dressings, and subsequently treated by delayed primary closure or be allowed to heal by secondary intention. Whenever in doubt, it is best to leave the wound open. A plan of wound closure in open fractures is shown in Flow chart-3.2.

**Fracture management:** In spite of the best debridement, an open fracture is a potentially infected fracture. Non-operative methods of treatment, as in closed fractures, usually give good results. In case an operative reduction of the fracture is considered necessary, it is safer to wait for the wound to heal before intervening. In cases where there is extensive damage to soft tissues, external fixation provides fixation of the fracture and allows good care of the wound. Some of the commonly used methods in the definitive care of an open fracture are as follows:

a) **Immobilisation in plaster:** For cases with moderate size wound, where a stable reduction of the fracture can be achieved, treatment by Plaster of Paris cast is as appropriate as for closed fractures. Care of the wound is possible through a window in the cast. Once the wound heals, the window is closed and the fracture treated on the lines of closed fractures.

b) **Pins and plaster:** For cases where the wound is moderate in size and is manageable through a window in a plaster cast, but reduction is unstable; the fracture can be stabilised by passing pins in the proximal and distal fragments, achieving reduction, and applying plaster cast with pins incorporated in it. This method is useful in open, unstable tibial fractures (Fig-3.6).
c) **Skeletal traction:** In cases where there is circumferential loss of skin or the wound is big, it may not be possible to treat them in plaster. In such cases, skeletal traction can be used to keep the fracture in good alignment until the wound heals. After healing of the wound, one can continue traction until the fracture unites, or change over to some other form of immobilisation such as plaster cast.

d) **External skeletal fixation:** It provides stability to fracture and permits access to virtually the whole circumference of the limb (see also page 33).

e) **Internal fixation:** Approach to management of open fractures has become very aggressive in last few years. In trauma centres in developed countries, more and more open fractures received early enough are treated with primary internal fixation. Closed methods of intramedullary fixation have been particularly useful. If everything goes well, the rehabilitation of the patient is highly accelerated. Such facilities are fast becoming available in most centres in India and other developing countries.

**Phase III - Rehabilitation**

Rehabilitation of a limb with an open fracture is along the lines of a simple fracture as discussed on page 19. It consists of joint mobilisation, muscle exercises during immobilisation, after removal of immobilisation, and advice regarding mobilisation of the injured limb.

**Further Reading**

- Gustilo RB, Kyle RF, Templeman D: *Fractures and Dislocations*, St. Louis: Mosby-Year Book Inc.
What have we learnt?

- Fracture has to be splinted as soon as possible.
- Fracture treatment consists of keeping the fracture in acceptable position till union. This can be done by non-operative methods or operative methods.
- Non-operative methods consists of bandaging, plaster application, use of brace, traction etc.
- Operative treatment consists of reduction of the fracture and holding it in position by internal fixation or external fixation.
- Open fractures are serious injuries, as they are more prone to complications. Adequate wound care and fracture treatment is required.

Additional information: From the entrance exams point of view

Securing the airway is the first step in treatment of polytrauma.
Splints and Traction

TOPICS
- Splints
- Traction

SPLINTS

OBJECTIVES
Splints are used for immobilising fractures; either temporarily during transportation or for definitive treatment. They are also used in other orthopaedic conditions like infection, congenital dislocation of the hip, etc.

TYPES
Some of the splints used in orthopaedic practice, and the conditions for which they are used are given in Table 4.1. The following are a few examples of common splints.

- **Cramer-wire splint**: This splint is used for temporary splintage of fractures during transportation. It is made up of two thick parallel wires with interlacing wires (Fig 4.1). It can be bent into different shapes in order to immobilise different parts of the body.

- **Thomas knee-bed splint (Thomas splint)**: It is one of the commonest splints used in orthopaedic practice. It was devised by H.O. Thomas, initially for immobilisation for tuberculosis of the knee. It is now commonly used for the immobilisation of hip and thigh injuries.

  Parts of a Thomas splint (Fig 4.2): A Thomas splint has a ring and two side bars joined distally. The ring is at an angle of 120° to the inside bar. The outside bar has a curvature near its junction with the ring to accommodate the greater trochanter.

Table 4.1: Common splints/braces and their uses

<table>
<thead>
<tr>
<th>Name</th>
<th>Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cramer-wire splint</td>
<td>Emergency immobilisation</td>
</tr>
<tr>
<td>Thomas splint</td>
<td>Fracture femur - anywhere</td>
</tr>
<tr>
<td>Bohler-Braun splint</td>
<td>Fracture femur - anywhere</td>
</tr>
<tr>
<td>Aluminium splint</td>
<td>Immobilisation of fingers</td>
</tr>
<tr>
<td>Dennis Brown splint</td>
<td>CTEV</td>
</tr>
<tr>
<td>Cock-up splint</td>
<td>Radial nerve palsy</td>
</tr>
<tr>
<td>Knuckle-bender splint</td>
<td>Ulnar nerve palsy</td>
</tr>
<tr>
<td>Toe-raising splint</td>
<td>Foot drop</td>
</tr>
<tr>
<td>Volkmann’s splint</td>
<td>Volkmann’s ischaemic contracture (VIC)</td>
</tr>
<tr>
<td>Four-post collar</td>
<td>Neck immobilisation</td>
</tr>
<tr>
<td>Aeroplane splint</td>
<td>Brachial plexus injury</td>
</tr>
<tr>
<td>SOMI brace</td>
<td>Cervical spine injury</td>
</tr>
<tr>
<td>ASHE (Anterior spinal hyper extension)</td>
<td>Dorso-lumbar spinal injury</td>
</tr>
<tr>
<td>Taylor’s brace</td>
<td>Dorso-lumbar immobilisation</td>
</tr>
<tr>
<td>Milwaukee brace</td>
<td>Scoliosis</td>
</tr>
<tr>
<td>Boston brace</td>
<td>Scoliosis</td>
</tr>
<tr>
<td>Lumbar corset</td>
<td>Backache</td>
</tr>
</tbody>
</table>

Fig 4.1 Cramer-wire splint
A patient in splint needs the following care:

a) The splint should be properly applied, well padded at bony prominences and at the fracture site.

b) The bandage of the splint should not be too tight as it may produce sores; nor too loose, lest it becomes ineffective.

c) The patient should be encouraged to actively exercise the muscles and the joints inside the splint as much as permitted.

d) Any compression of nerve or vessel, usually due to too tight a bandage or lack of adequate padding, should be detected early and managed accordingly.

e) Daily checking and adjustments, if required, should be made. Regular portable X-rays may be taken to ensure good position of the fracture.

Now-a-days, readymade braces are available for immobilising different joints. These are available in small to extra large (XL) sizes. Common ones in use are knee immobiliser, wrist immobiliser, and ankle support.

**CARE OF A PATIENT IN A SPLINT**

**TRACIONS**

**OBJECTIVES**

Traction is used for: (i) reduction of fractures and dislocations, and their maintenance; (ii) for immobilising a painful, inflamed joint; (iii) for the prevention of deformity, by counteracting the muscle spasms associated with painful joint conditions; and (iv) for the correction of soft tissue contractures by stretching them out.

**TYPES OF TRACTION**

For effectiveness of any traction, a counter-traction is necessary. Depending upon what acts as counter-traction, a traction can be fixed or sliding.

- **Fixed traction**: In this type, counter-traction is provided by a part of the body e.g., in Thomas splint fixed traction, the ring of the splint comes to lie against the ischial tuberosity and provides counter-traction (Fig-4.4a).
Splints and Traction

Sliding traction: In this type, the weight of the body acts as counter-traction; e.g., traction given for a pelvic fracture, where the weight of the body acts as counter-traction; made effective by elevating the foot-end of the bed (Fig. 4.4b).

METHODS OF APPLYING TRACTION
There are two methods of applying traction – skin and skeletal (Fig-4.4).

Skin traction: An adhesive strap is applied on the skin and traction applied. The traction force is transmitted from the skin through the deep fascia and intermuscular septae to the bone. These days, readymade foam traction kits are available for this purpose.

Skeletal traction: The traction is applied directly on the bone by inserting a K-wire or Steinmann pin through the bone.

Table 4.2: Comparison between skin and skeletal tractions

<table>
<thead>
<tr>
<th>Point</th>
<th>Skin traction</th>
<th>Skeletal traction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Required for</td>
<td>Mild to moderate force</td>
<td>Moderate to severe force</td>
</tr>
<tr>
<td>Age used for</td>
<td>Children</td>
<td>Adults</td>
</tr>
<tr>
<td>Applied with</td>
<td>Adhesive plaster</td>
<td>Steinmann pin, K-wire</td>
</tr>
<tr>
<td>Applied</td>
<td>On skin</td>
<td>Through bone</td>
</tr>
<tr>
<td>Common site</td>
<td>Below knee</td>
<td>Upper tibial pin traction</td>
</tr>
<tr>
<td>Weight permitted</td>
<td>Up to 3-4 kg</td>
<td>Up to 20 kg</td>
</tr>
<tr>
<td>Used for</td>
<td>Short duration</td>
<td>Long duration</td>
</tr>
</tbody>
</table>

Common traction systems used are given in Table 4.3.

Table 4.3: Traction systems and their uses

<table>
<thead>
<tr>
<th>Name</th>
<th>Use</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gallow’s traction</td>
<td>Fracture shaft of the femur in children below 2 years</td>
</tr>
<tr>
<td>Bryant’s traction</td>
<td>Same</td>
</tr>
<tr>
<td>Russell’s traction</td>
<td>Trochanteric fractures</td>
</tr>
<tr>
<td>Buck’s traction</td>
<td>Conventional skin traction</td>
</tr>
<tr>
<td>Perkin’s traction</td>
<td>Fracture shaft of femur in adults</td>
</tr>
<tr>
<td>90-90 traction</td>
<td>Fracture shaft of femur in children</td>
</tr>
<tr>
<td>Agnes-Hunt traction</td>
<td>Correction of hip deformity</td>
</tr>
<tr>
<td>Well-leg traction</td>
<td>Correction of adduction or abduction deformity of hip</td>
</tr>
<tr>
<td>Dunlop traction</td>
<td>Supracondylar fracture of humerus</td>
</tr>
<tr>
<td>Smith’s traction</td>
<td>Supracondylar fracture of humerus</td>
</tr>
<tr>
<td>Calcaneal traction</td>
<td>Open fractures of ankle or leg</td>
</tr>
<tr>
<td>Metacarpal traction</td>
<td>Open forearm fractures</td>
</tr>
<tr>
<td>Head-halter traction</td>
<td>Cervical spine injuries</td>
</tr>
<tr>
<td>Crutchfield traction</td>
<td>Cervical spine injuries</td>
</tr>
<tr>
<td>Halo-pelvic traction</td>
<td>Scoliosis</td>
</tr>
</tbody>
</table>

DAILY CARE OF A PATIENT IN TRACTION
A patient in traction can develop serious complications and needs the following care:

a) The traction should be as comfortable as possible.

b) Proper functioning of the traction unit must be ensured. Traction weights should not be touching the ground. See that the ropes are in the grooves of the pulleys. The foot of the patient or the end of the traction device should not be touching the pulley, as it makes traction ineffective.
c) One must see that terminal part of the limb in traction (hand or foot) is warm and of normal colour. Sensations over toes and fingers should be normal. Any numbness or tingling may point to a traction palsy of a nerve.

d) Any swelling over the fingers or toes may point to a tight bandage or slipped skin traction.

e) A pin tract infection in skeletal traction can be detected early by eliciting pain on gentle tapping at the site of the pin insertion.

f) The proper position of the fracture should be ensured by taking check X-rays in traction.

g) Physiotherapy of the limb in traction should be continued to minimise muscle wasting.

h) A watch must be kept on general complications of recumbency, i.e., bed sores, chest congestion, UTI, constipation etc.

i) Diversion therapy is important for any patient confined to bed for a long period of time. This may be done by suggesting the patient to do things he likes — such as reading, craft, games, watching television, net surfing, etc.

Further Reading

What have we learnt?
- Splints are useful and readily available methods of immobilising a limb. Due care is required while treating a patient in splint.
- There are two types of traction — fixed and sliding.
- There are two methods of applying traction — skin and skeletal.
- Skeletal traction is more convenient for giving traction for longer duration. Also, more weight can be applied by skeletal traction. Skin traction is suitable for short term traction only.
- How to take proper care of a patient in traction?

Additional information: From the entrance exams point of view

Skin traction is contraindicated in skin damage, deep vein thrombosis, significant vascular deficit and neurological deficit.
AO METHOD OF FRACTURE TREATMENT

The AO (Arbeitsgemeinschaft fur Osteosynthesefragen, a Swiss term meaning association for osteosynthesis) and its English counterpart, the ASIF (Association for the Study of Internal Fixation) advocated the internal fixation of fractures based on principles laid down by them. The basic guiding principle is that by achieving stable fixation of fractures, a limb can be mobilised early, thereby avoiding the disadvantages of immobilisation i.e., stiffness of joints, muscle wasting etc., all of which have been termed ‘fracture disease’ by them. The following principles are used to achieve stable fixation (Fig-5.1):

a) **Inter-fragmentary compression** i.e., achieving compression between different fracture fragments.

b) **Splinting** i.e., splinting the fracture internally or externally.

c) **Combination** of a and b.

INTER-FRAGMENTARY COMPRESSION

This means that the fracture fragments are not merely in contact with each other, but are compressed against each other. It improves the strength of the fixation. Compression between fragments can be produced at the time of surgery, and is called **static** compression. It can also occur between fragments as a result of muscle action as the limb is put to use, the so called **dynamic** compression.

**Methods of producing static compression:** It can be produced by the following methods:

a) **Lag screw fixation:** A screw is passed across the fracture site in such a way that, as the screw is tightened, the fracture surfaces are compressed against each other. In this technique, the fracture is reduced and held with a clamp. A hole is drilled across the fracture, as shown in Fig-5.1a. The proximal cortex is ‘overdrilled’. As the screw is passed, it slides through the proximal cortex (being overdrilled), and its threads catch the opposite cortex (Fig-5.1a). The head of the screw pushes the proximal cortex against the near cortex.

* Lag screw is no special screw. It is just the way a screw is used. Partially threaded screws are used to produce lag-screw effect. Any threaded screw can be used as a lag screw by ‘over drilling’ the near cortex.
SPLINTING

There are various methods of ‘splinting’ the fracture surgically. These do not provide ‘rigid’ fixation, but the fixation is good enough to support fracture healing. Following are some such methods (Fig-5.4).

a) **Intra-medullary splinting**: This is useful for fixation of fractures of the long bones, e.g., fracture of the shaft of the femur. A long hollow rod, called ‘nail’ is inserted in the medullary cavity of the long bone (Fig-5.4a).

b) **Extra-medullary splinting**: This can be done for any fracture by applying a plate on the surface which tends to ‘open up’ on use of the limb, because it is subjected to a distracting force. If this force is countered by some device fixed on the tension side, the fracture comes under compression. This is called *tension-band principle*. Dynamic compression can be achieved by the following methods:

a) **Tension-band wire** (Fig-5.3a): This can be used for producing dynamic compression in fractures of the patella and olecranon.

b) **Tension-band plate** (Fig-5.3b): This can be used for fractures of the humerus and tibia by applying the plate on tension surface.
of the bone. This plate, being used just to 'splint' the fracture, without producing compression, is called a neutralisation plate (Fig-5.4b). Sometimes, the plate may just be buttressing the fracture, without really fixing it (buttress plating), as done for fixing tibial condyle fractures (Fig-5.4c).

c) Outside the body: Pins are inserted through the skin into the bone, and the same are held outside with clamps and rod (external fixators). This is used to hold the fragments of an open fracture (Fig-5.4d).

![Fig-5.5 An example of use of combination of compression screws and a neutralisation plate, used for a spiral fracture](https://kat.cr/user/Blink99/)

**COMBINATION OF COMPRESSION AND SPLINTING**

A combination of these two principles is required in achieving stable fixation in most fractures e.g., a spiral fracture of the shaft of the femur can be stabilised by using inter-fragmentary screws across the fracture, and a neutralisation plate to add to fixation (Fig-5.5). Just inter-fragmentary screw fixation alone may not be strong enough fixation.

**CHANGING AO CONCEPTS**

Before 50's, fractures were treated primarily non-operatively. The healing of fractures occurred as per stages described in Chapter 2. Callus formation was an important stage in the healing of fractures. Non-operative treatment required immobilisation of the limb, which lead to stiffness of joints, wasting of muscles etc. (the so-called fracture disease). Operative techniques were primitive, and were associated with unacceptable complications.

**EARLY AO CONCEPTS**

In the late 50's, AO group brought revolution in the treatment of fractures. They proposed anatomical reduction of the fracture, which meant, putting each and every bone fragment back to where it belonged. The fragments were rigidly fixed using AO principles (as explained earlier). The aim of rigid fixation was to achieve bone-to-bone healing, without callus formation (primary bone healing). In an attempt to achieve anatomical reduction and rigid fixation, it became necessary to widely expose the fracture, and thus damage its blood supply. It was soon realised that, although this method is mechanically superior, it does not respect the biological environment of the fracture, and hence often resulted in problems in fracture healing.

**CURRENT AO CONCEPTS**

Hence, in the 90's, the concept changed from rigid fixation to 'stable fixation'. By stable fixation it meant that the fixation should be good enough to achieve union. In intra-articular fractures, stable fixation meant anatomical reduction and absolute stable fixation, the concept similar to early AO concept. But for diaphyseal fractures, anatomical reduction is not necessary. A more biological, less-than-rigid (relative stable) fixation is compatible with healing. For this, exact anatomical reduction is not considered necessary. Just functional reduction (achieving length and overall alignment) is okay. For example, recommendations in the past for a comminuted fracture of the femur was to reconstruct the femur, fixing each and every small piece of bone to where it belonged, using small screws and plates. It was more like solving a jigsaw puzzle. It caused a lot of damage to blood supply of individual fragments, and hence delayed union. Now, the same fracture is 'stabilized' by using a bridging plate (Fig-5.6), or a nail where the length

![Fig-5.6 Biological fixation: the fracture is stabilised without touching the small comminuted pieces of bone (bridging plate)](https://kat.cr/user/Blink99/)
of the bone and its alignment is restored, without exposing the comminuted segment of the bone. The 'relative stability' provided by these devices is good enough to allow early mobilisation, but since the blood supply of the bone is preserved, the chances of fracture healing are better.

In order to preserve the blood supply of the fracture, emphasis has shifted from direct reduction where fragments are exposed and directly reduced, to indirect reduction where reduction is achieved by manipulating the limb without touching the fracture.

In general, intra-medullary nailing has been considered a mechanically superior device compared to plating. This is because a nail is a load sharing device (the load is shared by nail and bone). Intra-medullary nailing is a preferred option for fractures of long bones. The only disadvantage of the conventional nailing was that it did not provide rotational stability to the fracture. For this, the conventional nail has been modified. Holes have been made at the two ends of the nail. After the nail is inserted into the medullary canal, it is locked in place with the help of two bolts (Fig-5.7). This gives rotational stability to the fixation, and results in improved stability. The technique of 'interlock nailing' is state-of-the-art treatment for fixation of long bone diaphyseal fractures. An image intensifier (Fig-5.8), a special fracture table and surgical experience are pre-requisites for this technically demanding procedure.

There has been a change in the design of the conventional plates too. The new plates are so designed that they are in contact with the bone at minimum surface (low contact dynamic compression plates – LCDCP). This allows better vascularization of the fracture under the plate.

The latest development in plating technique is Locking Compression Plate (LCP). This plate has a specially designed 'combination' screw hole. The screw hole has two halves; one half is like a conventional DCP hole, and the other half is threaded (Fig-5.9a). The screw head also has threads so that as the screw is tightened, the head gets 'locked' in the plate (Fig-5.9b). This provides a rigid plate-screw construct, which has been found
Recent Advances in the Treatment of Fractures

LCP can be used as a compression plate, as a neutralisation plate, as a buttress plate, as a bridging plate, and as a locked plate. It is particularly suitable for peri-articular fractures and fractures in osteoporotic bones.

Carbon plates and bio-degradable* implants are also being used. They do not show up on the X-rays, and get 'absorbed' after sometime. These have yet not been widely accepted.

FUNCTIONAL BRACING

Sarmiento (1973) popularized functional bracing technique for treating fractures. In principle, the technique consists of applying an external splint (called a brace) to a fractured limb. The brace provides adequate support to the fracture while permitting function of that limb, until the union is complete. Bracing is done if the reduction of the fracture is satisfactory, and the swelling has subsided – usually 2-3 weeks after the injury.

It seems likely that the brace works by supporting the soft tissues in a tight compartment (Fig-5.10). As the limb is put to use, the axial pressure on the limb (caused by weight bearing or due to muscle contraction) tends to shorten the bone segment, producing a ‘bulging effect’ on the soft tissues. Since the whole leg is enclosed in a confined space (the brace), there develops a sort of hydraulic pressure within the brace, which helps in maintaining the fracture alignment. In long series of cases treated this way, it was found that shortening and angulation was not a significant problem.

Bracing may not reduce the time taken by the fracture to unite, but it markedly reduces the stiffness and wasting of muscles caused by immobilisation in traction or plaster. It also drastically reduces the length of rehabilitation. Functional bracing is used for simple fractures where the reduction is stable. It is also useful for a fracture fixed internally where internal fixation is inadequate. In such cases, a guarded mobilisation may be done using a brace, and thus strain on the implant is avoided. In severely compound fractures, after initial treatment with external fixator or traction, brace can be used to allow functional use of the limb.

Overall, bracing is most popular for fractures of the shaft of the tibia, humerus, and femur.

ILIZAROV’S TECHNIQUE

Gavril A. Ilizarov, a Russian surgeon revolutionized the application of external fixation in the management of difficult non-unions and limb lengthening (Fig-5.11). These used to be some of the most difficult orthopaedic problems before the advent of Ilizarov’s technique.

The basic premise of Ilizarov’s technique is that osteogenesis requires dynamic state. The dynamic state means that the site of osteogenesis (e.g., a fracture) requires either a controlled distraction or a controlled compression. This dynamic force,

Fig-5.11 Ilizarov’s fixation applied to thigh and leg

* These are implants made of special plastic, which ‘dissolves’ over a period of time.
when properly applied, causes the dormant mesenchymal cells at the non-union site to differentiate into functioning osteoblasts. This results in bone synthesis and fracture healing. The concept that compression enhances bone healing, was known even prior to Ilizarov, but the concept of *distraction osteogenesis* was put on a sound footing by Ilizarov. According to his theory in wider perspective, *any living tissue when subjected to constant stretch under biological conditions, can grow to any extent*. The biological conditions are provided by: (i) aligning the fracture with minimal damage to its vascularity, and (ii) performing an ‘osteotomy’ of the bone (e.g., in limb lengthening surgeries), without damaging its periosteal and endosteal blood supply. Such an ‘osteotomy’ was termed *corticotomy* by Ilizarov.

The whole segment of the limb is stabilised by a specially designed fixation system called *ring fixator*. This protects the growing tissues from bending or shearing forces, but permits loading in the long axis of the limb. Distraction or compression can be applied at the fracture or corticotomy site by twisting nuts on the fixation system (Fig-5.12). Distraction or compression is carried out at the rate of 1 mm per day. This is done in four sittings—½ mm, four times a day. Ring fixator application consists of inserting thin (1.5 or 1.8 mm) stainless steel wires through the bone. Outside the limb, the wires are attached to steel rings with the help of bolts. Before fixing the wire to the ring, the wire is put under tension so as to make it ‘stiff’, and thus impart stability to the fixator. The rings are interconnected with the help of threaded rods with nuts on either ends. It is by twisting these nuts that the rings can be moved up or down, and the fracture distracted or compressed. Ilizarov technique is useful in the management of the following conditions:

a) Limb lengthening, especially when the shortening is associated with deformity. In this situation the fixator assembly is so designed that it corrects the deformity and produces lengthening at the same time. Massive limb lengthening (even up to 18 inches) have been performed by using this technique. The fixator provides a stable biological environment to the new bone at the site of lengthening (Fig-5.13).

b) Non-union, especially those resistant to conventional methods of treatment or those associated with deformity or shortening. By this method, non-union, deformity as well
shortening can be treated by one-stage fixator application. Non-unions requiring skin cover and bone grafting operations can be managed by using Ilizarov technique without subjecting the patient to staged surgeries.

c) Deformity correction, which may be congenital or acquired can be corrected by Ilizarov’s technique.

d) Osteomyelitis can be treated, as this technique offers the possibility of liberal excision of necrotic bone. The gap thus created can subsequently be made up by transporting a segment of bone from either end.

e) Arthrodesis can be performed by “crushing” the articular surfaces against each other, and thus stimulating union between opposite bones.

ADVANTAGES OF ILIZAROV’S TECHNIQUE
- Immediate load bearing
- A healthy viable bone in place of devascularized bone
- Correction of more than one problems by one-stage operation.

DISADVANTAGES OF ILIZAROV’S TECHNIQUE
- Inconvenience, as the external fixator hampers normal activity
- Long duration of treatment
- Pin tract infection
- Nerve palsy by pin insertion or traction
- Joint stiffness caused by transfixation of the soft tissues by the external fixator.

Further Reading

What have we learnt?
- The AO methods of internal fixation are based on achieving inter-fragmentary compression or splintage or both. Plates and screws can be used to achieve this. Nailing is a splintage device, interlock nailing being a recent modification.
- In the treatment of diaphyseal fracture, emphasis has shifted from rigid (absolute) stability to relative stability. Anatomical reduction has been replaced by functional reduction.
- Plate fixation has undergone changes to make it a more ‘biological’ implant. Locking compression plate (LCP) is a recent addition to plating techniques.
- Functional bracing is a non-operative method of treating fractures of the long bones.
- Ilizarov method is a type of external fixator system which is versatile in its application, particularly suited for difficult non-unions, malunions and limb length discrepancies.
While examining a case of injury to the musculo-skeletal system, answers to the following questions are sought:

- Is there a fracture*?
- Is it a closed or an open fracture?
- Is it a traumatic or a pathological fracture?
- Are there any complications associated with the fracture?

**CLINICAL EXAMINATION**

With widespread availability of X-ray facilities, diagnosis of fractures and dislocations has become easy. But the clinical examination still continues to be important, especially in the following situations:

a) To decide whether an X-ray examination is needed. This is particularly relevant when a patient has to travel to far off places for X-ray.

b) To ascertain whether the injury under consideration needs a special view. For example, an oblique view of the wrist best shows a scaphoid fracture.

c) To avoid making a wrong diagnosis, by correlating the clinical findings with the radiological findings. This way, some artifacts otherwise likely to be diagnosed as ‘fracture’, are recognised.

d) To detect complications associated with a fracture e.g., injury to the neurovascular bundle.

This will be missed if a clinical examination is not carried out.

Thus, a thorough clinical examination must precede an X-ray in all cases of musculo-skeletal injury. The following questions should be kept in mind while performing the clinical examination:

**IS THERE A FRACTURE?**

Most often a fracture can be diagnosed on the basis of history and clinical examination. The following points in clinical examination need to be considered:

**Age of the patient**: Fractures occur at all ages but dislocations are uncommon in children**. Some fractures are common in a particular age group, as shown in the Table–6.1.

<table>
<thead>
<tr>
<th>Table–6.1: Common fractures at different ages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age group</td>
</tr>
<tr>
<td>------------------</td>
</tr>
<tr>
<td>At birth</td>
</tr>
<tr>
<td>In children</td>
</tr>
<tr>
<td>Epiphyseal injuries</td>
</tr>
<tr>
<td>In adults</td>
</tr>
<tr>
<td>In elderly people</td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

**Mechanism of injury**: The mechanism by which the patient sustains the injury often gives an idea

* For ease of discussion, the term ‘fracture’ is used for both, fracture and dislocation.

** In children, force around a joint produces an epiphyseal injury through the epiphyseal plate, and not a dislocation.
Table–6.2: Mechanisms of injury and fractures/dislocations

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>Common injuries</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fall on an out-stretched hand</td>
<td>Fracture clavicle</td>
</tr>
<tr>
<td>Fall with spine forced in a particular direction</td>
<td>Fractures around the elbow</td>
</tr>
<tr>
<td>Slipping in the bathroom (trivial trauma)</td>
<td>Flexion injuries, Extension injuries etc.</td>
</tr>
<tr>
<td>Dashboard injury</td>
<td>Fractures neck of the femur</td>
</tr>
<tr>
<td>Fall onto the heel</td>
<td>Fracture calcaneum</td>
</tr>
<tr>
<td>Hit by a stick</td>
<td>Fracture ulna</td>
</tr>
</tbody>
</table>

about the expected fracture/dislocation. For example, in a fall from some height onto the heels, one is likely to sustain a fracture of the calcaneum, fracture of lumbar vertebrae and fracture of pubic rami. Some common injuries and mechanisms involved are shown in Table–6.2.

**Presenting complaints:** A patient with suspected fracture may present with the following complaints:
- **Pain:** It is the commonest presenting complaint in cases of musculo-skeletal injury. The severity of pain has no bearing on the diagnosis. Sprains and strains can be as painful as fracture.
- **Swelling:** Fractures are usually accompanied with swelling. The swelling may be slight if patient presents immediately after the injury; but in those presenting late, the whole limb may be swollen, mostly because of gravitational oedema.
- **Deformity:** A fractured bone may result in deformity of that part of the body.
- **Loss of function:** Following a fracture, the patient may* be unable to use the affected limb.

**Examination:** A proper exposure of the body part is crucial to an accurate examination. At times the findings are subtle, and comparison of the injured limb with the opposite normal extremity may be useful. Joints proximal and distal to the injured bone should always be examined. In a patient of road traffic accident with multiple injuries, it is wise to expose the patient completely and examine each body part in a systematic manner. One should look for the following signs:

* Ability to keep using the limb after an injury is not conclusive of no fracture, especially in children.

- **Swelling:** Though most fractures are accompanied with swelling at the site, it can be a misleading sign as there may be minimal visible swelling in the presence of a serious fracture (e.g., fracture of the neck of the femur); on the other hand, a massive swelling may be present in the absence of a fracture (e.g., in cases of ligament sprain and muscle injuries). The swelling may be due to a haematoma, prominence of the bone ends or passive oedema.
- **Deformity:** An obvious deformity of a body part is a very specific sign of a fracture or dislocation. So characteristic is the deformity in some fractures and dislocations that a diagnosis can be made just by looking at the deformity (Table–6.3). Deformity may be absent if there is an impacted fracture.

Table–6.3: Injuries with characteristic deformities

<table>
<thead>
<tr>
<th>Deformity</th>
<th>Injury</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flattening of shoulder</td>
<td>Shoulder dislocation (anterior)</td>
</tr>
<tr>
<td>Dinner fork deformity</td>
<td>Colles’ fracture</td>
</tr>
<tr>
<td>Mallet finger</td>
<td>Avulsion of the insertion of the extensor tendon from the distal phalanx</td>
</tr>
<tr>
<td>Flexion, adduction and internal rotation of the hip</td>
<td>Posterior dislocation of the hip</td>
</tr>
<tr>
<td>Abduction, external rotation of the hip</td>
<td>Anterior dislocation of the hip</td>
</tr>
<tr>
<td>External rotation of the leg</td>
<td>Fracture neck of femur</td>
</tr>
<tr>
<td></td>
<td>Trochanteric fracture</td>
</tr>
</tbody>
</table>

- **Tenderness:** Pain elicited by direct pressure at the fracture site or by indirect pressure may suggest a fracture.

**Direct Pressure:** A localised tenderness on a subcutaneous bone, elicited by gently running the back of the thumb (Fig-6.1a) may suggest an underlying fracture. The site of maximum tenderness helps in differentiating ligament injuries from that of fractures around a joint (e.g., ankle injuries). One may feel or hear a crepitus while eliciting tenderness.

**Indirect Pressure:** It may be possible to elicit pain at the fracture site by applying pressure at a site away from the fracture. Some examples are given below:

- **Springing test:** It may be possible to elicit pain from a fracture of the forearm bones
by pressing the two bones towards each other at a distance away from the fracture (Fig-6.1b).

- **Axial pressure:** An axial pressure along the second metacarpal may elicit pain in the scaphoid fossa, in a case of scaphoid fracture.
- **Bony irregularity:** It is possible to feel bony elevations and depressions in fractures of subcutaneous bones such as the tibia. This is a definite sign of fracture.
- **Abnormal mobility:** In any limb, movements occur only at joints. If one can elicit mobility at sites other than the joints (say in the middle of the arm), or an abnormal range of movement at a joint, a fracture or dislocation is definite. One may hear or feel a crepitus while doing this.
- **Absence of transmitted movements:** Normally, if a bone is moved holding it at one end, the movement can be felt at the other end. This transmitted movement will be absent in case of a displaced fracture. In case the fracture is undisplaced or impacted, the movement will be transmitted even in the presence of a fracture.

**IS IT AN OPEN FRACTURE?**

Whenever there is a wound in the vicinity of a fracture, it is important to ascertain whether the wound is communicating with the fracture or not, in which situation, it should be taken as an open fracture. It must be ascertained whether the compounding is internal or from outside.

**IS IT A PATHOLOGICAL FRACTURE?**

A pathological fracture must be suspected if: (i) the force producing the fracture is insignificant (trivial trauma); (ii) there is a history of pain or swelling in the affected bone prior to the occurrence of fracture; (iii) there is history suggestive of frequent fractures in the past (e.g., osteogenesis imperfecta); and (iv) the patient is suffering from a debilitating systemic illness known to weaken bones (e.g., rheumatoid arthritis).

**ANY ASSOCIATED COMPLICATION?**

Complications such as injuries to the nerves and vessels etc. may be associated with a fracture. These can be diagnosed on clinical examination (see Chapter 10).

**RADIOLOGICAL EXAMINATION**

A radiological examination helps in: (i) diagnosis of a fracture or dislocation; (ii) evaluation of displacements, if any; and (iii) studying the nature of the force causing fracture. The following are some of the points to be remembered in a radiological examination of a case of skeletal injury:

**ASKING FOR AN X-RAY**

Before asking for an X-ray, the following points should be kept in mind:

- **Both**, antero-posterior and lateral views should be requested in most situations.
- **Joint above and below** should be included in the X-ray.
- **Special views** show fractures better in some cases (Table–6.4).

**Table–6.4: Some commonly used special views**

<table>
<thead>
<tr>
<th>Special View</th>
<th>Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oblique view wrist</td>
<td>Scaphoid fracture</td>
</tr>
<tr>
<td>Judet view</td>
<td>Acetabular fracture</td>
</tr>
<tr>
<td>Merchant view</td>
<td>Patello-femoral joint</td>
</tr>
<tr>
<td>Skyline view</td>
<td>Calcaneum fracture</td>
</tr>
</tbody>
</table>

- **X-ray requisition must specify** the area of suspicion e.g., if an injury to D_{12} vertebra is suspected, ask for an X-ray of the dorso-lumbar spine focusing D_{12}.
- **X-ray of the pelvis with both hips** should be asked for in all cases of suspected pelvic injury. Major
Injuries of the thigh are often associated with fractures of the pelvis, hence an X-ray of the pelvis must be taken as a routine in cases with major fractures of the leg.

- For an X-ray evaluation of the hands and feet, antero-posterior and oblique views (not lateral) are required.

**Reading an X-ray**

An X-ray view box should be used in all cases. If a fracture is obvious, one must make a note of the following points:

- Which bone is affected?
- Which part of the bone is affected? e.g., shaft etc.
- At what level is the fracture? i.e., whether the fracture is in the upper, middle or lower third.
- What is the pattern of the fracture? i.e., whether the fracture is transverse, oblique etc.
- Is the fracture displaced? If yes, in what direction, i.e., whether it is a shift (antero-posterior, sideways), a tilt, or angulation in any direction, or a rotational displacement. Rotational displacement is sometimes not visible on X-rays, and can only be diagnosed clinically.
- Is the fracture line extending into the nearby joint?
- Does the underlying bone appear pathological? e.g., a cyst, abnormal texture of the whole bone, etc.

---

**Flow chart-6.1 Approach to a patient with limb injury**

- History of injury
  - is it fracture?**
    - (A) Yes
      - Is it a closed or open fracture?
      - Is the trauma sufficient to cause fracture? (i.e., is it a pathological fracture?)
      - Is there any distal neurovascular deficit?
      - First aid
      - X-ray examination
      - Fracture obvious
        - Is it the only fracture?
        - Is it a pathological fracture?
        - Is it an old fracture?
        - Definitive treatment
      - Fracture not obvious
        - Carefully trace cortices of all the bones visible
        - Fracture seen
        - No fracture
          - Treat as (B)
    - (B) No
      - First-aid
      - X-ray examination
      - No bony injury
      - Which soft tissue is injured?
        - Sprain (ligament)
          - What degree?
            - I
            - II
            - III
          - Treatment
        - Strain (muscle)
          - Partial or complete
        - Skin contusion
          - Subcutaneous haematoma
          - Drain, Rest

---

* History of injury may not be present in pathological fractures.
** The term ‘fracture’ includes dislocation (for ease of presentation).
• Is it a fresh or an old fracture? An X-ray of a fresh fracture shows a soft tissue shadow resulting from haematoma, and the fracture ends are sharp. An old fracture shows callus formation and disuse osteoporosis and the fracture ends are smoothened.

If the fracture is not obvious immediately, all the bones and joints seen on the X-ray must be examined systematically for a break in the cortex or loss of joint congruity.

X-ray findings should be correlated with clinical findings, so as to avoid error because of some artifacts which may mimic a fracture. Also one must ensure that the part under question is visible on X-ray. An X-ray of a bone must include the joints proximal and distal to the bone. Do look for an associated injury to all the other bones and joints visible on the X-ray.

One must be aware of some normal X-ray findings which are often misinterpreted as fracture e.g., epiphyseal lines, vascular markings on bones, accessory bones etc. A comparison with the X-ray of the opposite limb helps in clearing any doubt.

There are some injuries particularly liable to be missed by a novice (Table–6.5). Before an X-ray is passed as normal, one must carefully look for these injuries.

A diagrammatic presentation of approach to a patient with a limb injury is as shown in Flow chart-6.1.

Table–6.5: Fractures commonly missed

<table>
<thead>
<tr>
<th>Upper Limb</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Greater tuberosity</td>
<td>In AP view of the shoulder, fracture of greater tuberosity is missed, as the fragment gets displaced behind the head of humerus.</td>
</tr>
<tr>
<td>AC joint subluxation</td>
<td></td>
</tr>
<tr>
<td>Posterior dislocation of shoulder</td>
<td></td>
</tr>
<tr>
<td>Head of radius, neck of the radius</td>
<td></td>
</tr>
<tr>
<td>Capitulum</td>
<td></td>
</tr>
<tr>
<td>Scaphoid</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Lower Limb</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Fracture neck of femur (Impacted)</td>
<td></td>
</tr>
<tr>
<td>Acetabulum</td>
<td></td>
</tr>
<tr>
<td>Patella</td>
<td></td>
</tr>
<tr>
<td>Calcaneum</td>
<td></td>
</tr>
<tr>
<td>Dislocation of foot</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Other</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Epiphyseal injuries</td>
<td></td>
</tr>
</tbody>
</table>

OLD FRACTURE

After 2-3 weeks, signs of a fresh fracture like pain, soft tissue swelling, tenderness etc. diminish markedly. On X-ray examination, the fracture ends will not appear sharp. Callus may be present. When a patient with fracture presents late after injury, to decide further treatment, it is important to ascertain: (i) whether the position of the fracture fragments is acceptable; and (ii) at what stage of union is the fracture? e.g., united, uniting or non-union (Fig-6.2).

APPROACH TO A POLYTRAUMA PATIENT

An isolated skeletal injury rarely poses any threat, but in association with multiple injuries, musculo-skeletal injuries assume great significance in terms of morbidity and mortality. Proper, well-articulated, early management plays a vital role in improving the outcome of these patients. The following constitute the key points in the management.

FIELD TRIAGE

Ideally, the management of a multiple injured should begin at the scene of accident where the medical technician should be able to work in coordination with the doctors. He should be able to provide basic life support and help in transportation. He should be able to decide, on the basis of the level of injury, whether the victim needs to be transferred to the local hospital or a hospital with developed trauma services. In the present day context, trauma services are not so well
developed in countries like India, but this is something to aspire for.

**TREATMENT IN THE EMERGENCY DEPARTMENT**

Once the patient reaches the emergency department, management consists of the following:

- **Primary survey:** This constitutes rapid assessment of vitals of the patient and ensuring patent airway, adequate breathing, circulation and control of external bleeding.
- **Resuscitation:** This is conducted on established lines of ABCDE (please refer to book on Anaesthesia for details).
- **Secondary survey:** Once the patient is stable, proper neurologic assessment, thoraco-abdominal assessment, genito-urinary assessment and musculo-skeletal assessment is carried out. As is obvious from the above sequence, musculo-skeletal assessment is not high on priority. The only injuries related to musculo-skeletal system which are important at this stage are: (i) vascular injuries – so check all peripheral pulses and capillary circulation; (ii) nerve injuries – check peripheral nerves and correct malposition of the fractures to relieve pressure or stretch on the affected nerve; (iii) dislocation or subluxation – these need to be corrected early. All other orthopaedic injuries can be treated once other systems have been stabilised.
- **Definitive treatment:** From orthopaedic viewpoint, this consists of planning whether some fractures need internal fixation. Those being treated coservatively, need to be reduced and immobilised. Current trend is to internally fix as many fractures as possible in a polytrauma patient, often in one sitting. This has been shown to help in nursing care, and has resulted in decrease in mortality and morbidity.

**Further Reading**


**What have we learnt?**

- When evaluating a fracture, a correlation between clinical findings and X-ray findings is a must.
- Wherever possible take AP and lateral X-rays.
- Think of the possibility of a background pathological process which may have contributed to the fracture.
- Proper X-ray requisition can prevent a lot of problems.
- Comparative X-ray of the normal, opposite side may help in diagnosing a doubtful fracture.
Complications inevitably occur in a proportion of fractures. With early diagnosis and treatment, the disability caused by these complications can be greatly reduced.

**CLASSIFICATION**

Complications of fractures can be classified into three broad groups depending upon their time of occurrence. These are as follows:

a) *Immediate* complications – occurring at the time of the fracture.

b) *Early* complications – occurring in the initial few days after the fracture.

c) *Late* complications – occurring a long time after the fracture.

Some of the complications of the fractures seen in day-to-day practice are given in Table–7.1.

**HYPOVOLAEMIC SHOCK**

Hypovolaemic shock is the commonest cause of death following fractures of major bones such as the pelvis and femur. Its frequency is on the increase due to a rise in the number of patients with multiple injuries.

**Cause:** The cause of hypovolaemia could be external haemorrhage or internal haemorrhage. External haemorrhage may result from a compound fracture with or without an associated injury to a major vessel of the limb. Internal haemorrhage is more difficult to diagnose. It is usually massive bleeding in the body cavities such as chest or abdomen. Significant blood loss may occur in fractures of the major bones like the pelvis (1500–2000 ml), and femur (1000–1500 ml).

**Management:** This begins even before the cause can be ascertained. An immediate step is to put in at least two large bore intravenous cannulas (No. 16 or No. 14). If there is peripheral vasoconstriction, no time should be wasted in performing a cut down. 2000 ml of crystalloids (preferably Ringer lactate), should be infused rapidly, followed by colloids (Haemaccel) and blood. At the earliest opportunity, effort is made to localize the site of bleed – whether it is in the chest or in the abdomen. Needle aspiration from the chest, and diagnostic peritoneal lavage provide quick information to this effect. If possible, a plain X-ray chest, and X-ray abdomen may be done. A chest tube for chest bleeding, laparotomy for abdominal bleeding, may be required.

Excessive blood loss from fractured bone may be prevented by avoiding moving the patient from one couch to another. For fractures of the pelvis, temporary stabilisation with an external fixator has been found useful in reducing haemorrhage. In advanced trauma centres, an emergency angiography and embolisation of the bleeding.

<table>
<thead>
<tr>
<th>TOPICS</th>
<th>TOPICS</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Classification</td>
<td>• Injury to viscera</td>
</tr>
<tr>
<td>• Hypovolaemic shock</td>
<td>• Infection – osteomyelitis</td>
</tr>
<tr>
<td>• Adult respiratory distress syndrome</td>
<td>• Compartment syndrome</td>
</tr>
<tr>
<td>• Fat embolism syndrome</td>
<td>• Delayed and non-union</td>
</tr>
<tr>
<td>• Deep vein thrombosis</td>
<td>• Malunion</td>
</tr>
<tr>
<td>• Crush syndrome</td>
<td>• Shortening</td>
</tr>
<tr>
<td>• Injury to major blood vessels</td>
<td>• Avascular necrosis</td>
</tr>
<tr>
<td>• Injury to nerves</td>
<td>• Stiffness of joints</td>
</tr>
<tr>
<td>• Injury to muscles and tendons</td>
<td>• Sudeck’s dystrophy</td>
</tr>
<tr>
<td>• Injury to joints</td>
<td>• Myositis ossificans</td>
</tr>
</tbody>
</table>
A vessel is performed to control bleeding from deeper vessels.

ADULT RESPIRATORY DISTRESS SYNDROME

Adult respiratory distress syndrome (ARDS) can be a sequela of trauma with subsequent shock. The exact mechanism is not known, but it is supposed to be due to release of inflammatory mediators which cause disruption of microvasculature of the pulmonary system. The onset is usually 24 hours after the injury. The patient develops tachypnoea and laboured breathing. X-ray chest shows diffuse pulmonary infiltrates. Arterial PO$_2$ falls to less than 50. Management consists of 100 percent oxygen and assisted ventilation. It takes from 4-7 days for the chest to clear, and the patient returns to normal. If not detected early, patient's condition deteriorates rapidly, he develops cardio-respiratory failure and dies.

FAT EMBOLISM SYNDROME

This is one of the most serious complications, the essential feature being occlusion of small vessels by fat globules.

Causes: The fat globules may originate from bone marrow or adipose tissue. Fat embolism is more common following severe injuries with multiple fractures and fractures of major bones. The pathogenesis of the syndrome is not clear, but it seems likely that two events occur: (i) release of free fatty acids (by action of lipases on the neutral fat), which induces a toxic vasculitis, followed by platelet-fibrin thrombosis; and (ii) actual obstruction of small pulmonary vessels by fat globules.

Consequences: Symptoms are evident a day or so after the injury. Presenting features are in the form of two, more or less distinct types: (i) cerebral; and (ii) pulmonary. In the cerebral type, the patient becomes drowsy, restless and disoriented and gradually goes into a state of coma. In the pulmonary type, tachypnoea and tachycardia are the more prominent features. The other common feature of fat embolism is a patechial rash, usually on the front of the neck, anterior axillary folds, chest or conjunctiva. If untreated, and sometimes despite treatment, the patient develops respiratory failure and dies.

Diagnosis: In a case with multiple fractures, early diagnosis may be possible by strong suspicion. In addition to the classic clinical features, signs of retinal artery emboli (striate haemorrhages and exudates) may be present. Sputum and urine may reveal the presence of fat globules. X-ray of the chest may show a patchy pulmonary infiltration (snow storm appearance). Blood PO$_2$ of less than 50 mmHg may indicate impending respiratory failure.

Treatment: This consists of respiratory support, heparinisation, intravenous low molecular weight dextran (Lomodex-20) and corticosteroids. An intravenous 5 percent dextrose solution with 5 percent alcohol helps in emulsification of fat globules, and is used by some.
DEEP VEIN THROMBOSIS (DVT) AND PULMONARY EMBOLISM

Deep vein thrombosis (DVT) is a common complication associated with lower limb injuries and with spinal injuries.

**Cause:** Immobilisation following trauma leads to venous stasis which results in thrombosis of veins. DVT proximal to the knee is a common cause of life threatening complication of pulmonary embolism. DVT can be recognised as early as 48 hours after the injury. Embolism occurs, usually 4-5 days after the injury.

**Consequences:** DVT can be diagnosed early with high index of suspicion. The group of patients ‘at risk’ include the elderly and the obese patients. Leg swelling and calf tenderness are usual signs. The calf tenderness may get exaggerated by passive dorsiflexion of the ankle (Homan’s sign). Definitive diagnosis can be made by venography. One should keep a patient of DVT on constant watch for development of pulmonary embolism. This can be suspected if the patient develops tachypnoea and dyspnoea, usually 4-5 days after the accident. There may be chest pain or haemoptysis.

**Treatment** of DVT is elevation of the limb, elastic bandage and anticoagulant therapy. For pulmonary embolism, respiratory support and heparin therapy is to be done. Early internal fixation of fractures, so as to allow early, active mobilisation of the extremity is an effective means of prevention of DVT, and hence of pulmonary embolism.

CRUSH SYNDROME

This syndrome results from massive crushing of the muscles, commonly associated with crush injuries sustained during earthquakes, air raids, mining and other such accidents. A similar effect may follow the application of a tourniquet for an excessive period.

**Causes:** Crushing of muscles results in entry of myohaemoglobin into the circulation, which precipitates in renal tubules, leading to acute renal tubular necrosis.

**Consequences:** Acute tubular necrosis produces signs of deficient renal functions such as scanty urine, apathy, restlessness and delirium. It may take 2-3 days for these features to appear.

<table>
<thead>
<tr>
<th>Vessel injured</th>
<th>Trauma</th>
</tr>
</thead>
<tbody>
<tr>
<td>Femoral</td>
<td>Fracture lower third of femur</td>
</tr>
<tr>
<td>Popliteal</td>
<td>Supracondylar fracture of the femur</td>
</tr>
<tr>
<td>Posterior tibial</td>
<td>Dislocation of the knee, Fracture tibia</td>
</tr>
<tr>
<td>Subclavian</td>
<td>Fracture of the clavicle</td>
</tr>
<tr>
<td>Axillary</td>
<td>Fracture-dislocation of the shoulder</td>
</tr>
<tr>
<td>Brachial</td>
<td>Supracondylar fracture of the humerus</td>
</tr>
</tbody>
</table>

**Table-7.2: Vascular injuries and skeletal trauma**

**Treatment:** In a case with crushed limb, first aid treatment may necessitate the application of a tourniquet, which is gradually released, so that deleterious substances are released into the circulation in small quantities. If oliguria develops, the patient is treated as for acute renal failure.

INJURY TO MAJOR BLOOD VESSELS

Blood vessels lie in close proximity to bones, and hence are liable to injury with different fractures and dislocations (Table-7.2). The popliteal artery is the most frequently damaged vessel in musculo-skeletal injuries.

**Causes:** The artery may be damaged by the object causing the fracture (e.g., bullet), or by a sharp edge of a bone fragment (e.g., supracondylar fracture of the humerus). The damage to the vessel may vary from just a pressure from outside to a complete rupture.

**Consequences.** Obstruction to blood flow will not always lead to gangrene. Where the collateral circulation is good, the following may result:

- **No effect:** If collateral circulation of the limb around the site of vascular damage is good, there will be no adverse effect of the vascular injury.
- **Exercise ischaemia:** The collaterals are good enough to keep the limb viable but any further demand on the blood supply during exercise, causes ischaemic pain (vascular claudication).
- **Ischaemic contracture:** If the collaterals do not provide adequate blood supply to the muscles, there results an ischaemic muscle necrosis. This is followed by contracture and fibrosis of the necrotic muscles, leading to deformities (e.g., Volkmann’s ischaemic contracture, see page 102).
- **Gangrene:** If the blood supply is grossly insufficient, gangrene occurs.
**Diagnosis:** The pulses distal to the injury should be examined in every case of a fracture or dislocation. Some of the features which suggest a possible vascular injury of a limb are listed below:

a) *Signs at the fracture site:* The following signs may be present at the fracture site:
   - Rapidly increasing swelling
   - Massive external bleeding (in open fractures)
   - A wound in the normal anatomical path of the vessel

b) *Signs in the limb distal to the fracture:* The following signs may be present in the limb distal to the fracture (five P’s):
   - Pain – cramplike
   - Pulse – absent
   - Pallor
   - Paraesthesias
   - Paralysis

As a matter of rule, absent peripheral pulses in an injured limb should be considered to be due to vascular damage unless proved otherwise. The confirmation of obstruction to blood flow in a vessel and its site can be easily done by Doppler study. In the absence of such a facility, there is no need to waste crucial time by ordering an angiogram merely for confirmation of diagnosis.
An angiogram may be justified in cases with multiple fractures in the same limb, where it may help in localising the site of the vascular injury.

**Treatment:** Early diagnosis and urgent treatment are of paramount importance because of the serious consequences that may follow. Correct treatment at the site of first contact (Flow chart-7.1), followed by referral to a centre equipped with facilities to treat vascular injuries is essential. In case exploration of the vessel is required, the fracture should be suitably stabilised using internal or external fixation.

**INJURY TO NERVES**

Nerves lie in close proximity to bones, and hence are liable to damage in different fractures or dislocations (Table–7.3). The radial nerve is the most frequently damaged nerve in musculoskeletal injuries. Nerves and vessels lie together in limbs, and so are often injured together.

**Causes:** A nerve may be damaged in one of the following ways:

- By the agent causing the fracture (e.g., bullet).
- By direct pressure by the fracture ends at the time of fracture or during manipulation.
- Traction injury at the time of fracture, when the fracture is being manipulated or during skeletal traction.
- Entrapment in callus at the fracture site.

**Consequences:** Damage to the nerve may be neurapraxia, axonotmesis, or neurotmesis. It may result in a variable degree of motor and sensory loss along the distribution of the nerve (see Chapter 10, Peripheral Nerve Injuries).

**Treatment:** This depends upon the type of fracture, whether it is closed or open. When the nerve injury is associated with a closed fracture, the type of damage is generally neurapraxia or axonotmesis, and nerve recovery is good with conservative treatment. In case the fracture per se needs open reduction for other reasons, the nerve should also be explored. When associated with an open fracture, the type of nerve damage is often neurotmesis. In such cases, the nerve should be explored and repaired as per need, and the fracture fixed internally with nail, plate etc.

**INJURY TO MUSCLES AND TENDONS**

Some degree of damage to muscles and tendons occurs with most fractures. It may result from the object causing the fracture (e.g., an axe), or from the sharp edge of the fractured bone. Often these injuries are overshadowed by more alarming fractures, and are detected only late, when the joint distal to the fracture becomes stiff and deformed due to scarring of the injured muscle.

Rest to the injured muscle and analgesics is enough in cases with partial rupture. A complete rupture requires repair. Rarely, if rupture of a tendon or muscle is detected late, reconstruction may be required.

**INJURY TO JOINTS**

Fractures near a joint may be associated with subluxation or dislocation of that joint. This combination is becoming more frequent due to high-velocity traffic accidents. Early open reduction and stabilisation of the fracture to permit early joint movements has improved the results.

**INJURY TO VISCERA**

Visceral injuries are seen in pelvic and rib fractures. Their management is discussed in Surgery textbooks.

### Table–7.3: Nerve injuries and skeletal trauma

<table>
<thead>
<tr>
<th>Nerve</th>
<th>Trauma</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Axillary nerve</td>
<td>Dislocation of the shoulder</td>
<td>Deltoid paralysis</td>
</tr>
<tr>
<td>Radial nerve</td>
<td>Fracture shaft of the humerus</td>
<td>Wrist drop</td>
</tr>
<tr>
<td>Median nerve</td>
<td>Supracondylar fracture of humerus</td>
<td>Pointing index</td>
</tr>
<tr>
<td>Ulnar nerve</td>
<td>Fracture medial epicondyle humerus</td>
<td>Claw hand</td>
</tr>
<tr>
<td>Sciatic nerve</td>
<td>Posterior dislocation of the hip</td>
<td>Foot drop due to weakness of dorsiflexors of the foot</td>
</tr>
<tr>
<td>Common peroneal nerve</td>
<td>Knee dislocation</td>
<td>Foot drop</td>
</tr>
<tr>
<td></td>
<td>Fracture of neck of the fibula</td>
<td></td>
</tr>
</tbody>
</table>
INFECTION – OSTEOMYELITIS

Causes: Infection of the bone is an early complication of fractures. It occurs more commonly in open fractures, particularly in those where compounding occurs from outside (external compounding). The increasing use of operative methods in the treatment of fractures is responsible for the rise in the incidence of infection of the bone, often years later. Infection may be superficial, moderate (osteomyelitis), or severe (gas gangrene).

Treatment: Proper care of an open fracture can prevent osteomyelitis. Once infection occurs, it should be adequately treated.

COMPARTMENT SYNDROME

The limbs contain muscles in compartments enclosed by bones, fascia and interosseous membrane (Fig-7.1). A rise in pressure within these compartments due to any reason may jeopardize the blood supply to the muscles and nerves within the compartment, resulting in what is known as “compartment syndrome”.

Causes: The rise in compartment pressure can be due to any of the following reasons:
- Any injury leading to oedema of muscles.
- Fracture haematoma within the compartment.
- Ischaemia to the compartment, leading to muscle oedema.

Consequences: The increased pressure within the compartment compromises the circulation leading to further muscle ischaemia. A vicious cycle is thus initiated (Fig-7.2) and continues until the total vascularity of the muscles and nerves within the compartment is jeopardized. This results in ischaemic muscle necrosis and nerve damage. The necrotic muscles undergo healing with fibrosis,

![Fig-7.1 Osseo-fascial compartment](https://kat.cr/user/Blink99/)

leading to contractures. Nerve damage may result in motor and sensory loss. In an extreme case, gangrene may occur.

Diagnosis: Compartment syndrome can be diagnosed early by high index of suspicion. Excessive pain, not relieved with usual doses of analgesics, in a patient with an injury known to cause compartment syndrome must raise an alarm in the mind of the treating doctor. Injuries with a high risk of developing compartment syndrome are as follows:
- Supracondylar fracture of the humerus
- Forearm bone fractures
- Closed tibial fractures
- Crush injuries to leg and forearm.

Stretch test: This is the earliest sign of impending compartment syndrome. The ischaemic muscles, when stretched, give rise to pain. It is possible to stretch the affected muscles by passively moving the joints in direction opposite to that of the damaged muscle’s action. (e.g., passive extension of fingers produces pain in flexor compartment of the forearm).

Other signs include a tense compartment, hypoaesthesia in the distribution of involved nerves, muscle weakness etc. Compartment syndrome can be confirmed by measuring compartment pressure. A pressure higher than 40 mm of water is indicative of compartment syndrome. Pulses may remain palpable till very late in impending compartment syndrome, and should not provide a false sense of security that all is well.

Treatment: A close watch for an impending compartment syndrome and effective early preventive measures like limb elevation, active finger movements etc. can prevent this serious complication. Early surgical decompression

![Fig-7.2 Eaton and Green cycle for compartment syndrome](https://kat.cr/user/Blink99/)
is necessary in established cases. This can be performed by the following methods:

- **Fasciotomy:** The deep fascia of the compartment is slit longitudinally (e.g., in forearm).
- **Fibulectomy:** The middle third of the fibula is excised in order to decompress all compartments of the leg.

**DELAYED AND NON-UNION**

When a fracture takes more than the usual time to unite, it is said to have gone in *delayed union*. A large percentage of such fractures eventually unite. In some, the union does not progress, and they fail to unite. These are called *non-union*. Conventionally, it is not before 6 months that a fracture can be declared as non-union. It is often difficult to say whether the fracture is in delayed union, or has gone into non-union. Only progressive evaluation of the X-rays over a period of time can solve this issue. Presence of mobility at the fracture after a reasonable period is surely a sign of non-union. Presence of pain at the fracture site on using the limbs also indicates non-union. Non-union may be painless if pseudo joint forms between the fracture ends (pseudoarthrosis).

**Causes:** Some of the factors responsible for delayed union are given in Table-7.4. In any given case, there may be one or more factors operating.

**Types of non-union:** There are two main types of non-unions (Fig-7.3):

- Atrophic, where there is minimal or no attempt at callus formation.

<table>
<thead>
<tr>
<th>Table-7.4: Causes of delayed and non-union, and their common sites</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Causes related to the patient</strong></td>
</tr>
<tr>
<td>Age: Common in old age</td>
</tr>
<tr>
<td>Associated systemic illness: Malignancy, Osteomalacia</td>
</tr>
<tr>
<td><strong>Causes related to fracture</strong></td>
</tr>
<tr>
<td><em>Distraction at the fracture site</em></td>
</tr>
<tr>
<td>- Muscle pulling the fragments</td>
</tr>
<tr>
<td>- Fracture patella</td>
</tr>
<tr>
<td>- Fracture olecranon</td>
</tr>
<tr>
<td>- Gravity</td>
</tr>
<tr>
<td>- Fracture shaft of humerus</td>
</tr>
<tr>
<td><strong>Soft tissue interposition</strong></td>
</tr>
<tr>
<td>- Fracture shaft of humerus</td>
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<tr>
<td>- Fracture shaft of femur</td>
</tr>
<tr>
<td>- Fracture medial malleolus (abduction type)</td>
</tr>
<tr>
<td><strong>Bone loss at the time of fracture</strong></td>
</tr>
<tr>
<td>- Fracture tibia (open type)</td>
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<tr>
<td>- Fracture ulna (open type)</td>
</tr>
<tr>
<td><strong>Infection from an open fracture</strong></td>
</tr>
<tr>
<td>- Fracture tibia</td>
</tr>
<tr>
<td><strong>Damage to blood supply of fracture fragments</strong></td>
</tr>
<tr>
<td>- Fracture neck of femur</td>
</tr>
<tr>
<td>- Fracture lower third of tibia</td>
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<tr>
<td><strong>Pathological fracture</strong></td>
</tr>
<tr>
<td>- Fracture of the shaft of the femur</td>
</tr>
<tr>
<td>- Fracture of humerus</td>
</tr>
<tr>
<td><strong>Causes related to treatment</strong></td>
</tr>
<tr>
<td><em>Inadequate reduction</em></td>
</tr>
<tr>
<td>- Fracture shafts of long bones</td>
</tr>
<tr>
<td><em>Inadequate immobilisation</em></td>
</tr>
<tr>
<td>- Fracture shafts of long bones</td>
</tr>
<tr>
<td>- Fracture neck of femur</td>
</tr>
<tr>
<td><strong>Distraction during treatment</strong></td>
</tr>
<tr>
<td>- Fracture shaft of femur</td>
</tr>
</tbody>
</table>

- Hypertrophic, where though the callus is present, it does not bridge the fracture site.

**Common sites:** Sites where non-union occurs commonly are neck of the femur, scaphoid, lower third of the tibia, lower third of the ulna and lateral condyle of the humerus.

**Consequences:** Delayed and non-union can result in persistent pain, deformity, or abnormal mobility at the fracture site. A fracture in delayed union, if stressed, can lead to refracture.

**Diagnosis:** Delayed union is a diagnosis in relation to time. The fracture may not show any abnormal signs clinically, but X-rays may fail to show bony union. The following are some of the clinical findings which suggest delayed union and non-union:
• Persistent pain
• Pain on stressing the fracture
• Mobility (in non-union)
• Increasing deformity at the fracture site (in non-union).

The following are some of the radiological features suggestive of these complications:
• Delayed union: The fracture line is visible. There may be inadequate callus bridging the fracture site.
• Non-union: The fracture line is visible. There is little bridging callus. The fracture ends may be rounded, smooth and sclerotic. The medullary cavity may be obliterated.

It is sometimes very difficult to be sure about union of a fracture where internal fixation has been used. Evaluation of serial X-rays may help detect subtle angulation, non-progress of bridging callus, resorption of callus, loosening of screws and bending of the nail or plate. Excessive rotation may be the only abnormal mobility in a case with intramedullary rod in situ. Oblique views, done under fluoroscopy may show an unhealed fracture better than conventional AP and lateral X-rays. It may be possible to demonstrate mobility at the fracture by stress X-rays or weight bearing X-rays. 3-D CT scan is sometimes helpful in differentiating between delayed and non-union.

Treatment: Most fractures in delayed union unite on continuing the conservative treatment. Sometimes, this may not occur and the fracture may need surgical intervention. Bone grafting with or without internal fixation may be required. Treatment of non-union depends upon the site of non-union and the disability caused by it. The following possibilities of treatment should be considered, depending upon the individual cases.

• Open reduction, internal fixation and bone grafting: This is the commonest operation performed for non-union. The grafts are taken from iliac crest. Internal fixation is required in most cases.
• Excision of fragments: Sometimes, achieving union is difficult and time consuming compared to excision of one of the fragments. This can only be done where excision of the fragment will not cause any loss of functions. An excision may or may not need to be combined with replacement with an artificial mould (prosthesis). For example, the lower-end of the ulna can be excised for non-union of the fracture of the distal-end of the ulna without much loss. In non-union of fracture of the neck of femur in an elderly, the head of the femur can be replaced by a prosthesis (replacement arthroplasty).
• No treatment: Some non-unions do not give rise to any symptoms, and hence require no treatment, e.g., some non-unions of the fracture scaphoid.
• Ilizarov’s method: Prof. Ilizarov from the former USSR designed a special external fixation apparatus for treating non-union (see page 33).

MALUNION
When a fracture does not unite in proper position, it is said to have malunited. A slight degree of malunion occurs in a large proportion of fractures, but in practice the term is reserved for cases where the resulting disability is of clinical significance.

Causes: Improper treatment is the commonest cause. Malunion is therefore preventable in most cases by keeping a close watch on position of the fracture during treatment. Sometimes, malunion is inevitable because of unchecked muscle pull (e.g., fracture of the clavicle), or excessive comminution (e.g., Colles’ fracture).

Common sites: Fractures at the ends of a bone always unite, but they often malunite e.g., supracondylar fracture of the humerus, Colles’ fracture etc.

Consequences: Malunion results in deformity, shortening of the limb, and limitation of movements.

Treatment: Each case is treated on its merit. A slight degree of malunion may not require any treatment, but a malunion producing significant disability, especially in adults, needs operative intervention. The following treatment possibilities can be considered:

a) Treatment required: Malunion may require treatment because of deformity (e.g., supracondylar fracture of the humerus), shortening (e.g., fracture of the shaft of the femur) or functional limitations (e.g. limitation of rotations in malunion of forearm fractures). Some of the methods for treating malunion are as follows:
• **Osteoclasis** (refracturing the bone): It is used for correction of mild to moderate angular deformities in children. Under general anaesthesia the fracture is recreated, the angulation corrected, and the limb immobilised in plaster.

• **Redoing the fracture surgically**: This is the most commonly performed operation for malunion. The fracture site is exposed, the malunion corrected and the fracture fixed internally with suitable implants. Bone grafting is also performed, in addition, in most cases e.g., malunion of long bones.

• **Corrective osteotomy**: In some cases, redoing the fracture, as discussed above may not be desirable due to variety of reasons such as poor skin condition, poor vascularity of bone in that area etc. In such cases, the deformity is corrected by osteotomy at a site away from the fracture as the healing may be quicker at this new site, e.g. supra-malleolar corrective osteotomy for malunion of distal-third tibial fractures.

• **Excision of the protruding bone**: In a fracture of the clavicle, a bone spike protruding under the skin may be shaved off. Same may be required in a spikey malunion of fracture of the shaft of the tibia.

b) **No treatment**: Sometimes malunion may not need any treatment, either because it does not cause any disability, or because it is expected to correct by remodelling. Remodelling of a fracture depends on the following factors.

• **Age**: Remodelling is better in children.

• **Type of deformity**: Sideways shifts are well corrected by remodelling. Five to ten degrees of angulation may also get corrected, but mal-rotation does not get corrected.

• **Angulation in the plane of movement of the adjacent joint** is remodelled better than that in other planes e.g., posterior angulation in a fracture of the tibial shaft remodels better.

• **Location of fracture**: Fractures near joints remodel better.

**Cross union** is a special type of malunion which occurs in fractures of the forearm bones, wherein the two bones unite with each other. For details please refer to page 110.

**SHORTENING**

**Causes**: It is a common complication of fractures, resulting from the following causes:

• **Malunion**: The fracture unites with an overlap or marked angulation e.g., most long bone fractures.

• **Crushing**: Actual bone loss e.g., bone loss in gunshot wounds.

• **Growth defect**: Injury to the growth plate may result in shortening (see Salter-Harris classification of epiphyseal injuries, page 58).

**Treatment**: A little shortening in upper limbs goes unnoticed, hence no treatment is required. For shortening in lower limbs, treatment depends upon the amount of shortening.

• Shortening *less than 2 cm* is not much noticeable, hence can be compensated by a shoe raise.

• Shortening *more than 2 cm* is noticeable. In elderly patients, it may be compensated for by raising the shoe on the affected side. In younger patients, correction of angulation or overlap by operative method is necessary. Limb length equalisation procedure is required to correct shortening in an old, healed, remodelled fracture.

**AVASCULAR NECROSIS**

Blood supply of some bones is such that the vascularity of a part of it is seriously jeopardized following fracture, resulting in necrosis of that part.

**Common sites**: Some of the sites where avascular necrosis commonly occurs are given in Table–7.5.

**Consequences**: Avascular necrosis causes deformation of the bone. This leads to secondary osteoarthritis a few years later, thus causing painful limitation of joint movement.

---

**Table–7.5: Common sites of avascular necrosis**

<table>
<thead>
<tr>
<th>Site</th>
<th>Cause</th>
</tr>
</thead>
<tbody>
<tr>
<td>Head of the femur</td>
<td>Fracture neck of the femur.</td>
</tr>
<tr>
<td></td>
<td>Posterior dislocation of the hip</td>
</tr>
<tr>
<td>Proximal pole of</td>
<td>Fracture through the waist of the scaphoid</td>
</tr>
<tr>
<td>scaphoid</td>
<td></td>
</tr>
<tr>
<td>Body of the talus</td>
<td>Fracture through neck of the talus</td>
</tr>
</tbody>
</table>
Complications of Fractures

Diagnosis: Avascular necrosis should always be suspected in fracture where it is known to occur. Pain and stiffness appear rather late. Radiological changes as given below appear earlier (Fig-7.4).

- **Sclerosis of necrotic area:** The avascular bone is unable to share disuse osteoporosis as occurs in the surrounding normal bones. Hence, it stands out densely on the X-ray.
- **Deformity of the bone** occurs because of the collapse of necrotic bone.

Excision of the avascular segment of bone where doing so does not hamper functions e.g. fracture of the scaphoid.

Excision followed by replacement e.g., in fracture of the neck of the femur, the avascular head can be replaced by a prosthesis.

Total joint replacement or arthrodesis may be required once the patient is disabled because of pain from osteoarthritis secondary to avascular necrosis.

STIFFNESS OF JOINTS

It is a common complication of fracture treatment. Shoulder, elbow and knee joints are particularly prone to stiffness following fractures.

Causes: The following are some of the causes of joint stiffness:

- Intra-articular and peri-articular adhesions secondary to immobilisation, mostly in intra-articular fractures.
- Contracture of the muscles around a joint because of prolonged immobilisation.
- Tethering of muscle at the fracture site (e.g. quadriceps adhesion to a fracture of femoral shaft).
- Myositis ossificans (refer page 52).

Consequences: Stiff joints hamper normal physical activity of the patient.

Treatment: The treatment is heat therapy (hot fomentation, wax bath, diathermy etc.) and exercises. Sometimes, there may be a need for manipulating the joint under general anaesthesia. Surgical intervention is required in the following circumstances:

- To excise intra-articular adhesions, preferably arthroscopically.
- To excise an extra-articular bone block which may be acting as a ‘door stopper’.
- To lengthen contracted muscles.
- Joint replacement, if there is pain due to secondary osteoarthritis.

REFLEX SYMPATHETIC DYSTROPHY  
(SUDECK’S DYSTROPHY)

This is a term given to a group of vague painful conditions observed as a sequelae of trauma. The trauma is sometimes relatively minor, and hence symptoms and signs are out of proportion to the trauma.
Consequences: Clinical features consist of pain, hyperaesthesia, tenderness and swelling. Skin becomes red, shiny and warm in early stage. Progressive atrophy of the skin, muscles and nails occur in the later stage. Joint deformities and stiffness ensues. X-ray shows characteristic spotty rarefaction.

Treatment: It is a difficult condition to explain to the patient, and also the treatment is prolonged. Physiotherapy constitutes the principle modality of treatment. Further trauma in the form of an operation or forceful mobilisation is detrimental. In some cases, beta blockers have been shown to produce good response. In resistant cases, sympathetic blocks may aid in recovery. Prolonged physiotherapy and patience on the part of the doctor and the patient is usually rewarding.

MYOSITIS OSSIFICANS (POST-TRAUMATIC OSSIFICATION)

This is ossification of the haematoma around a joint, resulting in the formation of a mass of bone restricting joint movements, often completely.

Causes: It occurs in cases with severe injury to a joint, especially when the capsule and the periosteum have been stripped from the bones by violent displacement of the fragments. It is common in children because in them the periosteum is loosely attached to the bones. It is particularly common around the elbow joint. There is also a relatively high incidence in patients with prolonged or permanent neuronal damage from head injury, and in patients with paraplegia. Massage following trauma is a factor known to aggravate myositis.

Consequences: The bone formation leads to stiffness of the joint, either due to thickening of the capsule or due to the bone blocking movement. In extreme cases, the bone bridges the joint resulting in complete loss of movements (extra-articular ankylosis).

Radiologically, an active myositis and a mature myositis have been identified. In the former, the margins of the bone mass are fluffy (Fig-7.5); in the latter the bone appears trabeculated with well-defined margins.

Treatment: Massage following injury is strictly prohibited. In the early active stage of myositis the limb should be rested, and NSAID is given. In late stages, it is possible to regain movement by physiotherapy. In some cases, once the myositis mass matures, surgical excision of the bone mass may help regain movement.

Further Reading

What have we learnt?
- A neurovascular examination of the limb distal to the fracture is a must in every patient with fracture. It is a disastrous complication, if not attended to in time.
- Not all delayed union, non-union and malunion need surgical treatment. Some are quite compatible with normal functions. Treatment has to be tailored to patient’s need.
- Active joint mobilisation is necessary to prevent joint stiffness.
- No massage after a fracture or joint injury. It can lead to myositis ossificans.
Additional information: From the entrance exams point of view

- Basic pathology in myositis ossificans lies in the muscle fibres.
- Most common location for myositis ossificans is elbow, next common is hip.
- In myositis ossificans mature bone is seen in the periphery and immature bone in the centre.
- Myositis ossificans progressiva (fibrodysplasia): The life expectancy decreases and the most common cause of death is lung disease. It affects children before the age of 6 and involves deformities of spine, hands and feet.
- Bone scan (Tc99 three phase scan) is the most sensitive for early detection of heterotrophic ossification.
- Alkaline phosphatase and 24 hrs PGE2 urinary excretion are screening tests for heterotrophic ossification.
It is important to first understand the factors responsible for the stability of a joint in order to understand why a particular joint dislocates more often than another. Normally, a joint is held in position because of the inherent stability in its design, by the ligaments, and by the surrounding muscles, as discussed below:

The shape of a joint: The shape of the articulating surfaces in themselves may provide great security against displacement, e.g., the hip joint with its deep socket (the acetabulum) and an almost spherical ball (the femoral head) is a good design from the stability viewpoint. On the other hand, the shoulder joint with its shallow socket (the glenoid) and a large ball (the humeral head) is a poor design and therefore dislocates more easily than the hip joint.

The ligaments: These prevent any abnormal mobility of a joint and are called static stabilisers. The role of the ligaments in providing stability to a joint is variable. In some joints (e.g., the knee and finger joints), ligaments form the main stabilising structures, whereas in others (e.g., the hip or shoulder) they do not play an important role.

The muscles: A strong muscle cover around a joint gives it stability. Muscles may also provide a supporting function to the ligaments by reflexly contracting to protect the ligaments, when the latter come under harmful stresses. These are, therefore, called the dynamic stabilisers of a joint.

Dislocation: A joint is dislocated when its articular surfaces are completely displaced, one from the other, so that all contact between them is lost (see Fig-1.4, page 5).

Subluxation: A joint is subluxated when its articular surfaces are only partly displaced and retain some contact between them.

Dislocations and subluxations may be classified on the basis of aetiology into congenital or acquired. Congenital dislocation is a condition where a joint is dislocated at birth e.g., congenital dislocation of the hip (CDH). Acquired dislocation may occur at any age. It may be traumatic or pathological as discussed below.

<table>
<thead>
<tr>
<th>Common dislocations at different joints</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spine</td>
</tr>
<tr>
<td>Hip</td>
</tr>
<tr>
<td>Shoulder</td>
</tr>
<tr>
<td>Elbow</td>
</tr>
<tr>
<td>Wrist</td>
</tr>
<tr>
<td>MP joint</td>
</tr>
<tr>
<td>Knee</td>
</tr>
<tr>
<td>Patella</td>
</tr>
<tr>
<td>Ankle</td>
</tr>
<tr>
<td>Foot</td>
</tr>
<tr>
<td>Cervical spine</td>
</tr>
<tr>
<td>Posterior, anterior</td>
</tr>
<tr>
<td>Anterior (commonest overall), posterior</td>
</tr>
<tr>
<td>Posterior, postero-lateral</td>
</tr>
<tr>
<td>Lunate, perilunate</td>
</tr>
<tr>
<td>Dorsal (index finger)</td>
</tr>
<tr>
<td>Posterior</td>
</tr>
<tr>
<td>Lateral</td>
</tr>
<tr>
<td>Antero-lateral</td>
</tr>
<tr>
<td>Chopart’s dislocation</td>
</tr>
<tr>
<td>Lisfranc’s dislocation</td>
</tr>
</tbody>
</table>
Injury to Joints: Dislocation and Subluxation

Traumatic dislocation: Injury is by far the commonest cause of dislocations and sub-luxations at almost all joints (Table–8.1). The force required to dislocate a particular joint varies from joint to joint. The following are the different types of traumatic dislocations seen in clinical practice:

a) Acute traumatic dislocation: This is an episode of dislocation where the force of injury is the main contributing factor e.g., shoulder dislocation.

b) Old unreduced dislocation: A traumatic dislocation, not reduced, may present as an old unreduced dislocation e.g., old posterior dislocation of the hip.

c) Recurrent dislocation: In some joints, proper healing does not occur after the first dislocation. This results in weakness of the supporting structures of the joint so that the joint dislocates repeatedly, often with trivial trauma. Recurrent dislocation of the shoulder and patella are common.

d) Fracture-dislocation: When a dislocation is associated with a fracture of one or both of the articulating bones, it is called fracture-dislocation. A dislocation of the hip is often associated with a fracture of the lip of the acetabulum.

Pathological dislocation: The articulating surfaces forming a joint may be destroyed by an infective or a neoplastic process, or the ligaments may be damaged due to some disease. This results in dislocation or subluxation of the joint without any trauma e.g., dislocation of the hip in septic arthritis.

PATHOANATOMY

Dislocation cannot occur without damage to the protective ligaments or joint capsule. Usually the capsule and one or more of the reinforcing ligaments are torn, permitting the articular end of the bone to escape through the rent. Sometimes, the capsule is not torn in its substance but is stripped from one of its bony attachments (Fig-8.1). Rarely, a ligament may withstand the force of the injury so that instead of ligament rupture, a fragment of bone at one of its attachments may be chipped off (avulsed).

At the time of dislocation, as movement occurs between the two articulating surfaces, a piece of articular cartilage with or without its underlying bone may be ‘shaved off’ producing an osteochondral fragment within the joint. This fragment may lie loose inside the joint and may cause symptoms long after the dislocation is reduced (Fig-8.2).

DIAGNOSIS

Clinical examination: In most cases of dislocation, the clinical features are sufficiently striking and make the diagnosis easy. Never-theless, a dislocation or subluxation is sometimes overlooked, especially in a multiple injury case, an unconscious patient or in a case where the bony landmarks are obscured by severe swelling or obesity. Some dislocations, which are particularly notorious for getting overlooked are: (i) posterior dislocation of the shoulder especially in an epileptic; and (ii) dislocation of the hip associated with a fracture of the shaft of the femur on the same side. The classic deformity of a hip dislocation does not occur, and the attention is drawn on the more obvious injury – the femoral shaft fracture. Some of the salient clinical features of dislocation are as follows:

Fig-8.1 Pathoanatomy of dislocations

Fig-8.2 X-ray showing osteochondral fragment
Table–8.2: Typical deformities in dislocations

<table>
<thead>
<tr>
<th>Joint (dislocation)</th>
<th>Deformity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shoulder (anterior)</td>
<td>Abduction</td>
</tr>
<tr>
<td>Elbow (posterior)</td>
<td>Flexion</td>
</tr>
<tr>
<td>Hip</td>
<td>Flexion, adduction, internal rotation</td>
</tr>
<tr>
<td>Ankle</td>
<td>Varus</td>
</tr>
</tbody>
</table>

- **Pain:** Dislocations are very painful.
- **Deformity:** In most dislocations the limb attains a classic attitude (Table–8.2).
- **Swelling:** It is obvious in the dislocation of a superficial joint, but may not be so in a joint located deep.
- **Loss of movement** because of severe pain and muscle spasm and loss of articulation.
- **Shortening of the limb** occurs in most dislocations except in anterior dislocation of the hip where lengthening occurs.
- **Telescopy:** In this test, it is possible to produce an abnormal to and fro movement in a dislocated joint (see Annexure-III).

As with all limbs injuries, specific tests to establish the integrity or otherwise of major nerves and vessels of the extremity must be established in all cases of dislocation.

**Radiological examination:** In doubtful cases, the diagnosis must finally depend on adequate X-ray examination. The following principles should be remembered:
- X-ray should always be taken in two planes at right angles to each other, because a dislocation may not be apparent on a single projection.
- If in doubt, X-rays of the opposite limb may be taken for comparison. CT scan may also be of help.
- An associated fracture or an osteochondral fragment must always be looked for.

**Complications**

As with a fracture, complications following a dislocation can be immediate, early or late. Immediate complication is an injury to the neuro¬vascular bundle of the limb. Early complications are: (i) recurrence; (ii) myositis ossificans; (iii) persistent instability; and (iv) joint stiffness. Late complications are: (i) recurrence; (ii) osteoarthritis; and (iii) avascular necrosis.

**Treatment**

Treatment of a dislocation or subluxation depends upon its type, as discussed below:

**Acute traumatic dislocation:** In acute traumatic dislocation, an urgent reduction of the dislocation is of paramount importance. Often it is possible to do so by conservative methods, although sometimes operative reduction may be required.

a) **Conservative methods:** A dislocation may be reduced by closed manipulative manoeuvres. Reduction of a dislocated joint is one of the most gratifying jobs an orthopaedic surgeon is called upon to do, as it produces instant pain relief to the patient. Prolonged traction may be required for reducing some dislocations.

b) **Operative methods:** Operative reduction may be required in some cases. Following are some of the indications:
- Failure of closed reduction, often because the dislocation is detected late.
- Fracture-dislocation: (i) if the fracture has produced significant incongruity of the joint surfaces; (ii) a loose piece of bone is lying within the joint; and (iii) the dislocation is difficult to maintain by closed treatment.

**Old unreduced dislocations:** This often needs operative reduction. In some cases, if the function of the dislocated joint is good, nothing needs to be done. These are discussed in the respective chapters.

**Recurrent dislocations:** An individual episode is treated like a traumatic dislocation. For prevention of recurrences, reconstructive procedures are required. These are discussed in the respective chapters.

**Further Reading**


**What have we learnt?**

- Dislocation means complete loss of contact between articulating bones.
- Treatment of acute dislocation is an emergency.
- Shoulder is the joint to dislocate most often.
Fractures in Children

RELEVANT ANATOMY

Fractures in children are different from those in adults, mostly because of some anatomical and physiological differences between a child’s and an adult’s bone. Some of these are discussed below:

• **Growing skeleton:** Bones in children are growing. At each end of major long bones, and usually at only one end of short bones, there is a cartilaginous growth plate. This is a potential weak point giving rise to different types of epiphyseal injuries. In some injuries through the epiphyseal plate, the growth of the limb may be affected.

• **Springy bones:** Bones in children are more resilient and springy, withstanding greater deformation without fracture. This characteristic is responsible for ‘greenstick’ fractures in children. Such fractures do not occur in adults.

• **Loose periosteum:** The periosteum is attached loosely to the diaphysis in a child’s bone. This results in easy stripping of the periosteum over a considerable part following fracture. The haematoma soon gets calcified to become callus, therefore a child’s bone unites with a lot of callus.

• **Site of fractures:** Some fractures are more common in children than in adults as given in Table–9.1.

• **Healing of fractures:** Fractures unite quickly in children, taking almost half the time taken in adults.

---

**Table–9.1: Fractures* common in children**

- Forearm bones fractures
- Supracondylar fracture of the humerus
- Fracture of lateral condyle of the humerus
- Epiphyseal injuries
- Spiral fracture of tibial shaft

* Dislocations are uncommon in children. Fractures of hands and feet are also uncommon in children.

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**TOPICS**

- Relevant anatomy
- Types of fractures
- Diagnosis
- Treatment
- Complications

---

**TYPES OF FRACTURES**

Fractures in children can be conveniently considered under four headings: (i) birth fractures and related injuries; (ii) epiphyseal injuries; (iii) fractures of shafts of long bones in older children; and (iv) pathological fractures.

**Birth fractures:** Three types of fractures may occur in a newborn. These are as follows:

a) Fracture or epiphyseal separation *sustained during a difficult delivery:* These are the commonest fractures seen at birth. Fracture of the shaft of the humerus occurs most frequently; others are fracture of the shaft of the femur, fracture clavicle etc. Simple strapping of the fracture may be sufficient. Union occurs rapidly with a lot of
callus. Remodelling occurs during the first few years of life.

b) Multiple fractures associated with the congenital fragility of bones e.g., osteogenesis imperfecta (see page 316).

c) Pseudoarthrosis of tibia: This is a pathological entity, very different from a simple fracture or birth injury mentioned above. In this type, there is an inherent indolence of the fracture to unite.

**Epiphyseal injuries:** This is a group of injuries seen in a growing skeleton. An injury involving the growth plate may result in deformities due to irregular growth. Shortening may occur because of premature epiphyseal closure.

**Salter and Harris classification** (Fig-9.1): Epiphyseal injuries have been classified into 5 types based on their X-ray appearance. The higher the classification, the more severe the injury. The incidence of growth disturbance is common in types III, IV and V (Table-9.2).

**Shaft fractures in older children:** Although, fractures of the shaft of long bones have many similarities in children and adults, the following are some of the features peculiar to children:

a) **Displacement is less:** Fractures of the shaft of long bones in children often do not displace much. A special type called ‘greenstick fracture’ occurs only in children. In this type, the bones being resilient, do not break completely. The inner cortex bends, while the outer cortex breaks (Fig-9.2). Such fractures occur commonly in the shafts of forearm bones.

b) **Alignment:** Perfect, end-to-end alignment is not mandatory. Some amount of ‘mal-alignment’ is corrected with growth.

c) **Union:** Fractures unite faster in children.

**Table-9.2: Essential features of epiphyseal injuries (Salter and Harris classification)**

<table>
<thead>
<tr>
<th>Type</th>
<th>Example</th>
<th>Treatment</th>
<th>Prognosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Radial neck epiphysis separation</td>
<td>Closed reduction</td>
<td>Good</td>
</tr>
<tr>
<td>II</td>
<td>Lower end radius epiphysis</td>
<td>Closed reduction</td>
<td>Good</td>
</tr>
<tr>
<td>III</td>
<td>Medial malleolus epiphysis</td>
<td>Open reduction</td>
<td>Growth disturbance can occur</td>
</tr>
<tr>
<td>IV</td>
<td>Lateral condyle of humerus</td>
<td>Open reduction</td>
<td>Growth disturbance common</td>
</tr>
<tr>
<td>V</td>
<td>Lower tibial epiphysis injury</td>
<td>Conservative</td>
<td>Growth disturbance always</td>
</tr>
</tbody>
</table>
d) Treatment: Fractures in children can usually be treated by conservative methods. An operation is rarely necessary.

Pathological fractures: These are uncommon in children. However, there are some diseases which are particularly common in children and result in pathological fractures. These are: (i) fractures through infected bones; (ii) fractures through cysts; and (iii) fractures associated with osteogenesis imperfecta.

DIAGNOSIS
Diagnosis of fractures in children is often missed for the following reasons:

a) History of trauma is either concealed, or the child is not old enough to communicate.  
b) The more dramatic signs of fracture may be absent, especially in incomplete fractures. Thus, there may be no deformity, no abnormal mobility, no crepitus etc.  
c) Parents may attempt to conceal the fact that an infant has been injured, especially when there has been abuse (battered baby syndrome).  
d) Undisplaced fractures are often missed on X-ray, unless carefully looked for.

Therefore, irrespective of the history, possibility of an injury should always be considered whenever marked loss of function, pain and tenderness, and unwillingness to use a limb occurs in children. On the other hand, trauma may be falsely implicated as a cause, in some non-traumatic diseases; the episode of trauma being often days or weeks earlier.

treatment

Most fractures in children can be successfully treated by non-operative methods like plaster immobilisation, traction, sling etc.

Following are some facts about fractures in children
• Fractures in children heal faster.  
• Fractures close to the joint heal faster.  
• Sideways displacement will remodel.  
• Angulation in the plane of the adjacent joint is acceptable.  
• Rotational malalignment will never remodel.  
• Physis subjected to compressive forces inhibits growth.

Operative intervention is necessary in some fractures, as listed below:
• Fracture of the neck of the femur, displaced.  
• Fracture lateral condyle of the humerus (Type IV epiphyseal injury).  
• Fracture of the shaft of femur, in an adolescent.  
• Wherever operation is considered necessary for some other reason such as vascular injury, the fracture is also fixed internally.

With the availability of image intensifier and development of percutaneous methods of fixation, operative fixation of unstable fractures in children, using TENS nails or rush nails has become popular. This makes treatment of an unstable fracture more predictable.

COMPLICATIONS
Fractures in children are associated with few complications. Union of a fracture is generally not a problem; non-union being very rare. Some complications relatively important in children's fractures are:
• Growth disturbances in epiphyseal injuries.  
• Brachial artery injury in supracondylar fracture of the humerus.  
• Myositis ossificans in injuries around the elbow.  
• Avascular necrosis in fracture of the neck of the femur.

Further Reading
### What have we learnt?

- Fractures in children are easier to treat.
- A special category, i.e. epiphyseal injuries, occur in children, and can lead to growth disturbances.
- History of injury has to be carefully probed, as often non-traumatic problems such as infection or tumour, may be erroneously linked to an unrelated episode of ‘injury’.

### Additional information: From the entrance exams point of view

- Distal radius and ulna are the most common fracture locations in children followed by the clavicle.
- Most common bone fractured during birth is the clavicle.
- Injury to the perichondrial ring is type VI Salter Harris fracture or Rang’s injury.
- Multiple fractures at various stages of healing in a child, always consider battered baby syndrome.
- Epiphyseal enlargement seen in haemophilia.
- Epiphyseal dysgnesis seen in Hypothyroidism.
CHAPTER 10

Peripheral Nerve Injuries

TOPICS

• Relevant anatomy
• Diagnosis
• Pathology
• Electrodiagnostic studies
• Mechanism of injury
• Treatment
• Classification
• Prognosis

RELEVANT ANATOMY

Structure of a peripheral nerve: A peripheral nerve consists of masses of axis cylinders (axons), each with a neurilemmal tube (Fig-10.1). An individual nerve fibre is enclosed in a collagen connective tissue known as endoneurium. A bundle of such nerve fibres are further bound together by fibrous tissue to form a fasciculus. The binding fibrous tissue is known as perineurium. A number of fasciculi are bound together by a fibrous tissue sheath known as epineurium. An individual nerve, therefore, is a bundle of a number of fasciculi.

Formation of a peripheral nerve: These are formed from nerves arising from the spinal cord (spinal nerves). There are 31 pairs of spinal nerves in the body, each representing a segment of the spinal cord. These, either through direct branching or through a network of nerves (plexus), give rise to peripheral nerves. Peripheral nerves are mixed nerves carrying motor, sensory and autonomous supply to the limbs. The anatomy of individual nerves will be discussed in respective sections.

Motor innervation of limb muscles: A knowledge of motor innervation of different muscles in the limb is essential for diagnosis of a nerve injury. The following knowledge of anatomy is often required when dealing with a case of nerve injury, and is discussed subsequently in the sections on individual nerve injuries:

a) What is the nerve supply of a particular muscle?
b) What are the different muscles supplied by a nerve?

c) What is the action of a muscle and by what manoeuvre can one appreciate its action in isolation? Only such muscles, whose action can be elicited in isolation are suitable for testing.

**Sensory innervation of limbs:** The area of hypoaesthesia resulting from a nerve injury may be less than the area of skin innervated by that nerve because of the overlap of sensory supply by different nerves. A relatively small area supplied exclusively by a single nerve, called autonomous zone, is found in all nerve injuries (Fig-10.2). The sensory innervation by different nerves of the limbs is discussed in the section on individual nerve injuries.

**Anatomical features relevant to nerve injuries:** There are some features related to the anatomy of a nerve which make a particular nerve more prone to injury. These are as follows:

a) **Relation to the surface:** Superficially placed nerves are more prone to injury by external objects e.g., the median nerve at the wrist often gets cut by a piece of glass.

b) **Relation to bone:** Nerves in close proximity to a bone or a joint are more prone to injury e.g. radial nerve injury in a fracture of the shaft of the humerus.

c) **Relation to fibrous septae:** Some nerves pierce fibrous septae along their course. They may get entrapped in these septae (entrapment neuropathies).

d) **Relation to major vessels:** Nerves in close relation to a major vessel run the risk of ligation during surgery, or damage by an aneurysm.

e) **Course in a confined space:** A nerve may travel in a confined fibro-osseous tunnel and get compressed if there is a compromise of the space, e.g., median nerve compression in carpal tunnel syndrome.

f) **Fixation at points along the course:** Nerves are relatively fixed at some points along their course and do not tolerate the stretch they may be subjected to, e.g., the common peroneal nerve is relatively fixed over the neck of the fibula, and any stretching of the sciatic nerve often leads to isolated damage to this component of the nerve.

**PATHOLOGY**

**Nerve degeneration:** The part of the neurone distal to the point of injury undergoes secondary or Wallerian degeneration; the proximal part undergoes primary or retrograde degeneration up to a single node.

**Nerve regeneration:** As regeneration begins, the axonal stump from the proximal segment begins to grow distally. If the endoneural tube with its contained Schwann cells is intact, the axonal sprout may readily pass along its primary course and reinnervate the end-organ. The rate of recovery of axon is 1 mm per day. The muscle nearest to the site of injury recovers first, followed by others as the nerve reinnervates muscles from proximal to distal, the so-called motor march. If the endoneural tube is interrupted, the sprouts, as many as 100 from one axonal stump, may migrate aimlessly throughout the damaged area into the epineural, perineural or adjacent tissues to form an end-neuroma or a neuroma in continuity (Fig-10.3). An end-neuroma may form when the proximal-end is widely separated from the distal-end. A side neuroma usually indicates a partial nerve cut.

**MECHANISM OF INJURY**

Fractures and dislocations are the commonest cause of peripheral nerve injuries. Some other mechanisms by which a nerve may be damaged are: (i) direct injury – cut, laceration; (ii) infections – leprosy; (iii) mechanical injury – compression, traction, friction and shock wave; (iv) cooling and freezing – ‘frost bite’ etc.; (v) thermal injury; (vi) electrical injury – electric shock; (vii) ischaemic injury– Volkmann’s ischaemia; (viii) toxic agents – injection tetracycline resulting in radial nerve palsy; and (ix) radiation – for cancer treatment.
Peripheral Nerve Injuries

CLASSIFICATION

**Seddon’s classification:** Seddon classifies nerve injuries into three types: (i) neurapraxia; (ii) axonotmesis; and (iii) neurotmesis.

- **Neurapraxia:** It is a physiological disruption of conduction in the nerve fibre. No structural changes occur. Recovery occurs spontaneously within a few weeks, and is complete.

- **Axonotmesis:** The axons are damaged but the internal architecture of the nerve is preserved. Wallerian degeneration occurs. Recovery may occur spontaneously but may take many months. Complete recovery may not occur.

- **Neurotmesis:** The structure of a nerve is damaged by actual cutting or scarring of a segment. Wallerian degeneration occurs. Spontaneous recovery is not possible, and nerve repair is required.

Most nerve injuries are a combination of these. Table 10.1 compares the essential features of the three types of nerve injuries.

### Table 10.1: Seddon’s classification of nerve injuries

<table>
<thead>
<tr>
<th>Type of injury</th>
<th>Pathology</th>
<th>Degeneration</th>
<th>Neuroma</th>
<th>Prognosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neurapraxia</td>
<td>Physiological interruption, anatomically normal</td>
<td>Nil</td>
<td>Nil</td>
<td>Recovery complete within 6 weeks</td>
</tr>
<tr>
<td>Axonotmesis</td>
<td>Axons broken, nerve intact</td>
<td>Proximally + distally</td>
<td>Neuroma in continuity</td>
<td>Recovery +/-, Motor march +</td>
</tr>
<tr>
<td>Neurotmesis</td>
<td>Axons as well as nerve broken</td>
<td>Proximally + distally</td>
<td>End or side neuroma</td>
<td>Recovery poor</td>
</tr>
</tbody>
</table>

**Diagnosis:** In a case of peripheral nerve injury, the following information should be obtained by careful history and examination:

- **a)** Which nerve is affected?
- **b)** At what level is the nerve affected?
- **c)** What is the cause?
- **d)** What type of nerve injury (neurapraxia etc.) is it likely to be?
- **e)** In case of an old injury, is the nerve recovering?

**History:** A patient with a nerve injury commonly presents with complaints of inability to move a part of the limb, weakness and numbness. The cause of nerve injury may or may not be obvious.

In case the cause is obvious, say a penetrating wound along the course of a peripheral nerve (e.g., glass cut injury to the median nerve), the nerve affected and its level is easy to decide. Similarly, nerve injury may occur during an operation as a result of stretching or direct injury.

When the cause is not obvious, an inquiry must be made regarding any history of injection in the proximity of the nerve. Neurotoxic drugs such as quinine and tetracycline are known to damage nerves. Medical causes of nerve affection like leprosy, diabetes should be considered in patients who do not give a history of injury.

**Examination:** Often, the clinical findings in a case of nerve palsy are very few. Therefore, it is essential to perform a systematic motor and sensory examination of the involved limb. Classic deformities may not be present in an early case or in a case with partial nerve injury. A combination of nerve injuries and anatomical variation in the nerve supply may distort the clinical picture of a classic nerve lesion. The following observations must be made during examination:

**Which nerve is affected?**

**Attitude and deformity:** Patients with some peripheral nerve injuries present with a classic attitude and deformity of the limb. Some such attitudes in different nerve injuries are as follows:

- **Wrist drop:** The wrist remains in palmar flexion due to weakness of the dorsiflexors. It is seen in radial nerve palsy.

- **Foot drop:** The foot remains in plantar flexion due to weakness of the dorsiflexors. It occurs in common peroneal nerve palsy.

- **Winging of scapula:** The vertebral border of the scapula becomes prominent when the patient tries to push against a wall. It occurs in paralysis of the serratus anterior muscle in long thoracic nerve palsy.
Claw hand (Main-en-griffe): Claw hand means hyperextension at the metacarpo-phalangeal joints and flexion at the proximal and distal inter-phalangeal joints (Fig-10.4). This occurs due to paralysis of the *lumbricals*, which flex the metacarpo-phalangeal joints and extend the inter-phalangeal joints. Paradoxically, clawing is more marked in low ulnar nerve palsy than in high ulnar nerve palsy. This is because in the latter, flexors of the fingers (both profundus and superficialis), which cause clawing affect are also paralysed. In ulnar nerve palsy, only the medial two fingers develop clawing while all the four fingers develop clawing in combined median and ulnar nerve palsies. Clawing may not become apparent in the early post-injury period.

![Fig-10.4 Claw hand (hyperextension at the MP joint)](image)

Wasting of muscles: This will be obvious *some time* after the paralysis. It may be slight and become apparent only on comparing the affected limb with the sound limb. Some examples of this are given in Table–10.2.

**Table–10.2: Muscle wasting in nerve injuries**

<table>
<thead>
<tr>
<th>Muscle wasting</th>
<th>Nerve</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flat shoulder</td>
<td>Axillary nerve</td>
</tr>
<tr>
<td>Thenar eminence</td>
<td>Median nerve</td>
</tr>
<tr>
<td>Hypothenar eminence</td>
<td>Ulnar nerve</td>
</tr>
<tr>
<td>Hollowing between metacarpals</td>
<td>Ulnar nerve</td>
</tr>
<tr>
<td>Thigh wasting</td>
<td>Femoral nerve</td>
</tr>
<tr>
<td>Calf wasting</td>
<td>Sciatic nerve</td>
</tr>
</tbody>
</table>

Skin: The skin becomes dry (there is no sweating due to the involvement of the sympathetic nerves), glossy and smooth. In partial lesions, there may be vasomotor changes in the form of pallor, cyanosis, or excessive sweating. There may be trophic disturbances such as ridged and brittle nails, shiny atrophic skin, trophic ulcers etc.

Temperature: A paralysed part is usually colder and drier because of loss of sweating, best appreciated by comparing it with normal skin.

Sensory examination: The different forms of sensation to be tested in a suspected case of nerve palsy are touch, pain, temperature and vibration. The area of sensory loss may be smaller than expected. If it is so, look for sensation in the autonomous zone (Fig-10.2, page 61).

Reflexes: Reflexes in the area of nerve distribution are absent in cases of peripheral nerve injuries.

Sweat test: This is a test to detect sympathetic function in the skin supplied by a nerve. Sympathetic fibres are among the most resistant to mechanical trauma. The presence of sweating within an autonomous zone of an injured peripheral nerve reassures the examiner that complete interruption of the nerve has not occurred. Sweating can be determined by the starch test or ninhydrin print test. In these tests, the extremity is dusted with an
agent that changes colour on coming in contact with sweat.

**Motor examination:** For evaluation of motor functions, clear concepts about the anatomy, as to which nerve supplies which muscle is essential. The muscles which are exclusively supplied by a particular nerve are most suitable for motor examination. The tests are nothing but manoeuvres to make a muscle contract. One must carefully watch for trick movements—the movement produced by the adjacent muscles, often substituting for the paralysed muscle. The contraction of the muscle must be appreciated, wherever possible, by feeling its belly or its tendon getting taut. Motor examination conducted for different nerves is discussed below.

**RADIAL NERVE**

**Anatomy:** This nerve is a continuation of the posterior cord of the brachial plexus. In the axilla, it gives off a branch to the long head of triceps, and enters the arm.

**Course in the arm:** As it comes into the arm, the radial nerve gives off the posterior cutaneous nerve of the arm and a branch to the medial head of the triceps. It now travels infero-laterally into the groove for the radial nerve on the posterior surface of the humerus, winding spirally around the bone. In the groove, it gives branches to the lateral head of triceps and anconeus muscles, and cutaneous branches to the arm and forearm. After winding around the humerus, the nerve pierces the lateral intermuscular septum from behind, at the junction of the middle and lower-third of the arm. In the distal-third of the arm it comes to lie in the anterior compartment, between the brachialis muscle on the medial side and brachioradialis and extensor carpi radialis longus on the lateral side.

Before it crosses the elbow in front of the lateral condyle, it divides into two branches—superficial and deep. The **superficial branch** is primarily sensory and travels along side the radial artery into the forearm. The **deep branch** is primarily motor. It gives branches to the extensor carpi radialis brevis and the supinator. It then pierces the supinator and emerges in the posterior compartment of the forearm to become the **posterior interosseous nerve**, which divides immediately into branches supplying the extensor muscles of the forearm.

Branches of the radial nerve are as given in Table–10.3.

**Table–10.3: Major motor branches of radial nerve**

<table>
<thead>
<tr>
<th>Branch Description</th>
<th>Motor Supply</th>
</tr>
</thead>
<tbody>
<tr>
<td>Before the radial groove</td>
<td>Long and medial heads of triceps</td>
</tr>
<tr>
<td>After the radial groove, before crossing the elbow</td>
<td>Lateral head of triceps, anconeus, brachioradialis, extensor carpi radialis longus</td>
</tr>
<tr>
<td>After crossing the elbow, before piercing the supinator</td>
<td>Extensor carpi radialis brevis, the supinator</td>
</tr>
<tr>
<td>After piercing the supinator</td>
<td>Other extensor muscles of the forearm and hand</td>
</tr>
</tbody>
</table>

**Tests:** Various muscles supplied by the radial nerve will be affected according to the level of radial nerve injury i.e. high or low.

**a) High radial nerve palsy:** This occurs if the nerve is injured in the radial groove. In this type, all the muscles supplied by radial nerve except the triceps and anconeus are paralysed. Occasionally, the radial nerve may be injured still higher up, in which case even the triceps may be paralysed. This is called very high radial nerve palsy.

**b) Low radial nerve palsy:** This occurs if the nerve is injured around the elbow so that the muscles supplied by the radial nerve in the distal arm (brachioradialis, extensor carpi radialis longus and brevis) are spared.

From proximal to distal, the following muscles can be examined:

- **Triceps:** The patient is asked to extend his elbow against resistance applied by the examiner, whose other hand feels for triceps contraction.
- **Brachioradialis:** The patient is asked to flex the elbow from 90° onwards, *keeping the forearm midprone*. As he does so against resistance, the brachioradialis stands out, and can be felt.
- **Wrist extensors:** The patient with paralysed wrist extensors has ‘wrist drop’. In case the paralysis is partial, the contraction of the extensor carpi radialis and extensor carpi ulnaris muscle can be felt, though actual movement may not occur.
- **Extensor digitorum:** It causes extension at the metacarpo-phalangeal joints. The patient cannot do so if it is paralysed (finger drop). The examiner
should not be misled by the ability of the patient to ‘extend the fingers’ at the inter-phalangeal joints (function performed by the lumbricals).

- **Extensor pollicis longus**: This causes extension at the inter-phalangeal joint of the thumb. It is examined by stabilising the metacarpophalangeal joint of the thumb, while the patient is asked to extend the inter-phalangeal joint.

**MEDIAN NERVE**

**Anatomy**: This nerve is formed by the joining of branches from the lateral and medial cords of brachial plexus. In the arm, the median nerve descends adjacent to the brachial artery.

**Course in the forearm**: The nerve enters the forearm between the two heads of the pronator teres. It then passes deep to the tendinous bridge of the origin of the flexor digitorum superficialis, in the proximal-third of the forearm. In the mid-forearm it descends between the flexor digitorum superficialis and flexor digitorum profundus. About 5 cm above the wrist, it comes to lie on the lateral side of the flexor digitorum superficialis. It becomes superficial just above the wrist, where it lies between the tendons of the flexor digitorum superficialis and flexor carpi radialis.

**Course in the hand**: The nerve passes deep to the flexor retinaculum and enters the palm. Here a short and stout muscular branch from it supplies the muscles of the thenar eminence (abductor pollicis brevis, opponens pollicis and flexor pollicis brevis). The median nerve finally divides into 4 to 5 palmar digital branches supplying the area of skin shown in Fig-10.2. Also, motor branches are given to the first and second lumbrical muscles at this level. The nerve supply to various muscles by the median nerve along its course is given in Table–10.4.

**Tests**

- **High median nerve palsy** (injury proximal to the elbow): This will cause paralysis of all the muscles supplied by the median nerve in the forearm and hand. In addition, there will be sensory deficit in the skin of the hand.
- **Low median nerve palsy** (injury in the distal-third of the forearm): There will be sparing of the forearm muscles, but the muscles of the hand will be paralysed. In addition, there will be anaesthesia over the median nerve distribution in the hand.

From proximal to distal, the following muscles can be examined:

- **Flexor pollicis longus**: The patient is asked to flex the terminal phalanx of the thumb against resistance while the proximal phalanx is kept steady by the examiner.
- **Flexor digitorum superficialis and lateral half of flexor digitorum profundus**: If the patient is asked to clasp his hand, the index finger will remain straight, the so-called ‘pointing index’. This occurs because both the finger flexors, superficialis as well as the profundus of the index finger are paralysed; though the available medial-half of the flexor digitorum profundus (supplied by the ulnar nerve) makes flexion of the other fingers possible.
- **Flexor carpi radialis**: Normally, the palmar flexion at the wrist occurs in the long axis of the forearm. In a patient with paralysed flexor carpi radialis, the wrist deviates to the ulnar side while palmar flexion occurs. In addition, one cannot feel the tendon of the flexor carpi radialis getting taut.
- **Muscles of the thenar eminence**: Out of the three muscles of the thenar eminence, only two can be examined for their isolated action. These are as follows: (i) abductor pollicis brevis (Fig-10.5): The action of this muscle is to draw the thumb forwards at right angle to the palm. The patient is asked to lay his hand flat on the table with palm facing the ceiling. A pen is held above the thumb and the patient is asked to touch the pen with tip of his thumb. This is called the ‘pen test’; (ii) opponens pollicis: The function of this

<table>
<thead>
<tr>
<th>Table–10.4: Major motor branches of the median nerve</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>In the arm</strong></td>
</tr>
<tr>
<td><strong>In the forearm</strong></td>
</tr>
<tr>
<td>Proximal 1/3</td>
</tr>
<tr>
<td>Distal 1/3</td>
</tr>
<tr>
<td><strong>In the hand</strong></td>
</tr>
</tbody>
</table>

* The three muscles are flexor pollicis brevis, opponens pollicis and abductor pollicis. **Adductor pollicis is not supplied by median nerve.**
ULNAR NERVE

Anatomy: This nerve arises from the medial cord of the brachial plexus. In the arm, it lies on the medial side of the axillary artery. At the junction of the middle and lower-third of the arm, it pierces the medial intermuscular septum and comes to lie in the posterior compartment. It becomes more and more superficial as it approaches the elbow, where it lies behind the medial epicondyle.

Course in the forearm: The ulnar nerve enters the forearm between the two heads of the flexor carpi ulnaris, and descends along the medial side of the forearm. Here it lies anterior to the flexor digitorum profundus, along with the ulnar vessels.

Course at the wrist: It passes in front of the flexor retinaculum just lateral to the pisiform bone. On entering the palm, the ulnar nerve finally divides into superficial and deep terminal branches supplying the hand muscles. The nerve supply to various muscles by the ulnar nerve along its course are given in Table–10.5.

Tests: Various muscles supplied by ulnar nerve will be affected according to the level of ulnar nerve injury i.e., high or low.

a) High ulnar nerve palsy (injury proximal to the elbow): This will cause paralysis of all the muscles supplied by the ulnar nerve in the forearm and hand. In addition, there will be a sensory deficit in the skin of the hand.

b) Low ulnar nerve palsy (injury in distal-third of forearm): There will be sparing of forearm muscles but the muscles of the hand will be paralysed. Sensory deficit will be same as in high ulnar nerve palsy.

Individual muscles which could be examined in a case of ulnar nerve palsy are given below:

- Flexor carpi ulnaris: The patient is asked to palmar flex the wrist against gravity. In doing so, the hand deviates towards the radial side. The tendon of flexor carpi ulnaris just above the pisiform, does not stand out. On performing the same test against resistance, the tendon cannot be felt.

- Abductor digiti minimi: The patient is asked to abduct the little finger against resistance while keeping the hand flat on the table (in order to avoid action of flexors of the finger).

- Interossei: Palmar interossei do adduction (PAD), the dorsal interossei do abduction (DAB) of the fingers at metacarpo-phalangeal joints. These can be tested as follows:
  
  Egawa’s Test (Fig-10.6a): This is for dorsal interossei (abductors) of the middle finger. With the hand kept flat on a table palmar surface down, the patient is asked to move his middle finger sideways.

  Card Test (Fig-10.6b): This is for palmar interossei (adductors) of the fingers. In this test, the examiner inserts a card between two extended fingers and the patient is asked to hold it as tightly as possible while the examiner

---

**Table–10.5: Major motor branches of the ulnar nerve**

<table>
<thead>
<tr>
<th></th>
<th>In the arm</th>
<th>In the forearm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Proximal 1/3</td>
<td>Nil</td>
<td>Flexor carpi ulnaris, medial half of flexor digitorum profundus</td>
</tr>
<tr>
<td>Distal 1/3</td>
<td>Nil</td>
<td>Nil</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Superficial branch</th>
<th>Deep branch</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Hypothenar muscles</td>
<td>Adductor pollicis, all interossei and medial two lumbricals</td>
</tr>
</tbody>
</table>
tries to pull the card out. The power of adductors can thus be judged. In case of weak palmar interossei, it is easy to pull out the card. First dorsal interosseous muscle can be separately examined by asking the patient to abduct the index finger against resistance (Fig-10.6c).

- **The lumbricals:** These are mainly responsible for flexion at the metacarpo-phalangeal joints but their isolated action cannot be tested.

- **Adductor pollicis:** The patient is asked to grasp a book between the thumb and index finger. Normally, a person will grasp the book firmly with thumb extended, taking full advantage of the adductor pollicis and the first dorsal interosseous muscles. If the ulnar nerve is injured, the adductor pollicis will be paralysed and the patient will hold the book by using the flexor pollicis longus (supplied by median nerve) in place of the adductor. This produces flexion at the inter-phalangeal joint of the thumb. (Fig-10.7). This becomes more pronounced if the examiner tries to pull the book out while the patient tries to hold it. This sign is known as 'Froment's sign' or the 'book test.'

**ACCESSORY NERVE**
This supplies the trapezius muscle.

**Test:** The trapezius muscle is tested by asking the patient to elevate his shoulder against resistance. One can see and feel the trapezius belly stand out. Similarly, the patient is asked to brace his shoulder backward and depress it to examine middle and lower part of the muscle.

**LONG THORACIC NERVE**

**Anatomy:** The nerve arises from the ventral rami of C5, C6 and C7. It descends behind the brachial plexus on the lateral surface of the serratus anterior, which it supplies.

**Test:** The serratus anterior muscle can be examined by asking the patient to push against a wall with both hands. The medial border of the scapula on the affected side will become prominent (winging of scapula, Fig-10.8).

**AXILLARY NERVE**

**Anatomy:** The axillary nerve arises from the posterior cord of the brachial plexus and curves backwards on the lower border of the subscapularis. It crosses the quadrangular space and comes to lie on the medial side of the surgical neck of the humerus, medial and inferior to the capsule of the shoulder joint. Here it divides into anterior and posterior branches. The posterior branch supplies
the teres minor and posterior part of the deltoid and terminates as the cutaneous nerve which supplies the skin over the lower-half of the deltoid. The anterior branch continues horizontally between the deltoid and the surgical neck of the humerus, and supplies the rest of the deltoid.

**THE CAUSES OF INJURY**

Once it is decided on clinical examination, which nerve is affected and at what level, one must look for a tell tale signs along the course of the nerve for the cause. This may be in the form of an injury such as displaced bone fragments or a scar to suggest an old external injury. The nerve may be thickened (e.g. leprosy). If no such external evidence is present, the paralysis could be due to some medical cause such as neuropathy, myelopathy etc. (Refer to a Textbook of Medicine).

**THE TYPE OF INJURY**

Once the cause of nerve injury is established, one must make an attempt to evaluate the predominant type of nerve injury (Seddon’s classification). The nature of the causative factor, a period of observation and electrodiagnostic studies may help in deciding this.

**SIGNS OF REGENERATION**

Whenever a case of nerve injury is seen some time after the injury or following a repair, signs of regeneration of the nerve should be looked for during examination. These are as follows:

- **Tinel’s sign:** On gently tapping over the nerve along its course, from distal to proximal, a pins and needle sensation is felt in the area of the skin supplied by the nerve. A distal progression of the level at which this occurs, suggests regeneration.
- **Motor examination:** The muscle supplied nearest to the site of injury is the first to recover, noticed clinically by the ability of the muscle to contract. The muscles in the more distal area begin to contract as they are reinnervated one after another (motor march). This phenomenon is absent in neuropraxia where all muscles recover together.
- **Electrodiagnostic test:** This can help in predicting nerve recovery even before it is apparent clinically.

**SCIATIC NERVE**

**Anatomy:** The sciatic nerve consists of two anatomically distinct components – the tibial and common peroneal nerves. The common peroneal component is more frequently affected than the tibial. Complete lesion of the sciatic nerve is rare.

**Tests:** The common peroneal nerve supplies the extensors and the evertors of the foot. Paralysis of these muscles results in foot drop. The patient walks with a ‘high-step gait’, i.e. while walking he has to lift the foot high in order to clear the ground. The plantar flexors of the foot are normal.

The tibial nerve supplies the plantar flexors of the foot. One can test for weakness of these muscles by asking the patient to plantar-flex the ankle and toes. The function of the hamstring group of muscles, also supplied by the sciatic nerve, can be tested by flexing the knee against resistance.

**ELECTRODIAGNOSTIC STUDIES**

**Electromyography:** Electromyography (EMG) is a graphic recording of the electrical activity of a muscle at rest and during activity. **Normal muscle:** A normal muscle at rest shows no electrical activity. With voluntary contraction,
action potentials develop in the motor units. In a weak contraction, these may be recordable as single motor unit potentials in the vicinity of the recording electrode. In a strong contraction, impulses of a number of motor units firing simultaneously are superimposed, giving rise to an interference pattern (Fig-10.9).

c) Whether any regeneration occurring: The earliest evidence of reinnervation of a muscle is the appearance of reinnervation potentials on attempted voluntary contraction of the muscle. These potentials appear weeks before a contraction can be noticed clinically. The progress of a nerve recovery can thus be monitored.

d) Level of nerve injury: By performing an EMG of all the muscle supplied by a nerve, one can decide the level of nerve injury. Muscles supplied distal to the site of nerve injury would show changes of denervation.

**Strength-duration curve:** This is a graphic representation of the excitability of muscle and nerve tissue under test. A small strength of current can excite a normal muscle. This occurs by excitation of the muscle through neuromuscular junction, which needs a weaker current. In a denervated muscle, the excitation is possible only on direct stimulation of the muscle fibres, which need a higher strength of current.

A very low-strength current is given for a duration of 300 milliseconds and its response noted. The strength of the current is gradually increased until a minimal visible contraction of the muscle is observed. This minimal current strength, required to elicit muscle contraction, is called the **Rheobase**, and is measured in milliamperes. The **Chronaxie** is the duration of current required to excite a muscle with a current-strength of double the rheobase. It is measured in milliseconds. These two are the basic parameters of excitability of a muscle.

For knowing excitability of a muscle in relation to current-strength and its duration, the muscle is stimulated by reducing the duration of the current from 300 milliseconds, gradually to a 1 millisecond or even lower. A corresponding increase in strength of the current required is detected. A graph is plotted between current-duration and corresponding current-strength. This is called a strength-duration curve (Fig-10.10).

**Interpretation:** The pattern of the strength-duration curve of an innervated muscle is different from that of a denervated muscle or regenerating muscle, as discussed below:

- **Normal strength-duration curve:** A normal muscle will respond to stimuli varying in duration
Peripheral Nerve Injuries

from 300 milliseconds to as low as 3 or even 1 millisecond without any increase in the strength of the current. If the duration of current is decreased beyond it, a progressive increase in the strength of current is required in order to produce a contraction. A strength-duration curve plotted from such a muscle is termed a nerve curve, because the muscle contraction is caused by stimulation of the motor nerve entering the muscle.

• Denervated muscle: A totally denervated muscle will need current either of more strength or for a longer duration. A curve from such a muscle is termed a muscle curve.

• A partially denervated muscle: The curve of a partially denervated muscle or a muscle recovering after nerve injury lies between the normal and the curve of denervation, and is characterised by an upward kink. The kink denotes the superimposition of the two basic types of curves.

• Assessment of recovery by the strength-duration curve: If progressive recovery is occurring the curve will, on serial examination, become flatter with a shift to the left. On the other hand, if the process of denervation is progressive, the curve will become steeper and will shift to the right.

Nerve conduction studies: It is a measure of the velocity of conduction of impulse in a nerve. A stimulating electrode is applied over a point on the nerve trunk and the response is picked up by an electrode at a distance or directly over the muscle. The velocity of the conduction of the impulse between any two points of the nerve can be calculated. The normal nerve conduction velocity of motor nerve is 70 metres/second. This conduction study helps in the following:

• Whether a nerve injury is present: If a nerve injury is present there will be no conduction of the impulse across the suspected level.

• Whether it is a complete or partial nerve injury: Absence of any transmitted impulse across the suspected site is an indicator of a complete nerve injury.

• Compressive lesion: The conduction velocity may simply be delayed in compressive nerve lesions such as carpal tunnel syndrome, etc.

TREATMENT

General consideration: In fresh nerve injuries, the general condition of the patient must be evaluated before undertaking a nerve repair. Arterial, bone and joint repair takes precedence over nerve repair. The treatment of nerve injuries may be conservative or operative. Though conservative treatment yields good results, in selected cases an operation should not be delayed in the hope of spontaneous recovery.

CONSERVATIVE TREATMENT

This alone or in addition to operative treatment is required in all types of nerve injuries. The aim of conservative treatment is to preserve the mobility of the affected limb while the nerve recovers.

The following are the essential components of conservative treatment:

• Splintage of the paralysed limb: The first procedure to be adopted in every case of nerve injury is to splint the limb in the position which will most effectively relax the affected muscles. The type of splints used for common nerve injuries are as given in Table–10.6.

• Preserve mobility of the joints: Every joint of the affected limb must be put through full range of movement at least once every day.

• Care of the skin and nails: Since the skin is anaesthetic, it should be protected from trauma.

Table–10.6: Splints used for various nerve injuries

<table>
<thead>
<tr>
<th>Nerve injured</th>
<th>Splint</th>
</tr>
</thead>
<tbody>
<tr>
<td>Axillary nerve (deltoid paralysis)</td>
<td>Aeroplane splint</td>
</tr>
<tr>
<td>Radial nerve palsy (extensors of wrist &amp; MP joints paralysed)</td>
<td>Cock-up splint</td>
</tr>
<tr>
<td>Ulnar nerve palsy (lumbricals paralysis)</td>
<td>Knuckle-bender splint</td>
</tr>
<tr>
<td>Sciatic N. palsy or common peroneal N. palsy</td>
<td>Foot drop splint</td>
</tr>
</tbody>
</table>
burn or pressure sores. Trophic ulcers should be meticulously treated. Nails should be cleaned and cut with care.

- **Physiotherapy:** Physiotherapeutic measures consist of (i) massage of the paralysed muscles; (ii) passive exercises to the limb; (iii) building up of the recovering muscles; and (iv) developing the unaffected or partially affected muscles. Attempts were made in the past to preserve tone and functions of denervated muscles by electrical stimulation, but it has been found to be of no use.

- **Relief of pain:** Suitable analgesics are prescribed for relief of pain.

**OPERATIVE TREATMENT**

Operative procedures for nerve injuries consist of nerve repair, neurolysis, and tendon transfers.

**Nerve repair:** It may be performed within a few days of injury (primary repair) or later (secondary repair).

**Primary repair:** It is indicated when the nerve is cut by a sharp object, and the patient reports early. In such cases an immediate primary repair is the best. One needs experience in the use of the fine sutures and operative microscope for this kind of surgery. In case the wound is contaminated or the patient reports late, a delayed primary repair is better. In this, in the first stage, the wound is debrided and the two nerve ends approximated with one or two fine silk sutures so as to prevent retraction of the cut ends. This also makes identification of the cut ends easy at a later date. After two weeks, once the wound heals, a definitive repair is done. Some surgeons routinely perform a delayed primary repair because they feel that the epineurium gets thickened in two weeks and sutures hold better.

**Secondary repair:** It is indicated for the following cases:

a) **Nerve lesions presenting some time after injury:** Often nerve injuries are missed at the time of injury, or it may not have been possible to treat them early for reason, such as poor general condition of the patient.

b) **Syndrome of incomplete interruption:** If no definite improvement occurs in 6 weeks in cases with an apparently incomplete nerve injury, nerve exploration, and if required secondary repair should be carried out.

c) **Syndrome of irritation:** Cases with signs of nerve irritation need exploration and sometimes a secondary repair.

d) **Failure of conservative treatment:** If a nerve injury is treated conservatively and no improvement occurs within 3 weeks, one should proceed to electrodiagnostic studies, and if required, nerve exploration.

**Techniques of nerve repair:** Nerve repair can be either end-to-end or by using a nerve graft.

a) **Nerve suture:** When the nerve ends can be brought close to each other, they may be sutured by one of the following techniques (Fig-10.11):

- Epineural suture
- Epi-perineural suture
- Perineural suture
- Group fascicular repair

**Methods of closing nerve gaps:** Sometimes, the loss of nerve tissue is so much, that an end-to-end suture cannot be obtained. In such a situation, the following measures are adopted to gain length and achieve an end-to-end suture:

- Mobilisation of the nerve on both sides of the lesion.
- Relaxation of the nerve by temporarily positioning the joints in a favourable position.
- Alteration of the course of the nerve, e.g. the ulnar nerve may be brought in front of the medial epicondyle (anterior transposition).
- Stripping the branches from the parent nerve without tearing them.
- Sacrificing some unimportant branch if it is hampering nerve mobilisation.
b) **Nerve grafting:** When the nerve gap is more than 10 cm or end-to-end suture is likely to result in tension at the suture line, nerve grafting may be done. In this, an expandable nerve (the sural nerve) is taken and sutured between two ends of the original nerve as shown in Fig-10.11d.

**Neurolysis:** This term is applied to the operation where the nerve is freed from enveloping scar (perineural fibrosis). This is called external neurolysis. In many cases, the nerve sheath may be dissected longitudinally to relieve the pressure from the fibrous tissue within the nerve (intra-neural fibrosis). This is called internal neurolysis.

**Reconstructive surgery:** These are operations performed when there is no hope of the recovery of a nerve, usually after 18 months of injury. After this time even if the nerve recovers, transmission of impulses across the neuromuscular junction does not occur because the neuromuscular function itself has degenerated. Operations included in this group are tendon transfers, arthrodesis and muscle transfer. Rarely, an amputation may be justified for an anaesthetic limb or the one with causalgia.

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**Flow chart-10.1  Treatment plan for management of peripheral nerve injury**

- Nerve is injured
  - Is the nature of lesion known?
    - Yes
      - What is it?
        - Nerve in continuity
          - Wait for 3 weeks
            - Examine for return of activity (A)
              - Recovery occurs
                - Follow-up every 1-3 months
                  - Progressive recovery
                    - No progress
                      - Full recovery
                        - Do electrical studies
                          - EMG-FIBS,PSW,No VMP
                            - EMG-FIBS,PSW
                              - Number of VMP's
                                - EMG-No FIBS or PSW,VMP-absent
                                  - NCV-Motor & sensory conduction above & below the block normal
                                    - EMG-No FIBS or PSW,No VMP unless patient is ticked,NCV-Normal
                                      - EMG- Electromyography
                                        - NCV- Nerve conduction velocity
                                          - FIBS- Fibrillation potentials
                                            - PSW- Positive sharp waves
                                              - VMP- Voluntary motor potentials
    - No
      - Do electrical studies
        - EMG-FIBS,PSW, No VMP
          - NCV- No motor/sensory
            - Explore & surgically repair
              - Expect further return
                - Full return
                  - Patient is normal
                    - no nerve injury

DECISION MAKING IN NERVE INJURIES

It is often difficult to decide when to operate in a case of nerve injury. This is especially so if there is a partial lesion or partial recovery has taken place. Electrodiagnostic studies are helpful in these cases.

Unless the nature of the nerve injury is known before the operation, the appearance of the involved nerve is often the best guide to deciding on the type of operative procedure. Where the nerve ends are visibly apart, nerve repair is the only choice. Where the nerve is in continuity, it is often difficult to decide the further course. A fusiform thickening of a nerve (neuroma in continuity) indicates a partial cut. A nerve stimulator may be used to find if there is any continuity of the nerve. If there is a brisk response in the muscles supplied by the nerve on stimulating the nerve proximal to the neuroma, there is no need for nerve suture; neurolysis may suffice. In case there is little or no response, the neuroma should be excised and the nerve repaired.

A practical plan for management of nerve injury is shown in Flow chart-10.1.

PROGNOSIS

The following factors dictate recovery following a nerve repair:

a) **Age:** The lower the age, the better the prognosis.
b) **Tension at the suture line:** The more the tension, the poorer the prognosis.
c) **Time since injury:** After 18 months only sensory functions can be expected.
d) **Location of injury:** The more proximal the injury, the worse the prognosis.
e) **Type of nerve:** A primarily motor nerve, like radial nerve, has a better prognosis than a mixed nerve.
f) **Condition of the nerve ends:** The more the crushing and infection, the poorer the prognosis.
g) **Associated conditions:** Infection, ischaemia etc. indicate poor prognosis.

What have we learnt?

- There are three types of nerve injuries: Neurapraxia, Axonotmesis, Neurotmesis.
- Nerve recovers at the rate of 1 mm/day.
- How to diagnose a nerve injury, clinical signs & their explanation.
- How to monitor nerve regeneration, and role of electrodiagnostic studies.
- Methods of nerve repair.
- Decision making in nerve injury.
- How to predict whether a nerve will recover or not?

Additional informationL: From the entrance exams point of view

- Tinel’s sign indicates regeneration of nerves.
- Prognosis after nerve suturing: radial nerve > ulnar nerve > peroneal nerve> sciatic nerve> femoral nerve.
- Erb’s palsy is the most common neurological deficit in the upper limb.
- Erb’s palsy is injury to upper C5,6 roots of brachial plexus.
- Klumpke’s palsy is injury to the lower trunk C8 and partially T1.
- Crutch palsy is injury to the radial nerve.
- Claw hand due to leprosy is classified as Grade II WHO grading.
- Sudden hyperflexion of the thigh over the abdomen (McRobert’s procedure) done on the mother for delivery of babies with shoulder dystocia leads to injury to the lateral cutaneous nerve of thigh (meralgia paresthetica).
CAUSES

Deformities may arise from an abnormality in the bone (e.g., a malunited fracture), joint (e.g., tuberculosis of the knee), or soft tissues (e.g., clubfoot). These may be either congenital or acquired.

CONGENITAL DEFORMITIES

These are deformities or malformations present at birth (e.g., clubfoot). Some of these malformations, though present at birth, may become apparent only later in life (e.g., spina bifida). The deformity may be severe and incompatible with life (e.g., osteogenesis imperfecta congenita), and can only be found in stillborn infants. On the other hand, it may be very minor and have no practical significance.

The underlying causative factors may be: (i) a genetic abnormality (e.g., diaphysial aclasis, mongolism etc.); (ii) environmental factors (e.g., phocomelia); and (iii) combined – genetic and environmental factors (e.g., congenital dislocation of the hip, clubfoot).

ACQUIRED DEFORMITIES

Deformities acquired later in life may be divided into those arising at a joint or in a bone (Fig-11.1), as discussed below:

Deformities arising at a joint: A joint may become deformed because of any of the following factors:

a) Dislocations and subluxations: These may be traumatic (e.g., most dislocations and subluxations seen in day-to-day practice) or pathological (e.g., following acute septic arthritis). Classic deformities are produced in some subluxation or dislocation (Table–8.2, page 56).

b) Muscle imbalance: All joints are spanned by two opposing groups of muscles. Normally, these muscles maintain a balance so that the joint can be kept in any position. In some diseases, an unbalanced action of the muscles may hold the joint in a particular position. With time the other soft tissues around the joint (the capsule, ligaments etc.) also contract and prevent the joint from returning to its neutral position. The muscle imbalance may arise from paralysis of a group (e.g., polio) or overactivity (e.g., spasticity in cerebral palsy).

c) Tethering or contracture of muscles and tendons: Joint movement is associated with contraction of a group of muscles and elongation of opposing group. To and fro gliding of tendons also happens in this process. If by some disease, these functions are interrupted, the joint is prevented from moving full range. For example, the muscles or tendons, may get tethered to the underlying bone (e.g., tethering of the quadriceps to the femur in a fracture). The muscle may get contracture (e.g., Volkmann’s ischaemic contracture of the flexor muscles of forearm, leading to flexion deformity of the wrist and fingers).
d) **Contracture of soft tissues other than muscles:** Apart from muscles, contracture of other soft-tissues like skin, deep fascia etc. may account for the deformity. For example, contracture of palmar aponeurosis may pull the metacarpo-phalangeal and **proximal** inter-phalangeal joints of one or more fingers (Dupuytren's contracture). Similarly, contracture of the scarred skin on the flexor aspect of the elbow or knee following a burn, may result in a flexion deformity of the respective joint.

e) **Arthritis:** Joint deformity may result from arthritis. This may occur: (i) because of sustained spasm of a group of muscles in response to pain; or (ii) as a result of damage to important structures like ligaments, cartilage etc., by the arthritic process.

f) **Posture:** The habitual keeping of a joint in a deformed position may result in a deformity, for example, lateral deviation of the great toe (hallux valgus) is seen in women who wear narrow pointed high-heeled shoes.

g) **Unknown factors:** Some deformities of joints result from no apparent reason. For example, knock knees deformity (**genu valgum**) commonly seen in children, often has no cause.

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**Deformities arising in a bone:** Three major causes of deformity arising in a bone are fractures, bone diseases and abnormally growing bones.

a) **Fracture:** This is the commonest cause of deformity of a bone. This results when a fracture unites in a mal-aligned position. Some of the common deformities resulting from malunion of fractures are given in Table–II.1.

b) **Bone diseases:** Some diseases of the bone result in a softening and bending of the bones. Most of these are generalised disorders where several or all of the bones are affected. The following are some examples:

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**Table–II.1: Deformities due to fractures**

<table>
<thead>
<tr>
<th>Deformity</th>
<th>Fracture</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gun stock deformity</td>
<td>Supracondylar fracture of the humerus</td>
</tr>
<tr>
<td>Cubitus varus</td>
<td>Fracture of lateral condyle of humerus</td>
</tr>
<tr>
<td>Cubitus valgus</td>
<td>Colles' fracture</td>
</tr>
<tr>
<td>Dinner fork deformity</td>
<td>Avulsion of the extensor tendon from base of distal phalanx</td>
</tr>
<tr>
<td>Mallet finger</td>
<td>Inter-trochanteric fracture</td>
</tr>
<tr>
<td>Coxa vara</td>
<td>Condylar fractures of tibia (e.g., bumper fracture)</td>
</tr>
<tr>
<td>Genu valgum</td>
<td>Ankle injuries</td>
</tr>
<tr>
<td>Varus-valgus at ankle</td>
<td></td>
</tr>
</tbody>
</table>

---

**Fig-11.1 Causes of deformities at joints**

- Dislocation
- Muscle imbalance
- Muscle tethering
- Contracture of fascia
- Soft tissue contracture
- Pointed shoe
- Postural
• Metabolic disorders – rickets, osteomalacia.
• Endocrine disorders – parathyroid osteodystrophy, Cushing’s syndrome.
• Disorder of unknown aetiology – Paget’s disease, fibrous dysplasia, senile osteoporosis.

c) Abnormal bone growth: Bone deformity may result from uneven growth occurring at the epiphyseal plate. Unequal growth of one of the two bones in a part of the limb with two bones (e.g. forearm or leg), may result in deformity at the joint adjacent (e.g. wrist or ankle). The common causes of uneven growth at the epiphyseal plate are as follows:
• Crushing fracture involving the epiphyseal plate (Grade-V, Salter and Harris epiphyseal injury).
• Infection from a nearby osteomyelitis or arthritis, spreading to the epiphyseal plate, and damaging it.
• A tumour may retard the growth of a nearby epiphyseal plate (e.g., enchondroma as in Ollier’s disease). Occasionally, the tumour may stimulate uneven growth of the adjacent plate by causing local hyperaemia (e.g., haemangioma).
• Dysplasia: In some epiphyseal dysplasias, abnormal growth at the epiphysis results in joint deformities.

TREATMENT
Many deformities do not need treatment, as they are of no significant functional or cosmetic concern. A simple reassurance and watchful neglect may be appropriate in these cases. Most other deformities cause functional impediment or cosmetic concerns, and have to be corrected. Some deformities (e.g. bow legs), may not be of immediate functional concern, but may cause problem in long term, and thus may need to be corrected. The methods used for correction of deformities may be non-operative or operative.

NON OPERATIVE METHODS
Wherever possible, non-operative methods are attempted first. These are suitable for deformities due to soft tissue contracture. The method essentially consists of stretching the contracted soft tissue, and then maintaining the correction by splints. The disadvantage of this method is that the treatment is long drawn, and an equally prolonged effort at maintenance is required. Recurrence of deformity is common. Correction of deformity by non-operative methods is done by the following ways:

a) Manipulative correction: The contracture is gently manipulated, so as to stretch it. Once corrected, it is maintained in the corrected position in a plaster cast or splint. An example of use of this method is treatment of a club foot by manipulation and PoP.

b) Wedging cast: In this technique, a cast is applied on the limb with deformed joint. A wedge of plaster is cut out on the convex side of the deformity, the wedge closed by forcing the part, thus achieving correction.

c) Traction: Gradual traction can stretch out contracted soft tissues. The correction is subsequently maintained in a splint or calipers.

d) Splints: These are special splints which permit gradual stretching of the soft tissues, leading to the correction of deformities (e.g., turn-buckle splint for VIC see page 103).
In cases where the non-operative methods fail or the deformity is primarily bony, operative correction may be required. The following methods are used:

a) **Soft tissue release**: The contracted soft tissues are released. Tethering of soft tissues is removed.

b) **Osteotomy (Fig-11.2)**: It is used for correcting bony deformity. The deformed bone is cut and suitably realigned in a corrected position (e.g., for genu varum and genu valgum).

c) **Arthrodesis (fusion of joint)**: This method is adopted where a joint is not only deformed, but also its articulating surfaces damaged beyond repair. Arthrodesis is suitable for joints where loss of motion at the joint does not produce much functional disability (e.g., wrist). In other situations, such as hip and knee, joint replacement is a better option. With the availability and better longevity of artificial joints, arthrodesis has become less popular. There are situations where joint replacement cannot be done (e.g. joint infection, paralysed limb), and arthrodesis remains the only option. The procedure involves opening up the joint, removing its cartilage, and immobilizing it in functional position. The raw bone ends unite (as in a fracture), resulting in fusion.

d) **Arthroplasty**: The term arthroplasty means 'reconstructing a joint'. Reconstruction can be done by two methods: (i) by excising a part of the deformed joint, thereby relaxing the surrounding soft tissues, and thus correcting deformity or (ii) by replacing the joint with artificial components. The former is called excision arthroplasty, and is done for joints damaged due to infection. The latter is called replacement arthroplasty (joint replacement), and is done for most other damaged and deformed joints (e.g. osteoarthritis knee). See also Chapter 42.

e) **Correction of deformity by selective retardation of epiphyseal growth**: This is useful in cases where the cause of deformity is unequal epiphyseal growth, and the child has residual growth potential. Here, the faster growing side of the epiphysis is temporarily or permanently
stopped by surgical means (stapling, direct damage etc.). Over a period of time, the slower growing side keeps growing, while the stapled side does not, resulting in correction of the deformity. This is performed in selected cases of genu varum or valgum in a growing child. It is a minimally invasive operation, but a little unpredictable.

f) **Ilizarov’s technique:** This is a versatile technique of correcting deformity. Its utility is more when the deformity is associated with shortening, or if the deformity is in more than one plane. The apparatus provides an opportunity for correcting the deformity very accurately (see also page 33) A comprehensive plan of management of a deformed joint is given in Flow chart-11.1.

**Further Reading**

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**What have we learnt?**

- Deformities may be since birth, or develop later.
- The cause of deformity may be bone (e.g. malunion), joint (e.g. OA knee), or soft tissue contracture (e.g. Volkmann’s contracture).
- Initial treatment of deformity is by non-operative methods, and thereafter operative methods.
- Different operative methods of correction of deformity exist.
Orthopaedic treatment can be broadly divided into two types, non-operative and operative. At times a patient needs no definite treatment except reassurance. Whenever treatment is required, it is preferable to try non-operative methods first, though there are occasions when early or an immediate operation must be advised.

**NON-OPERATIVE METHODS OF TREATMENT**

**REST**
Since olden times, rest has been the mainstay of orthopaedic treatment. It helps in reducing inflammation and pain. The word ‘rest’ could mean complete inactivity or immobility, as is sometimes required in acute inflammatory conditions like acute osteomyelitis etc. But, more often than not, it means no more than ‘relative rest’, implying simply a reduction of activity and avoidance of strain. With advances in orthopaedic treatment, more and more methods have been devised by which the period of rest could be reduced significantly (e.g., by internal fixation for most fractures).

**SUPPORT**
A limb or a joint not capable of functioning because of inadequate muscle power needs support (e.g., a polio limb). Temporary support may be given with a splint made from Plaster of Paris or other plastic splinting material. A permanent or prolonged support may be required in some cases, in the form of life time appliances called *orthoses* (see page 332).

**PHYSIOTHERAPY**
This includes a variety of treatment modalities based on physical methods of treatment such as heat therapy, exercises etc. It may be aimed at alleviation of pain, restoration of functions, or both. It may be used as a primary treatment modality (e.g., for backache etc.) or in conjunction with other methods of treatment (e.g., post-operative physiotherapy).

When appropriately prescribed and adequately pursued under the supervision of a skilled physiotherapist, it can perform wonders. On the other hand, an unskilled physiotherapist by his over enthusiastic approach may retard rather than hasten patient’s recovery. The following are the common methods used in physiotherapy:

**Ice therapy:** Ice therapy is beneficial during the first 24-72 hours after injury. It causes relief in pain, reduces haematoma formation and reduces inflammation. The pain relieving effect of cold therapy is appreciated more after the application than during it.

**Heat therapy:** Heat produces a soothing effect on many aches and pains, probably by increasing the blood flow, or possibly by some other mechanism.
Heat application is done for 15 to 20 minutes 2 or 3 times a day. Heat must not be applied to insensitive or ischaemic skin, and if there is underlying acute infection or neoplastic tissue. Depending upon the depth of penetration of the heat, it can be either surface heat i.e., only the skin and subcutaneous tissues are heated, or deep heat i.e. deeper structures are heated.

a) **Surface heat**: This can be provided by (i) hot water bottle (rubber-bottle); (ii) warm bath; (iii) hot soaks or compresses; (iv) infra-red lamp; and (v) wax bath.

b) **Deep heat**: This can be provided by (i) short-wave diathermy—heat generated by a high frequency alternating current (frequency 27 mega cycles/second) using a short-wave diathermy emitter; (ii) ultrasonic therapy – these waves (a million cycles/second) are projected as a beam from a transducer; and (iii) microwave. The ultrasonic waves and microwaves penetrate to a considerable depth. When the waves strike the tissues, energy is converted into heat. It is most useful for localised tender fibrous nodules.

**Exercise therapy**: These are given for three purposes: (i) to mobilise joints; (ii) to strengthen muscles; and (iii) to improve coordination and balance.

a) **Joint mobilising exercises**: The following exercises may be advised:
   - Passive joint movements: These are used to preserve joint mobility when the patient is unable to move the joint himself (e.g., when the muscles are paralysed).
   - Active joint movements: The patient moves his joints actively so as to gain more and more range of movement. Sometimes, a patient's active efforts may be assisted by gentle pressure from the physiotherapist.
   - Continuous passive mobilisation (CPM): The joint is fitted in a machine which moves the joint slowly through a predetermined arc of motion. Since the motion produced is very slow, it is tolerated by the patient in even the very early post-operative period.

b) **Muscle strengthening exercises**: These are used to preserve or improve the strength of the muscles. These may be of the following types:
   - Static or isometric exercises i.e., the muscle contracts while its length remains the same e.g., muscle contraction while pushing a wall.
   - Dynamic or isotonic exercises i.e., the muscle contracts and produces movement. These exercises could be: (i) active — the patient does the movements himself; (ii) active-assisted — the patient does the movement while the physiotherapist helps; or (iii) active-resisted — the patient does the movement against resistance. The last is the most effective in gaining muscle strength.

c) **Exercises to improve coordination**: These are special exercises, useful in polio and cerebral palsy patients.

**Traction**: In physiotherapy, traction is applied: (i) to separate joint surfaces while giving passive movements to a joint; (ii) to obtain the relaxation of muscles which are in spasm (e.g., by giving cervical or lumbar traction); or (iii) to correct deformities by gentle continuous traction. For details please refer to Chapter 4.

**Massage**: This is a systematic and scientific manipulation of the skin and the underlying soft tissues which gives rise to relief of pain and the relaxation of muscles. Most massage and manipulations are soothing except for frictions which are painful, and are used to breakdown adhesions.

**Hydrotherapy**: The principles of buoyancy help to reduce pain by relaxation of the muscles, mobilisation of stiff joints, and thereby assist in the development of muscle power. This is useful as it produces a general sense of well-being.

**Occupational therapy**: Occupational therapy aims at enabling the person to become as independent as possible, inspite of the disability he may have. A person needs independence in following day-to-day activities:

a) **Activities of daily living (ADL)**: These constitute activities such as self care, bathing, eating, wearing clothes etc.

b) **Work related activities**: These constitute employment related and home management related activities.

c) **Leisure time activities**: These constitute sports and social activities.
Initial emphasis in rehabilitation is on restoring the abilities of the person by physiotherapy measures such as exercises, positioning etc. Even during this period, occupational therapy helps the patient to be as independent as possible. When it is not possible to achieve any further improvement, adaptation is done in the patient’s environment, so that he is able to maintain independence. Psychological adaptation constitutes an important part of this. Use of adaptive devices such as walking aids, adaptive clothing etc. is encouraged. The person is also trained to perform purposeful activities of daily living (ADL), and activities related to work and play environment. All these make him independent inspite of his disability.

DRUGS

Drugs have a limited role in orthopaedic disorders. Those used may be placed in five categories as given below:

a) **Analgesics anti-inflammatory**: These are the most important group of drugs. They are broadly divided into non-steroidal anti-inflammatory drugs (NSAIDs) and steroids. Depending upon the need, the choice varies from a primarily analgesic to a mainly anti-inflammatory drug. In long standing illnesses, it is desirable to use single daily-dose drugs e.g., Coxibs and slow release (SR) formulations.

b) **Antibacterial drugs**: These drugs are of immense value in acute infective conditions such as septic arthritis, acute osteomyelitis etc. It is important to start with a broad-spectrum drug and change over to specific drug after a culture-sensitivity report.

c) **Hormones**: The main drugs in this group are anabolic steroids, estrogens (for osteoporosis) and stilbesterol (for metastasis from prostate).

d) **Specific drugs**: These are used in certain specific disorders e.g., vitamin D for rickets, vitamin C for scurvy, etc.

e) **Cytotoxic drugs**: These are used in the treatment of malignant bone tumours.

f) **Local injections** of a depot preparation of hydrocortisone or methylprednisolone are used to control non-specific inflammation of a joint or an extra-articular lesion like tennis elbow.

**MANIPULATION**

This is a term used for a manoeuvre whereby passive movements of joints, bones or soft tissues are carried out with or without an anaesthetic, and often forcefully, as a deliberate step in treatment. It may be done for: (i) correction of deformity; (ii) improving the range of movement of a stiff joint; or (iii) relief of chronic pain in or about a joint.

a) **Manipulation for the correction of a deformity**: In this category, manipulation has its most obvious application in the reduction of fractures and dislocations. It is also used to correct a deformity due to contracted soft tissues, as in CTEV.

b) **Manipulation for stiff joints**: In the treatment of stiff joints, while efforts being made to achieve movements by joint mobilising exercises, manipulation under anaesthesia may speed up the process of recovery. The joint most amenable to manipulation is the knee. Manipulation is **strictly contraindicated** in the elbow because it may lead to increased stiffness due to ‘myositis’.

c) **Manipulation for the relief of pain**: The role of manipulation in some chronic painful conditions is empirical. It has been shown to be effective in some conditions like tennis elbow, low backache etc.

**RADIOThERAPy**

Radiotherapy is useful in the following orthopaedic disorders:

a) **Malignant tumours**: Ewing’s sarcoma is a highly radiosensitive malignant bone tumour. Radiotherapy is also used for other malignant tumours, either pre or post surgery.

b) **Benign tumours**: Giant cell tumours of the bone which are unsuitable for excision can be irradiated.

c) **Other conditions**: Recalcitrant ankylosing spondylitis.

**OPERATIVE METHODS OF TREATMENT**

An operation is a useful yet serious undertaking. A trained surgeon, modern operation theatre and adequate instruments are essential before one ventures to perform an orthopaedic operation.
Failing this, one may have to face serious complications like osteomyelitis etc. The following are some of the common orthopaedic operations:

**OSTEOTOMY**

It means the cutting of a bone (Fig-12.1). Indications for performing osteotomy are as follows:

- To correct excessive angulation, bowing or rotation of long bone.
- To correct mal-alignment of a joint.
- To permit elongation or shortening of a bone in cases of leg length inequality.
- Special indications where osteotomy is performed for purposes other than above e.g., McMurray’s osteotomy.

![Fig-12.1 Types of osteotomies](https://kat.cr/user/Blink99/)

**ARTHRODESIS**

In this operation, fusion is achieved between the bones forming a joint so as to eliminate any motion at the joint (Fig-12.2). Although fusing a joint has its disadvantages, this operation becomes necessary when the advantages of eliminating the joint are more than disadvantages of keeping a joint. For example a stiff, painful ankle may be more disabling than an arthrodesed (fused) ankle (stiff but painless). An arthrodesis is used most often for a painful, stiff joint. It is also performed for grossly unstable joints in polio etc.

**Types of arthrodesis:** An arthrodesis may be intra-articular, extra-articular or combined (Fig-12.3). In an *intra-articular arthrodesis*, the articulating surfaces are made raw and the joint immobilised in the position of optimum function until there is a bony union between the bones. In an *extra-articular*

Table–12.1 gives some of the commonly performed osteotomies and their indications.

**Table–12.1: Common osteotomies and their indications**

<table>
<thead>
<tr>
<th>Name</th>
<th>Indication</th>
</tr>
</thead>
<tbody>
<tr>
<td>McMurray’s osteotomy</td>
<td>Fracture neck of femur</td>
</tr>
<tr>
<td>Pauwel’s osteotomy</td>
<td>Osteoarthritis of the hip</td>
</tr>
<tr>
<td></td>
<td>Fracture neck of femur</td>
</tr>
<tr>
<td>High tibial osteotomy</td>
<td>Osteoarthritis of the knee</td>
</tr>
<tr>
<td>French osteotomy</td>
<td>Correction of cubitus varus deformaty</td>
</tr>
<tr>
<td>Spinal osteotomy</td>
<td>Ankylosing spondylitis</td>
</tr>
</tbody>
</table>

![Fig-12.2 X-rays of the knee, AP and Lateral views, showing arthrodesis of the knee](https://kat.cr/user/Blink99/)

![Fig-12.3 Types of arthrodesis](https://kat.cr/user/Blink99/)
arthrodèsis, an extra-capsular bridge of bone is
created between the articulating bones. This acts
as a block to movement. Triple arthrodesis (talocalcaneal, calcaneo-cuboid and talo-navicular) is
one of the most commonly performed arthrodesis.

**Position of arthrodesis of different joints:** The
best position of arthrodesis of a joint is the one that conforms to the requirements of the patient’s work.
Table–12.2 gives positions in which common joints are fused.

---

**Table–12.2: Position of arthrodesis of joints**

<table>
<thead>
<tr>
<th>Joint</th>
<th>Position</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Shoulder</strong></td>
<td>Flex. 25 , abd. 30 , int. rot. 45</td>
</tr>
<tr>
<td><strong>Elbow</strong></td>
<td></td>
</tr>
<tr>
<td>Single</td>
<td>Flex. 75</td>
</tr>
<tr>
<td>Both</td>
<td>One in flex. 70 , other in flex. 130</td>
</tr>
<tr>
<td><strong>Wrist</strong></td>
<td>Dorsiflex. 20</td>
</tr>
<tr>
<td><strong>Hip</strong></td>
<td>Flex. 15 , no add./abd.</td>
</tr>
<tr>
<td></td>
<td>neutral rotation</td>
</tr>
<tr>
<td><strong>Knee</strong></td>
<td>Flex. 5-10</td>
</tr>
<tr>
<td><strong>Ankle</strong></td>
<td></td>
</tr>
<tr>
<td>Males</td>
<td>Neutral position</td>
</tr>
<tr>
<td>Females</td>
<td>Plantar flexion for high heels</td>
</tr>
</tbody>
</table>

---

**Fig-12.4 Types of arthroplasties**

---

**Bone grafting**

Bone grafting is an operation whereby pieces of bone (bone grafts) taken from some part of a patient’s body are placed at another site. Bone grafts are sometimes taken from another person or another species. Bone grafting is usually required for stimulating bone formation in a case of non-union of a fracture, or for filling bone defects. The defect may have been caused by a disease or by a surgeon.

**Type of bone grafts:** There are three types of bone grafts (Flow chart-12.1); autograft (from the same person); allograft (from another person of the same species); xenograft (from a different species). Autografts are used most commonly.

**Autogenous grafting:** This is the ‘gold standard’ bone grafting technique. Human body has a lot of ‘spare’ bone for such use. Iliac crests are the *commonest* site for taking bone grafts. When the graft is required for osteogenic purpose (as in non-union), cancellous
bone grafts are preferred. It is available in plenty from iliac crests and upper end of tibia. When the graft is used for providing stability (as for filling bone gaps), cortical graft is used. Fibulae are the common source of cortical bone grafts.

The grafts described above are free grafts. These do not survive as it is. These primarily provide a scaffolding upon which the new bone is laid down. A bone stimulating protein called bone morphogenetic protein (BMP) is liberated from bone grafts. This helps in osteogenesis. Over a period of time the bone grafts are replaced by new, living bone.

Newer techniques of autogenous bone grafting are such that the vascularity of a graft is preserved while it is being placed on its receptor area. There are two ways of doing it. In one, the bone graft is taken along with a pedicle of muscle. The muscle (with its intact blood supply) continues to supply blood to the graft, and hence it is a vascularised graft. It is called muscle-pedicle bone graft. It is commonly used for treating non-union of fracture of the neck of the femur.

The other method of preserving blood supply of a bone graft is free vascularized bone grafting. In this, the bone (usually fibula) is taken along with the vessels supplying it. It is placed at the new site, and its vessels anastomosed to a nearby vascular bundle. This way, the graft gets its blood supply almost instantaneously. Such a graft, therefore, remains ‘as it is’, and gets incorporated with the parent bone much faster. Microsurgical techniques are required for free vascularized grafting.

Allogenous grafting: Allogenous bone grafts (Allografts) are usually required when enough bone is not available from the host e.g., where a big defect is created following a tumour resection. Such bone grafts could be obtained from another human being, living or dead. The latter is called cadaveric graft. Allogenous graft from live donors could be, for example, from the mother when larger amount of bone graft is required for a child. It could be bone from another person, which is preserved by different techniques. Some of the techniques of bone preservation are: deep freezing (at –70°C), freeze dried, preservation by decalyzing bone (decal bone), or by formalin preservation. Such preserved bone can be used for another patient at a later date. Hospitals performing tumour excision surgery in a big way have a regular department procuring bones from patients and cadavers, processing it and storing it. These are called bone banks.

Xenografting: Bone grafts from other species, usually bovine are now available off the shelf. These are available in tailor made sizes. Their use is not common yet.

Artificial bone: This is a material derived from corals. It is hydroxyapatite with porous structure. It is supposed to have osteo-conductive potential, and is being used in some countries.

Indications: Bone grafts are used mainly for three types of cases; (i) non-union of fractures – to promote union; (ii) arthrodesis of joints – to achieve fusion between joint surfaces; and (iii) filling of bone defects or cavities in a bone.
Technique: A graft may be used as a solid slab from a cortical bone (commonly a segment of the fibula), or as cancellous bone slivers or chips (commonly from the iliac crest).

TENDON TRANSFER OPERATIONS
A tendon transfer is an operation in which insertion of the tendon of a functioning muscle is moved to a new site, so that the muscle, henceforth, has a different action. The transfer operation is planned in such a way that loss of the transferred muscle’s original function does not cause problem.

Table–12.3: Principles of tendon transfers

<table>
<thead>
<tr>
<th>Donor tendon</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Should be expandable</td>
</tr>
<tr>
<td>• Minimum power 4/5</td>
</tr>
<tr>
<td>• Amplitude of excursion to match that of the recipient muscle</td>
</tr>
<tr>
<td>• Preferably a synergistic muscle</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Recipient site</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Range of movements of the joints on which the transferred muscle is expected to work should be good</td>
</tr>
<tr>
<td>• No scarring at the bed of the transferred tendon</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Technical considerations</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Transferred tendon should take a straight route</td>
</tr>
<tr>
<td>• It should be placed in subcutaneous space</td>
</tr>
<tr>
<td>• Fixation must be under adequate tension</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Patient considerations</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Age – minimum 5 years*</td>
</tr>
<tr>
<td>• The disease should be non-progressive</td>
</tr>
</tbody>
</table>

* Minimum age when a child can be trained in using the transferred muscle.

Indications: Tendon transfers have their main application in three group of conditions: (i) muscle paralysis – to restore or improve active control of a joint by utilising a healthy muscle to act in place of a paralysed one (e.g., in nerve palsy); (ii) muscle imbalance – to restore the balance between opposite groups of muscles in case one is weaker than the other (e.g., in polio); (iii) rupture of a tendon – in cases where direct suture is not practicable.

The tendon transfer operation is commonly done for radial nerve palsy (Jone’s transfer). Some of the basic principles of tendon transfer procedure are as given in Table–12.3.

EXCISION OF TUMOURS
Excision of tumours (Fig. 12.5) can be of the following types: (i) intra-lesional excision: The lesion is curetted from within, as done for a simple bone cyst; (ii) extra-lesional excision: The lesion is removed along with its wall, as done for lipoma; (iii) wide excision: The lesion is removed with a margin of normal tissue; and (iv) radical excision: The tumour is removed along with the whole compartment in which it lies.

AMPUTATIONS
Amputation is the term used when a part of the limb is removed through a bone. Disarticulation is the corresponding term, used when a limb is removed through a joint. These operations are commonly performed for tumour ablation. This topic is discussed in detail in Chapter 40.

Further Reading

What have we learnt?
- Orthopaedic treatment methods fall in two groups: Non-operative and Operative. Non-operative methods are mainly physiotherapy and medication. Operative methods are osteotomy, arthrodesis, arthroplasty, tendon transfer, etc.
- Different techniques of bone grafting can be used. The most common is autografting.
CHAPTER 13

Injuries Around the Shoulder, Fracture Humerus

TOPICS

- Relevant anatomy
- Dislocation of the shoulder
- Fracture of the clavicle
- Fracture of the surgical neck of the humerus
- Fractures of the scapula
- Fracture of the greater tuberosity of the humerus
- Dislocation of the sterno-clavicular joint
- Fracture of the shaft of the humerus
- Subluxation or dislocation of the acromio-clavicular joint

RELEVANT ANATOMY

Shoulder girdle: It comprises of the clavicle, the scapula and the humerus. These three bones articulate with one another to give the shoulder a unique feature of freedom of movement in all directions. This maximises the reach of the hand in all directions.

The clavicle is the only long bone with membranous ossification. The muscles attached to its medial and lateral thirds are responsible for displacement following a fracture. Medial end of the clavicle articulates with the sternum to form the sterno-clavicular joint, the stability of which is provided by thickened portions of the capsule. The lateral end of the clavicle articulates with the acromion process to form the acromio-clavicular joint. The stability of this joint depends partially upon the acromio-clavicular ligaments, and the more important stabilising structures, the coraco-clavicular ligaments. The latter connect the conoid tubercle on the undersurface of the lateral end of clavicle to the coracoid process. This ligament has two parts – conoid and trapezoid (Fig-13.1). These ligaments must be torn before the acromio-clavicular joint can widely displace following an injury.

The scapula is a flat bone, thickly covered by muscles. Such thick cover do not allow displacement of fractures of this bone. Also, because of its rich vascularity, scapular fractures usually unite.

The proximal end of the humerus consists of the head articulating with glenoid cavity of the scapula (the gleno-humeral joint or the shoulder joint proper). The head is separated from the greater and lesser tuberosities by anatomical neck. The region below the tuberosities where the globular upper end of the bone joins the tubular shaft of the bone is called the surgical neck. Fractures are more common at the surgical than the anatomical neck.

Fig-13.1 Anatomy of the shoulder
Shoulder (gleno-humeral) joint: This is a ball and socket joint, inherently unstable because the ‘ball’ is big and the ‘socket’ is small and shallow. Consequently, only about one-third of the humeral head is in contact with the glenoid cavity at any one time. The capsule of the shoulder joint is lax and permits freedom of movement. The strong muscles surrounding the joint contribute a great deal to the stability of this joint. The important among these are the ‘rotator-cuff’ muscles (supra-spinatus, infraspinatus, teres minor, sub-scapularis). The interval between subscapularis tendon and supraspinatus tendon is called **rotator interval**.

**FRACUTRE OF THE CLAVICLE**

This is a common fracture at all age groups. It usually results from a fall on the shoulder or sometimes on an outstretched hand.

**PATHOANATOMY**

The junction of the middle and outer-third of the clavicle is the commonest site; the other common site being the outer-third of the clavicle. This fracture is usually displaced. The outer fragment displaces medially and downwards because of the gravity and pull by the pectoralis major muscle attached to it (Fig-13.2). The inner fragment displaces upwards because of the pull by the sterno-cleidomastoid muscle attached to it.

**DIAGNOSIS**

Diagnosis is simple in most cases. There is a history of trauma followed by pain, swelling, crepitus etc. at the site of fracture. One must look for any evidence of neurovascular deficit in the distal limb. The diagnosis can be confirmed on an X-ray.

**TREATMENT**

Fractures of the clavicle unite readily even if displaced, hence reduction of the fragment is not essential. A *triangular sling* is sufficient in cases with minimum displacement. Active shoulder exercises should be started as soon as the initial severe pain subsides, usually 10-14 days after the injury. A *figure-of-8 bandage* may be applied to a young adult with a displaced fracture (Fig-13.3). It serves the purpose of immobilisation, and gives pain relief. Open reduction and internal fixation is required, either when the fracture is associated with neurovascular deficit, or in some severely displaced fractures, where it may be more of a cosmetic concern. In such cases, the fracture is fixed internally with a plate or a nail.

**COMPLICATIONS**

**Early complications:** The fractured fragment may injure the subclavian vessels or brachial plexus.

**Late complications:** Shoulder stiffness is a common complication, especially in elderly patients. It can be prevented by shoulder mobilisation as soon as the patient becomes pain free. Malunion and non-union (the latter being very rare) often cause no functional disability and need no treatment. Rarely, for a painful non-union of the clavicle, open reduction and internal fixation with bone grafting may be necessary.
FRACTURES OF THE SCAPULA
Fractures of the scapula are less common, and in most cases unimportant because patients recover well without much treatment. The scapula can break at four sites: (i) the body; (ii) the neck; (iii) the acromion process and (iv) the coracoid process. Most often the fracture is undisplaced because the fragments are held in position by the surrounding muscles.

TREATMENT
The mainstay of treatment is to restore shoulder mobility by active exercises as soon as the pain subsides. A triangular sling for the period of pain and swelling (usually 1 week – 10 days) is usually sufficient.

DISLOCATION OF THE STERNO-CLAVICULAR JOINT
This is a rare injury. Here, the medial end of the clavicle is displaced forwards, or rarely backwards. Diagnosis is easier clinically than radiologically, because it is difficult to visualise this joint on X-ray.

TREATMENT
This is by reduction using direct pressure over the dislocated end. Reduction is maintained by a figure-of-8 bandage. Recurrence is common, but causes no disability.

SUBLUXATION OR DISLOCATION OF THE ACROMIO-CLAVICULAR JOINT
This is an uncommon injury, caused by a fall on the outer prominence of the shoulder.

PATHOANATOMY
The injury may result in a partial or complete rupture of the acromio-clavicular or coraco-clavicular ligaments. Acromio-clavicular joint injuries are divided into three grades depending upon their severity (Table–13.1).

DIAGNOSIS
Pain and swelling localised to the acromio-clavicular joint indicates an injury to this joint. In a Grade III injury the lateral end of the clavicle may be unusually prominent. X-ray with the acromio-clavicular joints of both sides, for comparison, in the same film will show the subluxation or dislocation.

TREATMENT
Grades I and II injuries are treated by rest in a triangular sling and analgesics. Grade III injury in young athletic individuals is treated by surgical repair.

DISLOCATION OF THE SHOULDER
This is the commonest joint in the human body to dislocate. It occurs more commonly in adults, and is rare in children. Anterior dislocation is much more common than posterior dislocation.

Shoulder instability: This is a broad term used for shoulder problems, where head of the humerus is not stable in the glenoid. It has a wide spectrum - from minor instability or a ‘loose shoulder’ to a frank dislocation. In the former, the patient may present with just pain in the shoulder, more on using the shoulder. Pain occurs due to stretching of the capsule, as the head ‘moves out’ in some direction without actually dislocating. A patient with frank instability may present with an ‘abnormal’ movement of the head of the humerus. This could be partial movement (subluxation) which gets spontaneously reduced, or a dislocation. The instability may be in one direction (unidirectional) or more (bidirectional). It may be in multiple directions – anterior, inferior, posterior, where it is called multi-directional instability (MDI).

MECHANISM
A fall on an out-stretched hand with the shoulder abducted and externally rotated, is the common mechanism of injury. Occasionally, it results from a direct force pushing the humerus head out of the glenoid cavity. A posterior dislocation may result from a direct blow on the front of the shoulder, driving the head backwards. More often, however, posterior dislocation is the consequence of an electric shock or an epileptiform convulsion.

Table–13.1: Grades of acromio-clavicular injury

<table>
<thead>
<tr>
<th>Grade</th>
<th>Pathoanatomy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grade I:</td>
<td>Minimal strain to acromio-clavicular ligament and joint capsule</td>
</tr>
<tr>
<td>Grade II:</td>
<td>Rupture of acromio-clavicular ligament and joint capsule</td>
</tr>
<tr>
<td>Grade III:</td>
<td>Rupture of acromio-clavicular ligament, joint capsule and coraco-clavicular ligaments</td>
</tr>
</tbody>
</table>
PATHOANATOMY

Classification: Dislocations of the shoulder may be of the following types:

a) Anterior dislocation: In this injury, the head of the humerus comes out of the glenoid cavity and lies anteriorly. It may be further classified into three subtypes depending on the position of the dislocated head (Fig-13.4).
   - Preglenoid: The head lies in front of the glenoid.
   - Subcoracoid: The head lies below the coracoid process. Most common type of dislocation.
   - Subclavicular: The head lies below the clavicle.

b) Posterior dislocation: In this injury, the head of the humerus comes to lie posteriorly, behind the glenoid.

c) Luxatio erecta (inferior dislocation): This is a rare type, where the head comes to lie in the subglenoid position.

Pathological changes: The following pathological changes occur in the commoner, anterior dislocation (Fig-13.5):

a) Bankart’s lesion: Dislocation causes stripping of the glenoidal labrum along with the periosteum from the antero-inferior surface of the glenoid and scapular neck. The head thus comes to lie in front of the scapular neck in the pouch thereby created. In severe injuries, it may be avulsion of a piece of bone from antero-inferior glenoid rim, called bony Bankart lesion.

b) Hill-Sachs lesion: This is a depression on the humeral head in its postero-lateral quadrant, caused by impingement by the anterior edge of the glenoid on the head as it dislocates.

c) Rounding off of the anterior glenoid rim occurs in chronic cases as the head dislocates repeatedly over it.

d) There may be associated injuries: like fracture of greater tuberosity, rotator-cuff tear, chondral damage etc.

DIAGNOSIS

Presenting complaints: The patient enters the casualty with his shoulder abducted and the elbow supported with opposite hand. There is a history of a fall on an out-stretched hand followed by pain and inability to move the shoulder. There may be a history of similar episodes in the past.

On examination: The patient keeps his arm abducted. The normal round contour of the shoulder joint is lost, and it becomes flattened. On careful inspection, one may notice fullness below the clavicle due to the displaced head. This can be felt by rotating the arm. The following are some of the signs, associated with anterior dislocation mostly of academic significance:
   - Dugas’ test: Inability to touch the opposite shoulder.
• **Hamilton ruler test:** Because of the flattening of the shoulder, it is possible to place a ruler on the lateral side of the arm. This touches the acromion and lateral condyle of the humerus simultaneously.

The diagnosis is easily confirmed on an anteroposterior X-ray of the shoulder (Fig-13.6). An axillary view is sometimes required.

**Posterior dislocation** usually occurs following a convulsion. There are few symptoms and signs.

![Fig-13.6 X-ray of the shoulder, AP view, showing anterior dislocation of the shoulder](https://kat.cr/user/Blink99/)

This injury is often missed even on X-ray. A clinical examination eliciting loss of external rotation, and a careful look at the X-ray may help diagnose these cases. CT scan may be diagnostic.

**TREATMENT**

Treatment of acute dislocation is reduction under sedation or general anaesthesia, followed by immobilisation of the shoulder in a chest-arm bandage for three weeks. After the bandage is removed, shoulder exercises are begun.

**TECHNIQUES OF REDUCTION OF SHOULDER DISLOCATION**

**Kocher's manoeuvre:** This is the most commonly used method. The steps* are as follows: (i) traction—with the elbow flexed to a right angle steady traction is applied along the long axis of the humerus; (ii) external rotation—the arm is rotated externally; (iii) adduction—the externally rotated arm is adducted by carrying the elbow across the body towards the midline; and (iv) internal rotation—the arm is rotated internally so that the hand falls across to the opposite shoulder.

**Hippocrates manoeuvre:** In this method, the surgeon applies a firm and steady pull on the semi-abducted arm. He keeps his foot in the axilla against the chest wall. The head of the humerus is levered back into position using the foot as a fulcrum.

A fracture of the greater tuberosity, often associated with an anterior dislocation usually comes back to its position as the head is reduced, and needs no special treatment.

**COMPLICATIONS**

Complications can be divided into early and late.

**Early complications:** Injury to the axillary nerve may occur resulting in paralysis of the deltoid muscle, with a small area of anaesthesia over the lateral aspect of the shoulder. The diagnosis is confirmed by asking the patient to try to abduct the shoulder. Though shoulder abduction may not be possible because of pain, one can feel the absence of contraction of the deltoid. Treatment is conservative, and the prognosis is good.

**Late complications:** The shoulder is the commonest joint to undergo recurrent dislocation. This results from the following causes: (i) anatomically unstable joint e.g., in Marfan’s syndrome; (ii) inadequate healing after the first dislocation, or (iii) an epileptic patient.

**Treatment:** If the disability is troublesome, operation is required. The following operations may be considered:

a) **Putti-Platt operation:** Double-breasting of the subscapularis tendon is performed in order to prevent external rotation and abduction, thereby preventing recurrences.

b) **Bankart’s operation:** The glenoid labrum and capsule are re-attached to the front of the glenoid rim. This is a technically demanding procedure, but has become simpler with the use of special fixation devices called anchors.

c) **Bristow’s operation:** In this operation, the coracoid process, along with its attached muscles, is osteotomized at its base and fixed to lower-half of the anterior margin of the glenoid. The muscles attached to the coracoid provide a dynamic anterior support to the head of the humerus.

*To remember the steps of reduction, remember TEA-I (Traction, Ext. rotation, Adduction, Int. rotation).*
Arthroscopic Bankart repair (Fig-13.7): With the development of arthroscopic techniques, it has become possible to stabilise a recurrently unstable shoulder arthroscopically. Initially it was considered suitable for cases where number of dislocations has been less than 5. But, with present day arthroscopic techniques, it is possible to stabilise most unstable shoulders arthroscopically. Apart from being a more cosmetic option, the rehabilitation after arthroscopic repair is faster and better. It is a technically demanding operation, and the anchor sutures used for repair are expensive. This technique is available only in select centres.

**FRACTURE OF THE SURGICAL NECK OF THE HUMERUS**

Fracture through the surgical neck of the humerus occurs most often in elderly women. The fracture is usually caused by a fall on the shoulder. In the majority of cases, these fractures are impacted; sometimes they are widely displaced. The possibility of this fracture should be kept in mind in all elderly persons complaining of pain in the shoulder following a fall. Often the symptoms are minimal.

It is important to properly evaluate these fractures by AP and axial X-rays. Neer has classified these fractures into 4 types depending upon the construction of the fracture. He identified 4 parts in the upper end of the humerus – shaft, head, greater tuberosity and lesser tuberosity. Depending upon in how many parts the bone has fractured, he divided them into one to four part fracture. For example, a fracture where the head, the greater tuberosity, the lesser tuberosity and the shaft, all have separated, it will be called a four-part fracture. This classification helps in deciding the treatment and prognosis.

**TREATMENT**

In elderly persons, even with moderate displacements, it is generally adequate to immobilise the affected shoulder in a triangular sling. As soon as the pain subsides, shoulder mobilisation is started. In younger persons, if the fragments are widely displaced, they are reduced by manipulation under anaesthesia. Once reduced, the fracture can be stabilised by multiple K-wires passed percutaneously under image intensifier control. Often, open reduction and internal fixation may be required. A number of internal fixation devices have been in use; from simple K-wires to modern LCP based special plates (Fig-13.8). In badly comminuted fractures in an elderly, replacement arthroplasty is desirable. Axillary nerve palsy and shoulder stiffness are common complications.

**FRACTURE OF THE GREATER TUBEROSITY OF THE HUMERUS**

Fracture of the greater tuberosity of the humerus occurs in adults. The fracture is usually caused by a fall on the shoulder, and is undisplaced and comminuted. Sometimes, it is widely separated due to the pull by the muscle (supraspinatus) attached to it.
Injuries Around the Shoulder, Fracture Humerus

TREATMENT
For minimally displaced, comminuted fractures, rest in a triangular sling is enough. The shoulder is mobilised as soon as the pain subsides. For displaced fractures, reduction is achieved by either holding the shoulder abducted in a plaster cast, or by open reduction and internal fixation. Painful arc syndrome (see page 304) and shoulder stiffness are the usual complications.

FRACTURE OF THE SHAFT OF THE HUMERUS
This is a common fracture in patients at any age. It is usually sustained from an indirect twisting or bending force – as may be sustained in a fall on out-stretched hand or by a direct injury to the arm.

RELEVANT ANATOMY
The humerus is a typical long bone (see page 8). The upper-half of the shaft is roughly cylindrical, and begins to flatten in its lower-half in the antero-posterior direction. The deltoid muscle is inserted on the deltoid tuberosity on the antero-lateral surface of the bone just proximal to its middle-third. The posterior surface is crossed obliquely by a shallow groove for the radial nerve.

The humerus is surrounded by muscles. This has the following clinical relevance: (i) the incidence of compound fractures is low; (ii) the union of fractures occurs early because a bone so well surrounded by muscles has a rich periosteal blood supply; and (iii) some degree of malunion is masked by the thick muscle cover.

PATHOANATOMY
A humerus fracture can be considered a prototype fracture because it occurs in all patterns (transverse, oblique, spiral, comminuted, segmental etc.), may be closed or open, and may be traumatic or pathological.

Displacements are variable. It may be an undisplaced fracture, or there may be marked angulation or overlapping of fragments. Lateral angulation is common because of the abduction of the proximal fragment by the deltoid muscle (Fig-13.9). This angulation is further increased by the tendency of the patient to keep the limb by the side of his chest, resulting in adduction of the distal fragment. Often, distraction occurs at the fracture site because of the gravity.

DIAGNOSIS
Diagnosis is simple because the patient presents with the classic signs and symptoms of a fracture. There may be wrist drop, if the radial nerve is injured. An X-ray of the whole arm including the shoulder and elbow should be done.

TREATMENT
For minimally displaced, comminuted fractures, rest in a triangular sling is enough. The shoulder is mobilised as soon as the pain subsides. For displaced fractures, reduction is achieved by either holding the shoulder abducted in a plaster cast, or by open reduction and internal fixation. Painful arc syndrome (see page 304) and shoulder stiffness are the usual complications.

Conservative methods: The following conservative methods are useful in most cases:

a) U-slab (Fig-13.10a): This is a plaster slab extending from the base of the neck, over the shoulder onto the lateral aspect of the arm;
under the elbow to the medial side of the arm. It should be moulded on the lateral side of the arm in order to prevent lateral angulation. The U-slab is supported with a triangular sling. Once the fracture unites, the slab is removed (approximately 6-8 weeks) and shoulder exercises started.

b) **Hanging cast** (Fig-13.10b): It is used in some cases of lower-third fractures of the humerus. The weight of the limb and the cast is supposed to provide necessary traction to keep the fracture aligned.

c) **Chest-arm bandage:** The arm is strapped to the chest. This much immobilisation is sufficient for fracture of the humerus in children less than five years of age.

In adults, early mobilisation of the limb can be begun by using a cast-brace once the fracture becomes sticky.

**Operative method:** In cases where a reduction is not possible by closed manipulation or if the fracture is very unstable, open reduction and internal fixation is required. Most fractures can be fixed well with plate and screws. Intramedullary nailing is another method of internal fixation. Contaminated open or infected fractures are stabilised by using an external fixator.

**COMPLICATIONS**

1. **Nerve injury:** The radial nerve is commonly injured in a fracture of the humeral shaft. The injury to the nerve is generally a neurapraxia only. It may be sustained at the time of fracture, during manipulation of the fracture or while the fracture is healing (nerve entrapment in the callus). A special type of humerus fracture, where there is a spiral fracture at the junction of the middle and distal third, is commonly known to be associated with a radial nerve palsy. This is called Holstein Lewis fracture. The radial nerve injury results in paralysis of the wrist, finger and thumb extensors (wrist drop), brachioradialis and the supinator. There is a sensory change in a small area on the radial side of the back of the hand.

**Treatment:** For cases reporting early, treatment depends on the expected type of nerve injury (for details, refer to Chapter 10). In most closed fractures, the nerve recovers spontaneously. In open fractures, exploration is usually required. In neglected cases or when repair of a divided nerve is impractical, tendon transfers are needed. Modified Jone’s transfer is most popular. Here the muscles of the forearm, supplied by median and ulnar nerves, are used for substituting wrist extension, finger extension and thumb abduction-extension. The following tendons are used:

   - Pronator teres → Ext. carpi radialis brevis
   - Flex. carpi ulnaris → Ext. digitorum
   - Palmaris longus → Ext. pollicis longus

2. **Delayed and non-union:** Fractures of the shaft of the humerus, especially transverse fracture of the midshaft, often go into delayed or non-union. The causes of non-union are: inadequate immobilisation or distraction at the fracture site because of the gravity.

**Treatment:** Open reduction, internal fixation with a plate, and bone grafting is usually performed. In cases where the quality of bone is poor, an intra-medullary fibular graft may be used to enhance the fixation. The limb is suitably immobilised using a U-slab or a shoulder spica.

**Further Reading**

What have we learnt?

- Shoulder is a group of joints — sterno-clavicular, acromio-clavicular, gleno-humeral and scapulo-thoracic.
- Rotator cuff muscles are important for shoulder functions.
- Fracture of the clavicle is usually treated non-operatively.
- Anterior shoulder dislocation is a common injury. It frequently leads to recurrent dislocation.
- Treatment of humerus shaft fractures is essentially non-operative. Unstable fractures need to be operated. Non-union is treated with plating and bone grafting.

Additional information: From the entrance exams point of view

- Inferior capsule is the weakest portion of the shoulder joint.
- Tests for anterior glenohumeral instability are apprehension test, fulcrum test, crank test, Jobe’s relocation test and surprise test.
- Test for posterior glenohumeral instability is jerk test.
- Sulcus test done for multi-directional and inferior instability.
- Lift off test evaluates subscapularis muscle activity.
Injuries Around the Elbow

TOPICS

- Relevant anatomy
- Pulled elbow
- Supracondylar fracture of the humerus
- Fracture of the olecranon
- Fracture of the lateral condyle of the humerus
- Fracture of the head of the radius
- Intercondylar fracture of the humerus
- Fracture of neck of the radius
- Fracture of the medial epicondyle of the humerus
- Fracture of the capitulum
- Dislocation of the elbow joint

RELEVANT ANATOMY

The elbow joint is a hinge joint, formed by the articulation between the lower end of the humerus with the ulna (humero-ulnar joint), and with the head of the radius (humero-radial joint). The lower end of the humerus is enlarged to form the trochlea medially and capitulum laterally. Medial to the trochlea is a prominent process i.e., medial epicondyle, and lateral to the capitulum is the lateral epicondyle. The two epicondyles are continuation of the medial and lateral supracondylar ridges respectively. The lateral epicondyle and capitulum together constitute the lateral condyle.

Three bony points relationship: The three prominent bony points around the elbow i.e., the medial epicondyle, lateral epicondyle and tip of the olecranon are important landmarks in the diagnosis of injuries around the elbow. Normally, in an elbow flexed to 90°, these three bony points form a near-isosceles triangle (Fig-14.1), but they lie in a straight horizontal line in an extended elbow. The base of the triangle (between the two epicondyles) is the longest arm. The side between the medial epicondyle and olecranon tip is the shortest. The head of the radius, also considered the 4th bony point, can be palpated in a semi-flexed elbow, just distal to the lateral epicondyle. It can be better felt moving during supination-pronation of the forearm.

Carrying angle: When the elbow joint is fully extended and supinated, the forearm and the arm do not lie in a straight line, but form an angle (Fig-14.2). This is called the carrying angle. It disappears on flexing the elbow. The normal carrying angle is 11° in males and 14° in females. In injuries around the elbow this angle may decrease or increase.
**Stability of the elbow:** This mainly depends upon the inherent stability of the articulating surfaces of the elbow joint i.e., the olecranon and the trochlea. The strong capsule and collateral ligaments add to the stability. The head of the radius rotates within an annular ligament which encircles it. In children, the head can slip out of this ligament (pulled elbow).

**Ossification around the elbow:** Knowledge of the appearance and fusion of different ossification centres around the elbow is necessary because these are sometimes mistaken for a fracture. Fig-14.3 shows the time of appearance of these epiphyses.

**MECHANISM OF INJURY**
Injuries around the elbow may result from an indirect or direct violence.

**Indirect violence:** This is the commoner of the two mechanisms. The cause is generally a fall onto the out-stretched hand. The type of force to which the elbow is subjected determines which fracture will occur. The elbow may be forced into valgus, varus, or hyperextension resulting in different injuries (Table–14.1).

**Direct violence:** This results either from a fall on to the point of the elbow or a direct hit on the olecranon. The result may be: (i) an olecranon fracture or (ii) an intercondylar fracture of the humerus.

**SUPRACONDYLAR FRACTURE OF THE HUMERUS**
This is one of the most serious fractures in childhood as it is often associated with complications.

**MECHANISM**
The fracture is caused by a fall on an out-stretched hand. As the hand strikes the ground, the elbow is forced into hyperextension resulting in fracture of the humerus above the condyles.

**PATHOANATOMY**
The fracture line extends transversely through the distal metaphysis of humerus just above the condyles.

**Types:** A supracondylar fracture may be of extension or flexion type, depending upon the displacement of the distal fragment (Fig-14.4).

The extension type is the commoner of the two. In this, the distal fragment is extended (tilted backwards) in relation to the proximal fragment. In the flexion type, the distal fragment is flexed (tilted forwards) in relation to the proximal fragment. Subsequent text is limited to the commoner, extension type of supracondylar fracture.

**Table–14.1: Injuries around the elbow and their mechanisms**

<table>
<thead>
<tr>
<th>Indirect</th>
<th>Direct</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Valgus injury</td>
<td>• Fall on the point of elbow</td>
</tr>
<tr>
<td>– Fracture head of the radius</td>
<td>– Olecranon fracture</td>
</tr>
<tr>
<td>– Fracture neck of the radius</td>
<td>– Intercondylar fractures of humerus</td>
</tr>
<tr>
<td>– Avulsion fracture of medial epicondyle of the humerus</td>
<td></td>
</tr>
<tr>
<td>• Varus injury</td>
<td>• Valgus injury</td>
</tr>
<tr>
<td>– Fracture lateral condyle of the humerus</td>
<td>– Fracture head of the radius</td>
</tr>
<tr>
<td>• Hyperextension injury</td>
<td>– Hyperextension fracture of the humerus</td>
</tr>
<tr>
<td>• Axial force</td>
<td>– Axial force</td>
</tr>
<tr>
<td>– Fracture of the capitulum</td>
<td>– Fracture neck of the radius</td>
</tr>
<tr>
<td></td>
<td>– Dislocation of the elbow</td>
</tr>
</tbody>
</table>

**Fig-14.3 Epiphyseal centres around the elbow and their time of appearance**

**Fig-14.4 Types of supracondylar fractures**

https://kat.cr/user/Blink99/
**Displacements**: Commonly, a supracondylar fracture is displaced (Fig-14.5). The distal fragment may be displaced in the following directions: (i) posterior or backward shift; (ii) posterior or backward tilt; (iii) proximal shift; (iv) medial or lateral shift; (v) medial tilt; and (vi) internal rotation.

**Fig-14.5 Displacements in supracondylar fracture**

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**DIAGNOSIS**

**Presenting complaints**: The child is brought to the hospital with a history of fall, followed by pain, swelling, deformity and inability to move the affected elbow.

**On examination**: When presented early, before significant swelling has occurred, the following clinical signs may be observed:

- Unusual posterior prominence of the point of the elbow (tip of olecranon) because of the backward tilt of the distal fragment.
- Since the fracture is above the condyles, the three bony points relationship is maintained as in a normal elbow.

When presented late, gross swelling makes it difficult to appreciate these signs, thus making clinical diagnosis difficult. The possibility of interruption of the blood supply to the distal extremity because of an associated brachial artery injury, must be carefully looked for in all cases. Radial and ulnar pulses may be absent with or without signs of ischaemia (five p’s—page 36). One must look for an injury to the median nerve (pointing index) or the radial nerve (wrist drop).

**Radiological examination**: Most often, it is easy to diagnose the fracture because of wide displacement.

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**TREATMENT**

**Undisplaced** fractures require immobilisation in an above-elbow plaster slab, with the elbow in 90° flexion. In all **displaced** fractures, the child should be admitted to a hospital because serious complications can occur within the first 48 hours. The following methods of treatment are used in displaced fractures:

a) **Closed reduction and percutaneous K-wire fixation**: Most displaced fractures are easily reduced by closed reduction, but they often slip. Hence, it is best to fix them with one or two K-wires, passed percutaneously under image intensifier guidance. Where facility for image intensifier is not available, a close watch on the fracture position in plaster, is a must.

**Technique of closed reduction of a supracondylar fracture**: Closed reduction of a supracondylar fracture requires experience. It is carried out in the following steps (Fig-14.7).

- **Traction** with the elbow in 30-40° of flexion: Traction is applied for two minutes, with an assistant giving counter-traction at the arm. While in traction, the elbow is gradually...
extended and the forearm fully supinated. This manoeuvre corrects proximal displacement and medial-lateral displacements. If required, the ‘carrying angle’ of the elbow is corrected at this stage.

**Flexion in traction:** With one hand maintaining traction, the upper arm is grasped with the other hand, placing the fingers over the biceps, so that the thumb rests on the olecranon. The elbow is now flexed slowly, using the hand with which traction is being applied, so as to flex the elbow while continuous traction is maintained in the long axis of the forearm.

**Pressure over the olecranon:** While the above manoeuvre is continued, the thumb over the olecranon presses the olecranon (and with it the distal fragment) forward into flexion. Traction is maintained as the elbow is flexed to beyond 90°.

Throughout this manoeuvre the radial pulse is felt. If it is obliterated on flexion, the elbow is extended again until the pulse returns and a posterior slab is applied in whatever position achieved. Further treatment in such cases will depend upon the acceptability of reduction. If it is possible to flex the elbow beyond 90°, the fragments become locked. The intact periosteum and triceps on the dorsal aspect of fracture act as an ‘internal splint’ (Fig-14.8), thereby stabilising the reduction. A posterior slab is applied in this position for 3 weeks. It is necessary to make a check X-ray after 48 hours, and after 1 week in order to detect any redisplacement. In case no redisplacement occurs, the plaster is removed after 3 weeks.

**b) Open reduction and K-wire fixation:** In some cases, it is not possible to achieve a good position by closed methods, or the fracture gets redisplaced after reduction. In such cases, open reduction and K-wire fixation is necessary (Fig-14.9). This is also used as a first line of treatment in some open fractures, and in those requiring exploration of the brachial artery for suspected injury.

**c) Continuous traction:** This is required in cases presenting late with excessive swelling or bad
wounds around the elbow. The traction may be given with a K-wire passed through the olecranon (Smith’s traction) or a below-elbow skin traction (Dunlop’s traction). These methods are no longer used.

A general treatment plan for a supracondylar fracture is shown in Flow chart-14.1.

COMPLICATIONS

The supracondylar fracture is notorious for a number of serious complications. These can be:

(i) Immediate – occurring at the time of fracture;
(ii) Early – occurring within first 2-3 days;
(iii) Late – occurring weeks to months after the fracture.

Immediate Complications

1. Injury to the brachial artery: This is a complication commonly associated with a displaced supracondylar fracture. The brachial artery is usually injured by the sharp edge of the proximal fragment (Fig-14.10). The damage may vary from just a pressure on the artery to complete disruption. The affects of arterial occlusion at the elbow differ from case to case. Most often, enough blood gets
through the collaterals around the elbow to keep the hand alive, but flexor muscles of the forearm may suffer ischaemic damage leading to Volkmann’s ischaemia. At times, the vascular compromise may be severe enough to result in gangrene.

The treatment plan for a supracondylar fracture with an absent pulse is shown in Flow chart-14.2.

2. Injury to nerves: The median nerve is the most commonly injured nerve. Radial nerve is also sometimes affected. Spontaneous recovery occurs in most cases.

**Early Complications**

1. **Volkmann’s ischaemia:** This is an ischaemic injury to the muscles and nerves of the flexor compartment of the forearm. It is caused due to occlusion of the brachial artery by a supracondylar fracture.

*Pathophysiology:* Volkmann’s ischaemia is the result of diminished blood supply to the flexor muscles of the forearm. The muscles supplied by the anterior interosseous artery, a branch of brachial artery, are most susceptible to ischaemic damage because this artery is an end-artery*. Most commonly affected muscles are the flexor pollicis longus and medial half of flexor digitorum profundus. The muscle ischaemia leads to compartment syndrome (see page 47).

*Diagnosis:* Early diagnosis of Volkmann’s ischaemia is of extreme importance. The following are some of the early signs:

- The child complains of severe pain in the *forearm*. He is unable to move the fingers fully. Ischaemic pain is more severe than the pain due to the fracture. A child needing more than usual doses

* End artery is the one which doesn't have any collaterals joining it.

```
Flow chart-14.2 Treatment plan for supracondylar fracture with absent pulse

Supracondylar fracture*  
Radial pulse absent  
Immediate closed reduction  
Pulse return within 1 hour (A)  
Maintain in slab for 48 hours  
Proper reduction and treat as any other supracondylar fracture  
Carry on as in (A)  
Circulation improves  
Circulation deteriorates  
Carry on as in (B)  
Explore vessel and internally fix the fracture  
Pulse does not return  
Capillary circulation good  
Keep in slab under close supervision  
Proper reduction and treat as any other supracondylar fracture  
Carry on as in (A)  
Proper reduction and treat as any other supracondylar fracture  
Carry on as in (A)  
Proper reduction and treat as any other supracondylar fracture
```

* In an open supracondylar fracture with absent pulse, wound debridement and exploration of the vessel is performed, and the fracture fixed internally as a primary procedure.

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of analgesics may be developing a compartment syndrome.

- **Stretch pain:** The child complains of pain in the flexor aspect of the forearm when the fingers are extended passively.
- Swelling and numbness over the fingers occur rather late.
- There is tenderness on pressing the forearm muscles.

*Treatment:* Volkmann’s ischaemia is an emergency of the highest order. The following actions need to be taken urgently in a suspected case:

- Any external splints or bandages that might be causing constriction are removed.
- The forearm is elevated and the child encouraged to move fingers.
- If no improvement occurs within 2 hours, an urgent decompression of the tight compartment is necessary. This is done by a *fasciotomy* – an operation where the deep fascia covering the flexor muscles of the forearm is slit along its entire length.

**Late Complications**

1. **Malunion:** It is the commonest complication of a supracondylar fracture and results in a *cubitus varus* deformity. This is because the fracture unites with the distal fragment tilted medially and in internal rotation. Malunion may occur either because of failure to achieve good reduction, or displacement of the fracture within the plaster. The cubitus varus deformity is often termed the *Gun stock deformity* (Fig-14.11). Sometimes, the distal fragment unites with an excessive backward tilt, resulting in hyperextension at the elbow along with limitation of flexion – basically a change in the arc of movement at the elbow.

*Treatment:* Cubitus varus deformity is a cosmetic problem, usually without much functional impairment. Mild deformity may not require treatment, but a badly deformed elbow should be corrected. Treatment is a supracondylar corrective osteotomy (French osteotomy).

2. **Myositis ossificans:** This is an ectopic new bone formation around the elbow joint, resulting in stiffness. Massage following the injury, so commonly resorted to in some places, is a major factor responsible for it.

*Treatment:* In the early stages, the elbow is put to rest in an above-elbow slab for 3 weeks. Following this, gentle elbow mobilisation is started. In some late cases, excision of the myositic bone or excision arthroplasty of the elbow is required. Whatever treatment is undertaken, the chances of the elbow regaining full range of movement are little.

3. **Volkmann’s ischaemic contracture (VIC):** This is a sequel of Volkmann’s ischaemia. The ischaemic muscles are gradually replaced by fibrous tissue, which contracts and draws the wrist and fingers into flexion (Fig-14.12a). If the peripheral nerves are also affected, there will be sensory loss and motor paralysis in the forearm and hand.

*Fig-14.12 (a) Volkmann’s ischaemic contracture (VIC). (b) Volkmann’s sign*

**Clinical features:** There is marked atrophy of the forearm, with flexion deformity of the wrist and fingers. The skin over the forearm and hand is dry and scaly. The nails also show atrophic changes. *Volkmann’s sign* helps in deciding the cause of flexion deformity of the fingers. In this sign, it is possible to extend the fingers fully at the interphalangeal joints only when the wrist is flexed (Fig-14.12b). On extending the wrist, the fingers get flexed at the inter-phalangeal joints. This is because when the wrist is extended, the shortened flexor muscle-tendon unit is stretched over the front of the wrist, resulting in flexion of the fingers. There may be hypoaesthesia or anaesthesia of the hand.
Treatment: Mild deformities can be corrected by passive stretching of the contracted muscles, using a turn-buckle splint (Volkman’s splint). For moderate deformities, a soft tissue sliding operation, where the flexor muscles are released from their origin at the medial epicondyle and ulna, is performed (Maxpage operation). For a severe deformity, bone operations such as shortening of the forearm bones, carpal bone excision etc. may be required.

FRACTURE OF THE LATERAL CONDYLE OF THE HUMERUS
A common fracture in children, it results from a varus injury to the elbow.

PATHOANATOMY
The fracture fragment comprises of the capitulum and the lateral epicondyle. The fracture line runs obliquely upwards and laterally from the intercondylar area (Fig-14.13).

![Fig-14.13 Lateral condyle fracture](https://kat.cr/user/Blink99/)

In younger children, the greater part of the detached fragment may be cartilaginous, and therefore appears smaller on X-rays, than it is in reality. It is Salter and Harris type IV epiphyseal injury (ref. page 58).

Displacement: It is common and occurs due to the ‘pull’ of the common extensor muscles which take origin from the lateral epicondyle. The fragment is rotated outwards along its vertical and horizontal axis (Fig-14.14); sometimes even as much as 90°.

DIAGNOSIS
There is mild swelling and pain over the outer aspect of the elbow. This is associated with tenderness over the lateral epicondyle. The fracture is usually diagnosed on X-rays, as the symptoms are not much (Fig-14.15).

![Fig-14.14 Displacements in a lateral condyle fracture. (a) along horizontal axis. (b) along vertical axis](https://kat.cr/user/Blink99/)

TREATMENT
For the fracture, a type-IV epiphyseal injury, accurate reduction is important if normal growth of the elbow is to be expected. Treatment depends upon whether the fracture is displaced or not.

a) An undisplaced fracture (an uncommon situation) needs support in an above-elbow plaster slab for 2-3 weeks.

b) A displaced fracture is treated by open reduction and internal fixation using two K-wires.

COMPLICATIONS
1. Non-union: If unreduced, the fracture goes into non-union. This is either because of wide displacement of the fragment or a constant ‘pulling’ force of the extensor muscles attached to it.
result is a persistent pain or growth disturbance at the distal humeral epiphysis.

**Treatment:** If detected early (usually within 2 months), it is treated with open reduction and internal fixation. In late cases, it may not be possible to achieve any improvement even after open reduction. In such cases, it is better to accept the position and treat its consequences (deformity etc.).

2. **Cubitus valgus deformity:** Diminished growth at the lateral side of distal humerus epiphysis results in a *cubitus valgus* deformity (Fig-14.16). This may result in late ulnar nerve palsy (*tardy ulnar nerve palsy*) because of friction neuritis of the ulnar nerve as it moves over the medial epicondyle, everytime the elbow is flexed and extended.

3. **Osteoarthritis:** In cases where the articular surface is significantly disorganised, elbow osteoarthritis develops after many years. Pain and stiffness are presenting symptoms. Physiotherapy is rewarding in most cases.

**INTERCONDYLAR FRACTURE OF THE HUMERUS**

This is a common fracture in adults. It results from a fall on the point of the elbow so that the olecranon is driven into the distal humerus, splitting the two humeral condyles apart.

**PATHOANATOMY**

The fracture line may take the shape of a T or Y (Fig-14.17). The fracture is generally badly comminuted and displaced. When displaced, the two condyles fall apart and are rotated along their horizontal axis.

**DIAGNOSIS**

There is generally severe pain, swelling, ecchymosis and crepitus around the elbow. The diagnosis is confirmed on X-rays.

**TREATMENT**

It depends upon the displacement. An undisplaced fracture needs support in an above-elbow plaster slab for 3-4 weeks, followed by exercises. A displaced fracture is treated generally by open reduction and internal fixation (Fig-14.18). In cases with severe comminution, olecranon pin traction...
is given to reduce the fracture and maintain the reduction.

**COMPLICATIONS**

1. **Stiffness of the elbow**: This is a common complication because of the intra-articular nature of this fracture. There may be associated myositis ossificans. Treatment is by physiotherapy.
2. **Malunion**: The fracture usually unites, but may unite in a bad position. This leads to cubitus varus or valgus deformity. A corrective osteotomy may be required for severe deformities.
3. **Osteoarthritis**: See in section on supracondylar fracture.

**FRACTURE OF THE MEDIAL EPICONDYLE OF THE HUMERUS**

It is more commonly injured than the lateral epicondyle, because the epiphysis of the medial epicondyle appears early and fuses late with the main epiphysis of the lower humerus. Its displacement varies from minimal to displacement of the whole fragment into the elbow joint. This fracture is commonly associated with posterior dislocation of the elbow. It may be associated with an ulnar nerve injury.

**Treatment** is generally conservative, by immobilisation in an above-elbow slab. If displaced into the joint, it may require open reduction and internal fixation.

**DISLOCATION OF THE ELBOW JOINT**

Posterior dislocation is the commonest type of elbow dislocation. Other dislocations are posteromedial, posterolateral, and divergent*. It may be associated with fracture of the medial epicondyle, fracture of the head of the radius, or fracture of the coronoid process of the ulna.

Clinically, there is severe pain at the elbow. The triceps tendon stands prominent (bowstringing of triceps). The three bony points relationship is reversed. There is often an associated median nerve palsy. Diagnosis is easily confirmed on X-rays (Fig-14.19).

**Treatment**: It is by reduction under anaesthesia followed by immobilisation in an above-elbow plaster slab for 3 weeks. Elbow stiffness and myositis are common complications.

**PULLED ELBOW**

This condition occurs in children between 2-5 years of age. The head of the radius is pulled partly out of the annular ligament when a child is lifted by the wrist. The child starts crying and is unable to move the affected limb. The forearm lies in an attitude of pronation. There may be mild swelling at the elbow. It is not possible to see the subluxated head on an X-ray because it is still cartilaginous; X-rays are taken only to rule out any other bony injury.

**Treatment**: The head is reduced by fully supinating the forearm and applying direct pressure over the head of the radius. A sudden click is heard or felt as the head goes back to its place. The child becomes comfortable and starts moving his elbow almost immediately.

**FRACTURE OF THE OLECRANON**

This is usually seen in adults. It results from a direct injury as in a fall onto the point of the elbow.

**PATHOANATOMY**

The proximal fragment may be pulled proximally by the attached triceps muscle, thus creating a gap at the fracture site. The fracture may be one of the three types (Fig-14.20).
Essential Orthopaedics

Fig-14.21 X-ray of the elbow, Lateral view, showing tension-band wiring of the olecranon

Unlike usual elbow fractures, which are immobilised in 90° flexion, here it is done in 30° of flexion to relax the triceps, and thus avoid pulling the olecranon away.

Fig-14.20 Types of olecranon fractures
- **Type I**: Crack without displacement of fragments.
- **Type II**: Clean break with separation of fragments.
- **Type III**: Comminuted fracture.

**DIAGNOSIS**
Pain, swelling and tenderness are present at the point of the elbow. A crepitus or a gap between the fragments may be present. Active extension of the elbow is not possible in fractures with a gap. The diagnosis is confirmed on an X-ray (Fig-14.21).

**TREATMENT**
It depends upon the type of fracture:
- **Type I**: A crack without displacement is treated by immobilising the elbow in an above-elbow plaster slab in 30 degree* of flexion. After 3 weeks the plaster is removed and elbow exercises begun.
- **Type II**: A clean break with separation of the fragments is treated by open reduction and internal fixation using the technique of tension-band wiring (Fig-14.21). It is not possible to keep the fragments together in the plaster alone because of the constant pull exerted by the triceps.
- **Type III**: A comminuted fracture, if not separated, is treated in a plaster slab as in type-I, but if the fragments are separated, tension-band wiring or excision of the fragments may be required.

With improvement in methods of internal fixation, fracture of the olecranon, being an intra-articular fracture is treated by internal fixation wherever possible. This helps in early mobilisation of elbow, and hence achieving good range of movements.

**COMPLICATIONS**
1. **Non-union** is a common complication in cases with a gap at the fracture site which prevents the fracture from uniting. Treatment is by open reduction, internal fixation and bone grafting.
2. **Elbow stiffness** occurs in some cases. Treatment is physiotherapy. In selected cases surgical release of adhesions (arthrolysis) may be required. This can be now done arthroscopically.
3. **Osteoarthritis** occurs late, often after many years in some cases, because of the irregularity of the articular surface. Treatment is physiotherapy. In selected cases, elbow replacement may be required.

**FRACTURE OF THE HEAD OF THE RADIUS**
This is seen in adults, in contrast to fractures of the neck of the radius which occurs in children. It is a valgus injury.

**PATHOANATOMY**
The head is deformed because of scattering of fragments. Sometimes a fragment of bone becomes loose and lies inside the elbow joint. The fracture may be of the following three types (Fig-14.22):
- A crack only.
- A fragment of the head is broken off.
- Comminuted fracture (the commonest type).

**Fig-14.22 Types of fracture head of radius.**
(a) Undisplaced (b) Fragment < 1/3 (c) Fragment >1/3 (d) Comminuted
DIAGNOSIS
This fracture is often missed because of minimal symptoms. There is mild pain and swelling over the lateral aspect of the elbow. A localised tenderness over the head of the radius, located immediately distal to the lateral epicondyle in a semi-flexed elbow, and painful forearm rotation are useful signs.

TREATMENT
It depends upon the type of fracture as discussed below:

a) **A crack only**: The fracture is treated by immobilisation in an above-elbow plaster slab for 2 weeks with the elbow at $90^\circ$ of flexion and the forearm in mid pronation.

b) **A fragment of the head broken off**: If the fragment is less than $1/3$ the size of the head it can be treated as above. If it is more than $1/3$ in size, or if it is lying loose inside the joint, it needs excision.

c) **Comminuted fracture with displacement**: This is treated by excision of the head.

COMPLICATIONS
1. **Joint stiffness**: Limitations of supination-pronation is a common complication associated with this injury. Treatment is persistent physiotherapy.

2. **Osteoarthritis**: It is an uncommon complication, and occurs because of joint irregularity. It usually does not cause much disability.

FRACTURE OF THE NECK OF THE RADIUS
This fracture occurs in children. It is a valgus injury of the elbow. Displacements are usually mild, and immobilisation of such fracture in an above-elbow plaster slab for 2-3 weeks is generally sufficient. In some cases with severe angulation (usually more than $60^\circ$), it may be possible to achieve acceptable reduction by closed manipulation. Sometimes, open reduction and fixation with K-wire is required. Cubitus valgus deformity may occur in a malunited fracture.

FRACTURE OF THE CAPITULUM
This is an uncommon fracture, seen in adults. The chipped off capitulum may get displaced into the joint. Due to overlap of bones, the fracture fragment may go unnoticed on X-rays. If the fragment is small or comminuted, excision is carried out. If it is a big fragment, open reduction and internal fixation is performed.

**Further Reading**

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**What have we learnt?**
- Three bony point relationship has a diagnostic value in elbow fractures.
- Supracondylar fracture is a common injury in children, and is fraught with complications such as malunion, Volkmann’s ischaemia etc.
- Fracture of the lateral condyle of humerus is a type IV epiphyseal injury, and needs primary internal fixation in most cases.
- A displaced olecranon fracture needs early surgery, as it commonly leads to non-union.
- Fractures around the elbow, commonly missed on X-rays are: (a) fracture capitulum; (b) fracture medial epicondyle; (c) fracture head or neck of radius; and (d) fracture lateral condyle.

**Additional information: From the entrance exams point of view**
- Anconeus triangle formed by radial head, lateral epicondyle and the tip of the olecranon.
- The most common cause of Volkmann’s ischaemic contracture (VIC) in a child is supracondylar fracture of the humerus.
- Most common muscle involved in VIC is flexor digitorum profundus.
- Head of radius excision leads to valgus deformity at the elbow.
The radius and ulna are common sites for fracture in all age groups. These may result from direct or indirect injury. Frequently, these fractures are open, mostly from within. Common combinations of injury in this region are: (i) fracture of both bones of the forearm; (ii) Monteggia fracture-dislocation; and (iii) Galeazzi fracture-dislocation.

RELEVANT ANATOMY

Muscles controlling supination and pronation
Supination and pronation occur at the radio-ulnar joints. The muscles producing these movements are attached to the forearm bones, and are responsible for the rotational displacement of these fractures. The supinators of the forearm (biceps and the supinator) are attached to the radius in its proximal-third (Fig-15.1). The pronators (pronator teres and pronator quadratus) are attached to the middle and distal-thirds of the radius respectively. This means that the supinators control the proximal half of the forearm whereas the pronators control the distal-half.

Therefore, in fractures of the proximal-third of the forearm bones, the proximal half of the forearm has only supinators attached to it, and is supinated. The distal half on the other hand is pronated. In fractures of the middle-third, both the proximal and the distal halves of the forearm are in midpronation. This knowledge helps in predicting relative positions of the proximal and distal halves of the forearm after a fracture. Once this is known, the reduction can be obtained by realigning the distal part of the forearm in relation to the expected rotational position of the proximal part.

Radio-ulnar articulation: The radius and ulna articulate with each other by the proximal and distal radio-ulnar joints, and interosseous membrane. Hence, an injury to the forearm usually results in fractures of both the bones. In a case where there is a fracture of only one bone, and the fracture is displaced, there should be dislocation of the proximal or the distal radio-ulnar joint.
FRACTURES OF THE FOREARM BONES

The radius and ulna are commonly fractured together – termed fracture of ‘both bones of the forearm’. Sometimes, there may be a fracture of either of the bones without much displacement. The cause of fracture may be either an indirect force such as a fall on the hand, or a direct force such as a ‘lathi’ blow to the forearm.

DISPLACEMENTS

In children, these fractures are often undisplaced, or minimally displaced (greenstick fractures), but in adults they are notoriously prone to severe displacement. A combination of any of the following displacements may occur:

- Angulation – commonly medial and anterior
- Shift – in any direction
- Rotation – the proximal and distal fragments lie in different positions of rotations (e.g., the proximal fragment may be supinated and the distal pronated).

DIAGNOSIS

It is usually simple because of the obvious signs. Fractures in children are often undisplaced and may not have much signs.

TREATMENT

Conservative treatment is sufficient in most cases. For adults with displaced fractures, operative treatment is often required.

Conservative treatment: This consists of closed reduction by manipulation under general anaesthesia, and immobilisation in an above-elbow plaster cast.

Technique of closed manipulation: The elbow is flexed to 90°. The surgeon applies traction to the hand against counter-traction by an assistant grasping the upper arm. Angulation and displacement are generally corrected by traction alone. The distal part of the forearm can now be placed in the correct rotational alignment in relation to the proximal part, as judged from the site of the fracture (Refer to page 108, Fig-15.1). Once a fracture is reduced, an above-elbow plaster cast is applied. It is important to keep the two bones apart and maintain the interosseous space, by moulding the cast while it is setting (Fig-15.2). Weekly X-rays should be taken for 3 weeks, for early detection of redisplacement.

Open reduction and internal fixation: In a large proportion of cases, especially in adults, it is impossible to obtain satisfactory reduction by closed manipulation, or to maintain it in plaster. Open reduction and internal fixation has become a popular method now. The following points have to be kept in mind:

- The radius and ulna should be approached through separate incisions to avoid cross union.
- Compression plating is the preferred method.
- The other method is intra-medullary nailing.
- Additional bone grafting should be used in fractures older than three weeks.
- The limb should be mobilised depending upon rigidity of the fixation.
- External fixation is used in some compound fracture for ease of dressing.

Deciding plan of treatment: Main point is to decide whether it is a closed or an open fracture. If it is a closed fracture, as it commonly is, the plan of treatment is as shown in Flow chart-15.1.

COMPLICATIONS

1. Infection: An open fracture of both bones of the forearm may become secondarily infected, leading to osteomyelitis.

2. Volkmann's ischaemia: This occurs within 8 hours of injury, as a result of ischaemic damage to the muscles of the flexor compartment of the forearm (For details, please refer to page 47).

3. Delayed union and non-union: Fractures of shafts of both bones of the forearm are prone to delayed union, particularly that of ulnar shaft at the junction of the middle and lower-thirds. The cause of non-union is usually inadequate immobilisation. Partial impairment of the blood supply to one of the fragments is also a contributory factor in some cases.
Treatment: Treatment of non-union of these bones is open reduction and internal fixation using plates, and bone grafting. In a non-union involving the distal 5 cm of the ulna, good functions can be achieved by simply excising the short distal fragment.

4. Malunion: This results from failure to achieve and maintain a good reduction so that the bones unite in an unacceptable position, leading to deformity and limitation of movement – especially that of rotation of the forearm. Treatment is open reduction and internal fixation using plates, and bone grafting.

5. Cross union: When radius and ulna fractures are joined to each other by a bridge of callus, it is called a cross union. It is likely to develop in a case where the two fractures are at the same level. It result in a complete limitation of forearm rotations.

Treatment: If the cross union is in mid-pronation, the position most suitable for function, it is left as it is. If it occurs in excessive pronation or supination, operative treatment may be required. The cross union is undone, mal-alignment corrected, and the fracture internally fixed.

**MONTEGGIA Fracture-Dislocation**

This is a fracture of the upper-third of the ulna with dislocation of the head of the radius. It is caused by a fall on an out-stretched hand. It may also result from a direct blow on the back of the upper forearm.

**TYPES**

These fall into two main categories depending upon the angulation of the ulna fracture – extension and flexion type. The *extension type*, is the commoner of the two, where the ulna fracture angulates

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To remember, in Monteggia, medial side bone (i.e. ulna) is fractured.
anteriorly (extends) and the radial head dislocates anteriorly. The flexion type is where the ulna fracture angulates posteriorly (flexes) and the radial head dislocates posteriorly.

Fig-15.4 X-rays of the forearm, AP and Lateral views, showing Galeazzi fracture-dislocation. Note the dislocated distal radio-ulnar joint (arrow)

DIAGNOSIS
In a case with an isolated fracture of the ulna in its upper half, a dislocation of the head of the radius should be carefully looked for (Fig-15.3).

TREATMENT
This is a very unstable injury, frequently redispersing even if it has been reduced once. One attempt at reduction under general anaesthesia is justified. If reduction is successful, a close watch is kept by weekly check X-rays for the initial 3–4 weeks. In case, the reduction is not possible or if redisplacement occurs, an open reduction and internal fixation using a plate is performed. The radial head automatically falls into position, once the ulna fracture is reduced.

COMPLICATIONS
Malunion occurs because of displacement of the fragment. It results in deformity and limitation of supination and pronation.

GALEAZZI FRACTURE-DISLOCATION
This injury is the counterpart of the Monteggia fracture-dislocation. Here, there is a fracture of the lower third of the radius with dislocation or subluxation of the distal radio-ulnar joint. It commonly results from a fall on an outstretched hand.

DISPLACEMENT
The radius fracture is angulated medially and anteriorly (Fig-15.4). The distal radio-ulnar joint is disrupted, resulting in dorsal dislocation of the distal end of the ulna.

DIAGNOSIS
In an isolated fracture of the distal-half of the radius, the distal radio-ulnar joint must be carefully evaluated for subluxation or dislocation.

TREATMENT
Perfect reduction is essential for complete restoration of functions, particularly rotation of the forearm. It is difficult to achieve and maintain perfect reduction by conservative methods (except in children). Most adults require open reduction and internal fixation of the radius with a plate. The dislocated radio-ulnar joint may automatically fall back in place or may require open reduction.

COMPLICATIONS
Malunion occurs because of displacement of the fragment. It results in deformity and limitation of supination and pronation.

COLLES’ FRACTURE
This is a fracture at the distal end of the radius, at its cortico-cancellable junction (about 2 cm from...
the distal articular surface), in adults, with typical displacement (Fig-15.5). It is the commonest fracture in people above forty years of age, and is particularly common in women because of post-menopausal osteoporosis. It nearly always results from a fall on an out-stretched hand.

**RELEVANT ANATOMY**

The distal end of the radius articulates with the carpal bones (radio-carpal joint), and the distal end of the ulna (radio-ulnar joint). Normally, the distal articular surface of the radius faces ventrally and medially (Fig-15.6). The tip of the radial styloid is about 1 cm distal to the tip of the ulnar styloid.

**PATHOANATOMY**

Displacement: The fracture line runs transversely at the cortico-cancellous junction. In the majority of cases, one or more of the displacements described below occur; although in a few cases it may be a crack fracture without displacement. The following are the displacements seen in Colles' fracture (Fig-15.7):

- Impaction of fragments
- Dorsal displacement
- Dorsal tilt
- Lateral displacement
- Lateral tilt
- Supination

As the displacement occurs, some amount of comminution of the dorsal and lateral cortices, and that of the soft cancellous bone of the distal fragment occurs. Rarely, the whole of the distal fragment is broken into pieces. Some of the following injuries are commonly associated with Colles' fracture:

**DIAGNOSIS**

Clinical features: The patient presents with pain, swelling and deformity of the wrist. On examination, tenderness and irregularity of the lower end of the radius is found. There may be a typical ‘dinner fork deformity’ (Fig-15.8). The radial
styloid process comes to lie at the same level or a little higher than the ulnar styloid process.

**Radiological features:** It is important to differentiate this fracture from other fractures at the same site (e.g., Smith’s fracture, Barton’s fracture) by looking at the displacements.

The dorsal tilt is the most characteristic displacement. It can be detected by looking at the direction of the distal articular surface of the radius on a lateral X-ray. Normally it faces ventrally*. If after fracture it faces dorsally or becomes neutral, a dorsal tilt has occurred. Similarly, a lateral tilt can be detected on an antero-posterior X-ray. Normally the distal articular surface faces medially; if it faces laterally or becomes horizontal, a lateral tilt has occurred. Most displacements can be identified on X-ray.

**TREATMENT**
Treatment of Colles’ fracture is essentially conservative. For an undisplaced fracture, immobilisation in a below-elbow plaster cast for six weeks is sufficient. For displaced fractures, the standard method of treatment is manipulative reduction followed by immobilisation in Colles’ cast (Fig-15.9).

**Technique of closed manipulation** (Fig-15.10):
The muscles of forearm must be relaxed, either by general or regional anaesthesia. The surgeon grasps the injured hand as if he was ‘shaking hands’. The first step is to disimpact the fragments which have often been driven together. This is achieved by firm longitudinal traction to the hand against the counter-traction by an assistant who grasps the arm above the flexed elbow. Some displacements are corrected by traction alone. The surgeon now presses the distal fragment into palmar flexion and ulnar deviation using the thumb of his other hand. As this is done, the patient’s hand is drawn into pronation, palmar flexion and ulnar deviation. A plaster cast is applied extending from below the elbow to the metacarpal heads, maintaining the wrist in palmar flexion and ulnar deviation. This is Colles’ cast.

An X-ray is taken to check the success of the closed reduction. Besides displacements, it is important to look for correction of the dorsal tilts i.e., the distal articular surface of the radius must face ventrally (as in a normal case).

The patient is encouraged to move his fingers as soon as the plaster dries. In addition, the shoulder and elbow joints are moved through their full range several times in a day. It is important to make check X-rays every week for the first 3 weeks in order to detect re-displacement. The plaster is removed after six weeks and joint mobilising and muscle strengthening exercises started for the wrist and fingers.

In comminuted fractures treated by the above method, incidence of re-displacement is very high. In most elderly people, malunion is compatible with useful functions, and is acceptable. In young adults, particularly those where a dominant hand is involved, more aggressive approach of surgical stabilisation is followed. These fractures are sometimes transfixed percutaneously using two K-wires which are incorporated in the plaster.

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*Normal ventral tilt of the distal articular surface can be identified on lateral X-ray of the wrist by noting that the surface faces (i) toward the side of the thumb; and (ii) toward the thicker soft-tissues of the palm, both of which are structures on ventral side.
cast. In some centres, an external fixator is used to keep the fracture ‘distracted’, so that the stretched ligaments and periosteum keep the comminuted fragments in place (ligamentoraxis). A more recent addition to the surgical treatment is to fix these fractures with LCP (Locking compression plate).

**COMPLICATIONS**

Most patients progress rapidly to full functional recovery. Stiffness of the fingers and malunion are common complications. Other complications seen occasionally are – Sudeck’s osteodystrophy, carpal tunnel syndrome, and rupture of the extensor pollicis longus tendon.

1. **Stiffness of joints:** Finger stiffness is the commonest complication; the shoulder, wrist and elbow are the other joints which commonly get stiff. This occurs because of lack of exercise, and can be prevented by actively moving these joints. The joints which are out of plaster should be moved several times a day.

2. **Malunion:** A Colles’ fracture always unites, but malunion occurs in a large proportion of cases. The cause of malunion is redisplacement of the fracture within the plaster so that a ‘dinner fork’ deformity results. There may be a limitation of wrist movement and forearm rotation.

   **Treatment:** Not always does a malunited Colles’ fracture need treatment. Often, the only disadvantage is the ugly deformity, which does not hamper the day-to-day activities of the patient. In some active adults, the deformity and impairment of functions may be severe enough to justify correction by an osteotomy.

3. **Subluxation of the inferior radio-ulnar joint:** Shortening of the radius because of the impaction of the distal fragment leads to subluxation of the distal radio-ulnar joint. The head of the ulna becomes unduly prominent. Wrist movements, especially ulnar deviation and forearm rotations are painful and restricted.

   **Treatment:** A minor degree of displacement, especially in an elderly person may be accepted. In selected cases, excision of the lower end of the ulna (Darrach’s resection) is worthwhile.

4. **Carpal tunnel syndrome:** This uncommon complication, occurs a long time after the fracture unites. The median nerve is compressed in the carpal tunnel, which is encroached by the fracture callus. **Treatment** is decompression of the carpal tunnel.

5. **Sudeck’s osteodystrophy:** Colles’ fracture is the commonest cause of Sudeck’s dystrophy in the upper limb. It is noticed after the plaster is removed. The patient complains of pain, stiffness and swelling of the hand. The overlying skin appears stretched and glossy. **Treatment** is by intensive physiotherapy. Full recovery takes a long time, but eventually occurs.

6. **Rupture of the extensor pollicis longus tendon:** This is an extremely rare complication and occurs a long time after the fracture has united. It is either due to loss of blood supply to the tendon at the time of fracture (a tiny vessel supplying blood to a part of the tendon is severed), or due to friction the tendon is subjected to everytime it moves over a malunited fracture. **Treatment** is by tendon transfer (extensor indicis to extensor pollicis longus).

**SMITH’S FRACTURE (Reverse of Colles’ Fracture)**

This uncommon fracture is seen in adults and in elderly people. Its importance lies in differentiating it from the commoner Colles’ fracture which occurs at the same site. It differs from Colles’ fracture in that the distal fragment displaces ventrally and tilts ventrally. **Treatment** is by closed reduction and plaster cast immobilisation for 6 weeks. Complications are similar to those in Colles’ fracture.

**BARTON’S FRACTURE**

This is an intra-articular fracture of the distal radius. Here, the fracture extends from the articular surface of the radius to either its anterior or posterior cortices. The small distal fragment gets displaced and carries with it, the carpals (Fig-15.11).

Depending upon the displacement, there is a volar Barton’s fracture (anterior type), and a dorsal Barton’s fracture (posterior type). **Treatment** is closed manipulation and a plaster cast. Open reduction and internal fixation with plate may be required in those cases where closed reduction fails. It may be considered as a primary choice in young adults with significantly displaced fractures.

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Fig-15.11 Barton’s fracture: a) Anterior, b) Posterior
SCAPHOID FRACTURE

A scaphoid fracture is more common in young adults. It is rare in children and in elderly people. Commonly, the fracture occurs through the waist of the scaphoid (Fig-15.12). Rarely, it occurs through the tuberosity. It may be either a crack fracture or a displaced fracture.

DIAGNOSIS

Clinical features: Pain and swelling over the radial aspect of the wrist following a fall on an outstretched hand, in an adult, should make one suspect strongly the possibility of a scaphoid fracture. On examination, one may be able to elicit tenderness in the scaphoid fossa* (anatomical snuff box). A force transmitted along the axis of second metacarpal may produce pain in the region of the scaphoid bone.

Radiological features: Whenever suspected, an oblique view of the wrist, in addition to the antero-posterior and lateral views, is essential. Sometimes, it is just a crack fracture (Fig-15.13) and is not visible on initial X-rays. If a fracture is strongly suspected, X-rays should be repeated after 2 weeks.

TREATMENT

The treatment of a scaphoid fracture is essentially conservative. The affected hand is immobilised in a scaphoid cast for 3-4 months. Sometimes, the fracture may not be visible on the initial X-ray. All such cases with clinical suspicion of a scaphoid fracture should be treated in a scaphoid cast for 2 weeks. After two weeks the X-ray is repeated. Sometimes, the fracture becomes visible at this stage, because of resorption of the fracture ends in two weeks time. If no fracture is seen even at 2 weeks, no further treatment is required.

Scaphoid cast: This is a cast extending from below the elbow to the metacarpal heads, includes the thumb, up to the inter-phalangeal joint. The wrist is maintained in a little dorsiflexion and radial deviation (glass holding position). In widely displaced fractures, open reduction and internal fixation using a special compression screw (Herbert's screw) is required.

COMPLICATIONS

Fractures of the scaphoid bone are potentially troublesome. The incidence of complications, inspite of the best treatment, is high. The most important complications are as follows:

1. Avascular necrosis: The blood supply of the scaphoid is precarious. In fractures through the waist, there is high probability of the proximal fragment becoming avascular. The patient complains of pain and weakness of the wrist. On the X-ray one finds non-union of the fracture with sclerosis and crushing of the proximal pole of the scaphoid.

Treatment: It is a difficult problem to treat. If the patient is symptomatic, the avascular segment of the bone is excised. In some cases, the wrist develops osteoarthritis, and is treated accordingly (as discussed later).

2. Delayed and non-union: A high proportion of cases of fractures of the scaphoid go into delayed or non-union. More than one factor contributes to this. It may be because of imperfect immobilisation, the synovial fluid hindering the formation of fibrinous bridge between the fragments, or impaired blood supply to one of the fragments. The diagnosis is made on the X-ray. In delayed union, the fracture, line may persist on

* It is the fossa between tendons of extensor pollicis longus and brevis, at the wrist.
X-ray even after 4-6 months. In non-union, distinct radiological features present are: (i) rounding of the fracture surfaces; (ii) the fracture becomes rather sharply defined; and (iii) cystic changes occur in one or both fragments. In a late case of non-union, changes of wrist osteoarthritis such as joint space reduction, osteophyte formation may also be seen.

Treatment: The treatment of delayed union and non-union depends largely on the severity of the symptoms. In a case where functions are not much impaired, nothing needs to be done. In a case where there is wrist pain and weakness of grip, operative intervention is necessary. For delayed union, bone grafting is sufficient. For non-union, the type of operation depends upon the presence of associated osteoarthritis of the radio-carpal joint. Once this happens, it is too late to expect relief by aiming at fracture union alone. An excision of part of the radio-carpal joint, or its fusion may be required.

3. Wrist osteoarthritis: In some cases of scaphoid fractures, osteoarthritis of the wrist develop as a result of avascular necrosis or non-union. Treatment depends upon the symptoms. Conservative treatment with hot fomentation and physiotherapy is sufficient in most cases. In some, excision of the styloid process of the radius; or in extreme cases, wrist arthrodesis may be required.

LUNATE DISLOCATIONS

These are rare dislocations of the wrist. These are of two types: lunate dislocation and peri-lunate dislocation (Fig-15.14). In lunate dislocation the lunate dislocates anteriorly but the rest of the carpals remain in position. In peri-lunate dislocation, the lunate remains in position and the rest of the carpal bones dislocate dorsally. The former type is commoner. Treatment is usually by open reduction. Severe loss of wrist movements is inevitable. Avascular necrosis of the lunate is a common complication.

Further Reading

What have we learnt?
- Forearm fractures are common, usually displaced. Open reduction has become a method of choice in unstable, displaced fractures; and in those associated with radio-ulnar joint dislocations (Monteggia and Galeazzi).
- Colles’ fracture occurs at the cortico-cancellous function of the distal radius. It is often comminuted and displaced. It usually malunites, but this does not cause much functional disability. Colles’ fracture should be differentiated from less common Smith’s fracture and Barton’s fracture.

Additional information: From the entrance exams point of view

Fractures of both bones of the forearm, above the insertion of the pronator teres is immobilised in supination, below the insertion of the pronator teres is immobilised in mid-neutral position.
The hand is an important functional unit of the upper limb without which the whole of the upper limb becomes almost useless. This calls for adequate treatment of all hand injuries, however minor they may appear. The following discussion includes only the important hand injuries.

**Bennett's Fracture-Dislocation**

It is an oblique intra-articular fracture of the base of the first metacarpal with subluxation or dislocation of the metacarpal [Fig-16.1(a)]. It is sustained as a result of a longitudinal force applied to the thumb.

**TREATMENT**

Accurate reduction and restoration of the smooth joint surface is important. This is because, being an intra-articular fracture, if not reduced accurately, it will lead to incongruity of the articular surfaces.

This would increase the chances of developing osteoarthritis. The following methods of treatment are used:

a) **Closed reduction and percutaneous K-wire fixation** under an image intensifier, is a good technique. K-wire is used and incorporated in a plaster cast (Fig-16.2).

b) **Open reduction and internal fixation** with a K-wire or a screw may be necessary in some cases.

**Complications**

Osteoarthritis develops if the joint surface is left irregular. It may cause persistent pain and loss of grip, so the patient is disabled when attempting heavy work. Excision of the trapezium may be required in particularly painful arthritis cases.
ROLANDO’S FRACTURE
This is a complete articular, ‘T’ or ‘Y’ shaped fracture of the first metacarpal [Fig-16.1(b)] Perfect reduction is not as important as in Bennett’s fracture-dislocation. Treatment is by accurate reduction and fixation with ‘K’ wires and immobilisation in a thumb spica for 3 weeks.

FRACTURES OF THE METACARPALS
Fractures of the metacarpal shaft are common at all ages. The common causes are: (i) a fall on the hand, (ii) a blow on the knuckles (as in boxing) and (iii) crushing of the hand under a heavy object. Fracture of one or more metacarpals may occur. The fracture may be classified, according to the site, as follows:

a) Fracture through the base of the metacarpal, usually transverse and undisplaced.

b) Fracture through the shaft – transverse or oblique. These fractures are usually not much displaced because of the splinting effect of the interosseous muscles and adjacent metacarpals. When more than one metacarpal shaft is fractured, this “auto-immobilisation” advantage is lost. Such fractures are unstable and require operative treatment.

c) Fracture through the neck of the metacarpal – It commonly affects the neck of the fifth metacarpal. The distal fragment is tilted forwards. It is usually sustained when a closed fist hits against a hard object (Boxer’s fracture).

TREATMENT
Conservative treatment is sufficient in most cases. It consists of immobilisation of the hand in a light dorsal slab for 3 weeks. A minimal displacement is acceptable, but in cases with severe displacement or angulation, reduction is necessary. This is achieved in most cases by closed reduction; in some, particularly those with multiple metacarpal fractures, internal fixation with K-wires or mini plates may be required.

Mallet finger (Baseball finger) results from the sudden passive flexion of the distal interphalangeal joint so that the extensor tendon of the distal interphalangeal (DIP) joint is avulsed from its insertion at the base of the distal phalanx. Sometimes it takes a fragment of bone with it. Clinically, distal phalanx is in slight flexion. Treatment is by immobilising the DIP joint in hyperextension with the help of an aluminium splint or plaster cast.

DISLOCATION OF THE METACARPO-PHALANGEAL JOINTS
These are uncommon injuries, resulting from hyperextension of the metacarpo-phalangeal (MP) joint, so that the head of the metacarpal
button-holes through the volar capsule. The MP joint of the index finger is affected most commonly. Open reduction is required in most cases.

**AMPUTATION OF FINGERS: PRINCIPLES OF TREATMENT**

1. Every effort should be made to save as much length of the thumb as possible.
2. Amputations in children are more conservative.
3. Finger tip amputations need reconstruction in such a way that full-thickness skin covers the tip.
4. In amputations at the level of the distal phalanx, replantation is not possible.
5. Replantation is not performed in the elderly persons, or sometimes in labourers who do not need delicate functions of the hand. In such cases, rather the finger is amputated and the stump closed.
6. Thumb reconstruction is possible using microsurgical technique by: (i) replantation; (ii) pollicisation of the finger (one of the fingers is made into a thumb); or (iii) transfer of a toe with its neurovascular bundle using microsurgery.

**TENDON INJURIES OF THE HAND**

Flexor tendons of the fingers are commonly injured by sharp weapons. Extensor tendons are injured less commonly.

**DIAGNOSIS**

Often these injuries are missed. The reason is that an apparently ‘small’ cut wound in the hand is sutured as it is, without examining for the underlying tendon injury which goes unnoticed. Hence, whenever confronted with a wound over hand or wrist (or foot), one must visualise the tendons and nerves underlying that wound, and test for their function.

**Testing for flexor tendons:** For this, we must know the action of each and every flexor tendon in the hand.

*Flexor carpi radialis and flexor carpi ulnaris:* These are flexors of the wrist. To test for these, the patient is asked to palmar-flex the wrist. Normally, this motion occurs in the long axis of the forearm. In case tendon of one of the flexor carpi (radialis or ulnaris) is cut; the wrist while being flexed will deviate in the direction of the muscle whose tendon is intact. For example, if flexor carpi radialis is cut, on asking the patient to palmar-flex the wrist, one will see that the hand goes towards the ulnar side. The tendon of the muscle which is working can also be felt as it gets taut when the muscle contracts.

*Flexor digitorum:* There are two groups of these tendons, the flexor digitorum superficialis (FDS) and flexor digitorum profundus (FDP). The FDS flexes the proximal interphalangeal (PIP) joint; the FDP flexes primarily the distal interphalangeal (DIP) joint. But, since FDP runs across the PIP joint also, it causes flexion at this joint as well. To test FDP, the PIP joint of the respective finger is stabilized (Fig-16.4a), and the patient asked to flex the DIP joint. It will not be possible if FDP of that finger is cut. FDS is tested by looking at flexion at PIP joint. But, in the presence of an intact FDP, even if FDS of that finger is cut, it will be possible to flex the PIP joint by the action of the FDP (which works on both PIP & DIP joints). This makes testing of cut FDS a little tricky. To be able to test the FDS of a finger, the action of FDP of that finger is to be eliminated. This is done by hyperextending the other fingers (Fig-16.4b). By doing so, the FDP of the finger being tested is kept taut. This is because profundus tendons of all the fingers are interconnected by fibrous strands.

The arrangement of tendons of finger flexors is such at the wrist, that tendons of some fingers are cut more often than those of the others (Fig-16.5). Small digital nerves and vessels run alongside flexor tendons, and are commonly cut along with flexor tendons.
CRUSH INJURY TO THE HAND

With industrialisation, the incidence of crush injury to the hand is on the rise. In developing countries, farm injuries, machine injuries and road traffic accidents constitute a majority of such injuries. The purpose of treatment in such injuries is to restore function of the hand. With advances in microsurgical techniques and powerful antibiotics, a lot of ‘badly crushed hands’, which were not considered salvageable in the past, can now be rehabilitated to useful function.

CONSIDERATIONS FOR AMPUTATION

The most demanding aspect of treatment of a crushed hand is the assessment of the injury. The first question faced by the treating doctor is whether the hand or its part is salvageable. The only indication for a primary amputation is an irreversible loss of blood supply to the part. In the absence of such an indication, a number of factors must be considered in deciding whether an amputation is advisable. These are discussed as follows:

a) **Age of the patient:** In children, amputation is indicated only when the part is totally non-viable. However, in persons over 50 years of age, amputation of one or two digits, except the thumb, may be indicated when both digital nerves and both flexor tendons are severed.

b) **Cause of crushing:** The severity of crushing can be judged from the history of injury. High speed, machine injuries produce more crushing than those caused by fall of a heavy object onto the hand. The causative factor also determines the extent of contamination, and thereby chances of infection; which in turn influences the decision to salvage the hand or not.

c) **Time since injury:** In developing countries, often a patient reaches the hospital after considerable delay, without proper first-aid. In such situations, there is increased risk of infection and poor tissue viability, which may tilt the balance in favour of an amputation.

d) **Severity of crushing:** A systematic examination of the hand, with a viewpoint to evaluate the five tissue areas (skin, tendon, nerve, bone and joint) helps in judging the severity of crushing. When three or more of these require special procedures...
such as grafting of skin, tendon suture, alignment of bone and joint, amputation should be strongly considered.

e) **The part of the hand affected:** Every effort should be made to salvage as much of thumb and index finger as possible. One should be hesitant in amputating a finger when other fingers are also injured.

f) **Other considerations:** In some cases, the expected ultimate function of the part may not be good enough to warrant the time and effort required of the patient in not amputating the part. For example, a person engaged in manual labour may be served better by amputating a severely crushed finger, and putting him back to work, than subjecting him to a series of operations only to produce a ‘cosmetic’ finger.

**PRINCIPLES OF TREATMENT**

Once it has been decided that the crushed part of the hand can be salvaged, the purpose of treatment is to restore functions. Following basic principles guide the surgeon:

a) **Assessment of the injury:** A detailed history and thorough clinical examination is most important for accurate assessment of the injury. It is done in two stages: (i) soon after the patient is seen, and (ii) again prior to the operation. The purpose of first examination is to assess whether the injury needs care in a specialised hand unit. The basic principle guiding the assessment is that each one of the deeper structures must be considered damaged until proved otherwise. An orderly examination is helpful. Attention is first directed to the skin and then to bones, tendons and nerves.

b) **Treatment priorities:** The first priority is thorough cleaning and debridement of the wound. Next is stabilisation of fractures and dislocations, and after that is wound closure with or without skin graft, skin flaps etc. Nerves and tendons may be repaired in the primary phase of the care, but this is of secondary importance.

c) **Individual tissue considerations:** Even debridement of a crushed hand needs sufficiently experienced surgeon. Skin should be excised conservatively. Any enlargement of the skin wound must not cross a skin crease. Skeletal stabilisation is performed if fracture or dislocation is unstable. Joshi’s fixator (JESS system) is a versatile fixator for stabilising all types of fractures of the bones of the hand, with the possibility of adequate soft tissue care. Small K-wires can also be used for this purpose. Primary repair of the extensor tendons, if ends can be visualised, is usually possible. Repair of the flexor tendons must not be attempted if the wound is grossly contaminated or if extensive dissection is required to find its ends. Cut ends of the tendons are either tagged to each other or to the surrounding tissues in order to prevent retraction. Secondary suture or grafting can be carried out 3-6 weeks later in such cases. Dead muscles, and those with doubtful viability are excised with care to avoid nerves. Digital nerves can be repaired primarily in a clean wound, or they can be repaired after 3-6 weeks.

d) **Proper splintage (Fig-16.6):** Proper splintage of the hand during treatment is necessary, otherwise the ligaments at MP and IP joints shorten, causing stiffness. The ideal position of immobilisation is with the MP joints in 90° of flexion and IP joints in extension (Jame’s position). In this position, the collateral ligaments of these joints are kept. If possible, the finger tips are left visible to evaluate circulation from time to time.

e) **Supportive care:** The following supportive care is required:
   - Elevation of the hand for first 3-4 days to avoid oedema

![Fig-16.6 Position of immobilisation of a hand (Note MP joints in 90° flexion, IP joints in extension)](https://kat.cr/user/Blink99/)
• Finger movements to avoid oedema and stiffness
• Antibiotics, prophylaxis against tetanus and gas gangrene
• Suitable analgesics
• Dressings as necessary

f) **Rehabilitation:** In the initial period, this consists of exercises, wax bath and splintage. Later, various appliances may be designed to help the patient perform better. Once maximum benefit has been obtained by physiotherapy, secondary operations may be considered for further improvement in functions.

**Further Reading**

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**What have we learnt?**

• Intra-articular fracture of lst metacarpal has to be accurately reduced, as it causes disabling arthritis.
• Minimally displaced metacarpal fractures can be treated by splintage.
• Phalangeal fractures need accurate reduction, sometimes surgically.
• Tendon injuries around the hand are often missed. A thorough clinical examination of each and every tendon is the key to diagnosis.
• Crushed hand is a serious injury. Prognosis depends upon accurate initial assessment, good first-aid and splintage, and early referral to specialised facility.

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**Additional information: From the entrance exams point of view**

• The proximal fragment of a scaphoid fracture is more prone to avascular necrosis due to retrograde blood flow to the proximal fragment.
• Lunate dislocation can lead to median nerve injury.
• The incidence of injury in carpal bones is scaphoid > triquetral > trapezium > lunate.
• Bennett’s fracture is difficult to maintain in a reduced position due to the pull of the Abductor pollicis longus.
• Skier’s thumb/Gamekeeper’s thumb is an injury to the ulnar collateral ligament of the metacarpo-phalyngeal joint. It is injured during skiing, holding a catch and twisting the neck of small animals. An incomplete rupture is treated conservatively with a thumb spica or functional cast brace. A complete rupture is treated by surgical repair.
• Stener lesion occurs when the adductor pollicis aponeurosis becomes interposed between the retracted ligament, and this hinders healing.
The incidence of pelvic fractures is on the rise following the increased number of vehicular accidents. It is commonly found as one of the fractures in a patient with multiple injuries. Often this fracture is not a serious management problem in itself, but may become so, because of the visceral complications so often associated with it. These fractures occur in all age groups but are most common in young adults.

RELEVANT ANATOMY

**Pelvic ring:** The pelvis is a ring shaped structure joined in the front by the pubic symphysis and behind by the sacro-iliac joints. There are projecting iliac wings on either side, a frequent site of fractures. The pelvic ring is formed, in continuity from the front, by pubic symphysis, pubic crest, pectineal line of pubis, arcuate line of the ilium, and ala and promontory of the sacrum (Fig-17.1). Fractures in the anterior half of the ring may have an associated injury in the posterior half. Such injuries make the pelvic ring unstable.

**Stability of the pelvis:** The stability of the pelvic ring depends, posteriorly on the sacro-iliac joints and anteriorly on the symphysis pubis. The sacro-iliac joints are bound in front and behind by the strong, band-like, sacro-iliac ligaments (Fig-17.2). The pubic symphysis is reinforced by ligamentous fibres above and below it. Accessory ligaments of the pelvis, such as ilio-lumbar ligament, sacrotuberous ligaments and sacro-spinous ligaments provide additional stability to the ring.

**Nerves in relation to the pelvis:** The obturator nerve and the sacral plexus pass over the ala of the sacrum, and cross the pelvic brim. These are likely to suffer injury in fractures in this region.

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**Fig-17.1** The pelvic ring

**Fig-17.2** Transverse section through the pelvic ring
CLASSIFICATION

Marvin Tile (1988) classifies pelvis fractures on the basis of stability into three types: Types A, B and C (Table–17.1). Type A, the minimally displaced stable fractures, were previously known as ‘isolated’ fractures. Types B and C, the unstable fractures, were previously known as ‘pelvic ring disruption’ injuries.

TYPE A: Stable, minimally displaced fractures: In this type, the pelvic ring is stable and displacement is insignificant. These are avulsion fractures of the parts of pelvis and fractures of the iliac wing, pubic rami fractures and undisplaced fractures of the acetabulum. These are generally treated conservatively, and have good prognosis.

TYPE B: Unstable fractures - rotationally unstable but vertically stable: In this type of injury, the pelvis is unstable. Rotational displacement can occur but no vertical displacement can occur. Open-book injury is an example of this type where an antero-posterior force causes disruption of symphysis pubis, and thus tends to open up the pelvis (Fig-17.3). There is no vertical displacement.

Table–17.1: Classification of pelvic injuries (Tile, 1988)

<table>
<thead>
<tr>
<th>TYPE A: Stable</th>
</tr>
</thead>
<tbody>
<tr>
<td>• A1 – Fractures of the pelvis not involving the ring</td>
</tr>
<tr>
<td>• A2 – Stable, minimally displaced fractures of the ring</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TYPE B: Rotationally unstable, vertically stable</th>
</tr>
</thead>
<tbody>
<tr>
<td>• B1 – Open-book type</td>
</tr>
<tr>
<td>• B2 – Lateral compression – ipsilateral</td>
</tr>
<tr>
<td>• B3 – Lateral compression – contralateral (Bucket-handle type)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>TYPE C: Rotationally and vertically unstable</th>
</tr>
</thead>
<tbody>
<tr>
<td>• C1 – Unilateral</td>
</tr>
<tr>
<td>• C2 – Bilateral</td>
</tr>
<tr>
<td>• C3 – Associated with acetabular fracture</td>
</tr>
</tbody>
</table>

TYPE A INJURY — ISOLATED FRACTURES

This is the commonest injury but the least serious of the three types. Any part of the pelvis may be affected. The essential feature being that the pelvis remains stable. Complications are uncommon in these relatively minor fractures of the pelvis. The following are some of the fractures included in this group:

Ischio-Pubic Rami Fracture

These are the commonest of pelvic fractures. One or more rami may be fractured on one or both sides; the latter is called as straddle fracture. Displacement is usually minimal. The fracture of rami may extend into the acetabulum. There may be an associated injury to the urethra or bladder.

Clinically the patient presents with pain and tenderness over the fracture site. Sometimes, a patient with multiple injuries may not have any complaint referring to this fracture, and it is detected by the routine pelvic compression test (see page 124).

Radiologically, once an ischio-pubic rami fracture is detected, one must carefully rule out an associated fracture in the posterior half of the pelvic ring (i.e. fracture through sacrum, sacro-iliac joint or ilium). It is only after this is done that a diagnosis of ‘isolated’ pubic rami fracture can be made.

Treatment: These fractures pose no problems in successful union. Treatment is basically for relief of pain. Bed rest for 1-3 weeks is usually sufficient.
Iliac Wing Fracture
This is a relatively uncommon fracture resulting from direct injury to the wing of the ilium (e.g. in a road traffic accident). Sometimes, these patients may lose so much blood from ‘vascular’ iliac wings that they develop hypovolaemic shock. The fractures are otherwise without complications, and unite in 4-6 weeks with rest and analgesics.

Avulsion Fracture of Anterior Inferior Iliac Spine
The straight head of the rectus femoris muscle takes its origin from the anterior inferior iliac spine. Sometimes, due to a violent contraction of this muscle, as may occur during a jump, the anterior inferior iliac spine may be pulled off (avulsed). The fracture unites quickly in 3-4 weeks without any complications.

Acetabular Fractures
Some of the undisplaced or minimally displaced fractures of the acetabulum can be considered in this group of relatively ‘benign’ fractures. These fractures usually unite without any complications. Late, secondary osteoarthritis develops in some cases because of the irregularity of the articular surface following the injury.

TYPES B AND C INJURIES (RING DISRUPTION INJURIES)
These are uncommon but more important injuries because of the higher incidence of associated complications. Road traffic accidents are the commonest cause of such injuries.

PATHOANATOMY
If a portion of the pelvic ring is broken, and the fragments displaced, there must be a fracture or dislocation in another portion of the ring. The following combinations of fracture and dislocation in anterior and posterior halves of the pelvis may occur:

- Anterior
  - Fracture of superior and inferior pubic rami
  - Disruption of pubic symphysis

- Posterior
  - Fracture through ala of sacrum
  - Dislocation through SI joint
  - Fracture through ilium

Displacements: It is generally slight. The type of displacement depends upon the force causing the fracture. The following displacements may occur:

a) External rotation of the hemi-pelvis (open-book type): The pelvic ring is opened up from the front like a book. There may be a pubic symphysis disruption or rami fractures in front and damage to the sacro-iliac joint behind.

b) Internal rotation of hemi-pelvis: This may result from a lateral compression force. There may be an overlap anteriorly with or without a posterior lesion.

c) Rotation superiorly (bucket-handle type): The hemi-pelvis rotates superiorly along a horizontal antero-posterior axis.

d) Vertical displacement: This results from a vertical force causing upward displacement of half of the pelvis.

DIAGNOSIS
Clinical examination: Pelvic fractures are major injuries, often with little or no clinically obvious deformity. It may be one of the fractures in a seriously injured patient where the surgeon’s attention may be diverted to other injuries with more obvious manifestations. A pelvic fracture must be carefully looked for in all cases of road accident, especially in those with multiple injuries, those associated with hypovolaemic shock, and those with major lower limb fractures (fracture of the femur etc.). The pelvic compression test is a useful screening test in all such cases.

Pelvic compression test: The patient lies supine on the couch. The examiner compresses both iliac crests of the patient’s pelvis towards each other. Any pain during this manoeuvre or a ‘springy’ feeling, is an indicator of pelvic fracture. A pelvic distraction test may reveal similar findings.

In displaced pelvic fractures there may be shortening of one of the lower limbs. The limb may lie in external rotation. There may be a haematoma in the region of pubic symphysis or at the back, in the region of sacro-iliac joints. Palpation may reveal a localised tenderness or crepitus. A gap at the symphysis pubis is occasionally felt. There may be signs due to associated injury to the urethra, bladder or intestine etc., as discussed on page 127. There may be anaesthesia or weakness of one leg due to injury to the sciatic plexus.

Radiological examination: Pelvis with both hips-AP is the basic X-ray required for screening.
purposes. In case there is a pelvic injury, special views (inlet/outlet views) are sometimes necessary. CT scan may be needed for better evaluation in cases where operative intervention is contemplated. With current CT scan machines 3-dimensional reconstruction is possible. This helps in better evaluation of the fracture.

**TREATMENT**

The importance of treatment of pelvic fractures lies in identifying the possibility of life threatening hypovolaemic shock and associated visceral injuries. The patient should be moved as little as possible, as movement at the fracture site may result in further bleeding or fat embolism.

Once the patient is stabilised, an assessment regarding the nature of the injury is made by suitable X-ray examination. Further treatment of the pelvic fracture depends on the type of fracture and presence of associated complications. In case a complication like urethral injury etc. is present, emergency treatment for the same is executed. A pelvic fracture may fall into one of the following three categories from the treatment viewpoint:

a) **An injury with minimal or no displacement:**
   The patient is advised absolute bed rest for 3-4 weeks. Once the fracture becomes ‘sticky’ and the pain subsides, gradual mobilisation and weight bearing is permitted. It takes from 6-8 weeks for the patient to be up and about.

b) **An injury with anterior opening of the pelvis** (open-book injury): A minimal opening up (less than 2.5 cm) does not need any special treatment, and is treated on the lines of (a). Reduction is needed if the opening is more than 2.5 cm. This is done by manual pressure on the two iliac wings so as to ‘close’ the pelvic ring. The reduction thus achieved is maintained by one of the following methods:
   - **External fixator:** This is a reliable and comfortable method. Two or three pins threaded at the tip (Schanz pin) are inserted in the anterior part of the wing of the iliac bone on each side. After reduction of the displacement by manual pressure, the pins are clamped to a metal rod or frame placed transversely over the front of the pelvis (Fig-17.5b).
   - **Internal fixation:** The pubic symphysis disruption may be reduced and internally fixed with a plate.
   - **Hammock-sling traction** (Fig-17.5a): It was a popular method in the past but poses nursing problems. The patient requires prolonged hospitalisation.

c) **Injuries with vertical displacement:** These are the most difficult pelvic injuries to treat. These are treated by bilateral upper tibial skeletal traction. A heavy weight (upto 20 kg) may be required to achieve reduction. After 3 weeks,

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Fig-17.5 Treatment of ‘open-book’ injury. (a) Hammock-sling, (b) External fixator
the weight is reduced to about 10 kg to maintain the position. The traction is removed after 6-8 weeks, and the patient mobilised.

Flow chart-17.1 shows a treatment plan for pelvic ring disruption injuries.

Recently, a critical review of patients with pelvic fractures (Tile, 1988) has shown that late sequelae like persistent pain etc., occur in patients treated by non-operative methods. There is now a trend towards treating these fractures by operative reduction and stabilisation. This method allows early mobility of the patient, and is most suitable for patients with multiple injuries. It requires adequate facilities and a surgeon well-versed in operative fixation of these fractures.

COMPLICATIONS

1. Rupture of urethra: This is commonly associated in cases where wide disruption of symphysis pubis and pubic rami fractures is present. The urethra in males is more commonly injured – membranous urethra being the commonest site. The rupture may be complete or incomplete, partial thickness or full thickness. Diagnosis may be made by three cardinal signs of urethral injury i.e., blood per urethra, perineal haematoma and distended bladder.

   Treatment: It may be possible to pass a catheter gently in a case with partial and incomplete urethral tear. In case this fails, the help of a uro-surgeon should be sought. Principles of treatment are: (i) drainage of the bladder by suprapubic cystostomy;
and (ii) micturating cysto-urethrogram after 6 weeks to assess the severity of urethral stricture, and treatment accordingly.

2. **Rupture of bladder:** The bladder is ruptured in pubic symphysis disruption or pubic rami fractures. In case the bladder is full at the time of injury, the rupture is usually extra-peritoneal, and urine extravasates into perivesical space. Diagnosis may be suspected if a patient has not passed urine for a long time after the fracture. Catheterisation may be successful but only a few drops of blood-stained urine come out. A cysto-urethrogram will distinguish between a bladder and a urethral rupture.

_Treatment:_ An urgent operation is required, preferably by a urologist. The principles of treatment are: (i) to repair the rent in the bladder; (ii) drainage of the bladder by an indwelling catheter, and (ii) to drain the urine in the prevesical space.

3. **Injury to rectum or vagina:** There may be disruption of the perineum with damage to the rectum or vagina. General surgeons and gynaecologists suitably manage these injuries.

4. **Injury to major vessels:** This is a rare but serious complication of a pelvic fracture. The common iliac artery or one of its branches may be damaged by a spike of bone. Aggressive management is crucial. If facilities are available, embolisation of the bleeding vessel under X-ray control is a good procedure. In other cases, the vessel is explored surgically and ligated or repaired.

5. **Injury to nerves:** In case of major disruption of the pelvic ring with marked vertical displacement of half of the pelvis, it is common for the nerves of the lumbo-sacral plexus to be injured. The damage may be caused by a fragment pressing on the nerves, or by stretching. _Treatment_ is conservative. Recovery occurs in some cases, but in most the injury is irreversible and the consequent paralysis permanent.

6. **Rupture of the diaphragm:** A traumatic rupture of the diaphragm sometimes occurs in cases with severely displaced pelvic fractures. It is worthwhile getting an X-ray of the chest in case a patient with pelvic fracture complains of breathing trouble or pain in the upper abdomen. _Treatment_ is by surgical repair.

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**What have we learnt?**

- Pelvic fractures are serious, potentially life threatening injuries.
- These are often associated with visceral injuries.
- Prolonged hospitalization and in-bed immobilisation becomes necessary for treatment of these fractures.
- Operative fixation, resulting in early mobilisation has become a desired treatment now.

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**Additional information: From the entrance exams point of view**

- In a pelvic fracture, the blood loss is 4-8 units.
- Jumper’s fracture is a type of pelvic fracture.
- Kocher-Langenbeck approach is for posterior caudal exposure.
- Ilioinguinal approach is for internal or anterior approach.
- Extended iliofemoral approach is to expose both the anterior and posterior columns.
These constitute some of the most difficult injuries of the musculo-skeletal system, from treatment point of view. The following injuries will be discussed in this chapter: (i) dislocations of the hip; (ii) fractures of neck of the femur (intra-capsular); and (iii) inter-trochanteric fractures (extra-capsular).

RELEVANT ANATOMY
The hip joint is a ball and socket joint with inherent stability, largely as a result of the adaptation of the articulating surfaces of the acetabulum and femoral head to each other. The capsule and ligaments of the joint provide additional stability.

The acetabulum faces an angle of 30° outwards and anteriorly. The normal neck-shaft angle of the femur is 125° in adults, with 15° of anteversion. The neck is made up of spongy bone with aggregation of bony trabeculae along the lines of stress. The most important of these is the medial longitudinal trabecular stream. These run from the lesser trochanter, along the medial cortex of the neck to the postero-medial quadrant of the head (Fig-18.1). A thin vertical plate of bone springs from the compact medial wall of the shaft, and extends into the spongy bone of the neck. This is called the calcar femorale.

Blood supply of the femoral head: This comes from three main sources (Fig-18.2): (i) the medullary vessels from the neck; (ii) the retinacular vessels entering from the lateral side of the head; and (iii) the foveal vessel from the ligamentum teres. The most important of these, (i) and (ii) are generally cut off following a fracture of the neck of the femur, and sometime result in avascular necrosis of the head.

The abductor mechanism of the hip: When a person stands on one leg, the body weight tends to tilt the pelvis down on the other side. The ipsilateral hip acts as a fulcrum in this.
abductors of the hip on the side on which one is standing, contract to counter this. This helps in keeping the pelvis horizontal. This *abductor mechanism* (hip joint – neck of the femur – abductor muscles) is called *abductor lever arm*, as it acts like a lever (Fig-18.3). This has great clinical relevance in biomechanics of the hip, and is disrupted in conditions like dislocation of the hip, fracture of the neck of the femur etc.

**DISLOCATIONS OF THE HIP**

**CLASSIFICATION**

There are three main types of dislocations of the hip: (i) posterior dislocation (the commonest); (ii) anterior dislocation; (iii) central fracture-dislocation. All of these may be associated with fracture of the lip of the acetabulum.

**POSTERIOR DISLOCATION OF THE HIP**

The head of the femur is pushed out of the acetabulum posteriorly. In about 50 per cent of cases, this is associated with a chip fracture of the posterior lip of the acetabulum, in which case it is called a fracture-dislocation.

**MECHANISM OF INJURY**

The injury is sustained by violence directed along the shaft of the femur, with the hip flexed. It requires a moderately severe force to dislocate a hip, as often occurs in motor accidents. The occupant of the car is thrown forwards and his knee strikes against the dashboard. The force is transmitted up the femoral shaft, resulting in posterior dislocation of the hip. It is, therefore, also known as *dashboard injury*.

**DIAGNOSIS**

**Clinical features:** An isolated posterior dislocation of the hip is easy to diagnose. The patient presents with a history of severe trauma followed by pain, swelling and deformity (flexion, adduction and internal rotation). This is associated with a shortening of the leg. One may be able to feel the head of the femur in the gluteal region. The injury is sometimes missed, especially when associated with other more obvious injuries such as fracture of the shaft of the femur. It may go unnoticed in an unconscious patient. It is wise to X-ray the pelvis in *all* patients with fracture of the femur to avoid this mistake.

**Radiological features:** The femoral head is out of the acetabulum. The thigh is internally rotated so that the lesser trochanter is not seen. Shenton’s line is broken. One must look for any bony chip from the posterior lip of the acetabulum or from the head. A comparison from the opposite, normal side may be useful. CT scan may be necessary, in cases where an associated fracture is suspected.

**TREATMENT**

Reduction of a dislocated hip is an *emergency*, since longer the head remains out, more the chances of it becoming avascular. In most cases it is possible to reduce the hip by *manipulation under general anaesthesia*. The chip fracture of the acetabulum, if present, usually falls in place as the head is reduced. *Open reduction* may be required in cases where: (i) closed reduction fails, usually in those presenting late; (ii) if there is intra-articular loose fragment not allowing accurate reduction; and (iii) if the acetabular fragment is large and is from the weight bearing part of the acetabulum. Such a fragment makes the hip unstable.

**Technique of closed reduction:** The patient is anaesthetised and placed supine on the floor. An assistant grasps the pelvis firmly. The surgeon flexes the hip and knee at a right angle.

* It is an imaginary semi-circular line joining medial cortex of the femoral neck to lower border of the superior pubic ramus. See also Fig-26.7 on page 222.
Injuries Around the Hip

In Fig-18.4, and exerts an axial pull. Usually one hears a ‘sound’ of reduction, after which it becomes possible to move the hip freely in all directions. The leg is kept in light traction with the hip abducted, for 3 weeks. After this, hip mobilisation exercises are initiated.

**COMPLICATIONS**

1. **Injury to the sciatic nerve**: The sciatic nerve lies behind the posterior wall of the acetabulum. Therefore, it may be damaged in a posterior dislocation of the hip; more so if the dislocation is associated with a large bony fragment from the posterior lip of the acetabulum. 

   **Treatment**: Injury is a neurapraxia in most cases and recovers spontaneously. In cases where the fragment of the posterior lip is not reduced by closed method, open reduction of the fracture, and nerve exploration may be required. If the sciatic nerve is severely damaged at this level, prognosis is poor.

2. **Avascular necrosis of the femoral head**: In some 15-20 per cent of cases of dislocation, the femoral head undergoes avascular necrosis. The changes of avascular necrosis appear on X-rays generally 1-2 years after the injury. The avascular head appears dense, and gradually collapses—wholly or in part. The patient complains of pain in the hip after a seemingly painless period following treatment for a dislocated hip. Over a period of a few years, changes of osteoarthritis become apparent, clinically and radiologically. Such cases eventually need hip replacement.

3. **Osteoarthritis**: This is a late complication of hip dislocation, occurring a few to many years after the injury. The underlying cause may be an avascular deformed head, or an incongruous acetabulum and femoral head. The treatment is initially conservative. In some cases an operation may be necessary. Commonly, a total hip replacement is required (see also page 339).

4. **Myositis ossificans**: This occurs a few weeks to months after the injury. The patient complains of persistent pain and stiffness of the hip. X-rays shows a mass of fluffy new bone around the hip. It is particularly common in patients with head injury. Treatment is rest and analgesics (for details see page 52).

**ANTERIOR DISLOCATION OF THE HIP**
This is a rare injury, usually sustained when the legs are forcibly abducted and externally rotated. This may occur in a fall from a tree when the foot gets stuck and the hip abducts excessively, or in a road accident. Clinically, the limb is in an attitude of external rotation. There may be true lengthening, with the head palpable in the groin. Treatment and complications are similar to that of posterior dislocation.

**CENTRAL FRACTURE-DISLOCATION OF THE HIP**
In this common injury, the femoral head is driven through the medial wall of the acetabulum towards the pelvic cavity (Fig-18.5). The displacement of the head varies from the minimal to as much as the whole head lying inside the pelvis. Joint stiffness and osteoarthritis are inevitable. Therefore, the aim of treatment in these cases is to achieve as congruous an articular surface as possible. For this, skeletal traction is applied distally and laterally. If the
fragments fall in place and reasonably reconstitute the articular margins, the traction is continued for 8-12 weeks. In some young individuals, in whom the fragments do not fall back in place by traction, surgical reconstruction of the acetabular floor may be necessary.

**COMPLICATIONS**

Hip stiffness, myositis and osteoarthritis are common complications of this injury.

**FRACTURE OF NECK OF THE FEMUR**

There are two types of fractures of neck of the femur: intra-capsular and extra-capsular. As a matter of convention, the term ‘fracture of the neck of the femur’ is used for intra-capsular fracture of the neck (Fig-18.6). The extra-capsular fracture is usually called inter-trochanteric fracture. The necessity to differently classify these fractures in rather close proximity, is due to the fact that they behave differently in terms of outcome. Whereas, fracture of neck of the femur almost never unites, inter-trochanteric fracture unites readily. Other differences between these two 'neighbours' are as given in Table 18.1 on page 134.

**PATHOANATOMY**

Most of these fractures are displaced, with the distal fragment externally rotated and proximally migrated (Fig-18.7). These displacements also occur in inter-trochanteric fracture in which these are more marked. This is because in an intra-capsular fracture, the capsule of the hip joint is attached to the distal fragment. This capsule prevents extreme rotation and displacement of the distal fragment (and with it, the fragment "Neck" of femur)

**CLASSIFICATION**

Fractures of the neck of the femur can be classified on different basis as discussed below:

a) **Anatomical classification** (Fig-18.8): On the basis of anatomical location of the fracture, it can be classified as: (i) subcapital – a fracture just below the head; (ii) transcervical – a fracture in the middle of the neck; or (iii) basal – a fracture at the base of the neck. The more proximally the fracture is located, the worse is the prognosis.

b) **Pauwel’s classification**: This classification is based on the angle of inclination of the fracture in relation to the horizontal plane (Pauwel’s angle, Fig-18.9). The fractures are divided into three types (type I–III). The more the angle,
the more unstable is the fracture, and worse the prognosis.

c) **Garden’s classification:** This is based on the degree of displacement of the fracture (mainly rotational displacement). The degree of displacement is judged from change in the direction of the medial trabecular stream of the neck, in relation to the bony trabeculae in the weight bearing part of the head and in the corresponding part of the acetabulum (Fig-18.10).

- **Stage 1:** The fracture is incomplete, with the head tilted in postero-lateral direction, so that there is an obtuse angle laterally at the trabecular stream. This is also called an impacted or abducted fracture.
- **Stage 2:** The fracture is complete but undisplaced, so that there is a break in the trabecular stream with little angulation.
- **Stage 3:** The fracture is complete and partially displaced. As the distal fragment rotates externally, it causes internal rotation of the head. One can make out rotation of the head and displacement of the fracture by carefully following the medial trabecular stream. The trabecular stream at the fracture site is broken and displaced. Alignment between the trabeculae of the head and the acetabulum is also lost because of the rotation of the head in relation to the acetabulum.

- **Stage 4:** The fracture is complete and fully displaced. As the distal fragment rotates further outwards, it looses contact with the head, which springs back to its original position. Therefore, whereas there is a total loss of contact between the head and neck trabecular streams, those between the head and the acetabulum are normally aligned.

Though Garden’s classification is scientifically more appealing, but unless one can make proper assessment on good quality X-rays, it is difficult to decide the stage of the fracture.
MECHANISM
In elderly people, the fracture occurs with a seemingly trivial fall. Osteoporosis is considered an important contributory factor at this age. In young adults, this fracture is the result of a more severe injury. The fracture is uncommon in children.

DIAGNOSIS
Clinical features: Occasionally, a patient with an impacted fracture may arrive walking; the only complaint being a little pain in the groin. More often, the patient is an elderly, brought to the casualty department with complaints of pain in the groin and inability to move his limb or bear weight on the limb following a ‘trivial’ injury like slipping on the floor, missing a step etc. There is little pain or swelling. Often, the injury is trivial, and pain and swelling almost absent. In such cases, the fracture diagnosis is missed for days or weeks. Careful examinations reveals the following:
- External rotation of the leg, the patella facing outwards.
- Shortening of the leg, usually slight.
- Tenderness in the groin.
- Attempted hip movements painful, and associated with severe spasm.
- Active straight leg raising not possible.
Clinically, it is possible to differentiate this fracture from an inter-trochanteric fracture which presents with similar signs. In general, signs and symptoms are more in an inter-trochanteric fracture (Table–18.1).

Radiological features (Fig-18.11): It is useful to ask for X-ray of pelvis with both hips, rather than that of the affected hip alone. This helps in comparing the two sides. The following features should be noted:
- Break in the medial cortex of the neck.
- External rotation of the femur is evident; the lesser trochanter appearing more prominent*.
- Overriding of greater trochanter, so that it lies at the level of the head of the femur.
- Break in the trabecular stream.
- Break in Shenton’s line (page 222).

In impacted fracture, the only radiological finding is bending of the trabeculae. There is no clear cut

Table–18.1: Differences between fracture neck of the femur and inter-trochanteric fracture

<table>
<thead>
<tr>
<th>Point</th>
<th>Fracture neck femur</th>
<th>Inter-trochanteric fracture</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>After 50 yrs</td>
<td>After 60 yrs</td>
</tr>
<tr>
<td>Sex</td>
<td>F &gt; M</td>
<td>M &gt; F</td>
</tr>
<tr>
<td>Injury</td>
<td>Trivial</td>
<td>Significant</td>
</tr>
<tr>
<td>Ability to walk</td>
<td>May walk in impacted fracture</td>
<td>Not possible</td>
</tr>
<tr>
<td>Pain</td>
<td>Mild</td>
<td>Severe</td>
</tr>
<tr>
<td>Swelling</td>
<td>Nil</td>
<td>Severe</td>
</tr>
<tr>
<td>Ecchymosis</td>
<td>Nil</td>
<td>Present</td>
</tr>
<tr>
<td>Tenderness</td>
<td>In Scarpa’s triangle</td>
<td>On the greater trochanter</td>
</tr>
<tr>
<td>Ext. rotation deformity</td>
<td>Less than 45°</td>
<td>More than 45°</td>
</tr>
<tr>
<td>Shortening</td>
<td>Less than 1 inch</td>
<td>More than 1 inch</td>
</tr>
<tr>
<td>Treatment</td>
<td>Int. fixation always</td>
<td>Can be managed in traction</td>
</tr>
<tr>
<td>Complications</td>
<td>Non-union</td>
<td>Malunion</td>
</tr>
</tbody>
</table>

* The lesser trochanter is situated postero-medially on the shaft. So if the leg is externally rotated, the lesser trochanter appears more prominent on the X-ray.
TREATMENT

This fracture is rightly termed an ‘unsolved fracture’ because of the high incidence of complications. Two factors which make the treatment of this fracture particularly difficult are: (i) the blood supply to the proximal fragment (head) is cut off; and (ii) it is difficult to achieve reduction and maintain it because the proximal fragment is too small. Because of these factors, the fracture invariably needs operative treatment. There are numerous controversies in the treatment of this fracture. Discussed below is a balanced approach followed in most hospitals.

Impacted fracture: An impacted fracture can be treated in all age groups by conservative methods. Some surgeons fix these fractures internally with screws for fear of displacement. In children, a hip spica, and in adults immobilisation in a Thomas splint are preferred methods.

Unimpacted or displaced fractures: The aim of treatment in patients up to 60 years* of age is to achieve union. For this, internal fixation is usually required. Since incidence of failure of this type of treatment is reasonably high, in the elderly, it is preferable to excise the head of the femur and replace it by a prosthesis. In some younger patients presenting late, to achieve closed reduction of the fracture may be difficult. In such cases, an open reduction of the fracture is done. An accurate reduction and good fixation is important for a good result. Operations such as McMurray’s osteotomy, which were popular in the past, when facility of image intensifier was not available, are no longer

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* It is the physiological age which is considered. It means that the appearance, activities and health of the patient is that of a healthy person at 60 years of age. Often a patient may be more than 60 years, but physiologically younger, or vice versa.
done. Flow chart-18.1 shows the treatment plan of these fractures in adults.

**Internal fixation:** Any of the following implants may be used for internal fixation:

- Multiple cancellous screws – most commonly used.
- Dynamic hip screw (DHS) – used sometimes.
- Multiple Knowle’s pins/Moore’s pins used in children.

**The technique of internal fixation:** The technique being described here is, use of multiple cancellous screws. The screws used are partially threaded (Fig-18.11), the threaded part holds in the head, whereas the smooth part permits controlled collapse of the fracture, which helps in union. Facility of image intensifier is a must. Under anaesthesia, the patient is fixed to a special operating table (fracture table). The fracture is reduced by closed manipulation. The reduction is checked on image intensifier. The base of the greater trochanter is exposed and a guide-wire inserted in the centre of the femoral neck. If the position of the guide-wire is satisfactory, a cannulated screw is threaded over it. Minimum three screws, preferably parallel, are necessary. Cannulated screws make the operation simple. After the operation, no external immobilisation is required. The patient is allowed to sit up in bed, and be out of bed with crutches (non weight bearing) in the early post-operative period. Gradual weight bearing is permitted as the fracture shows evidence of union (usually 2-3 months).

**McMurray’s osteotomy:** This was a popular operation in yesteryears. This is an oblique osteotomy at the inter-trochanteric region. The **direction** of osteotomy is medially upwards, beginning at the base of the greater trochanter and ending just above the lesser trochanter (Fig-18.12). Once the osteotomy is made, the distal fragment is displaced medially and is abducted. The position is held by an external support (hip spica) or by internal fixation with plate and screws. The osteotomy converts the shearing stresses at the fracture site into compressive forces, thus enhancing fracture union. The line of weight bearing after the osteotomy passes from the head to the distal fragment. It therefore bypasses the fracture site. By this osteotomy the head is supported by the distal fragment (arm-chair effect). This helps in limb taking the weight on walking. This operation is done rarely now-a-days.

**Hemiarthroplasty** (Fig-18.13): This is a procedure used for elderly patients. In this, the head of the femur is excised and replaced by a prosthesis. There are two types of prosthesis commonly in use: unipolar and bipolar. Unipolar prosthesis have a ‘head’ with an attached stem. The stem is introduced inside the medullary canal of the femur, and the head sits over the neck of the femur. In bipolar prosthesis, the head has two parts: a smaller head, and a mobile plastic cup on top of it. Hence, when the prosthesis is fitted on the neck, there is movement at two planes – one between the acetabulum and the plastic cup, and other between the plastic cup and the head. This is why it is called ‘bipolar, and is supposed to be mechanically better (For details see pages 338).
Injuries Around the Hip

The patient presents either with a fracture not treated at all, or a fracture which has failed to unite even after treatment. The main complaint is pain and inability to bear weight on the affected limb. The limb is short and externally rotated. Active straight leg raising is not possible. The pain may be minimal and hip movements may be increased because of free mobility at the fracture site (pseudarthrosis). Trendelenburg’s test will be positive. Telescopy will be present.

In a patient who has been treated by internal fixation, non-union should be suspected if there is renewed pain after seemingly normal progress. Often, there is sudden deterioration in these cases; with acute pain, external rotation of the limb, shortening, and inability to walk. This is because the implant, which was ‘holding’ an ununited fracture, finally gives way (implant failure).

Treatment: The treatment of non-union depends upon the age of the patient, and on whether or not there is avascular necrosis of the femoral head. In patients beyond the age of 60 years, replacement arthroplasty is performed. In younger individuals, attempt is made to preserve the head of the femur by one of the following methods:

a) Neck reconstruction: The fracture is exposed from behind, the ends freshened and the fracture stabilized with multiple screws and vascularised muscle-pedicle bone graft taken from the femoral attachment of the quadratus femoris muscle. This operation is used in treating the fractures presenting late or those with significant comminution at the fracture site. It is also used for non-union of the femoral neck fractures.

b) Pauwel’s osteotomy: This is a valgus osteotomy at the level of the lesser trochanter (Fig. 18.15). A valgus effect so created at the fracture site

Treatment of cases presenting late: Patient with fracture of the neck of the femur often present late, either because the fracture is not diagnosed in time or the facilities for treatment were not available. These present a difficult problem because after 2–3 weeks, closed reduction is not possible, and opening the fracture is associated with complications such as avascular necrosis. In patients above 60 years of age, replacement arthroplasty is the best option. In younger patients, treatment depends upon whether the head of the femur is ‘vascular’ or not. This can be assessed by bone scanning or MRI. If the head is vascular, the hip should be reconstructed by either osteotomy (McMurray’s or Pauwel’s), or reconstruction procedures such as Meyers’ operation. If the head is not vascular, and the patient is young, bipolar prosthesis is preferred.

COMPLICATIONS

1. Non-union: It occurs in approximately 30 to 40 per cent of intra-capsular fractures. It is due to inadequate immobilisation of the fracture even with internal fixation, and its poor blood supply.
results in converting the shearing forces at the fracture site into compression forces. The osteotomy is fixed with double-angle blade plate. It is a technically demanding operation.

2. **Avascular necrosis:** After a fracture through the neck, all the medullary blood supply and most of the capsular blood supply to the head are cut off. The viability of the femoral head may therefore depend almost entirely on the blood supply through the ligamentum teres. If this blood supply is insufficient, avascular necrosis of a segment or whole of the head occurs. This may, in addition, be a cause of non-union. The avascular head may collapse and become deformed. These changes may not become evident early. It is only after a few months to as long as 2 years, that one can diagnose avascular necrosis on X-rays. MRI is the best investigation for this purpose. Deformation of the head results in osteoarthritis after a few years.

_Treatment:_ A fracture of the neck of the femur with avascular necrosis of the head is a very difficult problem to treat. In _young_ patients treatment options are between arthrodesing the hip, bipolar arthroplasty (a special type of prosthesis), Meyer’s procedure or rarely, total hip replacement (THR). In _elderly_ patients, a hemi-replacement arthroplasty is performed. In cases where there is an associated damage to the hip, a total hip replacement may be preferred.

3. **Osteoarthritis:** It develops a few years following fracture of the neck of femur. It may be because of: (i) avascular deformation of the head; or (ii) union in faulty alignment. The patient presents with pain and stiffness of the joint. Initially the pain is intermittent, but later it persists.

_Treatment:_ It depends upon the age and functional requirement of the patient. Younger patients are treated by either an inter-trochanteric osteotomy or arthrodesing of the hip. For an elderly patient, total hip replacement is the best option.

**INTER-TROCHANTERIC FRACTURES**

Fractures in the inter trochanteric region of the proximal femur, _involving either_ the greater or the lesser trochanter or both, are grouped in this category (Fig-18.16). In the elderly, the fracture is normally sustained by a sideway fall or a blow over the greater trochanter. In the young, it occurs following violent trauma, as in a road traffic accident.

![Fig-18.16 X-ray of the hip, AP view, showing inter-trochanteric fracture](image)

**PATHOANATOMY**

The distal fragment rides up so that the femoral neck-shaft angle is reduced (coxa vara). The fracture is generally comminuted and displaced. Very rarely, it can be an undisplaced fracture.

**DIAGNOSIS**

_Clinical features:_ As for fractures of the neck of the femur, the patient is brought in with a history of a fall or road accident, followed by pain in the region of the groin and an inability to move the leg. There will be swelling in the region of the hip, and the leg will be short and externally rotated. There is tenderness over the greater trochanter. The physical findings in such a case are _more marked_ compared to those in a fracture of the neck of the femur.

_Radiological features:_ Diagnosis is easy on an X-ray. Presence of comminution of the medial cortex of the neck, avulsion of the lesser trochanter and extension of the fracture to the subtrochanteric region indicate an _unstable_ fracture, and a poor prognosis.

**TREATMENT**

Contrary to fracture of the neck of the femur, trochanteric fractures _unite readily_. The main objective of treatment is to maintain a normal femoral neck-shaft angle during the process of union. This can be done by conservative means (traction) or by internal fixation. In elderly patients, internal fixation is preferred because in them prolonged bed rest (as much as 3-4 months) in traction may cause complications related to recumbency i.e., bed sores, pneumonia etc.
Conservative methods: There are a number of tractions described for an inter-trochanteric fracture. Those used most frequently are Russell’s traction (Fig-18.17) and skeletal traction in a Thomas splint. With the success of operative methods, whereby, early mobilisation is possible, conservative methods are used less often.

Operative methods: The fracture is reduced under X-ray control and fixed with internal fixation devices. The most commonly used ones are: (i) Dynamic Hip Screw (DHS) (Fig-18.18); (ii) Ender’s nails and (iii) Nails such as gamma nail, Proximal femoral nail (PFN). External fixation is useful for patients with bed sores, and for those who are unfit for a major operation.

COMPLICATIONS
1. Malunion: As discussed earlier, inter-trochanteric fractures almost always unite, but because of possible failure in keeping the fragments aligned, these often malunite. Malunion gives rise to coxa vara (decreased femoral neck-shaft angle), shortening and the leg in external rotation.

Treatment: In elderly patients, malunion does not cause a great deal of disability, except a limp while walking, and shortening. Compensation for this shortening, by giving a suitable shoe raise, suffices in most cases. In young people with severe coxa vara and shortening, correction may be required. This is achieved by an inter-trochanteric osteotomy whereby the neck-shaft angle is corrected and held in the proper position by internal fixation devices.

2. Osteoarthritis: Due to changes in the hip biomechanics following trochanteric fractures, osteoarthritis of the hip develops after a few years. The patient complains of pain and stiffness in the hip after a reasonably symptom free period following union of the fracture. An X-ray confirms changes of osteoarthritis in the hip joint.

Treatment: In the early stages, treatment is by physiotherapy. Later, a trochanteric osteotomy (in younger patient) or a total hip replacement (in elderly patient) may be required.

What have we learnt?
- Injury around the hip is common in elderly.
- Fracture neck of femur is notorious for complications such as non-union.
- Treatment of fracture neck of femur in younger patients is by internal fixation (head preservation), and in the elderly by replacement (head replacement).
- Inter-trochanteric fractures are cousins of fracture neck femur, but very different behaviour-wise. Union usually occurs, though malunion is common. These can be treated by conservative methods, but internal fixation is preferred.
Additional information: From the entrance exams point of view

- Main blood supply to the head of the femur in adults is the lateral ascending cervical or retinacular and epiphyseal branches of the medial circumflex femoral artery.
- The commonest hip injury in the elderly patient is intertrochanteric (extracapsular fractures).
- Occult fracture neck of femur is best diagnosed by MRI.
- Maximum chances of avascular necrosis in subcapital fractures.
- Fracture head of femur classified by Pipkin classification.
- Femoral head palpable on per rectal examination in central dislocation of hip.
- Paralysis of gluteus medius/minimus supplied by the superior gluteal nerve causes Trendelenburg’s gait.
A fracture of the shaft of the femur is usually sustained by a severe violence, as may occur in a road accident. The force causing the fracture may be indirect (twisting or bending force) or direct (traffic accidents).

**PATHOANATOMY**

The fracture may occur at any site and is almost equally common in the upper, middle and lower thirds of the shaft. It may be a transverse, oblique, spiral or comminuted fracture depending upon the nature of the fracturing force.

**Displacements:** In children, the fracture does not displace a great deal; but in adults, more often than not, there is marked displacement. The proximal fragment is flexed, abducted and externally rotated by the pull of the muscles attached to it (Fig-19.1). The distal fragment is adducted because of attachment of adductor muscles. The unsupported fracture end of the distal fragment sags because of the gravity. There is proximal migration (overriding) of this fragment because of the pull by the muscles going across the fracture.

**DIAGNOSIS**

**Clinical features:** The patient presents with a history of severe violence followed by classic signs of fracture in the region of the thigh (pain, swelling, deformity, abnormal mobility etc.). Diagnosis is not difficult.

**Radiological examination:** X-rays done for a femoral shaft fracture must include the whole femur. In addition, an X-ray of the pelvis should be done. It is common that a patient with fracture of the femur has an associated injury in the pelvis.

**TREATMENT**

Fracture of the shaft of the femur occurs in so many different forms that practically all methods of fracture treatment discussed in Chapter 3 may be applicable. The treatment methods can be conservative or operative.

**Conservative methods:** This consists of the following:

a) **Traction:** A fracture of the shaft of the femur can be treated by traction, with or without a splint. Usually a Thomas splint is used. Skin traction is sufficient in children, but skeletal traction is required in adults. Skeletal traction is given by a Stenmann pin passed through the upper-end of tibia.
b) **Hip spica**: This is a plaster cast incorporating part of the trunk and the limb. It may be a single spica (involving only the fractured limb) or one-and-a-half as shown in Fig-19.2. It can be safely used for immobilising these fractures in children. It may also be used for treating fractures in young adults, once the fracture becomes ‘sticky’.

With wider availability of operative methods, more and more fractures of the shaft of femur, even in children, are now treated operatively.

![Fig-19.2 Hip spica (1½)](image)

**Operative methods**: Intra-medullary nailing is the preferred method. The fracture may be reduced by closed or open methods. Plating is preferred in cases where good hold is not possible by a nail. The following are some of the commonly used methods of operative treatment:

a) **Closed Interlock nailing**: This is the preferred method of treatment of most femoral shaft fractures. In this, the fracture is reduced under X-ray control (Image intensifier), without opening it. The nail (a kind of rod) is introduced into the medullary canal from the greater trochanter under monitoring by image intensifier. This is called **closed nailing**. It is a minimally invasive counterpart of conventional nailing done by opening the fracture (**open nailing**). The recent addition to closed nailing is **interlock nailing**. In this, two horizontal screws are passed through two holes at the ends of the nail. This locks the nail in place. It is a technically demanding operation, and an image intensifier is necessary for this.

![Fig-19.3 X-rays showing intra-medullary nailing done for fracture of femoral shaft](image)

b) **Plating (fixing with a thick strip of metal)**: For fractures where medullary canal is too wide for a nail to provide a good hold, or for a comminuted fracture, plating may be used (Fig-19.4). Minimum of 16 cortex hold (8 screws) is desirable. AO heavy duty plates with or without compression may be used. Special condylar blade-plate may be used for fractures closer to either end of the bone. With the advent of interlock nailing, the fractures which were unsuitable for simple nailing, can be satisfactorily stabilized by interlock nailing. Therefore, there is a trend towards nailing the fractures of the shaft of the femur rather than plating.

![Fig-19.4 X-rays showing plating done for fracture of distal third of the shaft of the femur](image)
DECIDING TREATMENT PLAN

The treatment depends primarily upon the age of the patient, location of the fracture, type of the fracture (transverse, oblique etc.) and presence of a wound. In general, an open fracture is treated conservatively; in bad cases an external fixator may be used.

In children, treatment is mostly by non-operative methods. The technique of traction varies in different age groups.

**Fig-19.5 Gallow's traction**

a) **From birth to 2 years:** These fractures are treated by Gallow’s traction (Fig-19.5). In this, the legs of the child are tied to a overhead beam. The hips are kept a little raised from the bed so that the weight of the body provides counter-traction and the fracture is reduced. This is continued till sufficient callus forms (3-6 weeks).

b) **From 2 years to 16 years:** The treatment at this age is essentially conservative. Different methods of traction are used to keep fragments in proper alignment. Once the fracture becomes ‘sticky’, further immobilisation can be provided in a hip spica. Older the child, more difficult it becomes to keep the fracture reduced for required period. It is therefore, sometimes preferred to internally fix the fracture in older children (more than 10 yrs. of age). TENS (Titanium Elastic Nail System) nails are used for this.

In adults and in the elderly, as far as possible, and if proper facilities are available, the treatment of these fractures is by operation. It allows the patient to be up and about, out of bed with the help of crutches very early.

COMPLICATIONS

The complications following a fracture of the femoral shaft can be divided into early and late.

**EARLY COMPLICATIONS**

1. **Shock:** In a closed fracture of the shaft of the femur, on an average, 1000-1500 ml of blood is lost. Such sudden loss of blood can result in hypovolaemic shock. Hence, all patients with this fracture should be on I.V. line, with blood arranged, in case the need arises. A close watch should be kept on pulse and blood pressure during the early post-injury period.

2. **Fat embolism:** Patient shows signs and symptoms of fat embolism after 24-48 hours of the fracture. Frequent shifting of the patient without proper splintage of the fracture should be avoided.

3. **Injury to femoral artery:** Rarely, a sharp edge of the bone may penetrate the soft tissues and damage the femoral artery. This occurs most commonly in fractures at the junction of middle and distal-third of the femoral shaft. Unless the continuity of the vessel is restored by immediate operation, the viability of the limb is in danger.

4. **Injury to sciatic nerve:** It may be damaged by a sharp bone end or by traction. The severity of damage varies from neurapraxia to complete severance of the nerve. Treatment is discussed in Chapter 10.

5. **Infection:** In cases with open fractures, wound contamination with consequent infection, can lead to osteomyelitis. The risk is maximum in fractures with extensive wounds, and those with gun shot wounds.

**LATE COMPLICATIONS**

1. **Delayed union:** Although, there is no definite time period beyond which the union of a fracture is said to be delayed, but if union is still insufficient to allow unprotected weight bearing after 5 months, it is considered delayed. X-ray will show evidence of union, but not solid enough to allow weight bearing.

   **Treatment:** It needs experience to decide whether continuation of conservative treatment would lead to fracture union, or an operative intervention is required. It is better to cut short the uncertainty by resorting to bone grafting, especially in an elderly person.
2. **Non-union:** It occurs when the fracture surfaces become rounded and sclerotic. A persistent mobility at the fracture site in a fracture fixed internally, not yet united, sometimes leads to fatigue fractures of the plate or nail (implant failure). Clinically, there may be frank mobility, pain on stressing or tenderness at the fracture site. *Treatment* is by internal fixation and bone grafting. A nail or a plate may be used for fixation.

3. **Malunion:** If a fracture of the shaft of the femur is not kept in proper position, or if it redisplaces, it may unite in an unacceptable position. The deformity is generally lateral angulation and external rotation. There may be significant shortening due to overlap of the fragments.

   *Treatment:* This depends upon the degree of malunion and age of the patient. In an elderly patient, if the disability is not much, tendency is toward accepting the deformity. Shortening may be compensated by giving a shoe raise. In younger patients, correction of the deformity is done by operative means. After redoing the fracture or by osteotomy, the deformities are corrected and the fracture fixed with internal fixation devices. Bone grafting is done in addition. In children, mild deformities get corrected by the process of remodelling. Significant deformities require corrective surgery.

4. **Knee stiffness:** Some amount of temporary knee stiffness occurs in most cases of fracture of the shaft of the femur. It is possible to regain full movements with physiotherapy. At times, the stiffness persists. The following could be the reasons: (i) intra-articular and peri-articular adhesions; (ii) quadriceps adhering to the fracture site; (iii) an associated, often undetected, knee injury.

   *Treatment:* Cases where a conscientious treatment by exercises has not been rewarding, a proper assessment of the contributing factor and its treatment is required. Intra-articular adhesions can be released by arthroscopic technique (arthrolysis), or by gentle manipulation under GA. Quadriceps adhesion may require release, and contracted quadriceps may need to be ‘lengthened’ (Quadricepsplasty).

### Further Reading

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**What have we learnt?**

- Fracture of the femur is a major, disabling injury.
- It is treated by conservative methods in children. Most fractures in adults and elderly are treated by operation.
- Nailing and plating are the two methods of internal fixation.
- Interlock nailing is the current choice of treatment. Non-union, delayed union and malunion are common complications.

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**Additional information: From the entrance exams point of view**

- Upper one-third shaft of femur most commonly fractured at birth.
- Maximum shortening of lower limb is seen in fracture shaft of femur and posterior dislocation of hip.
The knee joint is the most frequently injured joint. The following injuries will be discussed in this chapter: (i) condylar fractures of the femur; (ii) fracture of the plateau; (iii) tibial plateau fractures; (iv) injuries to the ligaments of the knee; (v) injuries to the menisci of the knee; (vi) miscellaneous knee injuries.

**RELEVANT ANATOMY**

The knee is a hinge joint formed between the tibia and femur (tibio-femoral). The patella glides over the front of femoral condyles to form a patello-femoral joint. The stability of the knee depends primarily upon its ligaments. The functions of different ligaments of the knee are given in Table-20.1.

**Table-20.1: Functions of the knee ligaments**

<table>
<thead>
<tr>
<th>Ligament</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medial collateral</td>
<td>Prevents medial opening up</td>
</tr>
<tr>
<td>Lateral collateral</td>
<td>Prevents lateral opening up</td>
</tr>
<tr>
<td>Anterior cruciate</td>
<td>Prevents anterior translation of the tibia on the femur</td>
</tr>
<tr>
<td>Posterior cruciate</td>
<td>Prevents posterior translation of the tibia on the femur</td>
</tr>
</tbody>
</table>

**Extensor apparatus of the knee:** It is constituted from proximal to distal, by quadriceps muscle, quadriceps tendon, patella with patellar retinaculae on the sides, and the patellar tendon (Fig-20.1). Failure of any of these results in inability to actively extend the knee, called extensor lag.

**MECHANISM OF KNEE INJURIES**

The knee joint is subjected to a variety of forces during day-to-day activities and sports. The nature of the forces may be direct or indirect. An indirect force on the knee may be: (i) valgus; (ii) varus; (iii) hyperextension; or (iv) twisting (Fig-20.2). Most often it is a combination of the above forces.

**CONDYLAR FRACTURES OF THE FEMUR**

Condylar fractures of the femur are of three types (Fig-20.3): (i) supracondylar fractures; (ii) intercondylar fractures – T or Y types; and (iii) unicondylar fractures – medial or lateral. These fractures commonly result from a direct trauma to the lower end of the femur. An indirect force more often results in unicondylar (by a varus/valgus bending force) or supracondylar fracture (by a hyperextension force).

**DIAGNOSIS**

Diagnosis of these fractures is suggested by pain, swelling and bruising around the knee. These
Essential Orthopaedics

Fractures are often missed, when associated with more severe injuries, such as a fracture of the shaft of the femur. Diagnosis is made on X-rays. A careful assessment of the intra-articular extension of the fracture and joint incongruity must be made.

**TREATMENT**

**Unicondylar fractures:** If undisplaced, a long leg cast is given for 3-6 weeks, followed by protected weight bearing. If displaced, open reduction and internal fixation with multiple cancellous screws is performed. A buttress plate may be required in some cases.

**Intercondylar fractures:** The aim of treatment is to restore congruity of the articular surface as far as possible. In displaced T or Y fracture with minimal comminution, the joint is reconstructed by open reduction and internal fixation. Condylar blade-plate, DCS and LCP are popular implants. Comminuted fractures are difficult to accurately reconstruct, but well done open reduction and internal fixation permits early knee mobilisation and thus better functions. In selected comminuted fractures, conservative treatment in skeletal traction may be the best option, and give acceptable results.

**Supracondylar fractures:** It is best to treat displaced supracondylar fractures with internal fixation. This could be done by closed or open techniques. Nail or plate may be used.

**COMPLICATIONS**

1. **Knee stiffness:** Residual knee stiffness sometimes remains because of dense intra- and peri-articular adhesions. A long course of physiotherapy is usually rewarding. Arthrolysis may be required in resistant cases.

2. **Osteoarthritis:** Fractures with intra-articular extension give rise to osteoarthritis a few years later.
3. **Malunion**: A malunion may result in varus or valgus deformities, sometimes requiring a corrective osteotomy.

**FRACTURES OF THE PATELLA**

This is a common fracture. It may result from a direct or an indirect force. In a direct injury, as may occur by a blow on the anterior aspect of the flexed knee, usually a comminuted fracture results. The comminution may be limited to a part or whole of the patella. The latter is also called a *stellate fracture* (Fig-20.4a). Sometimes, a sudden violent contraction of the quadriceps, gives rise to a fracture with the fracture line running transversely across the patella, dividing it into two; the so-called *two-part fracture*. Most often, both of these mechanisms are at play simultaneously, so that once the fracture occurs by a direct violence, a simultaneous contraction of the quadriceps pulls the fragments apart, and results in a separated fracture of the patella with some comminution.

**PATHOANATOMY**

The fracture may remain undisplaced because the fragments are held in position by intact pre-patellar expansion of the quadriceps tendon in front, and by patellar retinaculæ on the sides. If the force of the quadriceps contraction is strong, it will pull the fragments apart and will result in rupture of patellar retinaculæ (Fig-20.4b).

**CLINICAL FEATURES**

**Presenting complaints**: The patient complains of pain and swelling over the knee. In an undisplaced fracture the swelling and tenderness may be localised over the patella. A crepitus is felt in a comminuted fracture. In displaced fractures, one may feel a gap between the fracture fragments. The patient will not be able to lift his leg with the knee in full extension; it remains in a position short of full extension (*extensor lag*) because of disruption of the extensor apparatus. There may be bruises over the front of the knee – a tell tale sign of direct trauma. The knee may be swollen because of haemarthrosis.

**Radiological examination**: Antero-posterior and lateral X-rays of the knee are sufficient in most cases. In some undisplaced fractures, a ‘skyline view’ of the patella (Fig-20.5) may be required. A fracture with wide separation of the fragments is easy to diagnose on a lateral X-ray. Often it is not possible to visualise comminution on the X-ray; it becomes obvious only during surgery (Fig-20.6).

**TREATMENT**

It depends upon the type of fracture, and in some cases on the age of the patient. The following groups may be considered:

a) **Undisplaced fracture**: Treatment is aimed primarily at relief of pain. A plaster cast extending from the groin to just above the malleoli, with the knee in full extension (cylinder cast) should be given for 3 weeks, followed by physiotherapy.
b) **Clean break with separation of fragments (two-part fracture):** The pull of the quadriceps muscle on the proximal fragment keeps the fragments apart, hence an operation is always necessary. The operation consists of reduction of the fragments, fixing them with *tension-band wiring (TBW)* and repair of extensor retinaculae. The knee can be mobilised early following this operation. In cases where it is not possible to achieve accurate reduction of the fragments, it is better to excise the fragments (patellectomy) and repair the extensor retinaculae. In cases where one of the fragments constitutes only one of the poles of the patella, it is excised. The major fragment is preserved and the extensor retinaculae repaired (partial patellectomy). Such operations on the patella are followed by support in a cylinder cast for 4-6 weeks.

c) **Comminuted fracture:** In comminuted fractures with displacement, it is difficult to restore a perfectly smooth articular surface, so excision of the patella (patellectomy) is the preferred option. This takes care of any future risk of osteoarthritis at the patello-femoral joint. With improvement in fixation techniques, more and more comminuted fractures of the patella are being reconstructed (patella saving operations).

**Complications**

1. **Knee stiffness:** It is a common complication after a fracture of the patella, mostly due to intra- and peri-articular adhesions. Treatment is by physiotherapy. Sometimes, an arthroscopic release of adhesions may be required.

2. **Extensor weakness:** This results from an inadequate repair of the extensor apparatus or due to quadriceps weakness.

3. **Osteoarthritis:** Patello-femoral osteoarthritis occurs a few years after the injury.

**INJURIES TO THE LIGAMENTS OF THE KNEE**

With increasing sporting activities, injuries to the knee ligaments are on the rise. The type of injury depends upon the direction of force and its severity.

**Mechanism**

Knee ligaments are injured most often from indirect, twisting or bending forces on the knee. The various mechanisms by which knee ligaments are injured are given below:

- **Medial collateral ligament:** This ligament is damaged if the injuring force has the effect of abducting the leg on the femur (*valgus force*). It ruptures most commonly from its femoral attachment.

- **Lateral collateral ligament:** This ligament is damaged by a mechanism just the reverse of above i.e., adduction of the tibia on the femur (*varus force*). Commonly, the ligament is avulsed from head of the fibula with a piece of bone. Lateral collateral ligament injuries are uncommon because the knee is not often subjected to varus force (the knee is not likely to be hit from the inside).

- **Anterior cruciate ligament:** This ligament is *most commonly* ruptured, often in association with the
tears of medial or lateral collateral ligaments. Commonly, it occurs as a result of twisting force on a semi-flexed knee. Often the injury to medial collateral ligament, medial meniscus and anterior cruciate ligament occur together. This is called O’Donoghue triad.

- **Posterior cruciate ligament**: This ligament is damaged if the anterior aspect of the tibia is struck with the knee semi-flexed so as to force the tibia backwards on to the femur.

### PATHOANATOMY

The ligament may tear at either of its attachment. Sometimes, it takes a chip of bone from its attachment. The ligament may be torn in its substance (mid-substance tear). The severity of the tear varies from a rupture of just a few fibres to a complete tear (see classification of ligament injury on page 5).

It may be an 'isolated' ligament injury, or more than one ligaments may be injured. The combination depends upon the direction and severity of the force. Rarely, in a very severe injury, the knee may get dislocated and a number of ligaments injured.

### DIAGNOSIS

**Clinical examination**: Pain and swelling of the knee are the usual complaints. Often, the patient is able to give a history of having sustained a particular type of deforming force at the knee (valgus, varus etc.), followed by a sound of something tearing. The pain may be localised over the torn ligament (in cases of injury to collateral ligaments), but there is vague pain in cruciate ligament injuries. The swelling (haemarthrosis) is variable, but appears early after the injury.

Damage to the medial and lateral collateral ligaments can be assessed clinically by stress tests* (see page 6).

Pain at the site of the torn ligament and/or an abnormal opening up of the joint indicate a tear. Cruciate ligaments prevent anterior–posterior gliding of the tibia. The anterior cruciate prevents anterior glide, and the posterior cruciate prevents posterior glide. This property is made use in detecting injury to these ligaments.

**Anterior drawer test**: This is a test to detect injury to the anterior cruciate ligament. A similar test in which anterior glide of the tibia is judged with the knee in 10-15 degrees of flexion is called Lachmann test.

**Posterior drawer test**: This is a test to detect injury to the posterior cruciate ligament. A posterior sagging of the upper tibia may be obvious, and indicates a posterior cruciate tear.

Essential features of common knee ligament injuries are given in Table 20.2.

**Radiological examination**: A plain X-ray may be normal, or a chip of bone avulsed from the ligament attachment may be visible. It may be possible to demonstrate an abnormal opening-up of the joint on stress X-rays. MRI is a non-invasive method of diagnosing ligament injuries, and may be of use in doubtful cases.

**Other investigation**: Arthroscopic examination may be needed in cases where doubt persists.

A patient with **acute knee haemarthrosis** may have sustained: (i) an intra-articular fracture of femur, tibia or patella; (ii) ligament injury; (iii) meniscus tear and (iv) patellar subluxation or dislocation. An X-ray is usually done, where a fracture can be easily diagnosed. Of all cases with ‘no bony injury’, an accurate clinical examination may lead to diagnosis of a ligament injury or patellar subluxation. In doubtful cases, an MRI is asked for. In some cases, an examination under anaesthesia and arthroscopy may be required to come to a diagnosis. In young females presenting with

### Table 20.2: Essential features of knee ligament injury

<table>
<thead>
<tr>
<th>Name of the ligament</th>
<th>Mechanism of injury</th>
<th>Pain</th>
<th>Swelling</th>
<th>Tenderness</th>
<th>Tests</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medial collateral</td>
<td>Valgus force</td>
<td>Medial</td>
<td>Medial side</td>
<td>Medially on femoral condyle</td>
<td>Valgus stress + at 30° knee flexion</td>
</tr>
<tr>
<td>Lateral collateral</td>
<td>Varus force</td>
<td>Lateral</td>
<td>Lateral side</td>
<td>Laterally on fibular head</td>
<td>Varus stress + at 30° knee flexion</td>
</tr>
<tr>
<td>Anterior cruciate</td>
<td>Twisting extension</td>
<td>Diffuse</td>
<td>Haemarthrosis</td>
<td>Vague</td>
<td>Anterior drawer test + Lachmann test +</td>
</tr>
<tr>
<td>Posterior cruciate</td>
<td>Backward force on tibia</td>
<td>Diffuse</td>
<td>Haemarthrosis</td>
<td>Vague</td>
<td>Posterior drawer test +</td>
</tr>
</tbody>
</table>
acute knee haemarthrosis, a self-reduced patellar subluxation must be thought of.

TREATMENT
Treatment of ligament injuries is a controversial subject. Conventionally, these injuries have been treated by non-operative methods. With availability of newer techniques, better results have been achieved by operative reconstruction. Therefore, operative treatment has become more popular in high demand athletic individuals, particularly for anterior cruciate ligament tear.

Conservative method: The haematoma is aspirated and the knee is immobilised in a cylinder cast or commercially available knee immobiliser. Most cases of grade I and II injuries can be successfully treated by this method. After a few weeks, the swelling subsides, and adequate strength can be regained by physiotherapy.

Operative methods: These are indicated in multiple ligament injured knee, especially in young athletes. The operation is usually performed 2-3 weeks after injury after the acute phase subsides. It consists of the following:

a) Repair of the ligament: It is performed for fresh, grade III collateral ligament injuries. In cases presenting after 2-3 weeks, an additional reinforcement is provided by a fascial or tendon graft.

b) Reconstruction: This is done in cases of ligament injuries presenting late with features of knee instability. A ligament is ‘constructed’ using patient’s tendon or fascia lata. A tendon or fascia taken from another person (allograft) or a synthetic ligament has also been used.

Knee ligaments are torn more often than they are diagnosed. Unfortunately, since this injury is not detected on X-rays, it gets neglected. Patients usually present late with symptoms of knee giving-way (instability). The treatment at this stage depends upon activity level of the patient. For a patient with sedentary lifestyle, adequate stability is achieved with physiotherapy alone. In active patients, ligament reconstruction is necessary. The ACL is the commonest to be ruptured. The treatment of choice is arthroscopic ACL reconstruction. In this, the torn ligament is replaced with a tendon graft. This is done endoscopically (arthroscopic surgery), without opening the joint. The joint is first examined by a 4 mm telescope (arthroscope). A tendon graft taken from patellar tendon or hamstring tendons is introduced into the knee through bone tunnels. The graft is fixed at both ends with screws or other devices. Bio-absorbable screws are now being used. Arthroscopic surgery has advantages of being minimally invasive, and results in quick return to function with minimal risks.

COMPLICATIONS
1. Knee instability: An unhealed ligament leads to instability. The patient ‘loses confidence’ on his knee, and the knee often “gives-way”. Surgery is usually required.

2. Osteoarthritis: A neglected ligament injury may result in further damage to the knee in the form of meniscus tear, chondral damage etc. This eventually leads to knee osteoarthritis.

TIBIAL PLATEAU FRACTURES
These are common fractures sustained in two wheeler accidents when one lands on the knee. Either or both condyles of tibia are fractured. The mechanism of injury is:

(a) an indirect force causing varus or valgus force on the knee or (b) a direct hit on the knee.

Types of fracture: These fractures commonly occur in six patterns (Schatzker types). Type I-IV involve only one condyle, lateral or medial. Type V and VI are more complex intercondylar fractures.

Symptoms and signs: The patient complains of pain and swelling, and inability to bear weight. Often crepitus is heard or felt. Diagnosis can be made on X-rays. CT scan may be required for accurate evaluation.

Treatment: Like most fractures, both conservative and operative methods can be used. Conservative methods are used for minimally displaced fractures, and those in elderly people. There is more and more trend towards accurate reduction and early mobilisation of these fractures. Surgical treatment of these fractures is a technically demanding procedure, and needs variety of equipment and lot of experience.

MENISCAL INJURIES OF THE KNEE
These constitute a common group of injuries peculiar to the knee, frequently being reported with increasing sporting activity.
MECHANISM
The injury is sustained when a person, standing on a semi-flexed knee, twists his body to one side. The twisting movement, an important component of the mechanism of injury, is possible only with a flexed knee. During this movement the meniscus is 'sucked in' and nipped as rotation occurs between the condyles of femur and tibia. This results in a longitudinal tear of the meniscus. The meniscus may be torn with a minor twisting, as may occur while walking on uneven surface. A degenerated meniscus in the elderly may get torn by minimal or no injury. The medial meniscus gets torn more often because it is less mobile (being fixed to the medial collateral ligament).

PATHOANATOMY
The meniscus is torn most commonly at its posterior horn. With every subsequent injury, the tear extends anteriorly. The meniscus, being an avascular structure, once torn does not heal*. If left untreated, it undergoes many more subtears, and damages the articular cartilage, thus initiating the process of osteoarthritis.

Types of meniscal tear: The bucket-handle tears (Fig-20.7) are the commonest type; others are radial, anterior horn, posterior horn and complex tears. Some underlying pathological changes in the meniscus make it prone to tear. These are discoid meniscus (the meniscus, unlike the normal semilunar shape, is shaped like a disc), degenerated meniscus (in osteoarthritis), and a meniscal cyst.

CLINICAL FEATURES
Presenting complaints: The patient is generally a young male actively engaged in sports like football, volleyball etc. The presenting complaint is recurrent episodes of pain, and locking** of the knee. At times, the patient complains of a 'jhatka', a sudden jerk while walking, or 'something flicking over' inside the joint. This may be followed by a swelling, appearing after a few hours and lasting for a few days. After some time, the pain becomes persistent but with little or no swelling.

On tracing back the symptoms to their origin, one often finds a history of a classic twisting injury to the knee, followed by a swelling appearing overnight*** as effusion collects. After the effusion subsides, the knee may remain in about 10 degrees of flexion, beyond which the patient is unable to extend his knee (locking). This is because the torn portion of the meniscus gets interposed between the femoral and the tibial condyles. Locking may be missed because the attention is drawn to more obvious signs of pain and swelling. The displaced fragment sometimes returns to its original position spontaneously and thus the original episode of locking may never be noticed. Every successive episode of locking may be either spontaneously corrected or may need manipulation by the patient or a physician. The history of sudden locking and unlocking, with a click located in one or other joint compartment, is diagnostic of a meniscus tear.

On examination: In a typical episode presenting after injury, the knee may be swollen. There may be tenderness in the region of the joint line, either anteriorly or posteriorly. The knee may be locked. Gentle attempts to force full extension produces a sensation of elastic resistance and pain, localised to the appropriate joint compartment. In between the episodes, the knee may not have any finding except wasting of the quadriceps. The manoeuvres carried out to detect a hidden meniscus tear are McMurray’s and Apley’s test (see Annexure-III).

* The peripheral meniscal tears, being in a vascular area, often heal.

** True locking is an inability to extend the knee for the last few degrees. It is different from ‘pseudo-locking’ where the knee catches temporarily in one position due to sudden pain. The latter may occur in cases with a loose body. It may also occur due to hamstring muscles spasm, thus not allowing the knee to get extended.

*** The swelling in a case of meniscus tear is due to synovial reaction, hence appears after a few hours. This is unlike the swelling in other knee injuries, where haemarthrosis results in early swelling.
Often it is difficult to diagnose the cause of knee symptoms on history and clinical examination. Such non-specific symptom-complex is termed as internal derangement of the knee (IDK).

**RADIOLOGICAL EXAMINATION**
With meniscal tears there are no abnormal X-ray findings. X-rays are taken to rule out any associated bony pathology. MRI is a non-invasive method of detecting meniscus tears. It is a very sensitive investigation, and sometimes picks up tears which are of no clinical significance.

**Arthrography:** It is a technique where X-rays are taken after injecting radiopaque dye into the knee. The dye outlines the menisci, so that a tear, if present, can be visualised. Being an invasive technique, it is no longer used.

**ARTHROSCOPY**
This is a technique where a thin endoscope, about 4-5 mm in diameter – the arthroscope, is introduced into the joint through a small stab wound, and inside of the joint examined (Fig-20.8). For details see page 334.

**TREATMENT**

**Treatment of acute meniscal tear:** If the knee is locked, it is manipulated under general anaesthesia. No special manoeuvre is needed. As the knee relaxes, the torn meniscus falls into place and the knee is unlocked. The knee is immobilised for 2-3 weeks, followed by physiotherapy. In a case where locking is not present, immobilisation in a knee immobiliser is sufficient. With this, a small number of peripheral tears will heal. Rest of the tears may produce recurrent symptoms.

**Treatment of a chronic meniscal tear:** Once the diagnosis is established clinically, the treatment is to excise the displaced fragment of the meniscus. Now-a-days, it is possible to excise a torn meniscus arthroscopically (arthroscopic surgery). By this technique, once the fault is detected (e.g., a loose meniscal flap), the same is corrected using fine cutting instruments introduced from another puncture wound. This technique is a significant advancement as it can be done as a day care procedure. Since it is a minimally invasive technique, early return to work is possible. Recent research has shown that menisci are not ‘useless’ structures as was thought earlier. Hence, wherever possible the trend is to preserve the meniscus by suturing. The state-of-the-art is arthroscopic meniscus suturing.

**RARE INJURIES AROUND THE KNEE**

**Dislocation of the knee:** This rare injury results from severe violence to the knee so that all of its supporting ligaments are torn. It is a major damage to the joint, and is often associated with injury to the popliteal artery. Treatment is by reduction followed by immobilisation in a cylinder cast. Recent studies have shown superior results by operative treatment of these severe knee injuries, by multiple ligament reconstruction.

**Disruption of extensor apparatus:** Injury from sudden quadriceps contraction most often results in fracture of the patella. Sometimes, it may result in tearing of the quadriceps tendon from its attachment on the patella, or tearing of the attachment of the patellar tendon from the tibial tubercle. In either case, operative repair of the tendon is required.

**Dislocation of the patella:** The patella usually dislocates laterally. It can be one of three types: (i) acute dislocation; (ii) recurrent dislocation; and (iii) habitual dislocation.

**Acute dislocation of the patella** results from a sudden contraction of the quadriceps while the knee is flexed or semi-flexed. The patella dislocates laterally and lies on the outer side of the knee. The patient is unable to straighten the knee. The
medial condyle of femur appears more prominent. Sometimes, the dislocation reduces spontaneously but one can elicit marked tenderness antero-medially as a result of the rupture of the capsule at that site. Treatment consists of reduction and immobilisation in a cylinder cast or knee immobiliser for 3 weeks. A piece of bone covered with articular cartilage (osteochondral fragment), may be shaved off from the patella or the femoral condyle at the time of dislocation (Fig-20.9). This results in repeated episodes of pain, swelling and sensation of a loose body. Arthroscopic removal may be required.

Recurrent dislocation of the patella: After the first episode of dislocation, generally during adolescence, the dislocation tends to recur with more and more ease. The reason for recurrence may be laxity of the medial capsule or some underlying defect in the anatomy of the knee. These could be: (i) excessive joint laxity; (ii) a small patella; (iii) a patella alta (i.e., the patella is high-lying in the shallower part of intercondylar groove); and (iv) genu valgum. Treatment consists of operative reconstruction where the insertion of the patellar tendon on the tibial tuberosity is shifted medially and downwards so that the line of pull of the quadriceps shifts medially (Hauser’s operation). Prof. Baksi from Calcutta has described a useful operation for this condition whereby pes anserinus is transferred to lower pole of the patella to provide a ‘checkrein’ effect.

The current trend is to precisely find the cause of the recurrence, which may be bony (genu valgus, increased Q-angle etc.), or soft tissue (ruptured medial patello-femoral ligament). The corrective surgery is aimed at correcting the underlying cause. The operations are done arthroscopic-assisted.

Habitual dislocation of the patella: It means that the patella dislocates laterally everytime the knee is flexed. The patient presents early in childhood. Underlying defects are very similar to those in recurrent dislocation. In addition, a shortened quadriceps (vastus lateralis component) may result in an abnormal lateral pull on the patella when the knee is flexed. Treatment is by release of the tight structures on the lateral side and repair of the lax structures on the medial side. An additional ‘checkrein’ mechanism of some sort is created to prevent re-dislocation.

What have we learnt?
- Knee injuries are commonly sustained in scooter accident and sports.
- Fractures around the knee are difficult injuries as they commonly lead to knee stiffness. Hence, open reduction and internal fixation is the more popular method of treatment.
- Internal derangement of the knee (IDK) is a term used to group all the other, non-bony injuries of the knee. These consist of ligament injuries, meniscus injury and patello-femoral problems.
- MRI is an important investigation for diagnosis of ligament and meniscus injuries.
- Arthroscopic surgery has become a standard method of treating meniscus and ligament injuries of the knee.
Additional information: From the entrance exams point of view

- In 90° flexion of the knee, the tibial tuberosity is in line with the centre of the patella, on extension, it moves towards the lateral border due to the screw home mechanism.
- People with anterior cruciate deficient knees have a problem climbing downhill.
- Dial test, tests posterolateral corner and the posterior cruciate ligament.
  - Posterolateral corner deficiency positive at 30° flexion.
  - Posterior cruciate ligament positive at both 30° and 90° flexion.
- Physiological locking occurs with internal rotation of the femur over a fixed tibia by the quadriceps, unlocking refers to the lateral rotation of the femur over a stabilized tibia by the popliteus.
- Rotation force is most important in causing a meniscal injury.
FRACTURES OF SHAFTS OF TIBIA AND FIBULA

RELEVANT ANATOMY

The tibia is the major weight bearing bone of the leg. It is connected to the less important bone, the fibula, through the proximal and distal tibio-fibular joints. Like fractures of forearm bones, these bones frequently fracture together, and are referred to as ‘fracture both bones of leg’. The following are some of the characteristics of these bones.

a) A subcutaneous bone: This is responsible for the large number of open tibial fractures; also, often there is loss of bone through the wound.

b) Fractures in this region are often associated with massive loss of skin, necessitating care by plastic surgeons, early in the treatment.

c) Precarious blood supply: The distal-third of tibia is particularly prone to delayed and non-union because of its precarious blood supply. The major source of blood supply to the bone is the medullary vessels. The periosteal blood supply is poor because of few muscular attachments on the distal-third of the bone. The fibula, on the other hand is a bone with many muscular attachments, and thus has a rich blood supply.

d) Hinge joints proximally and distally: Both, the proximal and distal joints (the knee and ankle) are hinge joints. So, even a small degree of rotational mal-alignment of the leg fracture becomes noticeable. This is unlike a fracture of the femur or humerus, where some degree of rotational mal-alignment goes unnoticed because of the polyaxial ball and socket joint proximally.

MECHANISM

The tibia and fibula may be fractured by a direct or indirect injury.

Direct injury: Road traffic accidents are the commonest cause of these fractures, mostly due to direct violence. The fracture occurs at about the same level in both bones. Frequently the object causing the fracture lacerates the skin over it, resulting in an open fracture.

Indirect injury: A bending or torsional force on the tibia may result in an oblique or spiral fracture respectively. The sharp edge of the fracture fragment may pierce the skin from within, resulting in an open fracture.

PATHOANATOMY

The fracture may be closed or open, and may have various patterns. It may occur at different levels (upper, middle or lower-third). Occasionally, it may be a single bone fracture i.e., only the tibia or fibula is fractured. Displacements may be sideways, angulatory or rotational. Occasionally, the fracture may remain undisplaced.

CLINICAL FEATURES

The patient is brought to the hospital with a history of injury to the leg followed by the classic features of a fracture i.e., pain, swelling, deformity etc.
There may be a wound communicating with the underlying bone.

**RADIOLOGICAL FEATURES**

The diagnosis is usually confirmed by X-ray examination. Evaluation of the anatomical configuration of the fracture on X-ray helps in reduction.

**TREATMENT**

For the purpose of treatment, fractures of the tibia and fibula may be divided into two types: closed or open.

**Closed fractures:** Treatment of closed fractures, both in children and in adults, is by closed reduction under anaesthesia followed by an above-knee plaster cast. In children, it is possible to achieve good alignment in most cases, and the fracture unites in about 6 weeks. In adults, the fracture unites in 16-20 weeks. Sometimes, reduction is not achieved, or the fracture displaces in the plaster. In both these cases open reduction and internal fixation is required.

The trend is changing with the availability of minimally invasive techniques such as closed nailing. More and more unstable tibial fractures are being treated with closed interlock nailing.

**Open fractures:** The aim in the treatment of open fractures is to convert it into a closed fracture by judicious care of the wound, and maintain the fracture in good alignment. Following methods can be used for treating the fracture, depending upon the grade of open fracture:

- **Grade I:** Wound dressing through a window in an above-knee plaster cast, and antibiotics.
- **Grade II:** Wound debridement and primary closure (if less than 6 hours old), and above-knee plaster cast. The wound may need dressings through a window in the plaster cast.
- **Grade III:** Wound debridement, dressing and external fixator application. The wound is left open.

The trend is changing, from primarily conservative treatment to operative treatment, in care of open tibial fractures. More and more open fractures in grade I and II are being fixed internally. In a number of other cases, a delayed operation (ORIF) is done once the wound is taken care of.

**Technique of closed reduction:** Under anaesthesia, the patient lies supine with his knees flexed over the end of the table. The surgeon is seated on a stool, facing the injured leg. The leg is kept in traction using a halter, made of ordinary bandage, around the ankle (Fig-21.1). The fracture ends are manipulated and good alignment achieved. Initially, a below-knee cast is applied over evenly applied cotton padding. Once this part of the plaster sets, the cast is extended to above the knee.

**Wedging:** Sometimes, after a fracture has been reduced and the plaster applied, check X-ray shows a little angulation at the fracture site. Instead of cutting open the plaster and reapplying it, it is better to wedge the plaster as shown in Fig-21.2. In this technique, the plaster is cut circumferentially at the level of the fracture, the angulation corrected by forcing open the cut on the concave side of the angulation, and the plaster reinforced with additional plaster bandages.
Once the fracture becomes 'sticky' (in about 6 weeks), above-knee plaster is removed and below-knee PTB (patellar tendon bearing) cast is put. Use of modern, synthetic casting tapes (made of plastic polymer) has made ‘plaster’ treatment more convenient. Once the fracture has partly united, the cast can be replaced by removable plastic supports (braces), and the joints mobilised.

**Role of operative treatment:** Open reduction and internal fixation is necessary when it is not possible to achieve a satisfactory alignment of a fracture by non-operative methods. The internal fixation device used may be a plate or an intra-medullary nail depending upon the configuration of the fracture. Interlock nailing provides the possibility of internally fixing a wide spectrum of tibial shaft fractures. With the availability of facilities, operative treatment has now become a method of preference.

**Deciding the plan of treatment:** It depends on whether the fracture is closed or open. A practical plan of treatment is as shown in Flow chart-21.1.

**Flow chart-21.1  Treatment plan of tibial shaft fractures**

https://kat.cr/user/Blink99/
COMPLICATIONS

1. Delayed union and non-union: Fractures of the tibia sometimes take unusually long to unite; more so the ones in the lower-third. In some cases, clear signs of non-union become apparent on X-rays. The most important factor responsible for delayed and non-union is the precarious blood supply of the tibia; others being frequent compounding with loss of fracture haematoma, wound infection, etc. Failure of union results in pain and inability to bear weight on the leg.

Treatment: Treatment of delayed union and non-union is essentially by bone grafting, with or without internal fixation. Following treatment options are available:

a) Nailing with Bone Grafting: This is indicated in cases of non-union, where the alignment is not acceptable, or there is free mobility at the fracture site. Some surgeons prefer plating and bone grafting.

b) Phemister Grafting: This is a type of bone grafting done for selected cases which fulfill the following criteria:
   - There is minimal or no mobility at the fracture site (fibrous union).
   - The fracture has an acceptable alignment.
   - The knee joint has a good range of movement.

In this technique, grafting is performed without disturbing the sound fibrous union at the fracture site. The aim is to stimulate bone formation in the ‘fibro-cartilaginous tissue’ already bridging the fracture. Cancellous bone grafts are placed after raising the osteo-periosteal flaps around the fracture (Fig-21.3). In addition, petalling (lifting slivers of cortical bone attached at base) is carried out around the fracture. This results in bony union in about 3-4 months.

c) Ilizarov’s Method: This method is useful in treatment of difficult non-unions of tibia. These are non-unions with bone gap, infection, or those with bad overlying skin. (details on page 33)

2. Malunion: Some amount of angulation is acceptable in children as it gets corrected by remodelling. In adults, displacements especially angulations and rotations are not acceptable. These cause problems in walking and result in early osteoarthritis of the knee and ankle. Treatment requires correction of the deformity by redoing the fracture and fixing it by plating or nailing, and bone grafting.

3. Infection: Because of the frequency with which tibial fractures are associated with a communicating skin wound, contamination and subsequent infection is a common complication. Most often the infection is superficial and is controlled by dressing and antibiotics. Sometimes, the underlying bone gets infected, in which case more elaborate treatment on the lines of osteomyelitis may be necessary (see page 171). The fracture in such cases often does not unite. Ilizarov’s method is the treatment of choice in such infected non-unions.

4. Compartment syndrome (see page 47): Some cases of closed fracture of the tibia may be associated with significant crushing of soft tissues, leading to compartment syndrome. A compartment syndrome should be suspected if a fracture of the tibia is associated with excessive pain, swelling and inability to move the toes. Immediate operative decompression of the compartments is imperative.

5. Injury to major vessels and nerves: Occasionally a fracture of the tibia, especially in the upper-third of the shaft may be associated with injury to the popliteal artery or the common peroneal and tibial nerves. Therefore, examination of the neurovascular status of the limb in a fresh case is of vital importance to prevent serious complications.
like vascular gangrene etc. Treatment of these complications is as discussed in Chapter 7.

**ANKLE INJURIES**

The bones forming the ankle joint are a frequent site of injury. A large variety of bending and twisting forces result in a number of fractures and fracture-dislocation at this joint. All these injuries are sometimes grouped under a general title ‘Pott’s fracture’.

**RELEVANT ANATOMY**

The ankle joint is a modified hinge joint. The ‘socket’ is formed by the distal articular surfaces of the tibia and fibula, the intervening tibio-fibular ligament and the articular surfaces of the malleoli. These together constitute the **ankle-mortise** (Fig-21.4). The superior articular surface of the talus (the dome) articulates with this socket.

The strong tibio-fibular syndesmosis, along with the medial and lateral malleoli make the ankle a strong and stable articulation. Therefore, pure dislocation of the ankle is rare. Commonly, dislocation occurs only with fractures of the malleoli. The elongated posterior part of the distal articular surface of the tibia, often termed as **posterior malleous** gets chipped-off in some ankle injuries.

**Ligaments of the ankle:** The ankle joint has two main ligaments; the medial and lateral collateral ligaments (Fig-21.5).

*Medial collateral ligament* (deltoid ligament): This is a strong ligament on the medial side. It has a superficial (tibio-calcaneal) and a deep (tibio-talar) part.

*Lateral collateral ligament:* This is a weak ligament and is often injured. It has three parts: (i) anterior talo-fibular; (ii) calcaneo-fibular in the middle; and (iii) posterior talo-fibular.

**Some terms used in relation to ankle injuries** (Fig-21.6): Following are some of the terms used to
describe different forces the ankle may be subjected to:

a) **Inversion** (adduction): Inward twisting of the ankle.
b) **Eversion** (abduction): Outward twisting of ankle.
c) **Supination**: Inversion plus adduction of the foot so that the sole faces medially and in plantar-flexion.
d) **Pronation**: Eversion and abduction of the foot so that the sole faces laterally and in dorsiflexion.
e) **Rotation** (external or internal): A rotatory movement of the foot so that the talus is subjected to a rotatory force along its vertical axis.
f) **Vertical compression**: A force along the long axis of the tibia.

**CLASSIFICATION**

The Lauge-Hansen classification (Table–21.1) of ankle injuries is most widely used. It is based on the mechanism of injury. It is believed that a specific pattern of bending and twisting forces results in specific fracture pattern. Different types of ankle injuries have been classified on the basis of five basic mechanisms. These are as follows:

a) Adduction injuries.
b) Abduction injuries.
c) Pronation-external rotation injuries.
d) Supination-external rotation injuries.
e) Vertical compression injuries.

When a foot is subjected to these forces, different parts of the ankle-mortise are subjected to distraction and compression stress. The specific fracture pattern depends on the type of stress and its severity, as discussed below:

**Adduction injuries** (inversion): An inversion force with the foot in plantar-flexion results in a sprain of the lateral ligament of the ankle. It may be either a partial or complete rupture. A partial rupture is limited to the anterior fasciculus of the lateral ligament (talo-fibular component). In a complete rupture, the tear extends backwards to involve the whole of the lateral ligament complex. As complete rupture occurs, the talus tends to subluxate out of the ankle-mortise.

The inversion force on an ankle in neutral or dorsiflexed position results in a fracture of the medial malleolus, typically, a fracture with the fracture line running obliquely upwards from the medial angle of the ankle-mortise (Fig-21.7). On the lateral side, this may be associated with a low transverse (below the ankle-mortise) avulsion

**Table–21.1: Lauge-Hansen classification of ankle injuries**

<table>
<thead>
<tr>
<th>Type of injury</th>
<th>On medial side</th>
<th>Tibio-fibular syndesmosis</th>
<th>On lateral side</th>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Adduction injury</td>
<td>Med. malleolus fracture with an oblique fracture line</td>
<td>Normal</td>
<td>Avulsion fractures of lat. malleolus or Lat. coll. lig. injury</td>
<td>—</td>
</tr>
<tr>
<td>• Abduction injury</td>
<td>Avulsion fracture of med. malleolus (low) or Med. coll. lig. injury</td>
<td>Normal</td>
<td>Fracture of lateral malleolus at the level of ankle mortise with comminution of its lateral cortex</td>
<td>—</td>
</tr>
<tr>
<td>• Pronation-external rotation injury</td>
<td>Transverse fracture of med. malleolus at the level of ankle-mortise</td>
<td>Damaged</td>
<td>Spiral fracture of the fibula above the level of ankle-mortise or no fracture</td>
<td>Fracture of the posterior malleolus</td>
</tr>
<tr>
<td>• Supination-external rotation injury</td>
<td>Transverse fracture of med. malleolus at the level of ankle-mortise</td>
<td>Normal</td>
<td>Spiral fracture of the lat. malleolus at the level of ankle-mortise</td>
<td></td>
</tr>
<tr>
<td>• Vertical compression</td>
<td>Comminuted fractures of med. malleolus, distal end of the tibia and lateral malleolus.</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
fracture of the lateral malleolus, or a lateral ligament rupture.

**Abduction injuries** (eversion): In this type, the medial structures are subjected to a distracting force and the lateral structures to compressive force. This results in rupture of the deltoid ligament or a low-lying transverse fracture of the medial malleolus (avulsion fracture) on the medial side. On the lateral side, a fracture of the lateral malleolus at the level of the ankle-mortise with comminution of the outer cortex occurs (Fig-21.8). The talus, with both malleoli fractured, subluxates laterally.

![Fig-21.8 Abduction injury](image)

**Pronation-external rotation injuries:** When a pronated foot rotates externally, the talus also rotates outwards along its vertical axis. The first structures to give way are those on the medial side. There may occur a transverse fracture of the medial malleolus at the level of the ankle-mortise, or a rupture of the medial collateral ligament. With further rotation of the talus, the anterior tibiofibular ligament is torn. This is followed by a spiral fracture of the lower end of the fibula as the rotating talus hits the lateral malleolus. In a case, where the tibio-fibular syndesmosis is completely disrupted, the fracture occurs above the syndesmosis i.e., in the lower-third of the fibula (Fig-21.9).

![Fig-21.9 Pronation-external rotation injury](image)

The fracture may occur as high as the neck of the fibula (Massonaie’s fracture). Thus, a fracture of the fibula above the ankle-mortise, in an ankle injury, is an indication of disruption of the tibio-fibular syndesmosis.

**Supination-external rotation injuries:** With the foot supinated, the talus twists externally within the mortise. As the medial structures are lax, the first structure to give way is that on the lateral side, the head of the talus striking against the lateral malleolus, producing a spiral fracture at the level of the ankle-mortise. The next structure to break is the posterior malleolus. As the talus rotates further, it hits against the medial malleolus resulting in a transverse fracture (Fig-21.10).

![Fig-21.10 Supination-external rotation injury](image)

**Vertical compression injuries:** All the above injuries may become complex due to a component of vertical compression force. It may be primarily a vertical compression injury resulting in either an anterior marginal fracture of the tibia or a comminuted fracture of the tibial articular surface with a fracture of the fibula—Pilon fracture (Fig-21.11).

![Fig-21.11 Vertical compression injury](image)

*It is important to recognise disruption of tibio-fibular syndesmosis on a proper AP X-ray of the ankle, and reconstruct it.*
**CLINICAL FEATURES**

There is history of a twisting injury to the ankle followed by pain and swelling. Often the patient is able to describe exactly the way the ankle got twisted. On examination, the ankle is found to be swollen. Swelling and tenderness may be localised to the area of injury (bone or ligament). Crepitus may be noticed if there is a fracture. The ankle may be lying deformed (adducted or abducted, with or without rotation).

**RADIOLOGICAL EXAMINATION**

Antero-posterior and lateral X-rays of the ankle are sufficient in most cases (Fig-21.12). While examining an X-ray, it is important to make note of the following features:

- **Fracture line** of the medial and lateral malleoli should be studied in order to evaluate the type of ankle injury (Lauge-Hansen classification). Small avulsion fractures from the malleoli are sometimes missed. These often have attached to them the whole ligament.

- **Tibio-fibular syndesmosis**: All ankle injuries where the fibular fracture is above the mortice, the syndesmosis is bound to have been disrupted. In injuries where the fibular fracture is at the level of the syndesmosis, one must carefully look for any lateral subluxation of the talus; if it is so, width of the joint space between the medial malleolus and the talus will be more than the space between the weight bearing surfaces of tibia and talus.

- **Posterior subluxation** of the talus should be looked for, on the lateral X-ray.

- **Soft tissue swelling** on the medial or lateral side in the absence of a fracture, must arouse suspicion of a ligament injury. This should be confirmed or ruled out after thorough clinical examination and stress X-rays. MRI may help.

**TREATMENT**

**Principles of treatment:** The complexity of the forces involved produce a variety of combinations of fractures and fracture-dislocations around the ankle. The basic principle of treatment is to achieve anatomical reconstruction of the ankle-mortise. This helps in regaining good function and reducing the possibility of osteoarthritis developing later. In some cases, it is possible to do so by conservative methods. But in most, an operative reduction and internal fixation is required. Given below are some general principles:

**Fractures without displacement:** It is usually sufficient to protect the ankle in a below-knee plaster for 3-6 weeks. Good, ready-made braces can be used in place of rather uncomfortable plaster cast.

**Fractures with displacement:** Aim of treatment is to ensure anatomical reduction of the ankle-mortise. This means, ensuring anatomical reduction of medial and lateral malleoli, and reduction of the talus accurately within the mortice. Following modes of treatment may be useful:

a) **Operative methods:** More and more surgeons are now resorting to internal fixation for all displaced fractures of ankle without attempting closed reduction. This is done because by operative reduction, it is possible to achieve perfect alignment as well as stable fixation of fragments. This allows early motion of the ankle joint, thereby improving overall results. This approach is justified in hospitals where trained staff and all equipment necessary for such work is available.

**Internal fixation:** In general, operative reduction and internal fixation may be used in cases where closed reduction has not been successful, or the reduction has slipped during the course of conservative treatment. The following techniques
Injuries to the Leg, Ankle and Foot

COMPLICATIONS

Simple types of ankle injuries are almost free of complications. More serious fracture-dislocation may be complicated because of improper treatment. Sometimes, the nature of injury is such that perfect functions cannot be restored. The following complications may occur:

1. **Stiffness of the ankle:** Following immobilisation in plaster, stiffness occurs. In ankle injuries, recovery takes a long time because of the tendency for gravitational oedema which may hinder mobilisation exercises. It is most common in elderly persons. With persistent treatment, using limb elevation, crepe bandage and active toe movements, oedema subsides. It may be necessary to continue ankle exercises for a long period (6-8 months).

2. **Osteoarthritis:** Since most ankle fractures involve the articular surfaces, anything short of a perfect anatomical reduction with smooth and congruous joint surfaces will lead to wear and tear of the articular cartilage. This will start the process of degenerative osteoarthritis. Greater the irregularity of the articular surfaces, more rapidly will the degenerative changes occur. The patient will complain of persistent pain, swelling and joint stiffness. Once established, osteoarthritis cannot be reversed. In a case where the disability (pain, etc.) is severe, it may be required to eliminate the joint by fusing the talus to the tibia (ankle arthrodesis).

**SPRAINED ANKLE**

It is the term used for ligament injuries of the ankle. Commonly, it is an inversion injury, and the lateral collateral ligament is sprained. Sometimes, an eversion force may result in a sprain of the medial collateral ligament of the ankle.

**Conservative methods:** It is often possible to achieve a good reduction by manipulation under general anaesthesia. The essential feature of the reduction is to concentrate on restoring the alignment of the foot to the leg. By doing so the fragments automatically fall into place. Once reduced, a below-knee plaster cast is applied. If the check X-ray shows a satisfactory position, the plaster cast is continued for 8-10 weeks. The patient is not allowed to bear any weight on the leg during this period. Check X-rays are taken frequently to make sure the fracture does not get displaced. If everything goes well, the plaster is removed after 8-10 weeks and the patient taught physiotherapy to regain movement at the ankle.

**External fixation:** This may be required in cases where closed methods cannot be used e.g., open fractures with bad crushing of the muscles and tendons, with skin loss around the ankle.

Medial Malleolus Fracture
- Transverse fracture – compression screw, tension-band wiring
- Oblique fracture – compression screws
- Avulsion fracture – tension-band wiring

Lateral Malleolus Fracture
- Transverse fracture – tension-band wiring
- Spiral fracture – compression screws
- Comminuted fracture – buttress plating
- Fracture of the lower third of fibula – 4-hole plate

Posterior Malleolus
- Involving less than one-third of the articulating surface of the tibia – no additional treatment
- Involving more than one-third of the articulating surface of the tibia – internal fixation with compression screws
- Tibio-fibular syndesmosis disruption – needs to be stabilised by inserting a long screw from the fibula into the tibia

All major ligament injuries e.g., that of deltoid ligament, lateral ligament should be repaired.

b) **Conservative methods:** It is often possible to achieve a good reduction by manipulation under general anaesthesia. The essential feature of the reduction is to concentrate on restoring the alignment of the foot to the leg. By doing so the fragments automatically fall into place. Once reduced, a below-knee plaster cast is applied. If the check X-ray shows a satisfactory position, the plaster cast is continued for 8-10 weeks. The patient is not allowed to bear any weight on the leg during this period. Check X-rays are taken frequently to make sure the fracture does not get displaced. If everything goes well, the plaster is removed after 8-10 weeks and the patient taught physiotherapy to regain movement at the ankle.

* In comminuted fractures, it may not be possible to fix the fragments; in such situations, buttress plate keeps the fragments in place.
• Inversion of a plantar-flexed foot for anterior talo-fibular ligament sprain.
• Inversion in neutral position for complete lateral collateral ligament sprain.
• Eversion in neutral position for medial collateral ligament sprain.

Radiological examination: X-rays of the ankle (AP and lateral) are usually normal. In some cases, stress X-rays may be done to judge the severity of the sprain. A tilt of the talus greater than 20° on forced inversion or eversion indicates a complete tear of the lateral or medial collateral ligament respectively.

Treatment: It depends upon the grade of sprain:
• Grade I: Below-knee plaster cast for 2 weeks followed by mobilisation.
• Grade II: Below-knee cast for 4 weeks followed by mobilisation.
• Grade III: Below-knee cast for 6 weeks followed by mobilisation.

Current trend is to treat ligament injuries, in general, by ‘functional’ method i.e., without immobilisation. Treatment consists of rest, ice packs, compression, and elevation (RICE) for the first 2-3 days. The patient begins early protected range of motion exercises. Methods are devised by which during mobilisation, stress is avoided on ‘healing’ ligaments, and the muscles around the joint are built up. For this approach, a well-developed physiotherapy unit is required. For grade III ligament injury to the ankle, especially in young athletic individuals, operative repair is preferred by some surgeons.

CHRONIC ANKLE SPRAIN
Chronic recurrent sprain ankle is a disabling condition. If a course of physiotherapy and modification in shoe has not helped, a detailed evaluation with MRI and arthroscopy may be necessary. Pain in a number of these so-called chronic ankle sprains is in fact due to impingement of the scarred capsule or chondromalacia of the talus. Arthroscopy is a good technique for diagnosis and treatment of such cases.

FRACTURES OF THE CALCANEUM
RELEVANT ANATOMY
The calcaneum forms the bone of the heel. Its upper surface articulates with the talus, and the front surface with the cuboid. Its inferior surface is prolonged backwards as the tüber calcanei. Normally, the angle between the superior articular surface (between talus and calcaneum) and the upper surface of the tuberosity is 35° (tuber-joint angle, Fig-21.13). It is reduced in most fractures of the calcaneum.

Fig-21.13 Tuber-joint angle

PATHOANATOMY
Fractures of the calcaneum are caused by fall from height onto the heels, thus both heels may be injured at the same time. The fracture may be: (i) an isolated crack fracture, usually in the region of the tuberosity; or (ii) more often a compression injury where the bone is shattered like an egg shell. The degree of displacement varies according to the severity of trauma. The fracture may be of one of the following types (Fig-21.14).
• Undisplaced fracture resulting from a minimal trauma.

Fig-21.14 Types of fracture calcaneum
Injuries to the Leg, Ankle and Foot

- **Extra-articular fracture**, where the articular surfaces remain intact, and the force splits the calcaneal tuberosity vertically.

- **Intra-articular fracture**, where the articular surface of the calcaneum fails to withstand the stress. It is shattered and is driven downwards into the body of the bone, crushing the delicate trabeculae of the cancellous bone into powder. This is the *commonest* type of fracture.

**DIAGNOSIS**

**Clinical features**: The patient often gives a history of a fall from height, landing on their heels (e.g. a thief jumping from the first floor of a house). There is pain and swelling in the region of the heel. The patient is not able to bear weight on the affected foot. On examination, there is marked swelling and broadening of the heel. If first seen after a day or two, there will be ecchymosis around the heel and on the sole. Movement at the ankle is not appreciably impaired.

Many cases of compression fractures of the calcaneum are associated with a compression fracture of a vertebral body (usually in the dorso-lumbar region), fractures of the pubic rami, or an atlanto-axial injury. One must look for these injuries in a case of a fracture of the calcaneum.

**Radiological examination**: It is possible to diagnose most calcaneum fractures on a lateral X-ray of the heel. In some cases, an additional axial view of the calcaneum may be required. Very often, rather than a clear fracture extending through the calcaneum, there occurs crushing of the bone. This can be diagnosed on a lateral X-ray of the heel by reduction in the tuber-joint angle (Fig-21.15).

**TREATMENT**

**Undisplaced fracture**: Below-knee plaster cast for 4 weeks followed by mobilisation exercises.

**Compression fracture**: This is a serious injury which inevitably leads to permanent impairment of functions. Many different methods of treatment have been advocated with no appreciable difference in results. The following method is one used most widely.

The foot is elevated in a well padded below-knee plaster slab for 2-3 weeks. Once pain and swelling subside, the slab is removed and ankle and foot mobilisation begun. Leg elevation is continued, and a compression bandage (crepe bandage) applied for a period of 4-6 weeks in order to avoid gravitational oedema. Weight bearing is not permitted for a period of 12 weeks.

Trend is towards surgical reconstruction of these fractures, at centres where facilities are available.

**COMPICATIONS**

1. **Stiffness** of the subtalar and mid-tarsal joints: Some amount of stiffness of the subtalar joint, resulting in limitation to the inversion-eversion motion of the foot is inevitable in most compression fractures of the calcaneum. Stiffness can be kept to minimum by early physiotherapy.

2. **Osteoarthritis**. Because of the irreparable distortion of the subtalar joint surface, osteoarthritis is an expected complication. It results in pain and stiffness, most noticeable while walking on an uneven surface. A patient with a severe disability may require fusion of the subtalar joint (arthrodesis).

**FRACTURES OF THE TALUS**

Minor fractures in the form of a small chip from the margins of one of the articular surfaces of the talus are more common than the more serious fracture i.e. fracture of the neck of the talus.

**RELEVANT ANATOMY**

**Blood supply to the talus**: This is the only bone of the foot without any muscle attachment. The main blood supply to the talus is from the anastomotic ring of blood vessels, the osseous vessels entering its neck and running postero-laterally within the bone to supply its body. Therefore, blood supply to the body of the talus is often cut off following fractures occurring through the neck.

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*Fig-21.15 X-ray showing fracture calcaneum*
INJURIES OF THE TARSAL BONES

Fractures and dislocations of other tarsal bones are uncommon. Most of the fractures can be treated by a below-knee plaster cast. Most dislocations at any of the tarsal joints (subtalar, talo-navicular or inter-tarsal) can be treated by manipulation and immobilisation in a plaster cast. Sometimes, an open reduction and internal fixation with K-wires may be required.

FRACTURES OF THE METATARSAL BONES

Most metatarsal fractures are caused by direct violence from a heavy object falling onto the foot. A metatarsal fracture may be caused by repeated stress without any specific injury (march fracture). Some of the commoner types of metatarsal fractures are discussed below.

FRACTURE OF THE BASE OF 5TH METATARSAL (Jones’ fracture)

This is a fracture at the base of the 5th metatarsal, caused by the pull exerted by the tendon of the peroneus brevis muscle inserted on it. Clinically, there is pain, swelling and tenderness at the outer border of the foot, most marked at the base of the 5th metatarsal. Diagnosis is easily confirmed on X-ray (Fig-21.17). Treatment is by a below-knee walking plaster cast for 3 weeks.

DIAGNOSIS

Unless carefully examined on a lateral X-ray of the ankle, this fracture is frequently missed because of the overlapping of the tarsal bones.

TREATMENT

It depends upon the displacement. If undisplaced, a below-knee plaster cast for 8-10 weeks is sufficient. In a displaced fracture, open reduction and internal fixation of the fracture with a screw is required.

COMPLICATIONS

1. Avascular necrosis and non-union: Because of the poor blood supply, after a fracture through the neck, the body of the talus becomes avascular. The avascular fragment fails to unite with rest of the bone and gradually collapses, leading to deformation of the bone, and eventually osteoarthritis of the ankle.

2. Osteoarthritis: Besides avascular necrosis of the talus, an associated injury to its articular cartilage may lead to osteoarthritis of the ankle. The patient complains of pain and stiffness. Treatment is mostly by physiotherapy and fomentation. In severe cases, an ankle arthrodesis may be needed.

FRACTURE OF THE METATARSAL SHAFTS

One or more metatarsal shafts may be fractured, mostly following a crush injury. Treatment is by below-knee plaster cast for 3-4 weeks.

MARCH FRACTURE

It is a ‘fatigue’ fracture of third metatarsal, resulting from long continued or often repeated stress, particularly from prolonged walking or running.
in those not accustomed to it. Thus, it may occur in army recruits freshly committed to marching – hence the term ‘March fracture’. The fracture heals spontaneously, so treatment is purely symptomatic.

FRACTURES OF PHALANGES OF THE TOES
These are common injuries, most often resulting from fall of a heavy object, or twisting of the toes. The great toe is injured most commonly. Satisfactory general alignment is maintained in most cases and little or no treatment is required. The injured toe is covered with a soft woolly dressing and strapped to the toe adjacent to it.

Further Reading

What have we learnt?
• Fracture both bones of leg is one of the commonest fracture of lower extremity.
• These fractures are commonly open, hence associated with complications.
• Closed interlock nailing is a usual method of treating these fractures.
• Ankle injuries are common, operative stabilisation is the treatment of choice.
• Fracture of calcaneum occurs due to fall from height, is essentially treated by non-operative methods.

Additional information: From the entrance exams point of view
• Tuber-joint angle (Bohler’s angle), Gissane (crucial) angle and neutral angle are measured on a radiograph in a calcaneal fracture.
• Most common site for ligament injury in the body is the ankle.
• Most common mode of ankle injury is inversion and plantarflexion of the foot.
• Most common cause of insertional tendonitis and ruptured Achilles tendon is overuse.
Infections of Bones and Joints

TOPICS

- Acute osteomyelitis
- Secondary osteomyelitis
- Chronic osteomyelitis
- Garre’s osteomyelitis
- Brodie’s abscess
- Salmonella osteomyelitis
- Septic arthritis
- Gonococcal arthritis
- Syphilis of joints
- Fungal infections
- Leprosy and orthopaedics
- Salmonella osteomyelitis

Infection of the bone by micro-organisms is called osteomyelitis. Conventionally, an unqualified term ‘osteomyelitis’ is used for infection of the bone by pyogenic organisms. Osteomyelitis can be acute or chronic.

ACUTE OSTEOMYELITIS

This can be primary (haematogenous) or secondary (following an open fracture or bone operation). Haematogenous osteomyelitis is the commonest, and is often seen in children.

RELEVANT ANATOMY

Metaphysis of the long bones (Fig-22.1): It is a highly vascularised zone. From the diaphysis the medullary arteries reach up to the growth plate; the area of greatest activity, and branch into capillaries. The venous system begins in this area and drains toward the diaphysis. Thus, the vessels in this zone are arranged in the form of a loop (hair-pin arrangement). The blood stasis resulting from such an arrangement is probably responsible for the metaphysis being a favourite site for bacteria to settle, and thus a common site for osteomyelitis.

In most joints, the capsule is attached at the junction of the epiphysis with the metaphysis i.e., the metaphysis is extra-articular (Fig-22.2). In some joints, part of the metaphysis is intra-articular, so that the infection from the metaphysis can spread to the joint, resulting in pyogenic arthritis.

AETIOPATHOGENESIS

Staphylococcus aureus is the commonest causative organism. Others are Streptococcus and Pneumococcus. These organisms reach the bone via the blood circulation. Primary focus of infection is generally not detectable.
The bacteria, as they pass through the bone, get lodged in the metaphysis. Lower femoral metaphysis is the commonest site. The other common sites are the upper tibial, upper femoral and upper humeral metaphyses.

**Pathology:** The host bone initiates an inflammatory reaction in response to the bacteria. This leads to bone destruction and production of an inflammatory exudate and cells (pus). Once sufficient pus forms in the medullary cavity, it spreads in the following directions (Fig-22.3).

a) **Along the medullary cavity:** Pus trickles along the medullary cavity and causes thrombosis of the venous and arterial medullary vessels. Blood supply to a segment of the bone is thus cut off.

b) **Out of the cortex:** Pus travels along Volkmann’s canals and comes to lie sub-periosteally. The periosteum is thus lifted off the underlying bone, resulting in damage to the periosteal blood supply to that part of the bone. A segment of bone is thus rendered avascular (sequestrum). Dimensions of this segment vary from a small invisible piece to the whole diaphysis of the bone (Fig-22.4). Pus under the periosteum generates sub-periosteal new bone (periosteal reaction). Eventually the periosteum is perforated, letting the pus out into the muscle or subcutaneous plane, where it can be felt as an abscess. The abscess, if unattended, bursts out of the skin, forming a discharging sinus.

c) **In other directions:** The epiphyseal plate is resistant to the spread of pus. At times it may be affected by the inflammatory process. The capsular attachment at the epiphysismetaphysis junction prevents the pus from entering the nearby joint. In joints with an intra-articular metaphysis, pus can spread to the joint, and cause acute pyogenic arthritis e.g., in the hip, in the shoulder etc.

**DIAGNOSIS**
The diagnosis of acute osteomyelitis is basically clinical. It is a disease of childhood, more common in boys, probably because they are more prone to injury.

**Presenting complaints:** The child presents with an acute onset of pain and swelling at the end of a bone, associated with systemic features of infection like fever etc. Often the parents attribute the symptoms to an episode of injury, but the injury is coincidental. One may find a primary focus of infection elsewhere in the body (tonsils, skin, etc.).

**Examination:** The child is febrile and dehydrated with classic signs of inflammation – redness, heat, etc. localised to the metaphyseal area of the bone. In later stages, one may find an abscess in the muscle or subcutaneous plane. There may be swelling of the adjacent joint, because of either sympathetic effusion or concomitant arthritis.

**Investigations:** Investigations provide few clues in the early phase of the disease.

- **Blood:** There may be polymorphonuclear leucocytosis and an elevated ESR. A blood culture at the peak of the fever may yield the causative organism.

- **X-rays** (Fig-22.5): The earliest sign to appear on the X-ray is a periosteal new bone deposition
(periosteal reaction) at the metaphysis. It takes about 7-10 days to appear.

- **Bone scan:** A bone scan using Technetium-99 may show increased uptake by the bone in the metaphysis. This is positive before changes appear on X-ray. This may be indicated in a very early case where diagnosis is in doubt.

Indium-111 labelled leucocyte scan is most specific for diagnosis of bone infection.

**DIFFERENTIAL DIAGNOSIS**

Any acute inflammatory disease at the end of a bone, in a child, should be taken as acute osteomyelitis unless proved otherwise. Following are some of the differential diagnosis to be considered:

a) **Acute septic arthritis:** This can be differentiated from acute osteomyelitis by the following features in arthritis:
   - Tenderness and swelling localised to the joint rather than the metaphysis.
   - Movement at the joint is painful and restricted.
   - In case of doubt, joint fluid may be aspirated under strict aseptic conditions, and the fluid examined for inflammatory cells.

b) **Acute rheumatic arthritis:** The features are similar to acute septic arthritis. The fleeting character of joint pains, elevated ASLO titre and CRP values may help in diagnosis.

c) **Scurvy:** There is formation of sub-periosteal haematomas in scurvy. These may mimic acute osteomyelitis radiologically, but the relative absence of pain, tenderness and fever points to the diagnosis of scurvy. There may be other features of malnutrition.

d) **Acute poliomyelitis:** In the acute phase of poliomyelitis, there is fever and the muscles are tender, but there is no tenderness on the bones.

Parents often tend to relate an episode of injury to onset of symptoms in any musculo-skeletal pain. This may give a wrong lead, and a novice may make a diagnosis of a fracture or soft tissue injury. Often such a patient is immobilised in plaster cast, only to know later that the infection was the cause. *Any history of trauma, particularly in children must be thoroughly questioned.*

**TREATMENT**

Early, adequate treatment of acute osteomyelitis is the key to success. The child is admitted and investigated. Treatment depends upon the duration of illness after which the child is brought. Cases can be arbitrarily divided into two groups:

a) **If the child is brought within 48 hours of the onset of symptoms:** If a child is brought early, it is supposed that pus has not yet formed and the inflammatory process can be halted by systemic antibiotics. Treatment consists of rest, antibiotics and general building-up of the patient. The limb is put to rest in a splint or by traction. Choice of antibiotics varies from centre to centre. It broadly depends upon the age of the child and choice of the doctor. In children less than 4 months of age, a combination of Ceftriaxone and Vancomycin in appropriate dose is preferred. In older children, a combination of Ceftriaxone and Cloxacillin is given. Antibiotics are started after taking blood for culture and sensitivity. Antibiotics are changed to specific ones depending upon the culture and sensitivity report.

The child is adequately rehydrated with intravenous fluids. Response to the above treatment is evaluated by frequent assessment of the patient. A four hourly temperature chart and pulse record is maintained. It is a good idea to outline the area of local tenderness precisely, with the help of the back of a match stick over regular intervals. If the patient responds favourably, fever will start declining and local inflammatory signs will diminish. As the child improves, the limb can be mobilised. Weight bearing is restricted for 6-8 weeks.
After 2 weeks, antibiotics can be administered by oral route for 6 weeks. If the patient does not respond favourably within 48 hours of starting the treatment, surgical intervention is required.

b) **If the child is brought after 48 hours of the onset of symptoms:** If the child is brought late or if he does not respond to conservative treatment, it is taken for granted that there is already a collection of pus within or outside the bone. Detection of pus is often difficult by clinical examination because it may lie deep to the periosteum. An ultrasound examination of the affected part may help in early detection of deep collection of pus. Surgical exploration and drainage is the mainstay of treatment at this stage. A drill hole is made in the bone in the region of the metaphysis. If pus wells up from the drill hole, the hole is enlarged until free drainage is obtained. A swab is taken for culture and sensitivity. The wound is closed over a sterile suction drain. Rest, antibiotics and hydration are continued post-operatively. Gradually, the inflammation is controlled and the limb is put to use. Antibiotics are continued for 6 weeks.

**COMPLICATIONS**

This can be divided into two types, general and local:

- **General complications:** In the early stage, the child may develop septicaemia and pyaemia. Either complication, if left uncontrolled, may prove fatal.

- **Local complications:** It is unfortunate that a large number of cases of acute osteomyelitis in developing countries develop serious complications. Most of these are because of delay in diagnosis, and inadequate treatment. Some of the common complications are as follows:

  1. **Chronic osteomyelitis:** It is the commonest complication of acute osteomyelitis. There are hardly any radiological features in the early stage. A delay in diagnosis leads to sequestrum formation (Fig-22.6) and pent-up pus in the cavities inside the bone. Poor host resistance is another reason for the chronicity of the disease.

  2. **Acute pyogenic arthritis:** This occurs in joints where the metaphysis is intra-articular e.g., the hip (upper femoral metaphysis), the shoulder (upper humeral metaphysis), etc.

  3. **Pathological fracture:** This occurs through a bone which has been weakened by the disease or by the window made during surgery. It can be avoided by adequately splinting the limb.

  4. **Growth plate disturbances:** It may be damaged leading to complete or partial cessation of growth. This may give rise to shortening, lengthening or deformity of the limb.

**SECONDARY OSTEOMYELITIS**

This condition arises from a wound infection in open fractures or after operations on the bone. The incidence of these cases are on the rise because of increase in operative intervention in the treatment of fractures.

The constitutional symptoms are less severe than those in haematogenous osteomyelitis as the wound provides some drainage. The condition can be largely prevented by adequate initial treatment of open fractures, and adherence to sterile operating conditions for routine orthopaedic operations.

**CHRONIC OSTEOMYELITIS**

Conventionally, the term ‘chronic osteomyelitis’ is used for chronic pyogenic osteomyelitis. Although, its incidence is on the decline in developed countries, it continues to be an important problem in developing countries. The other causes of chronic osteomyelitis are tuberculosis, fungal infections etc. There are three types of chronic osteomyelitis:

a) Chronic osteomyelitis secondary to acute osteomyelitis.

b) Garre’s osteomyelitis.

c) Brodie’s abscess.
PATHOLOGY

Acute osteomyelitis commonly leads to chronic osteomyelitis because of one or more of the following reasons:

a) Delayed and inadequate treatment: This is the commonest cause for the persistence of an osteomyelitis. Delay causes spread of pus within the medullary cavity and subperiosteally. This results in the death of a part of the bone (sequestrum formation). Destruction of cancellous bone leads to the formation of cavities within the bone. *Such ‘non-collapsing’ bone cavities and sequestra are responsible for persistent infection.*

b) Type and virulence of organism: Sometimes, despite early, adequate treatment of acute osteomyelitis, the body’s defense mechanism may not be able to control the damaging influence of a highly virulent organism, and the infection persists.

c) Reduced host resistance: Malnutrition compromises the body’s defense mechanisms, thus letting the infection persist.

When infection persists because of the above reasons, the host bone responds by generating more and more sub-periosteal new bone. This results in thickening of the bone. The sub-periosteal bone is deposited in an irregular fashion so that the osteomyelitic bone has an irregular surface. Continuous discharge of pus results in the formation of a sinus. With time, the sinus tract gets fibrosed and the sinus becomes fixed to the bone.

*Sequestrum* is a piece of dead bone, surrounded by infected granulation tissue trying to ‘eat’ the sequestrum away. It appears pale and has a smooth inner and rough outer surface (Fig-22.7), because the latter is being constantly eroded by the surrounding granulation tissue.

Different types of sequestra seen in different conditions as shown in Table 22.1.

Table 22.1: Different types of sequestra

<table>
<thead>
<tr>
<th>Type</th>
<th>Disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tubular</td>
<td>Pyogenic</td>
</tr>
<tr>
<td>Ring</td>
<td>External fixator</td>
</tr>
<tr>
<td>Black</td>
<td>Actinomycosis</td>
</tr>
<tr>
<td>Coralliform</td>
<td>Perthe’s disease</td>
</tr>
<tr>
<td>Coke</td>
<td>Tuberculosis</td>
</tr>
<tr>
<td>Sandy</td>
<td>Tuberculosis</td>
</tr>
<tr>
<td>Feathery</td>
<td>Syphilis</td>
</tr>
</tbody>
</table>

*Involucrum* is the dense sclerotic bone overlying a sequestrum. There may be some holes in the involucrum for pus to drain out. These holes are called *cloacae*. The bony cavities are lined by infected granulation tissue.

DIAGNOSIS

Diagnosis is suspected clinically but can be confirmed radiologically by its characteristic features. The disease begins in childhood but may present later. The lower-end of the femur is the commonest site.

Presenting complaints: A chronic discharging sinus is the commonest presenting symptom. The onset of sinus may be traced back to an episode of acute osteomyelitis during childhood. Often sinususes heal for short periods, only to reappear with each acute exacerbation. Quality of discharge varies from seropurulent to thick pus. There may be a history of extrusion of small bone fragments from the sinus.

Pain is usually minimal but may become aggravated during acute exacerbations. Generalised symptoms of infection such as fever etc., are present only during acute exacerbations. A patient with chronic osteomyelitis may present with complications discussed subsequently (see page 174).

Examination: Some of the salient features observed on examination are as follows:

- **Chronic discharging sinus:** This is a characteristic feature of chronic infection. *A sinus fixed* to the underlying bone indicates that infection is coming from the bone. There may be sprouting granulation tissue at its opening, indicating a sequestrum within the bone. The sequestrum may be visible at the mouth of the sinus itself. The sinus may be surrounded by healed puckered scars, indicating previous healed sinuses.
• **Thickened, irregular bone**: This can be appreciated on comparing the girth of the affected bone with that of the bone on the normal side.

• **Tenderness** on deep palpation, usually mild, is present in some cases.

• **Adjacent joint** may be stiff, either due to excessive scarring in the soft tissues around the joint, or because of associated arthritis of the joint.

**INVESTIGATIONS**

**Radiological examination**: The following are some of the salient radiological features seen in chronic osteomyelitis:

• **Thickening and irregularity** of the cortices

• **Patchy sclerosis**

• **Bone cavity**: This is seen as an area of rarefaction surrounded by sclerosis.

• **Sequestrum**: This appears **denser** than the surrounding normal bone because the decalcification which occurs in normal bone, does not occur in dead bone. Granulation tissue surrounding the sequestrum gives rise to a radiolucent zone around it. A sequestrum may be visible in soft tissues.

• **Involucrum** and **cloaca** may be visible.

**Sinogram**: In this test, a sterile thin catheter is introduced into the sinus as far as it can go. Then, a radio-opaque dye is injected, and X-rays taken. The radio-opaque dye travels to the root of the infection, and thus helps localise it better. It is indicated in situations where one cannot tell on X-rays where the pus may be coming from.

**CT scan and MRI**: are sometimes indicated in patients where diagnosis is in doubt. CT scan is of particular use in better defining the cavities and sequestra, which sometimes cannot be seen on routine X-rays. Exact localisation of a cavity or sequestrum has bearing on surgical treatment.

**Blood**: A blood examination is usually of no help. ESR may be normal or mildly elevated. Total blood counts may be normal, may be increased during acute exacerbation only.

**Pus**: Pus culture may grow the causative organism. This should be taken from depth of the sinus after proper cleaning of the skin. If an organism is grown, it may be useful in controlling the acute phase of the disease. It may also help in selecting the pre-operative antibiotics as and when operation is performed.

**DIFFERENTIAL DIAGNOSIS**

A discharging sinus on a limb indicates deeper infection which could be from tissues, skin downward. A history of bone piece discharge from a sinus is **diagnostic** of chronic osteomyelitis. Other differential diagnosis to be considered in the absence of such a history are as follows:

a) **Tubercular osteomyelitis**: The discharge is often thin and watery. A tubercular sinus may show its characteristic features like undermined margins and bluish surrounding skin. Tubercular osteomyelitis is often multifocal. Patient may be suffering from or may have suffered from pulmonary tuberculosis.

b) **Soft tissue infection**: A longstanding soft tissue infection with a discharging sinus may mimic osteomyelitis. Absence of thickening of underlying bone, and absence of sinus fixed to the bone, may point towards the infection **not coming** from the bone. Absence of any radiological changes in the bone would help conform the diagnosis.

c) **Ewing’s sarcoma**: A child with Ewing’s sarcoma sometimes presents with a rather sudden onset pain and swelling, mostly in the diaphysis. Radiological appearance often resembles that of osteomyelitis. A biopsy will settle the diagnosis.

**TREATMENT**

**Principles of treatment**: *Treatment of chronic osteomyelitis is primarily surgical.* Antibiotics are useful only during acute exacerbations and during post-operative period. Aim of surgical intervention is: (i) removal of dead bone; (ii) elimination of dead space and cavities; and (iii) removal of infected granulation tissue and sinuses.

**Operative procedures**: Following are some of the operative procedures commonly performed:

a) **Sequestrectomy**: This means removal of the sequestrum. If it lies within the medullary cavity, a window is made in the overlying involucrum and the sequestrum removed. One must wait for adequate involucrum formation before performing sequestrectomy.
b) **Saucerisation**: A bone cavity is a ‘non-collapsing cavity’, so that there is always some pent-up pus inside it. This is responsible for the persistence of an infection. In saucerisation, the cavity is converted into a ‘saucer’ by removing its wall (Fig-22.8). This allows free drainage of the infected material.

c) **Curettage**: The wall of the cavity, lined by infected granulation tissue, is curetted until the underlying normal-looking bone is seen. The cavity is sometimes obliterated by filling it with gentamycin impregnated cement beads or local muscle flap.

d) **Excision of an infected bone**: In a case where the affected bone can be excised *en bloc* without compromising the functions of the limb, it is a good method e.g., osteomyelitis of a part of the fibula. With the availability of Ilizarov’s technique, an aggressive approach, i.e., excising the infected bone segment and building up the gap by transporting a segment of the bone from adjacent part has shown good results (Ref. page 34).

e) **Amputation**: It may, very rarely, be preferred in a case with a long standing discharging sinus, especially if the sinus undergoes a malignant change.

In most cases, a combination of these procedures is required. After surgery the wound is closed over a continuous suction irrigation system (Fig-22.9). This system has an inlet tube going to the medullary cavity, and an outlet tube bringing the irrigation fluid out. A slow suction is applied to the outlet tube. The irrigation fluid consists of suitable antibiotics and a detergent. The medullary canal is irrigated in this way for 4 to 7 days.

### COMPLICATIONS

1. **An acute exacerbation** or ‘flare up’ of the infection occurs commonly. It subsides with a period of rest, and antibiotics, either broad-spectrum or based on the pus culture and sensitivity report.

2. **Growth abnormalities**: Osteomyelitis may cause growth disturbances at the adjacent growth plate, in one of the following ways:
   - **Shortening**, when the growth plate is damaged.
   - **Lengthening** because of increased vascularity of the growth plate due to the nearby osteomyelitis.
   - **Deformities** may appear if a part of the growth plate is damaged and the remaining keeps growing.

3. **Pathological fracture** may occur through a weakened area of the bone. Treatment is by conservative methods.

4. **Joint stiffness** may occur because of scarring of soft tissues around the joint or due to the joint getting secondarily involved.

5. **Sinus tract malignancy** is a rare complication. It occurs many years after the onset of osteomyelitis. It is usually a squamous cell carcinoma. The patient may need amputation.

6. **Amyloidosis**: As with all other long standing suppurations, this is a late complication of osteomyelitis.
PROGNOSIS
To cure a bone infection is very difficult. Operative intervention may be useful if there is an obvious factor responsible for the persistence of the infection e.g., sequestrum, cavity etc.

GARRE’S OSTEOMYELITIS
This is a sclerosing, non-suppurative chronic osteomyelitis. It may begin with acute local pain, pyrexia and swelling. Pyrexia and pain subside but the fusiform osseous enlargement persists. There is tenderness on deep palpation. There is no discharging sinus. Shafts of the femur or tibia are the most commonly affected.

The importance of Garre’s osteomyelitis lies in differentiating it from bone tumours, which commonly present with similar features e.g., Ewing’s tumour or osteosarcoma.

Treatment is guarded. Acute symptoms subside with rest and broad-spectrum antibiotics. Sometimes, making a gutter or holes in the bone bring relief in pain.

BRODIE’S ABSCESS
It is a special type of osteomyelitis in which the body’s defense mechanisms have been able to contain the infection so as to create a chronic bone abscess containing pus or jelly-like granulation tissue surrounded by a zone of sclerosis (Fig-22.10).

Clinical features: The patient is usually between 11 to 20 years of age. Common sites are the upper-end of the tibia and lower-end of the femur. It is usually located at the metaphysis. A deep boring pain is the predominant symptom. It may become worse at night. In some instances, it becomes worse on walking and is relieved by rest. Occasionally, there may be a transient effusion in the adjacent joint during exacerbation of symptoms. An examination may reveal tenderness and thickening of the bone.

Radiological features: The radiological picture is diagnostic. It shows a circular or oval lucent area surrounded by a zone of sclerosis. The rest of the bone is normal.

Treatment is by operation. Surgical evacuation and curettage is performed under antibiotic cover. If the cavity is large, it is packed with cancellous bone chips.

SALMONELLA OSTEOMYELITIS
This occurs during the convalescent phase after an attack of typhoid fever. It is subacute type of osteomyelitis, usually occurring in the ulna, tibia, or vertebra. Often, multiple bones are affected, sometimes bilaterally symmetrical. The predominant radiological feature is a diaphyseal sclerosis. The disease occurs more commonly in children with sickle cell anaemia.

SEPTIC ARTHRITIS
This is an arthritis caused by pyogenic organisms. Typically, it presents as an acute painful arthritis, but it may present as subacute or chronic arthritis. Other terms often used to describe this condition are pyogenic arthritis, infective arthritis or suppurative arthritis.

AETIOPATHOGENESIS
It is more common in children, and males are more susceptible. Other predisposing factors are poor hygiene, poor resistance, diabetes etc. Staphylococcus aureus is the commonest causative organism. Other organisms are Streptococcus Pneumococcus and Gonococcus. The organisms reach the joint by one of the following routes:

a) Haematogenous: This is the commonest route. There may be a primary focus of infection in the form of pyoderma, throat infection, septicaemia etc.

b) Secondary to nearby osteomyelitis: This is a particularly common route in joints with intra-articular metaphysis e.g., the hip, shoulder etc.
c) **Penetrating wounds**: The knee, being a superficial joint, is often affected via this route.

d) **Iatrogenic**: This may occur following intra-articular steroid injections in different arthritis, and during femoral artery punctures for blood collection.

e) **Umbilical cord sepsis** in infants can travel to joints.

As the organism reaches the joint by one of the above routes, there begins an inflammatory response in the synovium resulting in the exudation of fluid within the joint. Joint cartilage is destroyed by inflammatory granulation tissue and lysosomal enzymes in the joint exudate. Outcome varies from complete healing to total destruction of the joint. The latter may result in a complete loss of joint movement (ankylosis).

**DIAGNOSIS**

Diagnosis is mainly clinical. The patient is usually a child. The knee is the commonest joint affected. Other joints commonly affected are the hip, shoulder, elbow etc.

**Presenting complaints**: In its typical acute form, a child with septic arthritis presents with a severe throbbing pain, swelling and redness of the affected joint. This is associated with high grade fever and malaise. The child is unable to use the affected limb. In its subacute form, the parents may notice that the child is not allowing anybody to touch the joint. He may not be moving it properly. In the lower limbs, a painful limp may be the first thing to draw attention. It may be associated with low grade fever.

**On examination**: The child is generally severely toxic with high temperature and tachycardia. The affected joint is swollen and held in the position of ease (Table-22.2). Palpation reveals increased temperature, tenderness and effusion. There is severe limitation in the joint movements in all directions. Any attempt at either passive or active movements causes severe pain and muscle spasms. In subacute forms, some amount of joint movement is possible.

**INVESTIGATIONS**

**Radiological Examination**: Diagnosis in early stage is crucial. X-rays are usually normal. A careful look at the X-ray may reveal increased joint space and a soft tissue shadow corresponding to the distended capsule due to swelling of the joint. *Ultrasound examination* is useful in detecting collection in deep joints such as the hip and shoulder. If found, one could aspirate the fluid and send for culturing the organism responsible for infection.

In the later stage, the joint space is narrowed. There may be irregularity of the joint margins. Occasionally, there may be a subluxation or dislocation of the joint.

**Blood** shows neutrophilic leucocytosis. ESR is markedly elevated. A blood culture may grow the causative organism.

**Joint aspiration** is the quickest and the best method of diagnosing septic arthritis. The fluid may show features of acute septic inflammation (Table-22.3). Gram staining provides a clue to the type of organism, till one gets the culture report.

**DIFFERENTIAL DIAGNOSIS**

A case with an acute septic arthritis should be differentiated from the following conditions:

a) **Other acute inflammatory conditions**: Diseases near a joint, such as acute osteomyelitis, acute lymphadenitis, acute bursitis etc. may mimic an arthritis because in some of these conditions, the joint is kept in a deformed position. Also, there may be pain and muscle spasm with attempted movements, but these signs are basically because the body is trying to prevent any motion in the vicinity of the inflamed part. Careful examination reveals that reasonably pain free movements are present at the joint, and the movements are not limited in every direction. The swelling may also be localised to one side of the joint.

b) **Other causes of acute arthritis**: An acute septic arthritis should also be differentiated from other causes of arthritis as discussed below:

<table>
<thead>
<tr>
<th>Joint</th>
<th>Position of ease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shoulder</td>
<td>Adduction, internal rotation</td>
</tr>
<tr>
<td>Elbow</td>
<td>Flexion, mid pronation</td>
</tr>
<tr>
<td>Wrist</td>
<td>Flexion</td>
</tr>
<tr>
<td>Hip</td>
<td>Flexion, abduction, external rotation</td>
</tr>
<tr>
<td>Knee</td>
<td>Flexion</td>
</tr>
<tr>
<td>Ankle</td>
<td>Plantar-flexion</td>
</tr>
</tbody>
</table>
Infections of Bones and Joints

with a suction drain. The same can be now done arthroscopically. As the inflammation is brought under control, general condition of the patient improves, fever and local signs of inflammation subside, the joint is then gradually mobilised. Antibiotics are continued for 6 weeks.

In late cases, with radiological destruction of the joint margins, subluxation or dislocation, it is not possible to expect joint movement. In such cases, after an arthrotomy and extensive debridement of the joint, it is immobilised in the position of optimum function, so that as the disease heals, ankylosis occurs in that position.

COMPLICATIONS

These can be divided into general and local, as for osteomyelitis. Inadequate early treatment leads to the following local complications.

1. Deformity and stiffness: The joint gets stiff due to intra-articular and peri-articular adhesions. In cases with advanced disease, the articular cartilage may be completely damaged, resulting in ankylosis. Bony ankylosis is the usual outcome of a neglected septic arthritis.

2. Pathological dislocation: As the joint gets filled with inflammatory exudate, the supporting ligaments and joint capsule get stretched. Muscle spasm associated with the disease may result in pathological dislocation of the joint. Posterior dislocation of the hip and triple displacement of the knee occur (Fig-22.11).

Rheumatic arthritis: Commonly a migratory polyarthritis, but may present with only one joint affected. The subsequent fleeting character of the arthritis, high C-reactive protein levels in the serum, and joint aspiration helps in its diagnosis.

Haemophilia: A past history of a bleeding disorder, especially in a boy with an acute painful joint, would suggest the diagnosis. Abnormal bleeding and clotting times are helpful for confirmation.

Tubercular arthritis: It may sometimes present in a rather acute form. A past or family history of tuberculosis may be present. Joint aspiration and AFB examination may help in its diagnosis.

TREATMENT

In its early stage, before any signs of joint destruction appear on X-ray, a correct diagnosis and aggressive treatment can save a joint from permanent damage. Whenever suspected, diagnosis of septic arthritis must be confirmed or ruled out by joint aspiration. Broad-spectrum antibiotics should be started by parenteral route. A combination of Ceftriaxone and Cloxacillin, in appropriate doses is usually given. These are subsequently changed to specific antibiotics as per aspirate culture and sensitivity reports. The joint must be put to rest in a splint or in traction.

Whenever pus is aspirated, the joint should be opened up (arthrotomy), washed and closed with a suction drain. The same can be now done arthroscopically. As the inflammation is brought under control, general condition of the patient improves, fever and local signs of inflammation subside, the joint is then gradually mobilised. Antibiotics are continued for 6 weeks.

In late cases, with radiological destruction of the joint margins, subluxation or dislocation, it is not possible to expect joint movement. In such cases, after an arthrotomy and extensive debridement of the joint, it is immobilised in the position of optimum function, so that as the disease heals, ankylosis occurs in that position.

COMPLICATIONS

These can be divided into general and local, as for osteomyelitis. Inadequate early treatment leads to the following local complications.

1. Deformity and stiffness: The joint gets stiff due to intra-articular and peri-articular adhesions. In cases with advanced disease, the articular cartilage may be completely damaged, resulting in ankylosis. Bony ankylosis is the usual outcome of a neglected septic arthritis.

2. Pathological dislocation: As the joint gets filled with inflammatory exudate, the supporting ligaments and joint capsule get stretched. Muscle spasm associated with the disease may result in pathological dislocation of the joint. Posterior dislocation of the hip and triple displacement of the knee occur (Fig-22.11).

---

Table-22.3: Synovial fluid examination

<table>
<thead>
<tr>
<th>Points</th>
<th>Normal</th>
<th>Non-inflammatory</th>
<th>Inflammatory</th>
<th>Septic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gross examination</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Volume (ml)</td>
<td>Often &lt; 3.5 ml</td>
<td>Often &gt; 3.5 ml</td>
<td>Often &gt; 3.5 ml</td>
<td>&gt; 3.5 ml.</td>
</tr>
<tr>
<td>Viscosity</td>
<td>High</td>
<td>High</td>
<td>Low</td>
<td>Variable</td>
</tr>
<tr>
<td>Colour</td>
<td>Colourless</td>
<td>Straw yellow</td>
<td>Yellow</td>
<td>Variable</td>
</tr>
<tr>
<td>Clarity</td>
<td>Transparent</td>
<td>Transparent</td>
<td>Translucent</td>
<td>Opaque</td>
</tr>
</tbody>
</table>

Examination in lab.

- PMN leucocytes: < 25% < 25% > 50% > 75%
- Culture: – – – +
- Mucin clot: Firm Firm Friable Friable
- Glucose level: Equal to blood glucose Nearly equal to blood glucose < 25 mg% of blood glucose > 25 mg% of blood glucose
- Crystal examination: Positive in Gout – Sodium biurate, Positive in Pseudogout – Ca pyrophosphate
Onset is sudden, similar to septic arthritis, but the general condition of the patient is well maintained in spite of severe local signs. This is typical of gonococcal arthritis. Knee is the commonest joint affected. Treatment is similar to that of septic arthritis. Penicillin is the drug of choice.

**SYMPHILIS OF THE JOINTS**

**CONGENITAL SYMPHILIS**

The joint may be affected early or late in congenital syphilis.

**Early:** During infancy, osteochondritis in the juxta-epiphyseal region results in breakdown of the bone and cartilage.

**Late:** A manifestation of congenital syphilis, ‘Clutton’s joints’ is a painless synovitis occurring at puberty. It most commonly affects the knee and elbow, mostly bilaterally.

**ACQUIRED SYMPHILIS**

The joints may be affected in the secondary and tertiary stages of acquired syphilis. In the secondary stage, transient polyarthritis and polyarthralgia involving the larger joints occur. In tertiary stage, gummatous arthritis occurs where the larger joints are most often involved. Neuropathic (Charcot’s) joint is an indirect consequence of syphilis. Please refer to a Medicine textbook for tests carried out for the diagnosis of syphilis.

**FUNGAL INFECTIONS**

Fungal infections of the bone occur usually in patients with depressed immunological status. The infection, particularly common in a rural population, is that of the foot, called ‘Madura foot’. As the infection results in a tumour-like mass, it is also called ‘Mycetoma’.

**MADURA FOOT**

This is caused by Maduromycosis. It starts as a nodular swelling over the dorsum or sole of the foot. The nodule bursts and discharges a thin pus. Gradually more nodules form and result in a swollen foot with a nodular surface and multiple discharging sinuses.

Pain is not a prominent feature, unless there is a secondary infection. The pus, characteristically contains small black granules, which on microscopic examination reveal the fungus. X-ray shows soft
tissue swelling around the foot bones. There may be multiple small sieve-like erosions in the bones of the foot.

**Treatment:** In early stages, the lesion responds to massive doses of penicillin or dapsone. In later stages, once the foot has become disorganised and there are multiple discharging sinuses, amputation may be necessary.

**LEPROSY AND ORTHOPAEDICS**

Leprosy is known in the society as a disease producing ugly deformities and mutilations. Deformities are seen in all types of leprosy, but are more common in tuberculoid and polyneuritic types.

**Mechanisms causing disability:** Nerve involvement leading to anaesthesia, dryness of the skin, and paralysis, is primarily responsible for deformity and disability of hands and feet. These factors predispose the affected limb to misuse, resulting in ulceration, scar formation and secondary infection. These, in turn, add to disability and create a vicious cycle whereby loss of deep tissue results. Flow chart-22.1 summarises the mechanism of disability.

Clinical manifestations of leprosy relevant from the viewpoint of orthopaedics are: (i) deformities; (ii) motor weakness and muscle atrophy; (iii) trophic ulcers; (iv) mutilations; and (v) neuritis.

**DEFORMITIES**

The primary factor responsible for deformities in leprosy is involvement of peripheral nerves, but secondary factors contribute to a large percentage of deformities. The latter are totally preventable, hence important. Some such factors are malpositioning of paralysed limbs, scarring and ulceration, self inflicted injuries to an anaesthetic part etc. Nerves commonly affected in leprosy are those in superficial locations. In order of frequency these are: ulnar nerve at the elbow, median nerve above the wrist and common peroneal nerve at the knee. Following are the common deformities seen:

- **Hand:** Common deformities in the hand are:
  - partial claw hand in ulnar nerve palsy, total claw hand in ulnar plus median nerve palsy, ape thumb deformity in median nerve palsy and wrist drop in radial nerve palsy.
- **Foot:** Foot drop occurs commonly due to involvement of common peroneal nerve.

**Treatment:** A great deal of deformities in leprosy are preventable, firstly by early detection of leprosy and adequate drug therapy; secondly by health education about hygienic care of anaesthetic foot, prevention of insect bites, proper splintage of paralysed part, and prompt and adequate care of trophic ulcers. Conservative treatment using splints, exercises and other physiotherapeutic measures are used in most cases. In some cases, surgical correction of the deformity is required. For details about methods of correction of deformities in general, please refer to Chapter 11.

**MOTOR WEAKNESS AND MUSCLE ATROPHY**

As a result of nerve involvement, commonly seen motor weakness are claw hand and wrist drop in the hand, and foot drop in the leg. Conservative treatment is by splints. Following reconstructive procedures may be performed in some cases.

- **For claw hand:** Paul Brand’s multi-tail tendon transfer.
- **For opponens weakness:** Opponensplasty, a tendon transfer operation where tendon of flexor digitorum superficialis of the ring finger is rerouted so that it passes through a pulley created at the flexor carpi ulnaris tendon, and is attached to the thumb.
- **For wrist drop:** Jone’s transfer.
- **For foot drop:** Transfer of tibialis posterior tendon on the dorsum of the foot.

**TROPHIC ULCERS**

These are found at anaesthetic sites, and are precipitated and perpetuated by recurrent injury or abnormal areas of pressure developing on paralysed hands and feet. Cause of injury could be mechanical, thermal etc. Common sites of trophic ulcers are heads of first and fifth metatarsals, heels and terminal phalanges of fingers. Early manifestation may be a spontaneous blister, a nodule or an injury at the anaesthetic site. This leads to ulcer formation, which may get secondarily infected. The ulcer may extend deep and affect soft tissues and bones, and become chronic and progressive. Causes responsible for chronicity of an ulcer are: (i) impeded vascular supply; (ii) repeated trauma to the ulcer; and (iii) superadded infection.

**Treatment:** Prevention of ulcer is most important, because once it occurs, healing takes a long time.
Treatment of an established ulcer consists of the following:

- **Eliminating stress caused by walking**, in acute stage, by resting the foot and in later stages by application of plaster cast.

- **Eradication of infection by**: (i) debridement; (ii) sequestrectomy; (iii) securing free drainage of the wound; (iv) antibiotics; and (v) occlusive dressings.

- **Other**: Besides debridement, the role of surgery is in using *plastic surgery procedures* to cover a large ulcer. *Amputation* may sometimes be considered necessary for a big, infected ulcer with osteomyelitis.

- **Prevention of recurrences** by protecting the foot and the scar from further injuries by good care of the part, proper footwear or splints, and careful use of the part (e.g., avoiding jumping, walking in cases with foot ulcers).

**MUTILATIONS**

Mutilations result from recurrent trophic ulcers, sequestration of bone and decalcification of bones. These are a result of ultimate neglect of a fairly treatable condition.
NEURITIS
Leprosy may result in acute or chronic neuritis. The patient complains of pain along the course of the nerve and later, neurological symptoms. In acute stage, rest to the part and anti-leprosy chemotherapy is given. Some surgeons prefer local or systemic corticosteroids in acute stage. In chronic cases, there is diffuse thickening of the nerves. Occasionally, a nerve abscess can be palpated. Indications for surgical intervention are: (i) abscess inside the nerve – in which case it is drained; and (ii) intractable pain in a person in whom paralysis in the distribution of the nerve is already present. Neurolysis has been attempted in these cases as no further harm can be done.

Further Reading

What have we learnt?
• *Staphylococcus aureus* is the commonest organism to cause bone and joint infection.
• Early diagnosis is crucial in acute osteomyelitis. Bone scan may be done in suspected cases.
• Early surgical drainage may prevent an acute osteomyelitis developing into chronic.
• Treatment of chronic osteomyelitis is essentially surgical. Giving prolonged antibiotics is of no use.
• In a case of suspected septic arthritis, aspiration of the joint is the best way to confirm the diagnosis. In case of a deep joint infection, ultrasound examination can help detect increased intra-articular fluid. Ultrasound guided aspiration can be done.
• In case of septic arthritis, early surgical drainage saves the joint from permanent damage.

Additional information: From the entrance exams point of view

- The earliest sign of osteomyelitis on X-ray is loss of soft tissue planes.
- The earliest bony change on X-ray is periosteal reaction.
- Evidence of osteomyelitis on X-ray occurs after 2 weeks of onset.
- Most common cause of post-surgical, post-traumatic and osteomyelitis of the spine is *Staphylococcus aureus*.
- Most common cause of osteomyelitis in drug abusers is *Pseudomonas aeruginosa*.
- Chondrolysis is seen in septic arthritis of infancy.
- Most common cause of bone and joint infection is haematogenous.

<table>
<thead>
<tr>
<th>Septic Arthritis</th>
<th>Transient synovitis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>0-5 yrs</td>
</tr>
<tr>
<td>ESR,WBC counts</td>
<td>Grossly elevated</td>
</tr>
<tr>
<td>Signs and Symptoms</td>
<td>More pronounced</td>
</tr>
</tbody>
</table>

https://kat.cr/user/Blink99/
AETIOPATHOGENESIS

Common causative organism is *Mycobacterium tuberculosis*. Bone and joint tuberculosis is *always* secondary to some primary focus in the lungs, lymph nodes etc. Mode of spread from the primary focus may be either haematogenous or by direct extension from a neighbouring focus.

Pathology: Tubercular infection of the bone and synovial tissue produces similar response as it produces in the lungs i.e., chronic granulomatous inflammation with caseation necrosis. The response may be proliferative, exudative or both;

a) *Proliferative response*: This is the commoner of the two responses. It is characterised by chronic granulomatous inflammation with a lot of fibrosis.

b) *Exudative response*: In some cases, particularly in immuno-deficient individuals, elderly people and people suffering from leukaemia etc., there is extensive caseation necrosis without much cellular reaction. This results in extensive pus formation. These are also termed *non-reactive* cases.

Natural history: Inflammation results in local trabecular necrosis and caseation. Demineralisation of the bone occurs because of intense local hyperaemia. In the absence of adequate body resistance or chemotherapy, the cortices of the bone get eroded, and the infected granulation tissue and pus find their way to the sub-periosteal and soft tissue planes. Here they present as *cold abscesses*, and may burst out to form sinuses. The affected bone may undergo a pathological fracture.
A tubercular osteomyelitis in the vicinity of a joint may result in the involvement of the joint.

Joint involvement is usually in the form of a low-grade synovitis, with thickening of the synovial membrane. Unlike pyogenic arthritis where proteolytic enzymes cause severe early destruction of the articular cartilage, tubercular infection causes slow destruction. Once the synovium is inflamed, it starts destroying the cartilage from the periphery. This inflammatory synovium at the periphery of the cartilage is called Pannus. Eventually, the articular cartilage is completely destroyed. The joint gets distended with the pus. Joint capsule and ligaments become lax, and the joint may get subluxated. Pus and tubercular debris burst out of the joint capsule to form a cold abscess, and subsequently a chronic discharging sinus.

Healing: It occurs by fibrosis, which results in significant limitation or near complete loss of joint movement (fibrous ankylosis). If considerable destruction of the articular cartilage has occurred, the joint space is completely lost, and is traversed by bony trabeculae between the bones forming the joint (bony ankylosis) as shown in Fig-23.1. Fibrous ankylosis is a common outcome of healed tuberculosis of the joints, except in the spine where bony ankylosis follows more often.

Clinical features depend upon the site affected. Patients of all ages and both sexes are affected frequently. The onset is gradual in most cases. Usual presenting complaints are pain, swelling, deformity and inability to use that part. Sometimes, the presentation is atypical. The following general principles will help in making a diagnosis:

a) High index of suspicion: Tuberculosis should be included in the differential diagnosis of any slow onset disease of the musculoskeletal system, particularly in countries where tuberculosis is still prevalent. Because of its slow onset and progress, the symptoms and signs are often minimal and non-specific. A high index of suspicion and a close watch over such symptoms in susceptible individuals, is the key to early diagnosis.

b) Fallacious history of trauma: Very often the patient assigns all his symptoms to an episode of injury. One should not get carried away by such information, as the injury may be coincidental. A detailed inquiry in such cases will reveal a symptom-free period between the episode of trauma and the beginning of symptoms, thus establishing the non-traumatic nature of the disease.

c) Lack of constitutional symptoms: Symptoms like fever, loss of appetite, weight loss etc. are present in only about 20 per cent cases. An active primary focus is detected only in about 15 per cent of cases at the time of diagnosis; in the rest it has already healed by the time the patient presents.

Specific signs and symptoms in patients with tuberculosis at major sites will be discussed in respective sections.

Investigations

Radiological examination: X-ray examination of the affected part, antero-posterior and lateral views, is the single most important investigation. Findings in the early stages may be minimal and are likely to be missed. A comparison with an identical X-ray of the opposite limb or with an X-ray repeated after some period, may be helpful. Following are some of the general radiological features of tuberculosis of the bones and joints:

TB osteomyelitis: A tubercular osteomyelitis presents as a well-defined area of bone destruction, typically with minimal reactive new bone formation. This is unlike a pyogenic infection, where reactive periosteal new bone formation is an important feature.
**TB arthritis:** In tubercular arthritis there is reduction of the joint space, erosion of the articular surfaces and marked peri-articular rarefaction. This is unlike many other causes of joint space reduction such as osteoarthritis, septic arthritis, etc., where there is subchondral sclerosis instead.

X-ray features specific to different sites will be discussed in their respective sections. A chest X-ray should be done routinely to detect any tubercular lesion in the lungs.

**Other investigations:** Some of the following investigations may be helpful in diagnosis:

- **Blood examination:** Lymphocytic leukocytosis, high ESR.
- **Mantoux test:** useful in children.
- **Serum ELISA test** for detecting anti-mycobacterium antibodies.
- **Synovial fluid aspiration** (see Table–22.3, page 177).
- **Aspiration of cold abscess and examination of pus for AFB.**
- **Histopathological examination** of the granulation tissue obtained by biopsy or curettage of a lesion.

**TREATMENT**

**Principles of treatment:** Treatment of tuberculosis of bones and joints consists of control of the infection and care of the diseased part. In most cases, conservative treatment suffices; sometimes operative intervention is required.

<table>
<thead>
<tr>
<th>Table–23.2: Common anti-tubercular drugs and their dosages</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Name/ Daily dose (max.)</th>
<th>Side effects</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Bactericidal</strong></td>
<td></td>
</tr>
<tr>
<td>Rifampicin (RF)</td>
<td>-Hepatotoxicity, pink coloured urine</td>
</tr>
<tr>
<td>10 mg/kg (600 mg)</td>
<td></td>
</tr>
<tr>
<td>Isoniazide (INH)</td>
<td>-Hepatotoxicity, Peripheral neuritis</td>
</tr>
<tr>
<td>5-10 mg/kg (300)</td>
<td></td>
</tr>
<tr>
<td>Streptomycin (SM)</td>
<td>-Vestibular damage, Nephrotoxicity Circumoral paraesthesia</td>
</tr>
<tr>
<td>30 mg/kg (1 gm)</td>
<td></td>
</tr>
<tr>
<td>Pyrazinamide (PZ)</td>
<td>-Hepatotoxicity</td>
</tr>
<tr>
<td>25 mg/kg (1.5 gm)</td>
<td></td>
</tr>
<tr>
<td><strong>Bacteriostatic</strong></td>
<td></td>
</tr>
<tr>
<td>Ethambutol (ETH)</td>
<td>-Optic neuritis, Colour blindness</td>
</tr>
<tr>
<td>25 mg/kg for 4 wks (1000 mg), thereafter</td>
<td></td>
</tr>
<tr>
<td>15 mg/kg (800 mg)</td>
<td></td>
</tr>
<tr>
<td>Cycloserine</td>
<td>-CNS toxicity -Headache, Tremor, Dysarthria</td>
</tr>
<tr>
<td>10 mg/kg (500 mg)</td>
<td></td>
</tr>
<tr>
<td>Ethionamide</td>
<td>-Anorexia, Nausea, Vomiting</td>
</tr>
<tr>
<td>25 mg/kg (750 mg)</td>
<td></td>
</tr>
<tr>
<td>Para-amino salicylate (PAS)</td>
<td>-Anorexia, Nausea, Vomiting</td>
</tr>
<tr>
<td>200-400 mg/kg (12 g)</td>
<td></td>
</tr>
</tbody>
</table>

**Control of infection:** It is brought about by potent anti-tubercular drugs, rest to the affected part and the building up of patient’s resistance.

a) **Anti-tubercular drugs:** Table–23.2 shows common anti-tubercular drugs, their dosage, route of administration and common side-effects. It is usual practice to start the treatment with 4 drugs — Rifampicin, INH, Pyrazinamide, Ethambutol for 3 months. In selected cases with multifocal tuberculosis, 5 drugs — RF, INH, PZ, ETH and Streptomycin, may be required for the initial period. The patient is monitored to detect any failure to respond or for any side-effects of the drugs.

b) **Rest:** The affected part should be rested during the period of pain. In the upper extremities this can be done with a plaster slab; in the lower extremities traction can be applied. In most cases of spinal tuberculosis bed rest for a short period is sufficient; in others, support with a brace may be necessary.

c) **Building up the patient’s resistance:** The patient should be given a high protein diet and exposed to fresh air and sunlight to build up his general resistance.

**Care of the affected part:** This consists of protection of the affected part from further damage, correction of any deformities and prevention of joint contractures. Once the disease is brought under control, exercises to regain functions of the joint are carried out. Care consists of the following:

a) **Proper positioning of the joint:** The joints should be kept in proper position so that contractures do not develop.

b) **Mobilisation:** As the disease comes under control and the pain reduces, joint mobilisation is begun. This prevents contractures and helps regain movement. In cases with extreme damage to the joint, it is best to expect ankylosis of the joint in the position of most useful function.

c) **Exercises:** As the joint regains movement, muscle strength building exercises are taught.

d) **Weight bearing:** It is started gradually as osteoporosis secondary to the disease is reversed.

* Multi-drug resistance is a serious upcoming problem. Anti-tubercular drugs, in proper combination, in proper dosages and under close supervision is the key to its prevention.
Operative intervention may be required in some cases. Following are some procedures commonly used:

a) Biopsy: For cases where the diagnosis is in doubt, a fine needle aspiration cytology (FNAC) may be performed from an enlarged lymph node or from a soft tissue swelling. An open biopsy may be necessary from a bony lesion, or in case FNAC fails to confirm the diagnosis.
b) Treatment of cold abscess: A small stationary abscess may be left alone as it will regress with the healing of the disease. A bigger cold abscess may need aspiration or evacuation (discussed in detail on page 191).
c) Curettage of the lesion: If the lesion is in the vicinity of a joint, infection is likely to spread to the joint. An early curettage of the lesion may prevent this complication.
d) Joint debridement: In cases with moderate joint destruction, surgical removal of infected and necrotic material from the joint may be required. This helps in the early healing of the disease, and thus promotes recovery of the joint.
e) Synovectomy: In cases of synovial tuberculosis, a synovectomy may be required to promote early recovery.
f) Salvage operations: These are procedures performed for markedly destroyed joints in order to salvage whatever useful functions are possible e.g., Girdlestone arthroplasty of the hip (page 198).
g) Decompression: In cases with paraplegia secondary to spinal TB, surgical decompression may be necessary.

Tuberculosis of the Spine
(Pott's disease)

The spine is the commonest site of bone and joint tuberculosis; the dorso-lumbar region being the one affected most frequently.

Relevant Anatomy

Development of a vertebra (Fig-23.2): A vertebra develops from the sclerotomes which lie on either side of the notochord. The lower-half of one vertebra and upper-half of the one below it, along with the intervening disc develop from each pair of sclerotomes and have a common blood supply. Therefore, infections via the arteries involve the

'Tuberculosis of Bones and Joints'
and spinal cord, the cord ends at the lower border of first lumbar vertebra. Beyond this, up to $S_2$, there is only the dural sac containing a bunch of nerve roots (cauda equina). The segment of the cord which corresponds to a given vertebra is therefore above the level of that vertebra. Relationship between the spinal segment and cord segment in different regions of the spinal column is as shown in Table–23.4, page 185.

**Pathology**

Like tuberculosis of the bones and joints elsewhere in the body, TB of the spine is always secondary. The bacteria reach the spine via the haematogenous route, from the lungs or lymph nodes. It spreads via the para-vertebral plexus of veins i.e., Batson's plexus, which has free communication with the visceral plexus of the abdomen, a common site of tuberculosis.

**Types of vertebral tuberculosis:** Lesions in the vertebrae may be of the following types (Fig-23.3):

a) **Parasidal:** This is the commonest type. In this, the contiguous areas of two adjacent vertebrae along with the intervening disc are affected.

b) **Central:** In this type, the body of a single vertebra is affected. This leads to early collapse of the weakened vertebra. The nearby disc may be normal. The collapse may be a ‘wedging’ or ‘concertina’ collapse (Fig-23.4); wedging being commoner.

c) **Anterior:** In this type, infection is localised to the anterior part of the vertebral body. The infection spreads up and down under the anterior longitudinal ligament.

d) **Posterior:** In this type, the posterior complex of the vertebra i.e., the pedicle, lamina, spinous process and transverse process are affected.

**Pathology:** Basic pathology is the same as that in other bone and joint tuberculosis. In the commoner paradiscal type, bacteria lodge in the contiguous areas of two adjacent vertebrae. Granulomatous inflammation results in erosion of the margins of these vertebrae. Nutrition of the intervening disc, which comes from the end-plates of the adjacent vertebrae is compromised. This results in disc degeneration, and as the process continues, complete destruction.

Weakening of the trabeculae of the vertebral body results in collapse of the vertebra. Type of collapse is generally a wedging, occurs early, and is severe in lesions of the dorsal spine. This is because, in the dorsal spine the line of weight bearing passes anterior to the vertebra, so that the anterior part of the weakened vertebra is more compressed than the posterior, resulting in wedging. In the cervical and
lumbar spines, because of their lordotic curvature (round forwards), wedging is less. Destruction occurs early, and is severe in children.

**Cold abscess:** This is a collection of pus and tubercular debris from a diseased vertebra. It is called a cold abscess because it is not associated with the usual signs of inflammation – heat, redness etc., found with a pyogenic abscess. The tubercular pus can track in any direction from the affected vertebra (Fig-23.5). If it travels backwards, it may press upon the important neural structures in the spinal canal. Pus may come out anteriorly (pre-vertebral abscess) or on the sides of the vertebral body (para-vertebral abscess). Once outside the vertebra the pus may travel along the musculo-fascial planes or neuro-vascular bundles to appear superficially at places far away from the site of lesion.

Healing: As healing occurs, the lytic areas in the bone are replaced by new bone. The adjacent vertebrae undergo fusion by bony-bridges. Whatever changes have occurred in the shape of the vertebral body are, however, permanent.

**CLINICAL FEATURES**

**Presenting complaints:** Clinical presentations of a case of TB of the spine is very variable – from a seemingly non-specific pain in the back to complete paraplegia. Following are some of the common presenting complaints:

- **Pain:** Back pain is the commonest presenting symptom. It may be diffuse; no more than a dull ache in the early stages, but later becomes localised to the affected diseased segment. It may be a ‘radicular’ pain i.e., a pain radiating along a nerve root. Depending upon the nerve root affected, it may present as pain in the arm (cervical roots), girdle pain (dorsal roots), pain abdomen (dorso-lumbar roots), groin pain (lumbar roots) or ‘sciatic’ pain (lumbo-sacral roots).

- **Stiffness:** It is a very early symptom in TB of the spine. It is a protective mechanism of the body, wherein the para-vertebral muscles go into spasm to prevent movement at the affected vertebra.

- **Cold abscess:** The patient may present the first time with a swelling (cold abscess) or problems secondary to its compression effects on the nearby visceral structures, such as dysphagia in TB of the cervical spine. A detailed examination in such cases reveals underlying TB of the spine.

- **Paraplegia:** If neglected, which is often the case in developing countries, a case of TB of the spine presents with this serious complication. For details see Pott's paraplegia, page 191.

- **Deformity:** Attention to TB of the spine may be attracted, especially in children, by a gradually increasing prominence of the spine – a gibbus.

- **Constitutional symptoms:** Symptoms like fever, weight loss etc., are rarely the only presenting symptoms.

**EXAMINATION**

The aim of examination is: (i) to pick up findings suggestive of tuberculosis of the spine; (ii) to localise the site of lesion; (iii) find skip lesions; and (iv) to detect any associated complications like cold abscesses or paraplegia. Following is the systematic way in which one should proceed to examine a case of suspected TB of the spine.

- **Gait:** A patient with TB of the spine walks with short steps in order to avoid jerking the spine. He may take time and may be very cautious while attempting to lie on the examination couch. In TB of the cervical spine, the patient often supports his head with both hands under the chin and twists his whole body in order to look sideways.

- **Attitude and deformity:** A patient with TB of the cervical spine has a stiff, straight neck. In dorsal spine TB, part of the spine becomes prominent (gibbus or kyphus*). Significant

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*Fig-23.5 Directions of tracking of tubercular pus from a vertebral focus*
deformity is generally absent in lumbar spine tuberculosis; there may just be loss of lumbar lordosis.

- **Para-vertebral swelling:** A superficial cold abscess may present as fullness or swelling on the back, along the chest wall or anteriorly. It is easy to diagnose because of its fluctuant nature. Sometimes, an abscess may be tense and it may not be possible to elicit fluctuation. A needle aspiration may be performed in such cases, to confirm the diagnosis. It is important to look for cold abscesses in not so obvious locations, depending upon the region of the spine affected (Table–23.5).

- **Tenderness:** It can be elicited by pressing upon the side of the spinous process in an attempt to rotate the vertebra.

- **Movement:** There is no necessity to examine for spinal movement in a patient with obviously painful spine. Spinal movement are limited in a case of TB of the spine, and can be tested, wherever considered suitable.

- **Neurological examination:** A thorough neurological examination of the limbs, upper or lower, depending on the site of tuberculosis should be performed. In addition to motor, sensory and reflexes examination, an assessment should be made of urinary or bowel functions. Aim of neurological examination is to find: (i) whether or not there is any neurological compression; (ii) level of neurological compression; and (iii) severity of neurological compression.

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**Table–23.5: Presentation of cold abscesses from different regions of the spine**

<table>
<thead>
<tr>
<th>Region of spine</th>
<th>Anteriorly</th>
<th>On the sides</th>
<th>Along musculo-facial plane</th>
<th>Along vascular bundle or arm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cervical spine</td>
<td>Retro-pharyngeal abscess</td>
<td>Para-vertebral abscess</td>
<td>At the posterior border of sterno-cleidomastoid muscle, in the posterior triangle of neck</td>
<td>To axilla, to arm along neuro-vascular bundle of the arm</td>
</tr>
<tr>
<td>Thoracic spine</td>
<td>Mediastinal abscess</td>
<td>Para-vertebral abscess</td>
<td>Trickles downward and enters either of the two lumbo-costal arches: • Lateral lumbo-costal arch –to present as lumbar abscess • Medial lumbo-costal arch –to present as psoas abscess</td>
<td>Along thoracic spinal nerves to present at • Anterior chest wall • Mid-axillary line • Posterior chest wall</td>
</tr>
<tr>
<td>Lumbar spine</td>
<td>Pre-vertebral abscess</td>
<td>Para-vertebral abscess</td>
<td>Lumbar abscess or psoas abscess lower</td>
<td>Along neurovascular bundle of the leg to present in groin or down in the leg</td>
</tr>
</tbody>
</table>

* There are three types of kyphotic deformities:
  (i) Knuckle – prominence of one spinous process
  (ii) Gibbus – prominence of two or three spinous processes
  (iii) Kyphus – diffuse rounding of the vertebral column

---

**Fig-23.6 X-ray findings in TB of the spine. (a) Early case, minimal loss of disc prolapse, (b) Complete loss of disc prolapse, (c) Destruction of vertebral bodies with loss of disc prolapse, (d) Advance destruction and wedging of vertebrae**
**General examination:** A general physical examination should be performed to detect any active or healed primary lesion. The patient may have some other systemic illness like diabetes, hypertension, jaundice etc., which may have a bearing on further treatment.

**Radiological investigations**

**X-ray examination:** One must specify the level of the suspected damage, when requisitioning an X-ray of the spine. Minimum of two views, AP and lateral, are necessary. A chest X-ray for primary focus or an X-ray of the abdomen – KUB, if a psoas abscess is suspected, may also be taken. Following are some of the important radiological features.

- **Reduction of disc space:** This is the earliest sign in the commoner, paradiscal type of tuberculosis (Fig-23.6a). In early stages, reduction in disc space may be minimal, and may be detectable only on comparing the height of the suspected disc with those above and below it. In advanced stages, disc space may be completely lost (Fig-23.6b). A lateral X-ray is better for evaluation of disc space. Reduction of disc space is an important sign because in other diseases of the spine e.g. secondaries in the spine, the disc space is well preserved.

- **Destruction of the vertebral body:** In early stages, the contiguous margins of the affected vertebrae may be eroded. The diseased, weakened vertebra may undergo wedging. In late stages, a significant part or whole of the vertebral body may be destroyed (Fig-23.6c), leading to angular kyphotic deformity. Severity of the deformity depends upon the extent of wedging and number of affected vertebrae (Table–23.6).

<table>
<thead>
<tr>
<th>Type</th>
<th>No. of vertebrae involved</th>
</tr>
</thead>
<tbody>
<tr>
<td>Knuckle</td>
<td>1</td>
</tr>
<tr>
<td>Gibbus</td>
<td>2-3</td>
</tr>
<tr>
<td>Angular kyphosis</td>
<td>3-4</td>
</tr>
<tr>
<td>Rounded kyphosis</td>
<td>&gt;4</td>
</tr>
</tbody>
</table>

- **Evidence of cold abscess:** Radiological evidence of a cold abscess is a very useful finding in diagnosing a case of suspected spinal TB. Following abscesses may be seen on X-rays:

- **Para-vertebral abscess:** A para-vertebral soft tissue shadow corresponding to the site of the affected vertebra in AP view indicates a para-vertebral abscess. It may be of the following types: (i) a fusiform para-vertebral abscess (bird nest abscess – an abscess whose length is greater than its width (Fig-23.7a); and (ii) globular or tense abscess – an abscess whose width is greater than the length (Fig-23.7b). The latter indicates pus under pressure and is commonly associated with paraplegia.

- **Widened mediastinum:** An abscess from the dorsal spine may present as widened mediastinum on AP X-ray.

- **Retro-pharyngeal abscess:** In cervical spine TB, a retro-pharyngeal abscess may be seen on a lateral X-ray. Normally, soft tissue shadow in front of the C3 vertebral body is 4 mm thick; an increase in its thickness indicates a retro-pharyngeal abscess (Fig-23.7c).
• Psoas abscess: In dorso-lumbar and lumbar tuberculosis, psoas shadow on an X-ray of the abdomen may show a bulge.

• Rarefaction: There is diffuse rarefaction of the vertebrae above and below the lesion.

• Unusual signs: In tuberculosis involving the posterior complex, there may be erosion of the posterior elements of pedicle, lamina etc. These are better visible on oblique X-rays of the spine. Anterior type of vertebral tuberculosis may show erosion of the anterior part of the body, much the same as that possibly seen sometimes in cases with aneurysm of aorta, thus termed aneurysmal sign. There may be lytic lesions in the ribs in the vicinity of the affected vertebra.

• Signs of healing: Once the disease starts healing, the density of the affected bones gradually improves. Areas surrounding the lytic lesion show sclerosis, and over a period of time these lesions are replaced by sclerotic bone. The adjacent vertebrae undergo bony fusion.

CT scan: It may detect a small para-vertebral abscess, not otherwise seen on plain X-ray; may indicate precisely the extent of destruction of the vertebral body and posterior elements; and may show a sequestrum or a bony ridge pressing on the cord (Fig-23.8). This is a very useful investigation in cases presenting as ‘spinal tumour syndrome’, where there may be no signs on plain X-rays.

MRI is the investigation of choice to evaluate the type and extent of compression of the cord (Fig-23.9). It also shows condition of the underlying neural tissues, and thus helps in predicting the prognosis in a particular case.

Myelography: This may be indicated in cases presenting with ‘spinal tumour syndrome’, or when the clinical level of neurological deficit does not correspond to the radiological level of the lesion.

Biopsy: CT guided needle biopsy, or an open biopsy may be required in a case with doubtful diagnosis.

Other general investigations: Investigations like ESR, Mantoux test, ELISA test for detecting anti-tubercular antibodies, chest X-ray, etc., to support the diagnosis of tuberculosis, may be carried out whenever required.

DIFFERENTIAL DIAGNOSIS

Cases with TB of the spine report fairly late in developing countries, so they present mostly with classic signs, symptoms and radiological features. In the early stages, and sometimes in some atypical presentations, diagnosis may be difficult. Some of the common differential diagnosis and their differentiating features are given in Table–23.7.

TREATMENT

Principles of treatment: Aim of treatment is: (i) to achieve healing of the disease; and (ii) to prevent, detect early, and treat promptly any complication like paraplegia etc. Treatment consists of anti-tubercular chemotherapy (page 184), general care (page 184), care of the spine, and treatment of the cold abscess. Only the latter two will be discussed here.

Care of the spine: This consists of providing rest to the spine during the acute phase, followed by guarded mobilisation.
Table–23.7: TB of the spine – Differential diagnosis

<table>
<thead>
<tr>
<th>Symptoms</th>
<th>Clinical features</th>
<th>Investigations</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Back pain</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Traumatic</td>
<td>History of trauma present</td>
<td>X-ray – disc height normal, Wedging of vertebrae present</td>
</tr>
<tr>
<td></td>
<td>No fever or abscess</td>
<td>No para-vertebral shadow seen</td>
</tr>
<tr>
<td>• Secondaries/Myeloma</td>
<td>History of ‘primary’ elsewhere or myeloma</td>
<td>Disc space normal</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Pedicles may be involved in secondaries</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Lesions in other bones present</td>
</tr>
<tr>
<td>• Prolapsed disc</td>
<td>Radiating pain</td>
<td>Normal X-rays</td>
</tr>
<tr>
<td></td>
<td>SLRT – positive</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Localised nerve root deficit</td>
<td></td>
</tr>
<tr>
<td>• Ank. spondylitis</td>
<td>Chronic back pain, starts in lower back</td>
<td>‘Bamboo spine’ appearance on X-rays</td>
</tr>
<tr>
<td></td>
<td>Diffuse morning stiffness</td>
<td>SI joints affected – hazy, fused</td>
</tr>
<tr>
<td></td>
<td>Chest expansion reduced</td>
<td></td>
</tr>
<tr>
<td><strong>Neurological deficit</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>• Spinal tumour</td>
<td>Present with gradually increasing neurologic deficit</td>
<td>X-ray – interpedicle space increased</td>
</tr>
<tr>
<td></td>
<td>No back pain, or other</td>
<td>Pedicle erosion present</td>
</tr>
<tr>
<td></td>
<td>findings on spine examination</td>
<td>CT myelogram confirms</td>
</tr>
<tr>
<td>• Traumatic</td>
<td>History of definite trauma present</td>
<td>X-ray suggestive of Fracture-dislocation</td>
</tr>
<tr>
<td></td>
<td>Weakness is sudden onset</td>
<td></td>
</tr>
<tr>
<td>• Secondaries in the spine</td>
<td>No history of trauma</td>
<td>X-ray shows erosion of vertebrae</td>
</tr>
<tr>
<td></td>
<td>Back pain present</td>
<td></td>
</tr>
<tr>
<td></td>
<td>History of ‘primary’ elsewhere</td>
<td></td>
</tr>
</tbody>
</table>

Rest: A short period of bed rest for pain relief may be sufficient during early stages of treatment. In cases with significant vertebral destruction, a longer period of bed rest is desirable to prevent further collapse and pathological dislocation of the diseased vertebrae. In children, a body cast is sometimes given, basically to force them to rest. Minerva jacket or a collar may be given for immobilising the cervical spine.

Mobilisation: As the patient improves, he is allowed to sit and walk while the spine is supported in a collar for the cervical spine, or an ASH brace for the dorso-lumbar spine. The patient is weaned off the brace once bony fusion occurs. He is advised to avoid sports for 2 years.

Treatment of cold abscess: A small cold abscess may subside with anti-tubercular treatment. Abscesses presenting superficially need treatment as discussed below;

Aspiration: A thick needle is required because often there is thick caseous material. It should be an anti-gravity insertion with the needle entering through a zig-zag tract.

Evacuation: In this procedure, the cold abscess is drained, its walls curetted, and the wound closed without a drain. This is unlike drainage of a pyogenic abscess, where a post-operative drain is always left. A psoas abscess can be drained extra-peritoneally using a kidney incision.

Medical Research Council of Great Britain conducted controlled trials to study various aspects of TB spine and published findings in four reports (1973-74). Their conclusions were that (i) bed rest is not necessary; (ii) Streptomycin is not necessary; (iii) PoP jacket offers no benefit; and (iv) debridement is not a good operation.

Complications
1. Cold abscess: This is the commonest complication of TB of the spine. Treatment is as discussed above.
2. Neurological compression: At times the patient presents as a case of spinal tumour syndrome; the first clinical symptom being a neurological deficit (discussed subsequently).

Pott's Paraplegia
(TB Spine with Neurological Involvement)

The incidence of neurological deficit has been reported to be 20 per cent. It occurs most commonly in tuberculosis of the dorsal spine because the spinal canal is narrowest in this part, and even
a small compromise can lead to a neurological deficit.

**PATHOLOGY**

This consists of pressure on the neural tissues within the canal by products from the diseased vertebrae. It could occur in the following ways:

- **Inflammatory oedema:** The neural tissues become oedematous because of vascular stasis in the adjacent diseased area.

- **Extradural pus and granulation tissue:** This is the commonest cause of compression on neural structures. The abscess formed around the diseased vertebrae may compress the neural structures from the front, much the same way as an extradural tumour.

- **Sequestra:** Devascularised bone and extruded disc material may be displaced into the canal.

- **Internal ‘gibbus’:** Angulation of the diseased spine may lead to formation of the bony ridge on the anterior wall of the spinal canal (Fig-23.10). This is called the internal gibbus.

**Types of Pott's paraplegia:** It can be divided into two types:

a) **Early onset paraplegia** i.e., paraplegia occurring during the active phase of the disease, usually within two years of onset of the disease.

b) **Late onset paraplegia** i.e., paraplegia occurring several years after the disease has become quiescent, usually at least two years after the onset of disease.

Pathology of the two types is different, as also is the prognosis. Table–23.8 gives the causes of neurological deficit in early and late onset paraplegia.

**CLINICAL FEATURES**

Neurological complications can occur in a known case of tuberculosis of the spine; or the case may present for the first time with a neurological deficit. In the latter, tuberculosis as the cause of paraplegia is detected only on examination and further investigation. Onset of paraplegia is gradual in most cases, but in some it is sudden. Tubercular paraplegia is usually spastic to start with. Clonus (ankle or patellar) is the most prominent early sign. Paralysis may pass with varying rapidity, through the following stages:

- **Muscle weakness**, spasticity and in-coordination due to pressure on the corticospinal tracts which are placed anteriorly in the cord and are probably more sensitive to pressure.

- **Paraplegia in extension:** Tone of the muscles is increased due to absence of normal corticospinal inhibition, resulting in paraplegia in extension.

- **Paraplegia in flexion:** Absence of paraspinal tract functions in addition to the corticospinal functions leads to paraplegia in flexion.

- **Complete flaccid paraplegia:** Paraplegia becomes completely flaccid once all transmission across the cord stops.

**Grades of Pott's paraplegia:** Potts' paraplegia has been graded on the basis of degree of motor involvement, into four grades (Goel, 1967):

**Grade I:** Patient is unaware of the neural deficit; the physician detects Babinski positive and ankle or patellar clonus on clinical examination.
Table–23.8: Causes* of paraplegia in TB of the spine

<table>
<thead>
<tr>
<th>Causes of paraplegia in TB of the spine</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Early onset paraplegia</strong></td>
</tr>
<tr>
<td>Inflammatory causes</td>
</tr>
<tr>
<td>• Abscess – commonest</td>
</tr>
<tr>
<td>• Granulation tissue</td>
</tr>
<tr>
<td>• Circumscribed tuberculous focus</td>
</tr>
<tr>
<td>• Posterior spinal disease</td>
</tr>
<tr>
<td>• Infective thrombosis of the spinal</td>
</tr>
<tr>
<td>blood supply</td>
</tr>
<tr>
<td>Mechanical causes</td>
</tr>
<tr>
<td>• Sequestrum in the canal</td>
</tr>
<tr>
<td>• Infected degenerated disc in the canal</td>
</tr>
<tr>
<td>• Pathological dislocation – a ridge of bone</td>
</tr>
<tr>
<td>pressing on the cord</td>
</tr>
<tr>
<td><strong>Late onset paraplegia</strong></td>
</tr>
<tr>
<td>• Recurrence of the disease</td>
</tr>
<tr>
<td>• Prominent anterior wall of the spinal canal in case of severe kyphosis (Internal gibbus)</td>
</tr>
<tr>
<td>• Fibrous septae following healing</td>
</tr>
</tbody>
</table>

* Although these several mechanisms have been described as acting separately to produce paraplegia, more than one cause may be responsible in a particular patient.

Grade II: Patient presents with complaints of clumsiness, in-coordination or spasticity while walking, but manages to walk with or without support.

Grade III: Patient is not able to walk because of severe weakness. On examination, he has paraplegia in extension. There may be partial loss of sensation.

Grade IV: Patient is unable to walk, and has paraplegia in flexion with severe muscle spasm. There is near complete loss of sensation with sphincter disturbances.

INVESTIGATIONS

It is usually possible to diagnose vertebral tuberculosis as a cause of paraplegia by typical radiological signs. In some cases, a MRI scan may be done to see: (i) type of vertebral destruction; (ii) presence of para-vertebral soft tissue abscess; and (iii) cause of paraplegia i.e., whether it is pus, sequestra etc. CT scan may be required in some cases to better evaluate the vertebral canal. MRI is the investigation of choice, wherever available.

TREATMENT

Principles of treatment: Aims of treatment are as follows:

a) To promote recovery of the affected neural tissues, by reversing the cause responsible for compression, either by drugs or by operation.

b) To achieve healing of the vertebral lesion, and to support the spine till the diseased segment becomes stable.

c) To undertake rehabilitative measures to prevent contractures, and to regain strength in the affected part.

Treatment of Pott’s paraplegia has been the topic of considerable study and discussion. Following is the treatment considered most acceptable in author’s opinion. Treatment may be divided into conservative and operative. All cases of Pott’s paraplegia must be treated under supervision, after admission to a hospital.

Conservative treatment: Anti-tubercular chemotherapy forms the mainstay of treatment. All patients are started on 4-drugs anti-tubercular chemotherapy as soon as the diagnosis is made. The spine is put to absolute rest by a sling traction for the cervical spine, and bed rest for the dorso-lumbar spine. The paralysed limbs are taken care of, as discussed in the Chapter 32. During treatment, repeated neurological examination of the limbs is carried out to detect any deterioration or improvement in the neurological status.

If paraplegia improves, conservative treatment is continued. Patient is allowed to sit in the bed with the help of a brace as soon as the spine has gained sufficient strength. Bracing is continued for a period of about 6 to 12 months.

Operative treatment: If paraplegia does not improve at a satisfactory rate, or if it actually deteriorates; surgical intervention is indicated. Following are the indications for surgery considered suitable in most centres.

Absolute indications

2. Paraplegia getting worse or remaining stationary despite adequate conservative treatment.
3. Severe paraplegia with rapid onset may indicate severe pressure from a mechanical accident or abscess.
4. Any severe paraplegia such as paraplegia in flexion, motor or sensory loss for more than six months, complete loss of motor power for one month despite adequate conservative treatment.

https://kat.cr/user/Blink99/
5. Paraplegia accompanied by uncontrolled spasticity of such severity that reasonable rest and immobilisation are not possible.

Relative indications
1. Recurrent paraplegia, even with paralysis that would cause no concern in the first attack.
2. Paraplegia with onset in old age: Indications for surgery are stronger because of the hazards of recumbency.
3. Painful paraplegia, pain resulting from spasm or root compression.
4. Complications such as urinary tract infection and stones.

Rare indications
1. Paraplegia due to posterior spinal disease.
2. Spinal tumour syndrome.
3. Severe paralysis secondary to the cervical disease.
4. Severe cauda equina paralysis.

Operative procedures for Pott’s paraplegia: The operative method aims at removal of the agents causing compression on the neural structures. The following operations are commonly performed:

a) **Costo-transversectomy** (Fig-23.11a): As the name suggests, this operation consists of the removal of a section of rib (about 2 inches), and transverse process. As this is done, sometimes liquid pus comes out under pressure. This is considered by some as a tense abscess relieved, and thus enough to decompress the neural tissues. It is indicated in a child with paraplegia, and when a tense abscess is visible on X-ray. In all other cases, it may not produce adequate decompression and an antero-lateral decompression may be necessary.

b) **Antero-lateral decompression (ALD):** This is the most commonly performed operation.

In this operation, the spine is opened from its lateral side and access is made to the front and side of the cord, thus it is called antero-lateral decompression. The cord is laid free of any granulation tissue, caseous material, bony spur or sequestrum pressing on it. Structures removed in order to achieve adequate exposure of the cord are; the rib, transverse process, pedicle and part of the body of the vertebra (Fig-23.11b). Lamina or facet joints are not removed, otherwise stability of the spine will be seriously jeopardized.

c) **Radical debridement and arthrodesis (Hongkong operation):** Wherever facilities are available, a radical debridement is performed by exposing the spine from front using trans-thoracic or trans-peritoneal approaches. All the dead and diseased vertebrae are excised and replaced by rib grafts. Advantage of this operation is early healing of the disease and no progress of the kyphosis.

d) **Laminectomy.** It is indicated in cases of spinal tumour syndrome, and those where paraplegia has resulted from posterior spinal disease.

Surgery for the cervical spine tuberculosis requires a separate technique; anterior decompression is preferable in this area.

**PROGNOSIS**

Prognosis of Pott’s paraplegia depends upon the following factors:
1. **Age:** Children respond to treatment better than adults.
2. **Onset:** Acute onset paraplegia has a better prognosis.
3. **Duration:** Long standing paraplegia has a worse prognosis.
4. **Severity:** Motor paralysis alone has a good prognosis. Sphincter involvement i.e., urinary or bowel incontinence are bad prognostic indicators.
5. **Progress:** Sudden progress of the paraplegia has a bad prognosis.

**TUBERCULOSIS OF THE HIP**

After spine, the hip is affected, most commonly. It usually occurs in children and adolescents, but patients at any age can be affected.
PATHOLOGY
The basic pathology is the same as that discussed on page 182. The usual initial lesion is in the bone adjacent to the joint i.e., either the acetabulum or the head of the femur (osseous tuberculosis). In some cases, the lesion may begin in the synovium (synovial tuberculosis), but quickly the articular cartilage and the bones are affected. A purely synovial tuberculosis, as seen in the knee joint, is uncommon in the hip. Common sites of initial bone focus in TB of the hip are as shown in Fig-23.12.

Natural history: The infected granulation tissue harbouring the bacilli, from the initial bony focus erodes the overlying cartilage or bone and reaches the joint. In early stage, this results in synovial hypertrophy and effusion. The pannus of hypertrophied synovium around the articular cartilage gradually extends over and under it. Cartilage is thus destroyed and the joint becomes full of pus and granulation tissue. Synovium gets thickened, oedematous, grey and ulcerated. Denuded of their protective cartilage, the bone ends become raw.

Multiple cavitation is typical of tuberculosis. Such cavities are formed in the femoral head and the acetabulum. Eventually, the head or the acetabulum gets partially absorbed. By the constant pull of the muscles acting on the hip, the remaining head of the femur may dislocate from the acetabulum onto the ilium, giving rise to the so-called wandering acetabulum (Fig-23.13). In later stages, pus bursts through the capsule and spreads in the line of least resistance. It may present as cold abscess in the groin or in the region of the greater trochanter. Pus may perforate the acetabulum and appear as a pelvic abscess.

Healing: If left untreated, healing may take place by fibrosis, leading to ankylosis of the hip usually in a deformed position (fibrous ankylosis).

CLINICAL FEATURES
Presenting complaints: The disease is insidious in onset and runs a chronic course. The child may be apathetic and pale with loss of appetite before definite symptoms pertaining to the hip appear. One of the first symptoms is stiffness of the hip, and it produces a limp. Initially, stiffness may occur only after rest, but later it persists all the time.

Pain may be absent in early stages, or if present, may be referred to the knee. The child may complain of ‘night cries’, the so called ‘starting pain’, caused by the rubbing of the two diseased surfaces, when movement occurs as a result of the muscle relaxation during sleep. Later, there may be cold abscesses around the hip or these may burst, resulting in discharging sinuses.

EXAMINATION
It should be carried out with the patient undressed. Following physical findings may be present:

- **Gait:** Lameness is one of the first signs. In the early stage, it is because of stiffness and deformity of the hip. Because of the flexion deformity at the hip, the child stands with compensatory exaggerated lumbar lordosis. While walking the hip is kept stiff. Forward–backward movement at the lumbar spine is used for propulsion of the lower limb. This is called the ‘stiff-hip gait’. Later the limp is exaggerated by pain so the child hastens to take the weight off the affected side. This is called the ‘painful or antalgic gait’.

https://kat.cr/user/Blink99/
• **Muscle wasting:** The thigh muscles and gluteal muscles are wasted.

• **Swelling:** There may be swelling around the hip because of a cold abscess.

• **Discharging sinuses:** There may be discharging sinuses in the groin or around the greater trochanter. There may be puckered scars from healed sinuses.

• **Deformity:** Gross deformities may be obvious on inspection. Minimal deformities are compensated for by pelvic tilt and can be made obvious by tests. Commonly it is flexion, adduction and internal rotation deformity of the hip. Method of measuring deformities is as discussed in Annexure-III.

• **Shortening:** There is generally a true shortening in TB of the hip, except in Stage I, in which an apparent lengthening occurs (Annexure III). Limb length discrepancy can occur at this joint not only because of actual shortening of the bones (true length) but also because of the adduction-abduction deformity, which results in pelvic tilt and thus affects the length of the limb (apparent length). The method of measuring true and apparent lengths is as discussed in Annexure-III. True and apparent length, and their relation to different deformities of the hip are given in Table-23.9.

• **Movements:** Both, active and passive movements are limited in all directions. An attempted movement is associated with muscle spasm.

There may be severe limitation of movements, both active and passive, in all directions in late cases of tuberculosis. This is called ankylosis of the hip. If there is no movement at all, it is bony ankylosis.

• **Abnormal position of the head:** In a dislocated hip, the head can be felt in the gluteal region.

• **Telescopy:** This test assesses the instability of the head if it is out of the acetabulum (details in Annexure-III).

**STAGES OF TB OF THE HIP**

TB of the hip has been arbitrarily divided into three stages in its clinical course (Fig-23.14).

**Stage I (stage of synovitis):** There is effusion into the joint which demands the hip to be in a position of maximum capacity. This is a position of flexion, abduction and external rotation. Since flexion and abduction deformities are only slight and are compensated for by tilting of the pelvis, these do not become obvious. The limb remains in external rotation. As the pelvis tilts downwards to compensate for the abduction deformity,
the affected limb appears longer (apparent lengthening), though on measuring true limb lengths, the two limbs are found to be equal. This stage is also called the *stage of apparent lengthening*. It lasts for a very short period. Very rarely does a patient present to the hospital in such an early stage of the disease.

**Stage II (stage of the arthritis):** In this stage, the articular cartilage is involved. This leads to spasm of the powerful muscles around the hip. Since the flexors and adductors are stronger muscle groups than the extensors and abductors, the hip takes the attitude of flexion, adduction and internal rotation. Flexion and adduction may be concealed by compensatory tilt of the pelvis but internal rotation of the leg is obvious. As the pelvis tilts upwards to compensate for the adduction, the affected limb appears shorter (apparent shortening), although on comparing the limb lengths in similar positions, the two limbs are equal. This is also called the *stage of apparent shortening*.

**Stage III (stage of erosion):** In this stage, the cartilage is destroyed and the head and/or the acetabulum is eroded. There may be a pathological dislocation or subluxation of the hip. Attitude of the limb is the same as that in Stage II i.e., flexion, adduction and internal rotation except for the fact that the deformities are exaggerated. There is *true shortening* of the limb because of the actual destruction of the bone. In addition, apparent length of the limb is further reduced because of the adduction deformity.

### Investigations

**Radiological examination:** An X-ray examination of the pelvis with both hips, AP and lateral views of the affected hip are essential. Inclusion of the normal hip in the same film on the AP view helps in comparing the joint spaces on the two sides. MRI scan and bone scan may be useful in early diagnosis. Some of the radiological signs in an established case of TB of the hip are as follows:

- **Haziness:** Haziness of the bones around the hip is the earliest sign. To appreciate it best, the affected hip is compared with the normal hip.
- **Lytic lesion:** There may be lytic lesions in the regions specified in Fig-23.12, on page 195).
- **Reduction of joint space:** This occurs because of destruction of the cartilage. It may be uniformly or irregularly diminished, better appreciated in the early stages on comparing it with the opposite side (Fig-23.15a).
- **Irregular outline:** The outline of the articular ends of the bone becomes irregular because of destruction by the disease process. In severe cases, a significant part of the head or acetabulum may be destroyed (Figs-23.15b and c).
- **Acetabular changes:** The head may be lying out of the acetabulum in a ‘pseudo’ acetabulum on the ilium – the *wandering acetabulum*. In some cases, the acetabulum simply gets enlarged and deepened with the deformed head shifted medially, giving the appearance of the ‘pestle and mortar’.
- **Signs of healing:** If the disease starts healing, there may be sclerosis around the hip.

![Radiological features of TB of the hip](https://kat.cr/user/Blink99/)

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*Fig-23.15  Radiological features of TB of the hip*
Other investigations: The investigations that can be carried out to confirm the diagnosis are as discussed on page 169.

Biopsy: It may be needed in some doubtful cases. This is done by exposing the hip by the posterior approach and taking a piece of the synovium for histopathological examination. It is possible to do an arthroscopic biopsy.

DIFFERENTIAL DIAGNOSIS

TB of the hip is the commonest cause of pain in the hip in children in countries where TB is still prevalent. Following differential diagnosis should be considered:

a) Other causes of monoarthritis of the hip:
Subacute low grade monoarthritis due to low grade septic infection or rheumatoid arthritis also presents with pain and stiffness of the hip. Lack of supportive evidence for TB (like positive family history, past history) and destruction and sclerosis on X-ray, favour a diagnosis of septic arthritis. It may sometimes be difficult to differentiate the two. In rheumatoid arthritis, joint space is uniformly reduced.

b) Inguinal lymphadenopathy or psoas abscess:
Patients with these extra-articular diseases often present with a flexion deformity of the hip because of spasm of the iliopsoas. An examination reveals that all movements of the hip except extension are pain free.

c) Other diseases of the hip presenting at that age: In a child presenting with a limp without much pain the following conditions should be considered:

- Congenital dislocation of the hip: The limp is painless. It can generally be detected at birth, but is often noticed only when the child starts walking. An abnormal femoral head can be felt in the gluteal region. Telescopy test is positive. X-rays are decisive.
- Congenital coxa vara: The limp is painless. The movements limited are abduction and internal rotation. In fact, adduction and external rotation may be increased. X-ray examination usually confirms the diagnosis.
- Perthes’ disease: This occurs in children in the age group of 5-10 years. The main complaint is a limp, which is generally painful. There is minimal limitation of movement, mainly of abduction and internal rotation. Little or no shortening is present. Typically, X-ray changes are out of proportion to the physical findings. The joint space, unlike in TB of the hip, may even be widened (for details see page 318).

d) Osteoarthritis: This occurs in older individuals. Hip movements are limited in all directions but only terminally. There is associated pain and crepitus. Most cases are of osteoarthritis secondary to some other pathology (see Chapter 35).

TREATMENT

Principles of treatment: It is to control the disease activity, and to preserve joint movement. In early stages (Stages I and II), it is possible to achieve this by conservative treatment. In later stages (Stage II and after), significant limitation of joint functions occur despite best treatment. Treatment may be conservative or operative.

Conservative treatment: It consists of antitubercular chemotherapy (page 184) and care of the hip.

- Care of the hip: The affected hip is put to rest by immobilisation using below-knee skin traction. In addition to providing pain relief, this also corrects any deformity by counteracting the muscle spasm.
- General care - Same as on page 184.

Operative treatment: The following operative procedures may be indicated in TB of the hip (see plan of treatment on page 200).

- Joint debridement: The joint is opened using posterior approach. Pus, necrotic tissue, inflamed synovium and dead cartilage are removed from the joint. Any cavities in the head of the femur or acetabulum are curetted. The joint is washed thoroughly with saline and the wound closed. Post-operatively the joint surfaces are kept apart by traction to the leg. After the wound heals, the joint is mobilised.
- Girdlestone arthroplasty: The hip joint is exposed using the posterior approach. Head and neck of the femur are excised (Fig-23.16). Dead necrotic tissues and granulation tissues are excised. Post-operatively, bilateral skeletal traction is given for 4 weeks, followed by mobilisation of the hip. It is possible to regain reasonable movement of the
Tuberculosis of Bones and Joints

hip by this procedure even in severely damaged joints.

- **Arthrodesis:** In selected cases, where a stiff hip in a functional position is more suitable considering day-to-day activities of the patient, it is produced surgically by knocking the joint out.

- **Corrective osteotomy:** Cases where bony ankylosis of the hip has occurred in an unacceptable position from the functional viewpoint, a subtrochanteric corrective osteotomy of the femur may be required.

- **Total hip replacement:** There is enough evidence now, that a total hip replacement is a useful operation in some patients with quiescent tuberculosis. But as of now in most Afro-Asian countries, where most cannot afford a total hip replacement, and where most patients want to be able to squat even at the cost of instability, an excision arthroplasty is a preferred option.

**Deciding the plan of treatment:** In early stages, ATT and skin traction is given. As the disease comes under control, as is evident from the relief of symptoms; joint mobilisation is begun. By physiotherapy good joint functions can be regained in most cases. In cases presenting in late stages, initial treatment is by ATT and below-knee skin traction. The traction keeps the hip in a functional position with the joint surfaces apart while healing occurs. As the disease activity comes under control and symptoms (pain etc.) subside, a decision has to be made whether useful hip functions can be regained, depending upon the X-ray appearance of the hip. In a case, where there is no or minimal destruction of the hip joint mobilisation is begun with the hope of regaining as much movement as possible. In a case where the X-ray picture suggests significant joint damage or subluxation, one expects that normal joint functions cannot be regained. In such a situation, the options before the surgeon are essentially these:

a) To provide a **painless, mobile but unstable joint** by an excision arthroplasty (Girdlestone arthroplasty). Though the hip becomes unstable and the limb short, one can expect that patient will be able to squat on the floor.

b) To provide a **painless, stable but fixed joint** by surgically fusing the joint (arthrodesis) or by conservative means. Though the hip is stable, lack of movement, and thus an inability to squat is the major problem.

In countries, where for most of day-to-day activities squatting is required, Girdlestone arthroplasty is still considered a suitable operation. In addition to providing a mobile hip, this operation enhances healing of the disease as physical removal of the infected bone, synovium etc. is done. In selected patients, joint debridement has resulted in a stable hip with reasonable mobility of the hip. A total hip replacement has also been advocated in some cases of healed TB.

Flow chart-23.1 shows a general plan of treatment for TB of the hip.

**Tuberculosis of the Knee**

The knee is a common site of tuberculosis. Being a superficial joint, early diagnosis is usually possible. A delay in diagnosis can severely compromise joint functions.

**Pathology**

The basic pathology is same as that described on page 182. The disease may begin in the bone (osseous tuberculosis), usually in the femoral or tibial condyles, or more rarely in the patella. More commonly, the disease begins in the synovial membrane (synovial tuberculosis), leading to hypertrophy of the synovium. In early stages, the disease may be confined to the synovium without significant damage to the joint.

**Natural history:** In later stages, the articular cartilage and bone are destroyed irrespective of the site of origin. In all types, there occurs synovial hypertrophy, synovial effusion and pus formation in the joint. The hypertrophied synovium spreads
under and over the cartilage and destroys it. The cartilage may become detached, leaving the bone exposed. Long standing distension of the joint and destruction of the ligaments produces subluxation of the tibia. The tibia flexes, slips backwards and rotates externally on the femoral condyles (triple subluxation). Pus may burst out of the capsule to present as a cold abscess, and subsequently a sinus. **Healing:** If untreated, nature’s attempt at healing may result in fibrosis, and thereby stiffness of the joint in a deformed position. Healing is by fibrosis (fibrous ankylosis).

**CLINICAL FEATURES**

**Presenting complaints:** The patient, usually in the age group of 10-25 years, presents with complaints of pain and swelling in the knee. It is gradual in onset without any preceding history of trauma. Subsequently, pain increases and the knee takes an attitude of flexion. The patient starts limping. There is severe stiffness of the knee.

**EXAMINATION**

Following findings may be present on examination:
• **Swelling**: The joint is swollen, which may be due to synovial hypertrophy or effusion. The same can be detected by tests, as discussed on pages 189 and 190.

• **Muscle atrophy**: Atrophy of the thigh muscles is more than what can be accounted for by disuse alone. This is an unexplained feature of joint tuberculosis.

• **Cold abscess**: There may be swelling due to a cold abscess, either around the knee or in the calf.

• **Sinus**: There may be discharging or healed sinuses.

• **Deformity**: In early stages, there is a mild flexion deformity of the knee because of effusion in the knee, and muscle spasm. Later, triple displacement (flexion, posterior subluxation and external rotation) occurs due to ligament laxity.

• **Movements**: The movements at the joint are limited. There is pain and muscle spasm on attempting movement.

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**INVESTIGATIONS**

**Radiological examination** (Fig-23.17): X-ray is essentially normal in a case of synovial tuberculosis, except for a soft tissue shadow corresponding to the distended knee. The joint space may be widened. There is diffuse osteoporosis of the bones around the joint. In osseous tuberculosis, one may see juxta-articular lytic lesions. The joint surfaces may be eroded. In later stages, joint space may be diminished or completely lost. In advanced stages, triple subluxation with cavitory bone lesions may be present.

**Other investigations** along the lines already discussed on page 183 may be carried out. A biopsy is sometimes required.

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**DIFFERENTIAL DIAGNOSIS**

Diagnosis is not difficult in a late case, but when the patient presents with synovitis, other causes of synovitis should be excluded before arriving at a diagnosis of TB of the knee. These include subacute pyogenic infection, mono-articular rheumatoid arthritis, chronic traumatic synovitis, rheumatic arthritis and haemophilic arthritis.

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**TREATMENT**

**Principles of treatment**: Aim of treatment is to achieve, wherever possible, a painless mobile joint. This is possible if a patient has come early for treatment. In later stages, some amount of pain and stiffness persist in spite of treatment.

**Conservative treatment**: This consists of antitubercular chemotherapy, general care and local care of the part affected. It is started in all cases and decision for surgery taken if indicated, as discussed later.

• **Care of the knee**: The knee is rested by applying below-knee skin traction or an above-knee PoP slab. This helps in the healing process, and also takes care of the associated muscle spasm which keeps the knee in a deformed position.

**Operative treatment**: Following operative procedures may be required in suitable cases:

• **Synovectomy**: It may be required in cases of purely synovial tuberculosis. Very often one finds ‘melon seed’ bodies within the joint.

• **Joint debridement**: This may be required in cases where the articular cartilage is essentially preserved. The pus is drained, the synovium excised, and all the cavities curetted.

• **Arthrodesis**: In advanced stages of the disease with triple subluxation and complete cartilage destruction, the knee is arthrodesed in
functional position, i.e., about 5-10° of flexion and neutral rotation. One popular method of knee arthrodesis is Charnley’s compression arthrodesis.

With the current state of development of surgery, all these operations can be performed by minimally invasive arthroscopic surgery.

**Plan of treatment:** A plan of treatment for a case of TB of the knee is similar to that of the hip and is shown in Flow chart-23.2.

**TUBERCULOSIS OF OTHER JOINTS**
Other joints uncommonly affected by tuberculosis are the elbow, shoulder and ankle joints. Clinical features are similar to tuberculosis of other joints. Diagnosis is generally possible by X-ray examination. Occasionally a biopsy may be required.

Shoulder joint tuberculosis, at times, may not produce any pus etc., and hence is called ‘caries sicca’ and should always be considered in

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**Flow chart-23.2 Treatment plan for TB knee**
subacute osteomyelitis of the shaft. Very often these are multifocal lesions. An important radiological feature is that the bone lysis is out of proportion to the new bone formation, unlike in pyogenic osteomyelitis. A biopsy may be necessary in some cases.

**Tuberculosis of short bones:** Tuberculosis of the small bones of the feet and hands is a rather common entity. These may occur as isolated lesions or as multiple ones. Diagnosis is easy with X-ray. Calcaneum is a common site (Fig-23.18). Treatment is by rest and ATT. In cases where diagnosis is in doubt, curettage of the lesion and histopathological examination of the curetted material may be required.

**Spina ventosa:** This is a name given to tuberculosis of the phalanges of hand. The affected phalanx swells up like a balloon. An X-ray typically shows a lytic lesion distending the phalanx, and a lot of new bone formation.

**Further Reading**

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**What have we learnt?**
- Bone and joint tuberculosis constitute 1-4 percent of total number of cases of tuberculosis.
- Systemic features of infection, such as fever, do not occur commonly in bone and joint tuberculosis.
- Spine is the commonest site of bone and joint tuberculosis.
- Dorsolumbar spine is the commonest region to get affected; Paradiscal being the commonest type.
- Reduction of disc space is the earliest radiological sign of TB spine.
- It is the neurological complications associated with spine TB, which are of serious concern.
- TB hip is common cause of monoarthritis in children. Early diagnosis can save the joint from developing ankylosis.
- TB knee is of two types: Synovial and articular.
- Joints gets affected early, if there is a juxta-articular tubercular osteomyelitis.
- Treatment is focussed on (a) control of the disease, (b) functional recovery.
- Healing of tuberculous arthritis occurs by fibrous ankylosis.
- TB osteomyelitis is common in small bones.
<table>
<thead>
<tr>
<th>Additional information: From the entrance exams point of view</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Tuberculosis most commonly affects the dorsolumbar region of the spine, mainly T12 L1 junction.</td>
</tr>
<tr>
<td>• The earliest feature of tuberculosis is pain.</td>
</tr>
<tr>
<td>• Tuberculosis of the spine affects the vertebral body first.</td>
</tr>
<tr>
<td>• Tuberculosis with polyarthritis is called Poncet’s disease.</td>
</tr>
<tr>
<td>• Hong Kong operation is done in TB.</td>
</tr>
<tr>
<td>• Poor prognostic factors in Pott’s paraplegia are acute onset, sudden progression and long standing paraplegia.</td>
</tr>
<tr>
<td>• Most common sequelae of TB spondylitis is bony ankylosis.</td>
</tr>
<tr>
<td>• First sign of TB spin is loss of curvature of the spine followed by reduction in intervertebral space radiologically.</td>
</tr>
<tr>
<td>• Most common cause of early onset paraplegia is cold abscess.</td>
</tr>
<tr>
<td>• Triple deformity is seen in TB knee and is characterized by posterior subluxation of the knee, external or lateral rotation of the tibia and flexion of the knee. It is treated by antitubercular drugs with replacement or arthrodesis.</td>
</tr>
</tbody>
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Infections of the Hand

**TOPICS**

- Classification
- Aetiopathology
- Acute paronychium
- Apical subungual infection
- Terminal pulp space infection

**CLASSIFICATION**

Infections of the hand can be classified into two broad categories; spreading infections and localised infections. **Spreading infections** are the ones which spread to involve a large area of the hand e.g., lymphangitis, cellulitis etc. **Localised infections** are those which are localised to an area of the hand because of certain anatomical factors. Infections of the hand are classified as given in Table–24.1.

**AETIOPATHOLOGY**

Hand infections are common in manual workers and housewives who frequently suffer small pricks or abrasions in the course of their work. *Staphylococcus aureus* is the causative organism in 80 per cent of cases; in others *Streptococcus* and other gram-negative bacteria are responsible. The organisms reach the tissue planes by direct implantation from outside or via the blood. They set up an acute inflammatory reaction, which in many cases progresses to suppuration. Without effective treatment, the infection may spread to adjacent tissue planes.

**ACUTE PARONYCHIUM**

Paronychium is an infection of the nail fold. It is the *commonest* infection of the hand, and usually results from careless nail paring or use of unsterile manicure instruments.

**Clinical Features:** There is pain, tenderness, redness and swelling at one or both sides of the nail fold, and at the base of the nail if suppuration has extended deep to the nail. There is a marked tenderness on pressing the nail.

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![Fig-24.1 Incision for draining acute paronychium](https://kat.cr/user/Blink99/)
**Treatment:** In its early stage, when no suppuration has occurred, conservative treatment may abort the infection. Once suppuration has occurred, the pus must be let out. For a mild infection, it is sufficient to raise the cuticle alone without incising, but better drainage is secured by vertical incision through the cuticle on one or both sides (Fig-24.1). When the pus extends beneath the nail, it is necessary to remove the proximal one-third of the nail for adequate drainage.

**Complications:** These are: (i) extension of the infection to the pulp space; and (ii) chronic paronychium.

**APICAL SUBUNGUAL INFECTION**

This is an infection of the tissues between the nail plate and the periosteum of the terminal phalanx. It results from a pin-prick or splinter beneath the nail. The lesion is excruciatingly painful with little swelling. Tenderness is maximum just beneath the free edge of the nail. The pus comes to the surface at the free edge of the nail.

**Treatment:** In the early stage, the infection can be aborted by conservative treatment, but once suppuration occurs, drainage is required. For drainage, a small V-shaped piece is removed from the centre of the free edge of the nail along with a little wedge of the full thickness of the skin overlying the abscess (Fig-24.2).

**Complications:** Pus may spread under the nail, and may lead to a chronic sinus. Occasionally, the tip of the phalanx becomes infected.

**TERMINAL PULP SPACE INFECTION**

(Whitlow or Felon)

**Surgical anatomy:** The terminal pulp space is the volar space of the distal digit. It is filled with compact fat, feebly partitioned by multiple fibrous septae (Fig-24.3). At its proximal end, this space is closed by a septum of deep fascia connecting the distal flexor crease of the finger to the periosteum just distal to the insertion of the profundus flexor tendon. The digital artery, before it enters the space gives a branch to the epiphysis at the base of the distal phalanx. On entering the space it divides into terminal branches.

**Clinical features:** This is the second most common infection of the hand, commonly resulting from a pin-prick. The index finger and the thumb are affected most often. The pulp is swollen, tense and tender. A severe throbbing pain and excruciating tenderness suggest suppuration.

**Treatment:** In its early stage, conservative treatment may abort the infection. In the later stage, when suppuration has occurred, drainage is required. This is achieved either by incising directly over the centre of the abscess where it is pointing, or by a lateral incision just in front of the plane of the terminal phalanx.

**Complications:** These are: (i) osteomyelitis of the terminal phalanx, often with necrosis and sequestration of its distal half. Thrombo-arteritis of the terminal branches of the digital vessels accounts for this. The basal plate of the epiphysis is rarely involved; (ii) pyogenic arthritis of the distal interphalangeal joint; and (iii) very rarely, infection spreads to the flexor tendon sheath (suppurative tenosynovitis).

**MIDDLE VOLAR SPACE INFECTION**

**Surgical anatomy:** The middle volar space extends from the proximal to the distal volar creases of the finger. It is filled with loosely packed fibro-fat tissue.
**Clinical features:** It commonly results from a pin-prick. Pain, swelling, and tenderness are maximally localised to this space. The finger is kept in semi flexion. Frequently a purulent blister appears in the distal flexor crease. In early cases, it may be difficult to distinguish this infection from infection of the underlying flexor tendon sheath. However, in the former, tenderness over the proximal end of tendon sheath – at the base of the finger, is lacking.

**Treatment:** In the early stage, conservative treatment is enough. In late stage, drainage of pus via a longitudinal incision on the lateral side is performed.

**Complications:** Infection may spread to the distal or the proximal volar spaces, into the inter-phalangeal joints, or into the synovial sheath of the flexor tendons.

**PROXIMAL VOLAR SPACE INFECTION**

**Surgical anatomy:** This space is well partitioned from the middle volar space, but it communicates freely with the corresponding web space.

**Clinical features:** It is usually a consequence of a pin-prick. Pain, swelling and tenderness are localised to the space. Often the swelling is asymmetrical because of the concomitant involvement of the web space.

**Treatment:** The abscess is drained by an incision on the lateral side or at the point of maximum tenderness.

**WEB SPACE INFECTION**

**Surgical anatomy:** The web space is the triangular space between the bases of adjacent fingers; the first one being between the thumb and the index finger.

**Clinical features:** The infection arises: (i) from a skin crack; (ii) from a purulent blister on the forepart of the hand; or (iii) from a proximal volar space infection which communicates with the web space through the lumbrical canal (the canal that carries the lumbrical tendon from the hand into the finger). In the early stage, before localisation of infection occurs, there is oedema over back of the hand. Although, the condition is strongly suspected by the location of the tenderness, a precise diagnosis is often difficult at this stage. Once localisation has occurred, signs of web space infection manifest themselves. The swelling at the base of the finger becomes obvious. In severe cases, the finger immediately adjacent to the space is ‘separated’ because of the mechanical effect of the abscess. Maximum tenderness is found in the web and on the volar surface of the base of the finger.

**Treatment:** In the early stage, conservative treatment may abort the infection. In late stage, drainage of the pus is required. The web space abscess is drained by a transverse incision on the palmar surface over the affected web space. Care needs to be taken to deepen the incision cautiously until the subcutaneous fat is reached. Only a few strands of palmar fascia need to be divided, and if pus does not flow, it is sought with a probe or a dissector. The edges of the wound are cut away so as to leave a diamond-shaped opening. When the abscess communicates with a dorsal pocket, a counter-incision is advisable on the dorsum of the hand.
**Complications:** Spread of the infection to the nearby spaces and tendon sheaths is a common complication.

**DEEP PALMAR ABSCESS**

An abscess beneath the palmar fascia is a serious but rare infection of the hand. It may be an infection in the thenar* or mid-palmar space.

**Surgical anatomy:** The deep palmar spaces of the hand lie in the hollow of the palm, deep to the flexor tendons and their synovial sheaths. The space is divided into two halves – a medial half (the mid-palmar space), and a lateral half (the thenar space). The posterior relation of the space is formed by the fascia covering the interossei and metacarpal bones on the medial side, and the adductor pollicis muscle on the lateral side. On the two sides of the space are the thenar and hypothenar muscles (Fig-24.4).

**Clinical features:** The infection can arise from a penetrating wound, via the blood stream, or as a complication of suppurative tenosynovitis. At an early stage, there is an intense throbbing pain, and deep tenderness in the palm. There is only little swelling on the volar aspect of the palm; rather it is severe on the dorsum of the hand, sometimes so great as to give rise to what is called ‘frog hand’. The fingers are kept flexed. Extension at the metacarpo-phalangeal joints is very painful, but painless at the inter-phalangeal joints. This distinguishes this condition from suppurative tenosynovitis where there is pain on extending the metacarpo-phalangeal as well as inter-phalangeal joints. Regional lymphadenopathy is commonly present.

As tension within the space mounts, the normal concavity of the palm becomes flattened. Subsequently, the pus erodes through the palmar fascia and the intense pain eases off. The palm now becomes slightly convex, and it may be possible to elicit fluctuation only at this stage of the infection.

**Treatment:** It is often difficult to diagnose deep palmar space infection in its early stages because of its deep location. In some cases, treatment by conservative methods may abort the infection, but more often suppuration follows. The pus, being in a deeper plane, is difficult to detect because of the lack of fluctuation. A strong suspicion and a throbbing pain are indications of deep seated pus requiring drainage. A needle aspiration may be helpful in confirming the presence of pus.

A central transverse incision is made in the line of the flexor crease, passing across the middle of the palm at the site of maximum tenderness. If pus is encountered beneath the aponeurosis, the floor of the abscess must be probed systematically for a sinus leading to a deeper plane. In order to ensure free drainage of the pus, the skin edges as well as those of the palmar fascia are trimmed.

**Complications:** Spread of infection to nearby spaces and tendon sheaths may occur. A chronic infection may result in a discharging sinus and stiffness of the hand.

**ACUTE SUPPURATIVE TENOSYNOVITIS**

This is a rare but important infection because prompt treatment is essential if the function of the finger is to be preserved.

**Surgical anatomy:** The flexor tendons of the hand are covered with fibrous and synovial flexor sheaths (Fig-24.5). The fibrous sheaths exist only up to the bases of the digits. A synovial sheath lines the fibrous sheaths. In the thumb and little finger, the synovial lining extends proximally through the palm and ends 2–3 cms above the wrist. The synovial sheaths of the index, middle

* Thenar space is not same as the space containing thenar muscles See Fig. 26.4

![Fig-24.5 Flexor tendon sheaths](image-url)
Infections of the Hand

flexor digitorum superficialis and flexor digitorum profundus.

It should be noted that there is normal but great anatomical variation in the arrangement of the synovial tendon sheaths. The ulnar and radial bursae may communicate. The tendon sheaths of the index, middle and ring fingers may communicate with the ulnar bursa.

**Clinical features:** The bacteria enter the tendon sheath with the point of a needle or other sharp objects penetrating the tendon sheath. *Exceptionally,* the sheath is infected by extension from the terminal pulp space infection.

The finger is swollen throughout its length, and is acutely tender over the flexor tendon sheath.

It is held semi-flexed, and active or passive extension at the inter-phalangeal joint is very painful. In tenosynovitis of the little finger, the ulnar bursa also becomes involved, giving rise to swelling of the palm and sometimes fullness immediately above the flexor retinaculum. The area of maximum tenderness in an ulnar bursa infection can be elicited over that part of the bursa lying between the transverse palmar creases. This is *Kanavel’s sign* (Fig-24.6). In infections of the radial bursa, there is more swelling over the thenar eminence and thumb. The other findings are similar to other tendon sheath infections.

**Treatment:** An aggressive conservative treatment is started at an early stage. Clinical re-examinations are carried out every 6 hours to assess improvement. Conservative treatment is continued only so long as there is good local and general response. Any delay in decompression leads to a spread of the infection proximally into the forearm.

**Complications:** These are: (i) permanent stiffness of the finger in semi-flexion because of necrosis of the tendons and adhesions between the tendon and the sheath; and (ii) spread of the infection to nearby structures such as the ulnar and radial bursae.

**Further Reading**

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**What have we learnt?**
- Hand infections are of two types: Superficial and Deep.
- Early surgical intervention is preferred.
- Stiffness is a common complication.

**Additional information:** From the entrance exams point of view
- Felon most commonly affects the thumb.
- Kanavel’s sign is seen in tenosynovitis, it consists of eliciting tenderness on percussion over the flexor tendon sheath of the finger, flexion posture of fingers with pain on hyprextension and uniform swelling involving the entire finger.
Chapter 25

Congenital Talipes Equino Varus (CTEV)

Topics

- Relevant anatomy
- Pathoanatomy
- Nomenclature
- Clinical features
- Aetiology
- Treatment

‘Clubfoot’ is a rather vague term which has been used to describe a number of different abnormalities in the shape of the foot, but over the years it has come to be synonymous with the commonest congenital foot deformity i.e., Congenital Talipes Equino Varus (CTEV). It occurs once in every 1000 live births.

Relevant Anatomy

The joints of the foot relevant to understanding of this chapter are: (i) the ankle joint between the tibia and the talus; (ii) the subtalar joint between the talus and the calcaneum; (iii) the talo-navicular joint; and (iv) the calcaneo-cuboid joint (Fig-25.1).

The ligaments related to the aetiology of clubfoot are as follows (Fig-25.2):

- Deltoid ligament: This is the medial collateral ligament of the ankle. It has a superficial and a deep component.
- Spring ligament: This is a ligament which joins the anterior end of the calcaneum to the navicular.
- Interosseous ligament: This ligament is between the talus and calcaneum, joining their apposing surfaces.
- Capsular ligaments: The thickened portions of the capsule of the talo-navicular, naviculo-cuneiform, and cuneiform-metatarsal joints, termed as the capsular ligaments, are important structures in pathology of CTEV.
Mechanical theory:

Genetic theory:

Ischaemic theory:

the meaning of terms understand various this wise it further, topic is discussing to Before

NOMENCLATURE

Before discussing this topic further, it is wise to understand the meaning of the various terms used to describe foot deformities (Fig-25.4). The following are some such terms:

- **Equinus**: (derived from ‘equine’ i.e., a horse who walks on toes). This is a deformity where the foot is fixed in plantar-flexion.
- **Calcaneus** (reverse of equinus): This is a deformity where the foot is fixed in dorsiflexion.
- **Varus**: The foot is inverted and adducted at the mid-tarsal joints so that the sole ‘faces’ inwards.
- **Valgus**: The foot is everted and abducted at the mid-tarsal joints so that the sole ‘faces’ outwards.
- **Cavus**: The longitudinal arch of the foot is exaggerated.
- **Planus**: The longitudinal arch is flattened.
- **Splay**: The transverse arch is flattened.

Invariably, the foot has a combination of above mentioned deformities; the commonest being equino-varus. The next most common congenital foot deformity is calcaneo-valgus.

AETIOLOGY

In the vast majority of cases, aetiology is not known, hence it is termed idiopathic. In others, the so called secondary clubfoot, some underlying cause such as arthrogryposis multiplex congenita (AMC) can be found.

**Idiopathic clubfoot**: Following are some of the theories proposed for the aetiology of idiopathic clubfoot:

a) **Mechanical theory**: The raised intrauterine pressure forces the foot against the wall of the uterus in the position of the deformity.

b) **Ischaemic theory**: Ischaemia of the calf muscles during intrauterine life, due to some unknown factor, results in contractures, leading to foot deformities.

c) **Genetic theory**: Some genetically related disturbances in the development of the foot have been held responsible for the deformity.

**Secondary clubfoot**: Following are some of the causes of secondary clubfoot:

a) **Paralytic disorders**: In a case where there is a muscle imbalance i.e., the invertors and plantar flexors are stronger than the evertors and dorsiflexors, an equino-varus deformity will develop. This occurs in paralytic disorders such as polio, spina bifida, myelodysplasia and Freidreich’s ataxia.
b) **Arthrogryposis multiplex congenita (AMC):**
   This is a disorder of defective development of the muscles. The muscles are fibroitic and result in foot deformities, and deformities at other joints.

**PATHOANATOMY**

All the tissues of the foot i.e., the bones, joints, ligaments and muscles have developmental abnormality.

**Bones:** Bones of the foot are smaller than normal. Neck of the talus is angulated so that the head of the talus faces downwards and medially. Calcaneum is small, and concave medially.

**Joints:** Deformities occur from the malpositioning of different joints:
- **Equinus** deformity occurs primarily at the ankle joint. Other tarsal joints also contribute to it.
- **Inversion** deformity occurs primarily at the subtalar joint. The inverted calcaneum takes the whole foot with it so that the sole faces medially.
- **Forefoot adduction** deformity occurs at the mid-tarsal joints, mainly at talo-navicular joint.
- **Forefoot cavus** deformity is the result of excessive arching of the foot at the mid-tarsal joints.

**Muscles and tendons:** Muscles of the calf are underdeveloped. As a result, the following muscles–tendon units are contracted:
- **Posteriorly**
  - Tendoachilles
- **Medially**
  - Tibialis posterior
  - Flexor digitorum longus
  - Flexor hallucis longus

**Capsule and ligaments:** All the ligamentous structures on the postero-medial side of the foot are shortened. Following are some of these structures:
- **Posterior (3 structures)**
  - Posterior capsule of the ankle joint
  - Posterior capsule of the subtalar joint
  - Posterior talo-fibular and calcaneo-fibular ligaments
- **Medial (3 ligaments)**
  - Talo-navicular ligament
  - Spring ligament
  - Deltoid ligament
- **Plantar**
  - Plantar fascia
  - Plantar ligaments
- **Others**
  - Interosseous ligament between the talus and calcaneum

**Skin:** The skin develops adaptive shortening on the medial side of the sole. There are deep creases on the medial side. There are dimples on the lateral aspect of the ankle and midfoot.

**Secondary changes:** These changes occur in the foot if the child starts walking on the deformed feet. Weight bearing exaggerates the deformity. Callosities and bursae develop over the bony prominences on the lateral side of the foot.

**CLINICAL FEATURES**

**Presenting complaints:** Though, the history dates back to birth, a child with CTEV may present some time after birth, often as late as adulthood. Following are some of the common presentations:

a) **Detected at birth:** At places where delivery is conducted by trained medical personnel, CTEV is detected at the time of routine screening of newborns for congenital mal-formations. At times, the deformity is very mild, the so-called postural equino-varus.

b) **Brought during early infancy:** At places where delivery is conducted at primary health centres, the child is generally brought to the hospital around the age of 3-6 weeks.

c) **Brought during late infancy and early childhood:** In these cases, the child has received treatment elsewhere, or the deformity has recurred, or it has never been corrected. Unfortunately, in developing countries, a large number of cases report to the hospital late.

d) **Brought during late childhood:** It is not uncommon in developing countries to have a grown up child, or sometimes an adult with clubfoot, reporting to the hospital for the first time. Ignorance, poverty and illiteracy are generally the reasons for such late presentations.

**EXAMINATION**

In addition to foot examination, a general examination should be carried out to detect associated malformations in other parts of the body.

**Foot examination:** Normally, the foot of a newborn child can be dorsiflexed until the dorsum touches the anterior aspect of the shin of the tibia (Fig-25.5). This is a good screening test for detecting the milder variety of clubfoot. The more classic one will have the following findings:

- **Bilateral** foot deformity in 60 per cent cases.
- **Size of the foot** smaller (in unilateral cases).
Table 25.1: Terminology to describe clubfoot

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Supple</td>
<td>Foot can be brought to normal position</td>
</tr>
<tr>
<td>Rigid</td>
<td>Foot deformities not correctable</td>
</tr>
<tr>
<td>Resistant</td>
<td>Foot deformities not responding to manipulation</td>
</tr>
<tr>
<td>Neglected</td>
<td>Not treated for 1 year</td>
</tr>
<tr>
<td>Relapsed</td>
<td>Got corrected but recurred</td>
</tr>
</tbody>
</table>

Table 25.2: Differences between primary and secondary club feet

<table>
<thead>
<tr>
<th>Differentiating features</th>
<th>Primary clubfoot</th>
<th>Secondary clubfoot</th>
</tr>
</thead>
<tbody>
<tr>
<td>Present since birth</td>
<td>Yes</td>
<td>Sometimes</td>
</tr>
<tr>
<td>Side affected</td>
<td>Bilateral common (60%)</td>
<td>Unilateral common</td>
</tr>
<tr>
<td>Foot size</td>
<td>Much smaller</td>
<td>Normal or small</td>
</tr>
<tr>
<td>Heel size</td>
<td>Small with fat++</td>
<td>Normal with fat</td>
</tr>
<tr>
<td>Skin</td>
<td>Chubby, creases present</td>
<td>Atrophic, creases absent</td>
</tr>
<tr>
<td>Neurological examination</td>
<td>Essentially normal</td>
<td>Motor and sensory loss present</td>
</tr>
<tr>
<td>Prognosis</td>
<td>Good</td>
<td>Poor</td>
</tr>
</tbody>
</table>

Fig-25.5 Foot of a newborn. It is possible to touch the dorsum of the foot to the shin

- Foot is in equinus, varus and adduction. This can be judged by the inability to bring the foot in the opposite direction. In late cases, in addition, cavus of the foot may also be present.
- Heel is small in size; the calcaneum may be felt with great difficulty.
- Deep skin creases on the back of the heel and on the medial side of the sole.
- Bony prominences felt on the lateral side of the foot, the head of the talus and lateral malleolus.
- Outer side of the foot is gently convex. There are dimples on the outer aspect of the ankle.

On attempted correction, one can feel the tight structures posteriorly (tendoachilles) and plantarwards (plantar fascia).

A child presenting late may have callosities over the lateral aspect of the foot. The calf muscles are wasted.

Terminology to describe clubfoot in different presentations is as shown in Table-25.1.

General examination: It is aimed at finding an underlying cause of the deformity as discussed on page 210. A patient of residual polio may present with equino-varus deformity, which may mimic clubfoot, but there will be paralysis of some other part of the limb. Presence of sensory deficit points to an underlying neurological cause. The presence of deformities at other joints indicate possible arthrogryposis multiplex congenita (AMC).

DIAGNOSIS

This is easy in cases presenting soon after birth. In those presenting late, secondary causes of talipes equino-varus deformity must be excluded (Table–25.2). X-rays of the foot are done (antero-posterior and lateral) with the foot in whatever corrected position possible. The talo-calcaneal angles*, in both, AP and lateral views, in a normal foot are more than 35°, but in CTEV these are reduced (Fig-25.6). X-rays are used by some

![Fig-25.6 X-rays showing reduced talo-calcaneal angles in clubfoot (normal 35°)](https://kat.cr/user/Blink99/)

* This is an angle between long axis of talus and calcaneum – also called as Kite’s angle.

**TREATMENT**

**Principles of treatment:** In principle, treatment consists of correction of the deformity, and its maintenance. Correction can be achieved by non-operative or operative methods. Maintenance is continued until the foot (and its bones) grows to a reasonable size, so that the deformity does not recur.

**METHODS OF CORRECTION OF DEFORMITY**

A deformity can be corrected by non-operative or operative methods.

**Non-operative methods:** Following are the non-operative methods of correcting deformities:

a) **Manipulation alone:** In a newborn, the mother is taught to manipulate the foot after every feed. The foot is dorsiflexed and everted as shown in Fig-25.7. While manipulating, sufficient pressure should be applied by the person so as to blanch her own fingers. This pressure should be maintained for about 5 seconds, and this is repeated several times, over a period of roughly 5 minutes. Minor deformities are usually corrected by this method alone. For major deformities, further treatment by corrective plaster casts is required.

b) **Manipulation and PoP:** In this method, the surgeon manipulates the foot after sedating the child. The foot is then held in the corrected position with plaster casts. There are two philosophies of treatment of clubfoot: Kite’s and Ponsetti’s.

- **Kite’s philosophy:** This has been a popular method for over 40 years. The foot is treated by manipulation and PoP, beginning at the age of 1 month. The deformities are corrected sequentially. Adduction deformity is corrected first followed by inversion deformity and then equinus deformity. A below-knee plaster cast is usually sufficient. The casts are changed every 2 weeks, and are continued until it is possible to ‘overcorrect’ all the deformities. Once this happens, the foot is kept in a suitable maintenance device (discussed subsequently). By this method, correction is achieved in 30% of cases, over a period of 6-9 months. The rest need surgical correction.

- **Ponsetti’s philosophy:** This philosophy is based on better understanding of the pathoanatomy of the deformed foot. According to Ponsetti, the calcaneo-cuboid-navicularexample is internally rotated (adducted) under the plantarflexed talus. Hence, the deformity can be corrected by bringing the complex back under the talus by gradually stretching the tight structures. This is done by putting thumb pressure over the talus head (and not over calcaneo-cuboid joint as in Kite’s method). By doing this, the calcaneo-cuboid-navicularexample is externally rotated under the talar head. Treatment is started within 1st week of life. The cavus aspect of the deformity is corrected first, followed by the adduction, then varus and lastly equinus. After every manipulation, an above-knee PoP cast is applied, which is changed every 5-7 days. It is usually possible to correct all components of the deformity within 6 weeks. The equinus deformity often remains undercorrected, and can be treated by percutaneous tenotomy of tendoachilles. The cut tendoachilles regenerates spontaneously.

**Operative methods:** In more severe deformities, which are not corrected by conservative methods, or in those that recur, operative treatment is required. Soft tissue release operations may be sufficient in younger children (younger than 3 years), but bony operations are required in older children. The following operations are performed:

a) **Postero-medial soft tissue release (PMSTR):** This operation consists of releasing the tight soft tissue structures (tendons, ligaments, capsule etc.) on the posterior and medial side of
Congenital Talipes Equino Varus (CTEV)

For cavus alone a plantar release.

- **Tendon transfers**: In some cases, the tibialis anterior and tibialis posterior (both invertors of the foot) may exert a deforming force against the weak peronei (evertors). This muscle imbalance may be corrected by transferring the tibialis anterior to the outer side of the foot, where it acts as an evertor. Minimum age for tendon transfers is 5 years.

- **Dwyer’s osteotomy**: This is an open-wedge osteotomy of the calcaneum, performed in order to correct varus of the heel (Fig-25.9). Minimum age at which this operation can be performed is 3 years, as prior to this the calcaneum is mainly cartilaginous. Some prefer a closed-wedge osteotomy on the lateral side.

- **Dilwyn Evan’s procedure**: This consists of a thorough soft tissue release (PMSTR) with calcaneo-cuboid fusion (Fig-25.10). It is used for a neglected or recurred clubfoot in children between 4-8 years. With fusion of the calcaneo-cuboid joint, the lateral side of the foot does not
grow as much as the medial side, thus resulting in gradual correction of the deformity.

f) **Wedge tarsectomy:** This consists of removing a wedge of bones from the mid-tarsal area (Fig-25.11). The wedge is cut with its base on the dorso-lateral side. Once the wedge is removed, the foot can be brought to normal (plantigrade) position. This operation is performed for neglected clubfeet between the age of 8-11 years.

g) **Triple arthrodesis:** This consists of the fusion of three joints of the foot (subtalar, calcaneo-cuboid and talo-navicular), after taking suitable wedges to correct the deformity (Fig-25.12). It is performed after the age of 12 years, because before this the bones are cartilaginous and it is difficult to achieve fusion. Of the three, talo-navicular joint fusion is most difficult to achieve.

h) **Ilizarov’s technique:** Using the principles of Ilizarov’s technique, different components of the deformity are corrected by gradual stretching, using an external fixator. Once correction is achieved, it is maintained by plaster casts. Ilizarov’s technique is indicated in neglected clubfeet, and in those in which it has recurred after previous operation. A simpler technique based on above principles has been popularized by Dr. B.B. Joshi from Bombay (JESS fixation).

**METHODS OF MAINTENANCE OF THE CORRECTION**

Correction once achieved, is maintained by the following methods:

a) **CTEV splints:** These are splints made of plastic, moulded in such a way that when tied with straps, it keeps the foot in corrected position.

b) **Denis-Brown splint (DB splint):** This is a splint to hold the foot in the corrected position (Fig-25.13). It is used throughout the day before the child starts walking. Once he starts walking, a DB splint is used at night and CTEV shoes during the day.

c) **CTEV shoes:** These are modified shoes, used once a child starts walking. The following modifications are made in the shoe (Fig-25.14):
Flow chart 25.1 Treatment plan for CTEV

Examination to rule out neuromuscular causes

**NO neuromuscular causes i.e. Idiopathic club foot**

Fresh case (A) At birth

Ponseti method of serial casting for 2-3 weeks

Percutaneous tenotomy of Tendoachilles

Correction achieved

Maintenance in D.B. splint till 1½ years

Splint at night, shoes during day after 1½ years

Follow up till 10-12 years

Progress well

Congratulations

Old, neglected case* (B)

Not corrected till 4-6 months

PMSTR**

Posterior release

Dwyer’s osteotomy

< 3 yrs — Soft tissue release

4-8 yrs — Evan’s operation

8-11 yrs — Wedge tarsectomy

> 12 yrs — Triple arthrodesis

Already — Ilizarov operated technique

Recurrence

Treat as (B)

** PMSTR – Postero-medial soft tissue release
• Straight inner border to prevent forefoot adduction.
• Outer shoe raise to prevent foot inversion.
• No heel to prevent equinus.

These shoes are used until the child is 5 years old.

**PLAN OF TREATMENT**

Most cases which are treated early, respond well to non-operative methods. Operative methods may be indicated in the following cases:

a) A child who does not respond to non-operative treatment (*resistant clubfeet*): These feet are generally severely deformed, ‘chubby’ or associated with underlying arthrogryposis multiplex congenita (AMC).

b) A child whose deformities have recurred (*recurrent clubfeet*): This usually happens if correction is not maintained. The *first* deformity to recur is the equinus.

c) A child who has presented late or has not been adequately treated (*neglected clubfeet*).

Once corrected a clubfoot has to be maintained in the corrected position by the methods described earlier. A comprehensive plan of treatment for CTEV is shown in Flow chart-25.1.

**Further Reading**


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**What have we learnt?**

- Idiopathic clubfoot is to be differentiated from secondary clubfoot, as the prognosis of the two is different.
- Treatment consists of manipulation and PoP, followed by maintenance in splints. Conservative treatment is successful in most cases.
- Different surgical procedures are indicated at different stages.
CONGENITAL DISLOCATION OF THE HIP (CDH)

This is a spontaneous dislocation of the hip occurring before, during or shortly after birth. In western countries, it is one of the commonest congenital disorders. It is uncommon in India and some other Asian countries, probably because of the culture of mothers carrying the child on the side of her waist with the hips of the child abducted (Fig-26.1). This position helps in reduction of an unstable hip, which otherwise would have dislocated. The general term “dysplastic hip” is sometimes used for these congenital malformations of the hip.

AETIOLOGY

Aetiology is not well understood, but the following factors appear to be important:

a) Hereditary predisposition to joint laxity: Hereditary related lax joints are predisposed to hip dislocation in some positions.

b) Hormone induced joint laxity: CDH is 3-5 times more common in females. This may be due to the fact that the maternal relaxin (a ligament relaxing hormone in the mother during pregnancy) crosses the placental barrier to enter the foetus. If the hormonal environment of the foetus is a female, relaxin acts on the foetus’s joints in the same way as it does on those of the mother. This produces joint laxity, and thus dislocation.

c) Breech malposition: The incidence of an unstable hip is about 10 times more in newborns with breech presentation than those with vertex presentation. It is possible that in breech presentation the foetal legs are pressed inside the uterus in such a way that if the hip ligaments are lax, dislocation may occur.

PATHOLOGY

Present evidence suggests that there are two distinct types of dysplastic hips; (i) those dislocated at birth (classic CDH); and (ii) those dislocatable after birth. The first are primarily due to a hereditary faulty development of the acetabulum, and are difficult to treat. The second are due to underlying joint laxity.
with a precipitating factor causing the dislocation. Following changes are seen in a dislocated joint (Fig-26.2):

- Femoral **head is dislocated upwards and laterally**; its epiphysis is small and ossifies late.
- Femoral neck is excessively **anteverted**.
- **Acetabulum is shallow**, with a steep sloping roof.
- Ligamentum teres is hypertrophied.
- Fibro-cartilaginous labrum of the acetabulum (limbus) may be folded into the cavity of the acetabulum (inverted limbus).
- Capsule of the hip joint is stretched.
- **Muscles** around the hip, especially the adductors, undergo adaptive shortening.

**DIAGNOSIS**
Diagnosis is easy in an older child; but may be very difficult in younger children, especially during infancy. This is because of subtle clinical findings and difficulties in interpreting X-rays of these children.

**CLINICAL FEATURES**
CDH is more common in first born babies, more on the left, more common in females (M:F=1:5), bilateral in 20% cases. CDH may be detected at birth or soon after; sometimes not noticed until the child starts walking. Following are the salient clinical features at different ages:
- **At birth**: Routine screening of all newborns is necessary. The examining paediatrician may notice signs suggestive of a dislocated or a dislocatable hip, as discussed subsequently.
- **Early childhood**: Sometimes, the child is brought because the parents have noticed an asymmetry of creases of the groin, limitation of movements of the affected hip, or a click everytime the hip is moved.
- **Older child**: CDH may become apparent once the child starts walking. Parents notice that the child walks with a ‘peculiar gait’ though there is no pain. On examination a CDH may be found to be the underlying cause.

**EXAMINATION**
A meticulous examination is the key to the early diagnosis of CDH. There may be limitation of hip abduction, asymmetry of groin creases or an audible click. Physical findings in a younger child may be little, and diagnosis may only be possible by special tests designed to elicit instability. These are as follows:

**Barlow’s test**: The test has two parts. In the first part, the surgeon faces the child’s perineum. He grasps the upper part of each thigh, with his fingers behind on the greater trochanter and thumb in front. The child’s knees are fully flexed and the hips flexed to a right angle (Fig-26.3). The hip is now gently **adducted**. As this is being done, gentle pressure is exerted by the examining hand in a proximal direction while the thumb tries to ‘push out’ the hip. As the femoral head rolls over the posterior lip of the acetabulum, it may, if dislocatable (but not, if dislocated) slip out of the acetabulum. One feels an abnormal posterior movement, appreciated by the fingers behind the greater trochanter. There may be a distinct ‘clunk’. If nothing happens, the hip may be normal or may already be dislocated; in the latter, second part of the test would be more relevant.
In the second part of the test, with the hips in 90° flexion and fully adducted, held as described above, thighs are gently abducted. The examiner’s hand tries to pull the hips while the fingers on the greater trochanter exert pressure in a forward direction, as if one is trying to put back a dislocated hip. If the hip is dislocated, either because of the first part of the test or if it was dislocated to start with, a ‘clunk’ will be heard and felt, indicating reduction of the dislocated hip. If nothing happens, the hip may be normal or it is an irreducible dislocation. In the latter case, there will be limitation of hip abduction. In a normal hip, it is possible to abduct the hips till the knee touches the couch.

**Ortolani’s test:** This test is similar to the second part of Barlow’s test. The hips and knees are held in a flexed position and gradually abducted. A ‘click of entrance’ will be felt as the femoral head slips into the acetabulum from the position of dislocation. In an older child, the following findings may be present:

- Limitation of abduction of the hip.
- Asymmetrical thigh folds (Fig-26.4).

**Ortolani’s test may be positive.**

**Trendelenburg’s test is positive:** This test is performed in an older child. The child is asked to stand on the affected side. The opposite ASIS (that of the normal side) dips down (details in Annexure-III).

**RADILOGICAL FEATURES**

In a child below the age of 1 year, since the epiphysis of the femoral head is not ossified, it is difficult to diagnose a dislocated hip on plain X-rays (Fig-26.6). Von Rosen’s view may help. Ultrasound

- The limb is short and slightly externally rotated. There is lordosis of the lumbar spine.
- **Telescopy positive:** In a case of a dislocated hip, it will be possible to produce an up and down piston-like movement at the hip. This can be appreciated by feeling the movement of the greater trochanter under the fingers (details in Annexure-III).
- A child with unilateral dislocation exhibits a typical gait in which the body lurches to the affected side as the child bears weight on it (Trendelenburg’s gait). In a child with bilateral dislocation, there is alternate lurching on both sides (waddling gait).
- Some hip pathologies mimicking CDH are: Coxa vara, posterior hip dislocation and paralytic hip dislocation and paralytic hip dislocation.
examination is useful in early diagnosis at birth. In an older child, the following are the important X-ray findings:

- Delayed appearance* of the ossification centre of the head of the femur.
- Retarded development of the ossification centre of the head of the femur.
- Sloping acetabulum.
- Lateral and upward displacement of the ossification centre of the femoral head.
- A break in Shenton’s line (Fig-26.7).

![Fig-26.7 Diagrammatic representation of the X-ray, showing break in Shenton’s line](image)

**TREATMENT**

**Principles of treatment:** Aim is to achieve reduction of the head into the acetabulum, and maintain it until the hip becomes clinically stable and a ‘round’ acetabulum covers the head. In most cases, it is possible to reduce the hip by closed means; in some an open reduction is required. Once the head is inside the acetabulum, in younger children, under the mould-like effect of the head, it develops into a round acetabulum. If reduction has been delayed for more than 2 years, acetabular remodelling may not occur even after the head is reduced for a long time. Hence, in such cases, surgical reconstruction of the acetabulum may be required.

**Methods of reduction:** Following methods of reduction may be used:

a) **Closed manipulation:** It is sometimes possible in younger children to reduce the hip by gentle closed manipulation under general anaesthesia.

b) In unilateral cases, reduction can be attempted till 10 years of age and till 8 years in bilateral cases

c) **Traction followed by closed manipulation:** In cases where the manipulative reduction requires a great deal of force or if it fails, the hip is kept in traction for some time, and is progressively abducted. As this is done, it may be possible to reduce the femoral head easily under general anaesthesia. An adductor tenotomy is often necessary in some cases to allow the hip to be fully abducted.

d) **Open reduction:** This is indicated if closed reduction fails. Reasons of failure of closed reduction could be the presence of fibro-fatty tissue in the acetabulum or a fold of capsule and acetabular labrum (inverted limbus) between the femoral head and the superior part of the acetabulum. In such situations, the hip is exposed, the soft tissues obstructing the head excised or released, and the head repositioned in the acetabulum.

**Maintenance of reduction:** Once the hip has been reduced by closed or open methods, following methods may be used for maintaining the head inside the acetabulum.

a) **Plaster cast:** A frog leg or Bachelor’s cast (Fig-26.8).

![Fig-26.8 Casts for CDH](image)

b) **Splint:** Some form of splint such as Von Rosen’s splint (Fig-26.9).

![Fig-26.9 Von Rosen’s splint](image)

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* Normally, epiphysis of the head of the femur appears at 1 year of age.
External splints can be removed once the acetabulum develops to a round shape. The hip is now mobilised, and kept under observation for a period of 2-3 years for any recurrence.

**Acetabular reconstruction procedures:** The available procedures are:

a) *Salter’s osteotomy:* This is an osteotomy of the iliac bone, above the acetabulum. The roof of the acetabulum is rotated with the fulcrum at the pubic symphysis, so that the acetabulum becomes more horizontal, and thus covers the head (Fig-26.10a).

![Salter’s osteotomy](https://kat.cr/user/Blink99/)

b) *Chiari’s pelvic displacement osteotomy:* The iliac bone is divided almost transversely immediately above the acetabulum, and the lower fragment (bearing the acetabulum) is displaced medially. The margin of the upper fragment provides additional depth to the acetabulum (Fig-26.10b).

c) *Pemberton’s pericapsular osteotomy:* A curved osteotomy as shown in the (Fig-26.10c) is made. The roof of the acetabulum is deflected downwards over the femoral head, with the fulcrum at the triradiate cartilage of the acetabulum.

In some cases, reduction of the hip may be possible only in extreme abduction or internal rotation of the thigh. In such cases a *varus derotation osteotomy* is done at the sub-trochanteric region. The distal fragment is realigned and the osteotomy fixed with a plate.

**Treatment plan:** Treatment varies according to the age at which the patient presents. For convenience of discussion, this has been divided into four groups on the basis of age of the patient:

- **Birth to 6 months:** The femoral head is reduced into the acetabulum by closed manipulation, and maintained with plaster cast or splint.
- **6 months to 6 years:** It may be possible up to 2 years to reduce the head into the acetabulum by closed methods. After 2 years, it is difficult and also unwise to attempt closed reduction. This is because, when the head has been out for some time, the soft tissues around the hip become tight. Such a hip, if reduced forcibly into the acetabulum, develops avascular necrosis of the femoral head. In these cases, reduction is achieved by open methods, and an additional femoral shortening may be required. In older children, an acetabular reconstruction may be performed at the same time or later. Salter’s osteotomy is preferred by most surgeons.
- **6-10 years:** The first point to be decided in children at this age is whether or not to treat the dislocation at all. No treatment may be indicated for children with bilateral dislocations because of the following reasons:
  - The limp is less noticeable.
  - Although having some posture and gait abnormalities, these patients tend to live normal lives until their 40’s or 50’s.
  - Results of treatment are unpredictable and a series of operations may be required.
  - In unilateral cases, an attempt at open reduction with reconstruction of the acetabulum may be made. A derotation osteotomy is needed in most cases.
- **11 years onwards:** Indication for treatment in these patients is pain. If only one hip is affected, a total hip replacement may be practical once adulthood is reached. Sometimes, arthrodesis of the hip may be a reasonable choice.

A general plan of treatment of a child with CDH is as shown in Flow chart-26.1.
OTHER CONGENITAL MALFORMATIONS

TRUNK AND SPINE

1. Klippel-Feil syndrome: Congenital short and stiff neck due to fused or deformed cervical vertebrae.

2. Sprengel’s shoulder: Failure of descent of the scapula, which is developmentally a cervical appendage, i.e. congenital high scapula.

3. Hemivertebra: Growth of only one half of a vertebra resulting in congenital scoliosis. This is common in the dorsal spine.

4. Block vertebra: The bodies of two vertebrae are joined together with no intervening disc space. This is common in the cervical spine.

5. Spondylolysis: A break in the pars inter-articularis of one of the lumbar vertebra, commonly L₅ (see page 285).
6. **Spondylolisthesis**: Displacement of one vertebra over the one below it, because of defective development, commonly L₅ over S₁ (see page 285).

7. **Diastematomyelia**: A longitudinal fibrous or bony septum dividing the spinal canal.

**UPPER LIMB**

8. **Phocomelia**: Lack of development of proximal part of the limb, the distal part being present (seal limb).

9. **Absence of radius**: The hand deviates to lateral side because of lack of normal support by the radius (radial club hand or manus valgus).

10. **Congenital radio-ulnar synostosis**: The forearm bones are joined together at the proximal end, thus preventing forearm rotation.

11. **Madelung’s deformity**: Defective growth of the distal radial epiphysis resulting in deformity of the distal end of the radius, and dislocation of the head of the ulna, dorsally.

12. **Syndactyly**: Webbing of two or more digits; the commonest being middle and ring fingers.

13. **Polydactyly**: More than five fingers; commonly an extra thumb.

**LOWER LIMB**

14. **Congenital dislocation of the hip**.

15. **Congenital coxa vara**: Reduced femoral neck-shaft angle due to a developmental defect in the growth of the proximal femur (see page 323).

16. **Congenital short femur**: Failure of development of the proximal half of the femur resulting in severe shortening.

17. **Congenital pseudarthrosis of the tibia**: A birth defect in the lower third of the tibia in children, whereby a fracture in this region fails to unite.

18. **CTEV or ‘Clubfoot’**: Congenital deformed foot (see page 210).

19. **Congenital vertical talus or ‘Rocker bottom foot’**: A vertically placed talus due to defective development (see page 326).

**Further Reading**


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**What have we learnt?**

- Congenital dislocation of the hip is uncommon in developing countries.
- Early diagnosis by ultrasound at birth, is useful.
- Aim of treatment is to put the dislocated head back in place, as soon as possible.
- Late cases need surgery, which essentially consists of reducing the head into acetabulum, and doing something to keep it there by reconstructing the acetabulum and/or femur.

**Additional information: From the entrance exams point of view**

- The best diagnostic modality for DDH is MRI.
- Screening for DDH done by ultrasound.
- Ultrasound-guided aspiration is the best way of differentiating septic arthritis and transient synovitis.
- Investigation of choice to diagnose early Perthe’s disease is MRI.
Poliomyelitis, commonly called polio, is an acute infectious disease caused by the poliovirus. In the majority of cases, the infection may manifest merely as an episode of diarrhoea; in others the virus may affect the anterior horn cells of the spinal cord and lead to extensive paralysis of the muscles. In extreme forms, the paralysis may involve the respiratory muscles, and may lead to death.

AETIOPATHOLOGY
The poliovirus enters the body either through the faeco-oral route or by inhalation of droplets. The infection occurs commonly in summer. Paralysis may be precipitated after strenuous physical activity, by an intramuscular injection or in a child on cortisone therapy. A tonsillectomy, adenoidectomy or tooth extraction predisposes to paralysis during polio epidemics.

Pathogenesis: The virus multiplies in the intestine. From here it travels to the regional lymph nodes and reticulo-endothelial structures, from where it enters the blood circulation. If the defense mechanism of the body is poor, the virus reaches the nervous system (mainly the anterior horn cells) via the blood or peripheral nerves. The neurons undergo varying degree of damage – some may permanently die, others may be only temporarily damaged, still others may undergo only functional impairment due to tissue oedema. The neurons, which are permanently damaged, lead to permanent paralysis; while the others may regenerate, so that partial recovery of the paralysis may occur. It is this residual paralysis (called post-polio residual paralysis – PPRP) which is responsible for the host of problems associated with a paralytic limb (deformities, weakness etc.).

CLINICAL FEATURES
Presenting complaints: Often, the patient is a child around the age of 9 months. The mother gives a history that the child developed mild pyrexia associated with diarrhoea, followed by inability to move a part or whole of the limb. The lower limbs are affected most commonly. Paralysis is of varying severity and asymmetrical in distribution. In extreme cases, the respiratory muscles may also be paralysed.

Often, the child is seen by a paediatrician in the early stages. When he is referred to an orthopaedic surgeon, the paralysis may already be on its
way to recovery. Recovery of power, if it occurs, may continue for a period of 2 years. Most of the recovery occurs within the first 6 months. Any residual weakness persisting after 2 years is permanent, and will not recover. For descriptive purposes, the disease is conveniently divided into five stages (Fig-27.1, page 226).

**EXAMINATION**

A patient of the paralytic polio may have the following features, on examination:

a) **In the early stage**, the child is febrile, often with rigidity of the neck and tender muscles. This may be associated with diffuse muscle paralysis. The following are some of the typical features of a paralysis resulting from polio:
   - It is asymmetric i.e., the involvement of the affected muscles is haphazard.
   - It occurs commonly in the lower limbs because the anterior horn cells of the lumbar enlargement of the spinal cord are affected most often.
   - The muscle affected most commonly is the quadriceps, although in most cases it is only partially paralysed.
   - The muscle which most often undergoes complete paralysis is the tibialis anterior.
   - The muscle in the hand affected most commonly is the opponens pollicis.
   - The motor paralysis is not associated with any sensory loss.
   - **Bulbar or bulbo-spinal polio**: This is a rare but life threatening polio, where the motor neurons of the medulla are affected. This results in involvement of respiratory and cardiovascular centres, and may cause death.

b) **In late stage (PPRP)**, the paralysis may result in wasting, weakness, and deformities of the limbs. The deformities result from imbalance between muscles of opposite groups at a joint, or due to the action of the gravity on the paralysed limb. The common deformity at the hip is flexion-abduction-external rotation. At the knee, flexion deformity is common; in severe cases *triple deformity* comprising of flexion, posterior subluxation and external rotation occurs. At the foot, equino-varus deformity is the commonest; others being equino-valgus, calcaneo-valgus and calcaneo-carus, in that order. In the upper limbs, polio affects shoulder and elbow muscles. Muscles of the hand are usually spared. The limb may become short. With time, the deformities become permanent due to contracture of the soft tissues and mal-development of the bones in the deformed position.

**DIAGNOSIS**

A diagnosis of poliomyelitis should be considered in an endemic area if a child presents with pyrexia and acutely tender muscles. At this stage, the poliomyelitis is usually confused with influenza, osteomyelitis, septic arthritis, scurvy etc. Once the paralysis sets in, other common conditions producing flaccid paralysis must be excluded. Some of these are as follows:

- Pyogenic meningitis: A lumbar puncture may reveal the diagnosis.
- Post-diphtheritic paralysis.
- Guillain Barre syndrome: This is to be considered if flaccid paralysis occurs later in life. In this syndrome, paralysis is symmetrical, and facial nerve involvement occurs early. Complete recovery usually occurs within 6 months.

In a patient presenting in the stage of residual paralysis, polio should be differentiated from other causes of flaccid paralysis. Some of these are myopathy, spina bifida, other spinal disorders producing paralysis, and peripheral neuropathy (Table–27.1).

**Table-27.1: Differential diagnosis of post-polio paralysis**

<table>
<thead>
<tr>
<th></th>
<th>Poliomyelitis</th>
<th>Myopathy</th>
<th>Spina bifida and other spinal disorders</th>
<th>Neuropathy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asymmetrical</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Lower motor neurone type</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>No sensory loss</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Improves with time, or is static</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Usually symmetrical, follows a pattern</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Lower motor neurone type</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>No sensory loss</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Deteriorates with time</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Usually symmetrical</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Motor+sensory loss</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Deteriorates with growth</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Usually bilateral, ‘Glove and stocking’ pattern</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Motor+sensory loss</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>May improve with treatment</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

**PROGNOSIS**

Of the total number of cases infected with the poliovirus, 50 per cent do not develop paralysis
at all (non-paralytic polio). 40 per cent develop paralysis of a varying degree (mild, moderate, or severe). 10 per cent patients die because of respiratory muscle paralysis. Of the patients with paralytic polio, 33 per cent recover fully, 33 per cent continue to have moderate paralysis, while another 33 per cent remain with severe paralysis.

**TREATMENT**

**Principles of treatment:** Polio can be prevented by immunisation. It is important to immunise patients even after an attack of acute poliomyelitis. This is because there are three strains of the virus, and the patient could still get paralytic polio by another strain. Once polio infection occurs, there is no specific treatment for it, and there is no way of preventing the paralysis or limiting its severity. Whatever recovery from paralysis occurs is spontaneous, and there is very little a doctor can do to enhance the recovery. The role of a doctor is: (i) to provide supportive treatment during the stage of paralysis or recovery; (ii) to prevent the development of deformities during this period; and (iii) to use, in a more efficient way, whatever muscles are functioning. The treatment appropriate to each stage of the disease is best considered stage by stage.

a) **Stage of onset:** It is generally not possible to diagnose polio at this stage. In an endemic area, if a child is suspected of having polio, intramuscular injections and excessive physical activity should be avoided.

b) **Stage of maximum paralysis:** In this stage, the child needs mainly supportive treatment. A close watch is kept for signs suggestive of bulbar polio. These are signs of paralysis of the vagus nerve, causing weakness of the soft palate, pharynx and the vocal cords – hence problem in deglutition, and speech. A respirator may be necessary to save life if the respiratory muscles are paralysed. Paralytic limbs may have to be supported by splints to prevent the development of contractures. All the joints should be moved through the full range of motion several times a day. Muscle pain may be eased by applying hot packs.

c) **Stage of recovery:** The patient should be kept under close supervision of a skilled physiotherapist. The principles of treatment during this stage are as follows:

- **Prevention of deformity** by proper splintage, and joint mobilising exercises.
- **Correction of the deformity** that may have already occurred (discussed in Chapter 11).
- **Retraining of muscles** that are recovering by exercises. Progress is judged by repeated examination of the motor power of the paralysed limb (muscle charting).
- **Encourage walking** with the help of appliances, wherever possible (Fig-27.2).

d) **Stage of residual paralysis:** It is the stage where more active orthopaedic treatment is required. It consists of the following:

- Detailed *evaluation of the patient:* Most patients with residual polio (PPRP) walk with a limp, with or without calipers. An assessment is made whether functional status of the patient can be improved. For this, an evaluation of the deformities and muscle weakness is made. Gait can be improved by the use of a caliper or by operations.

- **Prevention or correction of deformities:** The main emphasis is on prevention of deformity. This is done by splinting the paralysed part in such a way that the effect of muscle imbalance and gravity is negated (details in Chapter-11). An operation may sometimes be required to prevent the deformity. For example, in a foot with severe muscle imbalance between opposite group of muscles, a tendon transfer operation is done. This produces a more ‘balanced’ foot, hence less possibility of deformity. Commonly
performed operation for correction of deformities are as follows:

- For hip deformity (flexion-abduction-external rotation): Soutters’ release.
- For knee flexion deformity: Wilson’s release.
- For equinus deformity of the ankle: Tendoachilles lengthening.
- For cavus deformity of the foot: Steindler’s release.

**Tendon transfers:** The available muscle power is redistributed either to equalise an unbalanced paralysis, or to use the motor power for a more useful function (see page 86). It is not done before 5 years of age, as the child has to be manageable enough to be taught proper exercises. More commonly performed tendon transfers are as follows:

- Transfer of extensor hallucis longus (EHL) from the distal phalanx of great toe to the neck of the first metatarsal (modified Jone’s operation). This is done to correct first metatarsal drop in case of tibialis anterior muscle weakness.
- Transfer of peronius tertious and brevis muscles (evertors of the foot) to the dorsum of the foot. The transfer is required in a foot with dorsiflexor weakness. Evertors can be spared for more useful function of dorsiflexion of the foot.
- Hamstring (knee flexors) transfer to the quadriceps muscle to support a weak knee extensor.
- **Stabilisation of flail joints:** Joints with such severe muscle paralysis that the body loses control over them are called flail joints. Stabilisation of these joints is necessary for walking. This can be achieved by operative or non-operative methods. Non-operative methods consist of calipers, shoes etc. Operative methods consist of fusion of the joints (e.g., triple arthrodesis for stabilisation of the foot).
- **Leg length equalisation:** In cases where a leg is short by more than 4 cm, a leg lengthening procedure may be required.

**CEREBRAL PALSY (CP)**

This is defined as a non-progressive neuro-muscular disorder of cerebral origin. It includes a number of clinical disorders, mostly arising in childhood. The essential features of all these disorders is a varying degree of upper motor neurone type of limb paralysis (spasticity), together with difficulty in coordination (ataxia) and purposeless movements (athetosis).

**AETIOPATHOLOGY**

Birth anoxia and injuries are the commonest cause of CP in developing countries. Causes can be divided into prenatal, natal and postnatal (Table–27.2).

<table>
<thead>
<tr>
<th>Causes of cerebral palsy</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Prenatal causes</strong></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td><strong>Natal causes</strong></td>
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<td></td>
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<tr>
<td><strong>Postnatal causes</strong></td>
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</tbody>
</table>

* The commonest

**Pathology:** The pathology of this disorder is the degeneration of the cerebral cortex or basal ganglion, either because of their faulty development or because of damage caused by the various factors given in Table–27.2.

**CLINICAL FEATURES**

**Presenting complaints:** The clinical features vary according to the severity of the lesion, the site of the neurological deficit and the associated defects.

- **Severity of lesion:** The lesion may be mild in 20 per cent of cases, in which case the child may remain ambulatory without any help and may never require consultation. In the majority (almost 50 per cent of cases), the child requires help with ambulation. The usual presentation is a child less than one year old, in whom the parents have noticed a lack of control on the affected limb. There is a delay in the developmental milestones such as sitting up, standing or walking. In about 30 per cent of cases, the involvement is severe, and the child is bed-ridden.

- **Pattern of involvement:** The pyramidal tracts are involved in 65 per cent of cases, and they present with spasticity, exaggerated reflexes etc. One or all the limbs may be involved. The commonest pattern is a symmetrical spastic paresis of the
lower limbs, resulting in a tendency to flex and adduct the hips (scissoring), to keep the knees flexed and the feet in equinus. Less commonly, it may present as monoplegia, hemiplegia or quadriplegia. In the upper limb, there is typical flexion of the wrist and fingers with adduction of the thumb and pronation of the forearm. In 35 per cent of cases, extra-pyramidal symptoms such as ataxia, athetoid movements, dystonia predominate.

- **Associated defects:** These consist of speech defect, sensory defect, epilepsy, ocular defects and mental retardation. About 50 per cent of the patients are severely mentally retarded, 25 per cent have moderate mental retardation and 25 per cent have borderline mental retardation.

**EXAMINATION**

On examination, there may be weakness of muscles, the distribution of which is variable. This leads to marked muscle imbalance, resulting in deformities. The joints are stiff because of spasticity; hence when a steady pressure is applied, the muscle relaxes and the deformity is partially corrected. As the pressure is released, the spasm returns immediately. The tendon reflexes are exaggerated, and clonus may be present.

The patient exhibits a lack of voluntary control when asked to hold an object. As the patient tries to move a single group of muscles, other groups contract at the same time (athetoid movements). Mental deficiency may be present. There may also be defective vision and impaired hearing.

**TREATMENT**

**Principles of treatment:** The aim of treatment is to maintain and develop whatever physical and mental capabilities the child has. It consists of: (i) orthopaedic treatment; and (ii) speech and occupational therapy.

Orthopaedic treatment consists of the prevention and correction of deformities, and keeping the spasticity under check. Methods of controlling the spasticity are: (i) drugs – e.g., Diazepam, Beclofen; (ii) phenol nerve block; and (iii) neurectomy. Neurectomy may be required to control severe muscle spasm interfering with optimal rehabilitation. Obturator neurectomy is performed for spasm of adductors of the thigh. A number of other operative procedures may be necessary for improving selective functions.

Speech therapy and occupational therapy constitutes an important adjunct to the overall treatment of the child. Mild cases can be looked after at home, but specialised residential schools are required for severely handicapped children.

**PROGNOSIS**

Complete cure is impossible since an essential part of the brain is destroyed and cannot be repaired or replaced. All that can be hoped for is improvement. Depending upon the severity of the underlying damage, a child can be made independent enough to earn his own living in due course. A child formerly dependent on others for many daily activities may often become independent. This needs a great amount of patience and perseverance on the part of the parents and attendants of the child. In spite of all the treatment, there are a few in whom worthwhile improvement cannot be gained.

**SPINA BIFIDA**

**RELEVANT ANATOMY**

The vertebral bodies develop from the mesoderm around the notochord. From the centre of each body extend two projections which grow around the neural canal to form the vertebral arch (Fig-27.3). The two halves of the arch fuse in the thoracic region, from where the fusion extends up and down. Failure of fusion of these arches gives rise to spina bifida. It is often associated with maldevelopment of the spinal cord and the membranes.
with a major aberration in the development of the neural elements. Accordingly, there are two main types of spina bifida: (i) spina bifida occulta; and (ii) spina bifida aperta.

**Spina bifida occulta:** This is the mildest and the **commonest.** In this, the failure of the vertebral arches to fuse results in bifid spinous processes of vertebrae. The following are some of the important features:

- **Commonest site:** This is common in the lumbosacral spine; S1 being the commonest site.
- **Externally,** the skin may be normal or there may be tell tale signs in the form of a dimple in the skin, a lipomatous mass, a dermal sinus or a tuft of hair.
- **Neurological impairment** is not related to the severity of the bone defect. The commonest manifestation of neurological involvement is a muscle imbalance in the lower limbs with selective muscle wasting. This leads to foot deformities because of muscle imbalance; common ones being equinovarus or cavus. The cause of neural impairment may be: (i) tethering of the cord to the undersurface of the skin by a fibrous membrane (*membrana reuniens)*; (ii) tethering of the cord to the filum terminale; (iii) bifid cord, transfixed with an antero-posterior bone bar (*diastematomyelia*); or (iv) defective neural development (*myelodysplasia*).

**Treatment:** A symptomless patient, where the lesion is detected on an X-ray taken for some other problem, needs no treatment. Cases presenting with backache respond to physiotherapy. Cases presenting with a neurological deficit need to be evaluated regarding the cause and likelihood of worsening of the neurological deficit. MRI is the imaging modality of choice. Surgical treatment may be required in some cases. *Orthopaedic treatment* is the same as for a paralytic limb, i.e., (i) prevention and correction of deformities; (ii) using residual muscle power for more useful functions by tendon transfers and joint stabilisation; (iii) giving support for walking.

**Spina bifida aperta:** This developmental defect involves not only the vertebral arches but also the overlying soft tissues, skin, and often the meninges. In severe cases, the nerve tube itself may be exposed. The following are some of the important features:

**Commonest site:** The dorso-lumbar spine is affected most commonly. There is a variable structural defect of closure of the embryonal neural tube giving rise to the following (Fig-27.4):

a) **Meningocele** i.e., protrusion of meninges through a defect in the neural arch. This contains only CSF.

b) **Meningomyelocele** i.e., the protrusion of the meninges along with some neural elements (normally developed spinal cord or cauda equina).

c) **Syringomyelocele** i.e., the central canal of the cord is dilated (*syringomyelia*), and the cord lies within the protruded meningeal sac together with the nerves arising from it.

d) **Myelocele:** This results from an arrest in the development at the time of closure of the neural groove. An elliptical raw surface, representing the ununited groove is seen. At the upper end of this surface opens the central canal through which CSF may be seen leaking.

With the exception of spina bifida occulta, myelocele is the **commonest** type of spina bifida; though many of these cases are stillborn. If the child is born alive, death ensues within a few days from infection of the cord and meninges. The other types of spina
bifida with neural development defects where the patient often survives is meningocele and meningomyelocele. There may be mild to severe paralysis of the lower limbs. These children are often born with deformities, particularly flexion-adduction contracture of the hip, and deformities of the foot. These deformities are the direct result of muscle imbalance due to paralysis. There may be urinary and bowel incontinence.

_Treatment:_ Treatment of this condition consists of: (i) treatment of the basic defect i.e., spina bifida; (ii) orthopaedic treatment to prevent and correct the deformities, and to use the residual motor power in the best possible way; and (iii) urological treatment for bladder incontinence.

**DISORDERS OF THE MUSCLES**

These diseases are still incurable. Accurate diagnosis is important: (i) to rule out other treatable disorders; (ii) to act as a guide to optimal rehabilitation efforts; and (iii) to permit genetic counselling.

**DEFINITIONS**

Myopathy is a _generic term_ for somatic motor dysfunctions due to diseases of the skeletal muscles i.e., dysfunctions not otherwise attributable to lesions of the central nervous system, the lower motor neurone (LMN) or the neuro-muscular junction (NMJ).

Myopathy may be inherited or acquired. The underlying pathological process may be restricted to the muscles or may affect other organ systems as well. In an inherited myopathy, the defect may be: (i) abnormal cellular enzymes; (ii) abnormal structural proteins; or (iii) both. It is the inherited myopathy with structural protein as its underlying defect, which is designated _muscular dystrophy_ (Flow chart-27.1). It comprises of a group of heterogeneous disorders that share in common: (i) bilateral, usually symmetrical, topographically patterned loss of strength and muscle wasting; (ii) progression over several years; and (iii) largely non-specific laboratory and histologic evidence of myofibril necrosis. One type of dystrophy is distinguishable from another type by the following criteria:

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**Flow chart-27.1 Approach to a patient with motor dysfunction**

![Flow chart](image_url)
There are no proven pathognomonic clinical or laboratory tests to diagnose a muscular dystrophy. A confirmed family history is a very strong indicator of diagnosis. Classic syndromes present few diagnostic problems, but unfortunately incomplete or atypical forms are common. Diagnostic accuracy is directly proportional to the reliability of the neurologic data (the history, physical examination, and ancillary laboratory findings), and the care with which it is analysed (Flow chart-27.2). A systematic approach as suggested below helps in arriving at the correct diagnosis.

a) First localise the neuroanatomic site of the lesion. (Is it in the CNS, LMN, NMJ, or muscle?).

b) Establish its most likely pathogenetic mechanism (is it an inherited or acquired, myopathy?). If inherited, is it metabolic or structural? If acquired, is it infectious, metabolic, toxic drug induced, or dysimmune?

c) Identify a particular disease consistent with the site of the lesion and presumed pathogenesis.

**Treatment:** It is difficult. Most of the common types pursue a gradual course leading to severe muscular weakness. In selected cases, treatment along the lines of a paralytic limb may be used (refer to page 228).

**PERIPHERAL NEUROPATHIES**

This topic is discussed in detail in Medicine textbooks, and is discussed here since a paralysis secondary to neuropathy may present to an orthopaedic surgeon. Peripheral neuropathies are of two types – mononeuropathy, and polyneuropathy. Mononeuropathy is commonly due to
trauma and other causes as discussed in Chapter 10. The causes of polyneuropathy are as given in Table–27.3.

Nutritional deficiency, diabetes and infections constitute majority of the cases of polyneuropathy. Guillain Barre syndrome is an important treatable cause. The cause of neuropathy can be found in about 50-60 per cent of cases by clinical examination and investigations. The patient presents with bilateral involvement, complains of weakness of most distal group of muscles and paraesthesias in the distal parts of the extremities. There is loss of deep jerks in the affected extremities, and glove and stocking type of hypo-aesthesia. A detailed neurological examination should be performed in all cases.

**Table–27.3: Common causes of peripheral polyneuropathies**

<table>
<thead>
<tr>
<th>Toxic</th>
<th>Infections</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alcoholism</td>
<td>Leprosy, Diphtheria</td>
</tr>
<tr>
<td>Drugs: Nitrofurantoin, INH, Diphenylhydantoin</td>
<td>Guillain Barre syndrome</td>
</tr>
<tr>
<td>Metals: Pb, As, Hg</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Deficiency states</th>
<th>Genetic causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vit. B, deficiency</td>
<td>Peroneal muscular atrophy</td>
</tr>
<tr>
<td>Vit. B₁₂ deficiency</td>
<td>Progressive hypertrophic polyneuropathy</td>
</tr>
<tr>
<td>Multiple deficiencies</td>
<td></td>
</tr>
<tr>
<td>Malnutrition</td>
<td></td>
</tr>
<tr>
<td>Malabsorption</td>
<td></td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Metabolic diseases</th>
<th>Inflammatory causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diabetes mellitus</td>
<td>Polyarteritis nodosa (PAN)</td>
</tr>
<tr>
<td>Uraemia</td>
<td>Rheumatoid arthritis</td>
</tr>
<tr>
<td>Acute intermittent porphyria</td>
<td>Systemic lupus erythematosus (SLE)</td>
</tr>
<tr>
<td>Hepatic failure</td>
<td></td>
</tr>
</tbody>
</table>

**TREATMENT**

It consists of the following:

- Treatment of the underlying cause, if possible.
- Prevention of contractures by splintage and physiotherapy.
- Care of the anaesthetic limb by protecting it from injury.
- Treatment of neuropathic pain with analgesics and nerve blocks.

**Further Reading**


**What have we learnt?**

- It is the residue of polio, which needs treatment by orthopaedic surgery.
- In the initial stage, aim is to prevent occurrence of deformity. In late stages, reconstructive surgery is required to improve the functional capability of the affected part.

**Additional information: From the entrance exams point of view**

- Progression of congenital scoliosis is maximum in unilateral unsegmented bar with a hemi-vertebra.
- Progression of congenital scoliosis is least with a block vertebra.
The term ‘bone tumour’ is a broad term used for benign and malignant neoplasms, as well as ‘tumour-like conditions’ of the bone (e.g., osteochondroma). Metastatic deposits in the bone are commoner than primary bone tumours. Of the primary bone malignancies, multiple myeloma is the commonest. Osteochondroma is the commonest benign tumour* of the bone. Most primary malignant bone tumours occur in children and young adults; in whom these constitute one of the common malignant tumours. The nature of a bone tumour can be suspected based on the type of destruction seen on an X-ray (Table–28.1).

**TOPICS**

- Benign tumours
- Tumour-like conditions of bone
- Osteodastoma (GCT)
- Osteochondroma
- Primary malignant tumours
- Aneurysmal bone cyst
- Some uncommon malignant tumours
- Fibrous dysplasia
- Metastasis in bone

* Though it is not a true neoplasm.

Classification and nomenclature of bone tumours is given in Table–28.2.

**BENIGN TUMOURS**

**OSTEOMA**

This is a benign tumour composed of sclerotic, well-formed bone protruding from the cortical surface of a bone. The bones involved most often are the skull and facial bones. Generally, the tumour is of no clinical significance except that it may produce visible swelling. Sometimes, it may bulge into one of the air sinuses (frontal, ethmoidal or others), and cause obstruction to the sinus cavity, leading to pain.

**Treatment:** No treatment is generally required except for cosmetic reasons, where a simple excision is sufficient. It is not a pre-malignant lesion.

**OSTEOID OSTEOMA**

It is the commonest true benign tumour of the bone. Pathologically, it consists of a nidus of tangled arrays of partially mineralised osteoid trabeculae surrounded by dense sclerotic bone.

**Clinical presentation:** The tumour is seen commonly between the ages of 5-25 years. The bones of the lower extremity are more commonly affected; *tibia* being the commonest. The tumour is generally located in the diaphysis of long bones. Posterior elements of the vertebrae are a common site. The presenting complaint is a nagging pain, worst at night, and is relieved by salicylates. There are minimal or no clinical signs, except for mild tenderness at the site of the lesion, and a palpable swelling if it is a superficial lesion.

**Diagnosis:** It is generally confirmed on X-ray. The tumour is visible as a zone of sclerosis surrounding a radiolucent nidus, usually less than 1 cm in size (Fig-28.1a). In some cases, the nidus may not be seen on a plain X-ray because of extensive surrounding sclerosis, and may be detected on a CT scan.
Essential Orthopaedics

Treatment: Complete excision of the nidus along with the sclerotic bone is done. Prognosis is good. It is not a pre-malignant condition.

UNCOMMON BENIGN TUMOURS OF THE BONE

Osteoblastoma: This is a benign tumour consisting of vascular osteoid and new bone. It occurs in the jaw and the spine. If in long bones, it occurs in the diaphysis or metaphysis, but never in the epiphysis. It occurs in patients in their 2nd decade of life. The patient presents with an aching pain. Radiologically, it is a well-defined radiolucent expansile bone lesion 2-12 cm in size. There is minimal reactive new bone formation. Treatment is by curettage.

Chondroblastoma: This is a cartilaginous tumour containing characteristic multiple calcium deposits. It occurs in young adults, and is located around the epiphyseal plate. Bones around the knee are commonly affected. Radiologically, there is a well-defined lytic lesion surrounded by a zone of sclerosis. Areas of calcification within the tumour substance give rise to a mottled appearance (Fig-28.1b). Treatment is by curettage and bone grafting.

Haemangioma of the bone: This is a benign tumour of angiomatous origin, commonly affecting the vertebrae and the skull. It occurs in young adults. Common presenting symptoms are persistent pain and features of cord compression. At times the lesion is asymptomatic. Typically, one of the lumbar vertebrae is affected. Radiologically, it appears as loss of horizontal striations and prominence of

Table-28.2: Nomenclature and classification of bone tumours (WHO classification simplified)

<table>
<thead>
<tr>
<th>a) Bone forming tumours</th>
<th>b) Cartilage forming tumours</th>
<th>c) Giant cell tumours (GCT)</th>
<th>d) Marrow tumours</th>
<th>e) Vascular tumours</th>
<th>f) Others</th>
<th>g) Tumour-like lesions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benign: Osteoid osteoma, osteoma</td>
<td>Benign: Osteochondroma (exostosis)</td>
<td>Benign GCT</td>
<td>Malignant: Ewing’s sarcoma</td>
<td>Benign: Haemangioma</td>
<td>Benign: Neurilemmoma</td>
<td>Bone cysts – simple or aneurysmal</td>
</tr>
<tr>
<td>Osteoma</td>
<td>Benign: Enchondroma (chondroma)</td>
<td>Indeterminate GCT</td>
<td>Plasma cell tumour</td>
<td>Benign: Angiosarcoma</td>
<td>Benign: Neurofibroma</td>
<td>Fibrous dysplasia – mono or polyostotic</td>
</tr>
<tr>
<td>Osteoblastoma</td>
<td>Benign: Chondromyxoid fibroma</td>
<td>Malignant GCT</td>
<td>Multiple myeloma</td>
<td>Malignant: Malignant fibrous histiocytoma</td>
<td>Malignant: Liposarcoma</td>
<td>Reparative giant cell granuloma (e.g. epulis)</td>
</tr>
<tr>
<td>Indeterminate: Aggressive osteoblastoma</td>
<td>Benign: Chondroblastoma</td>
<td></td>
<td>Lymphoma</td>
<td>Malignant: Undifferentiated sarcoma</td>
<td></td>
<td>Fibrous cortical defect</td>
</tr>
<tr>
<td>Malignant: Osteosarcoma</td>
<td>Malignant: Chondrosarcoma</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Eosinophilic granuloma</td>
</tr>
</tbody>
</table>

Fig-28.1 Benign tumours of the bone
Bone Tumours

Giant cell tumour (GCT) is a common bone tumour with variable growth potential. Though generally classified as benign*, it tends to recur after local removal. Frankly malignant variants are also known.

PATHOLOGY
The cell of origin is uncertain. Microscopically, the tumour consists of undifferentiated spindle cells, profusely interspersed with multi-nucleate giant cells. The tumour stroma is highly vascular. These giant cells were mistaken as osteoclasts in the past, hence the name osteoclastoma.

CLINICAL FEATURES
The tumour is seen commonly in the age group of 20-40 years i.e., after epiphyseal fusion. The bones affected commonly are those around the knee i.e., lower-end of the femur and upper-end of the tibia. Lower-end of the radius is another common site. The tumour is located at the epiphysis**. It often reaches almost up to the joint surface. Common presenting complaints are swelling and vague pain. Sometimes, the patient, unaware of the lesion, presents for the first time with a pathological fracture through the lesion.

EXAMINATION
Examination reveals a bony swelling, eccentrically located at the end of the bone. Surface of the swelling is smooth. There may be tenderness on firm palpation. A characteristic ‘egg-shell crackling’ is often not elicited. The limb may be deformed if a pathological fracture has occurred.

DIAGNOSIS
GCT is one of the common cause of a solitary lytic lesion of the bone, and must be differentiated from other such lesions (Table–28.3). Following are some of the characteristic radiological features of this tumour:
• A solitary, may be loculated, lytic lesion.
• Eccentric location, often subchondral (Fig-28.1d).
• Expansion of the overlying cortex (expansile lesion).

Table–28.3: Differential diagnosis of a solitary bone lesion

<table>
<thead>
<tr>
<th>Features</th>
<th>Giant cell tumour</th>
<th>Simple bone cyst</th>
<th>Aneurysmal bone cyst</th>
<th>Fibrous dysplasia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>20-40 yrs.</td>
<td>&lt; 20 yrs.</td>
<td>10-40 yrs.</td>
<td>20-30 yrs.</td>
</tr>
<tr>
<td>Common bones</td>
<td>Lower femur,</td>
<td>Upper humerus</td>
<td>Tibia</td>
<td>Neck of the femur</td>
</tr>
<tr>
<td></td>
<td>Upper tibia</td>
<td>Upper femur</td>
<td>Humerus</td>
<td>Tibia</td>
</tr>
<tr>
<td></td>
<td>Lower radius</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Location</td>
<td>Epiphysis</td>
<td>Metaphysis</td>
<td>Metaphysis</td>
<td>Metaphysis</td>
</tr>
<tr>
<td>X-ray</td>
<td>Soap-bubble</td>
<td>Maximum width</td>
<td>Distending lesion,</td>
<td>Multi-loculated</td>
</tr>
<tr>
<td></td>
<td>appearance,</td>
<td>less than width</td>
<td>‘ballooning’ the bone</td>
<td>Ground-glass</td>
</tr>
<tr>
<td></td>
<td>eccentrically</td>
<td>of the growth plate</td>
<td></td>
<td>appearance</td>
</tr>
<tr>
<td></td>
<td>placed</td>
<td></td>
<td></td>
<td>Trabeculations++</td>
</tr>
<tr>
<td>Treatment preferred</td>
<td>Excision</td>
<td>Curettage and bone graft</td>
<td>Curettage and bone graft</td>
<td>Curettage and bone graft</td>
</tr>
</tbody>
</table>

* 1/3 are benign, 1/3 locally malignant and 1/3 frankly malignant.
** It is the area which was epiphysis before its fusion with the metaphysis.
‘Soap-bubble’ appearance – the tumour is homogeneously lytic with trabeculae of the remnants of bone traversing it, giving rise to a loculated appearance.

- No calcification within the tumour (Fig-28.2b).
- None or minimal reactive sclerosis around the tumour.
- Cortex may be thinned out, or perforated at places.
- Tumour usually does not enter the adjacent joint.

TREATMENT
Wherever possible, excision of the tumour is the best treatment. For sites like the spine, where excision is sometimes technically not possible, radiotherapy is done. Following treatment methods are commonly used:

a) **Excision**: This is the treatment of choice when the tumour affects a bone whose removal does not hamper with functions e.g., the fibula, lower-end of the ulna etc.

b) **Excision with reconstruction**: When excision of a tumour at some site may result in significant functional impairment, the defect created by excision is made up, usually partially, by some reconstructive procedure. For example, in tumours affecting the lower-end of femur, the affected part is excised en bloc, and the defect thus created made up by one of the following methods:

- **Arthrodesis by the Turn-o-Plasty procedure** (Fig-28.3a): In this technique, the required length of the tibia is split into two halves. One half is turned upside down and fixed with the stump of the femur left after excising the tumour. A similar procedure can be used for a tibial lesion by taking half of the femur.
- **Arthrodesis by bridging the gap** by double fibulae (Fig-28.3b), one taken from same extremity and the other from the opposite leg (Yadav, 1990).
- **Arthroplasty**: In this procedure, the tumour is excised, and an attempt is made to reconstruct the joint in some way (Fig-28.3c).

![Fig-28.2 Radiological features of giant cell tumour](image1)

![Fig-28.3 Methods of treating GCT around the knee](image2)
This can be carried out using an autograft (patella to substitute the articular defect), allograft (replacing the defect with the preserved bone of a cadaver), or an artificial joint (prosthesis).

c) **Curettage with or without supplementary procedures:** Curettage performed alone has the disadvantage of a high recurrence rate. This is because however thorough the curettage may be, some cells are always left along the walls of the cavity. Some supplementary procedures used with curettage have been reported to reduce recurrence. **Cryotherapy,** where liquid nitrogen is used to produce a freezing effect and thus kill the residual cells, and thermal burning of the residual cells using cauterization of the walls of the tumour are popular. Lately, thermal effect of bone cement has been used. The cavity is filled with ‘bone cement’, which by the heat it produces while setting, ‘kills’ the residual cells.

d) **Amputation:** For more aggressive tumours, or following recurrence, amputation may be necessary.

e) **Radiotherapy:** It is the preferred treatment method for GCT affecting the vertebrae. Treatment for GCT at commoner sites is as given in Table–28.4.

### Table–28.4: Treatment of GCT at common sites

<table>
<thead>
<tr>
<th>Site</th>
<th>Treatment of choice</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lower end of femur</td>
<td>Excision with Turn-o-Plasty</td>
</tr>
<tr>
<td>Upper end of tibia</td>
<td>Excision with Turn-o-Plasty</td>
</tr>
<tr>
<td>Lower end of radius</td>
<td>Excision with fibular* grafting</td>
</tr>
<tr>
<td>Lower end of ulna</td>
<td>Excision</td>
</tr>
<tr>
<td>Upper end of fibula</td>
<td>Excision</td>
</tr>
</tbody>
</table>

* Proximal end of the opposite fibula is preferred, since it matches the lower end of radius in shape.

**PROGNOSIS**

Recurrence following treatment is a serious problem. With every subsequent recurrence, the tumour becomes more aggressive.

**PRIMARY MALIGNANT TUMOURS**

**OSTEOSARCOMA (OSTEOGENIC SARCOMA)**

Osteosarcoma is the second most common, and a highly malignant primary bone tumour.

**Pathology:** An osteosarcoma can be defined as a malignant tumour of the mesenchymal cells, characterised by formation of osteoid or bone by the tumour cells.

**Classification:** This tumour has been subclassified on the basis of: (i) the clinical setting where it occurs; and (ii) its dominant histo-morphology.

a) On the basis of **clinical setting,** this tumour can be divided into primary and secondary. **Primary osteosarcoma,** the commoner, occurs in the age group of 15-25 years. There are no known pre-malignant conditions related to it. It is very much more malignant than the secondary one. The **secondary** osteosarcoma occurs in older age (45 years onwards). Some of the pre-malignant conditions often associated with it are Paget’s disease, multiple enchondromatosis, fibrous dysplasia, irradiation to bones, multiple osteochondroma etc.

b) On the basis of **dominant histo-morphology,** an osteosarcoma may be: (i) osteoblastic i.e., with a lot of new bone formation; (ii) chondroid i.e., with basic cell being a cartilage cell; (iii) fibroblastic i.e. the basic cell being a fibroblast; and (iv) telangiectatic or osteolytic type, a predominantly lytic tumour.

Whatever be the histo-morphologic characteristics and the site of origin, all osteosarcomas are aggressive lesions and metastasise widely through the blood stream, usually first to the lungs. Lymph node involvement, even local, is unusual. Osteolytic type is more malignant than the osteoblastic type. Despite its aggressiveness, osteosarcoma rarely penetrates the epiphysial plate. Most osteosarcomas fall into the primary conventional category, and have the following important features.

- **Age at onset:** These tumours occur between the ages of 15-25 years, constituting the commonest musculo-skeletal tumour at that age.
- **Common sites of origin:** In decreasing order of frequency these are: the lower-end of the femur; upper-end of the tibia; and upper-end of the humerus. However, any bone of the body may be affected.
- **Gross appearance** of the tumour depends upon its dominant histo-morphology. An **osteoblastic** tumour is greyish white, hard, and has a gritty feeling when cut. A **chondroid** type may appear opalescent and bluish grey. A **fibroblastic** type has a more typical fish flesh sarcomatous appearance. The highly malignant, **telangiectatic** type may have large areas of tumour necrosis.
and blood filled spaces within the tumour mass. Most tumours have mixed areas.

- **Histologically**, these tumours vary in the richness of the osteoid, cartilaginous, or vascular components; but common to all is a basically anaplastic mesenchymal parenchyma with tumour cells surrounded by osteoid.

**Clinical features:** Pain is usually the first symptom, soon followed by swelling. Pain is constant and boring, and becomes worse as the swelling increases in size. There may be a history of trauma, but more often it is incidental and just draws the attention of the patient to the swelling. Sometimes, the patient presents with a pathological fracture.

**Examination:** The swelling is in the region of the metaphysis. Skin over the swelling is shiny with prominent veins. The swelling is warm and tender. Margins of the swelling are not well-defined. Movement at the adjacent joint may be limited mainly because of the mechanical block by the swelling. The tumour may compress the neurovascular structures of the limb, and produce symptoms due to that. Regional lymph nodes may be enlarged, but are usually reactive.

**Investigations:** Following investigations may be carried out to confirm the diagnosis:

- **Radiological examination:** X-ray shows the following features (Fig-28.4):
  - An area of irregular destruction in the metaphysis, sometimes overshadowed by the new bone formation. The cortex overlying the lesion is eroded. There is new bone formation in the matrix of the tumour.
  - **Periosteal reaction:** As the tumour lifts the periosteum, it incites an intense periosteal reaction. The periosteal reaction in an osteosarcoma is irregular, unlike in osteomyelitis where it is smooth and in layers.
  - **Codman’s triangle:** A triangular area of subperiosteal new bone is seen at the tumour-host cortex junction at the ends of the tumour.
  - **Sun-ray appearance:** As the periosteum is unable to contain the tumour, the tumour grows into the overlying soft tissues. New bone is laid down along the blood vessels within the tumour growing centrifugally, giving rise to a ‘sun-ray appearance’ on the X-ray.

- **Serum alkaline phosphatase (SAP):** It is generally elevated, but is of no diagnostic significance. It has been considered a useful parameter for follow up of a case of osteosarcoma. A rise of SAP after an initial fall after tumour removal is taken as an indicator of recurrence or metastasis.

- **Biopsy:** An open biopsy is performed to confirm the diagnosis. Some pathologists have gained experience in diagnosing osteosarcoma by a small tissue sample obtained by a needle (core-biopsy), or by fine needle aspiration cytology (FNAC).

**Treatment:** The aim is to confirm the diagnosis, to evaluate spread of the tumour, and to execute adequate treatment.

a) **Confirmation of the diagnosis:** Histologically, tumour new bone formation is pathognomonic of osteosarcoma. In the absence of a classic appearance on histology, the clinical and radiological picture is taken into consideration.

b) **Evaluation of spread of tumour:** This consists of evaluation of the extent of involvement of the affected bone and that of spread of the tumour to other sites. Lung is the earliest site for metastasis. A chest X-ray should be done to detect the same. CT scan may be required in cases where the metastatic lesion is unclear on chest X-ray. It is important to know the extent of involvement of the affected bone by the tumour for the following reasons:
• **To plan amputation surgery:** Complete removal of local tumour is of vital importance in amputation surgery. The tumour may have slip areas in the medullary cavities, and can result in recurrence even after amputation.

• **To plan a limb saving operation:** In cases presenting early, a radical excision of the tumour is being performed these days (limb saving surgery), thus avoiding amputation.

• **Methods used for precise evaluation** of spread of the tumour locally are: bone scan for finding the intra-medullary spread (‘skip’ lesions), CT and MRI scans for finding the soft tissue spread. These investigations are indicated only if limb saving surgery or an amputation through the affected bone is contemplated.

c) **Treatment of the tumour:** Treatment consists of local control of the tumour, and control of the micro or macro-metastases.

• **Local control:** This is achieved by surgical ablation. Amputation remains the mainstay of treatment. It can be a *palliative* amputation, performed for advanced disease, aiming at pain relief and a better life. When done for a more *definitive* purpose, complete removal of the tumour must be ensured. In the past, high amputations or disarticulations through a joint proximal to the affected bone were done to avoid stump recurrence. With the availability of effective chemotherapy, stump recurrence can be prevented even if the amputation is performed through the affected bone, provided it is performed taking a safe margin beyond the tumour (usually 10 cm from the tumour margin). Table–28.5 gives the recommended levels of amputation in osteosarcoma at common sites.

Recently, with the possibility of early diagnosis, there is a trend towards limb saving surgery. After proper assessment of the local spread of the tumour, a radical excision is performed. Bone defect may be made up with bone grafts, or by using a prosthesis. Effective chemotherapy has played an important role in this change of approach.

**ROLE OF RADIOTherapy:** Radiotherapy is used for local control of the disease for tumours occurring at surgically inaccessible sites, or in patients refusing surgery. Routine pre-operative radiotherapy is no longer a preferred method.

• **Control of distant macro or micro-metastasis:** In the majority of cases, micro-metastasis has already occurred by the time diagnosis is made. These are effectively controlled by adjuvant chemotherapy, immunotherapy etc. A solitary lung metastasis may sometimes be considered suitable for excision.

**ROLE OF CHEMOTHERAPY:** Chemotherapy has revolutionised the treatment of osteosarcoma. It is given pre or post-operative; the basic principle being that the micro-metastases which are supposed to have occurred by the time diagnosis is made, can be effectively controlled. The drugs used are high dose Methotrexate, Citrovorum factor, Endoxan, and sometimes Cisplatinum. Many drug combinations and protocols are under trial. These drugs are highly toxic and should be given in centres where their side effects can be effectively managed.

**ROLE OF IMMUNOTHERAPY:** This is a new concept not yet practiced widely. In this technique, a portion of the tumour is implanted into a sarcoma survivor and is removed after 14 days. The sensitised lymphocytes from the survivor are infused into the patient. These cells then selectively kill the cancer cells.

d) **Follow up:** The patient is checked up every 6-8 weeks. Any evidence of recurrence of the primary tumour, or appearance of the secondary (usually in the chest) is diagnosed early and treated.

A practical plan for treatment management of a case of osteosarcoma is shown in Flow chart-28.1.

**Prognosis:** Without treatment, death occurs within 2 years, usually within 6 months of detection of
metastasis. 5-year survival with surgery alone is 20 per cent. With surgery and adjuvant chemotherapy, a 5-year disease free period is reported to be as high as 70 per cent. A primarily lytic type (telangiectatic) osteosarcoma has the worst prognosis.

SECONDARY OSTEOSARCOMA
This is an osteosarcoma developing in a bone affected by a pre-malignant disease. Some such diseases are as given in Table–28.6. The tumour is usually less malignant than the primary osteosarcoma. It is seen in the older age group (after 40 years). Treatment is along the lines of the conventional osteosarcoma.

PAROSTEAL OSTEOSARCOMA
This is a type of osteosarcoma, arising in the region of the periosteum. It is a slower growing tumour, seen in adults. The common site is lower-end of the femur. Treatment is on the lines of osteosarcoma. Prognosis is better.

Table–28.6: Pre-malignant bone lesions
- Paget’s disease
- Diaphysial aclasis
- Enchondromatosis
- Post-radiation
**EWING’S SARCOMA**

This is highly malignant tumour occurring between the age of 10-20 years, sometimes up to 30 years.

**Pathology:** Following are some of the important pathological features:

- **Bones affected:** It commonly occurs in long bones (in two-third cases), mainly in the femur and tibia. About one-third of cases occur in flat bones, usually in the pelvis and calcaneum. Occasionally, it is known to have a multicentric origin.
- **Site:** The tumour may begin anywhere, but diaphysis of the long bone is the most common site.
- **Gross pathology:** The tumour characteristically involves a large area, or even the entire medullary cavity. The tumour tissue is grey white. It is soft and may be thin, almost like pus. The bone may be expanded, and the periosteum elevated, with sub-periosteal new bone formation, often in layers. The tumour ruptures through the cortex early, and extends into the soft tissues.
- **Histopathology:** The tumour comprises of sheets of quite uniform, small cells, resembling lymphocytes. Often, the tumour cells surround a central clear area, forming a pseudo-rosette. The tumour grows fast and metastasises through the blood stream to the lungs and to other bones.

**Clinical features:** The tumour occurs between 10-20 years of age. The patient presents with pain and swelling. There may be a history of trauma preceding onset, but it is usually incidental. Often there is an associated fever, in which case it may be confused with osteomyelitis.

**Examination:** On examination, the swelling is usually located in the diaphysis and has features suggesting a malignant swelling.

**Radiological features:** In a typical case, there is a lytic lesion in the medullary zone of the midshaft of a long bone, with cortical destruction and new bone formation in layers – onion-peel appearance (Fig-28.5). In atypical presentations, the tumour may be located in the metaphysis, and may be confused with osteomyelitis. It may have a predominant soft tissue component with little cortical destruction, and may resemble a soft tissue sarcoma. In flat bones, it is primarily a lytic lesion with hardly any new bone formation.

**Differential diagnosis:** Ewing’s sarcoma can be differentiated from other bone tumours by features given in Table–28.7. From chronic osteomyelitis, it can be differentiated by the following features in the former:

- Sequestrum
- Well-defined cloacae and a rather smooth periosteal reaction
- Located at metaphysis

**Treatment:** This is a highly radio-sensitive tumour, melts quickly but recurs. In most cases, distant metastasis has occurred by the time diagnosis is made. Treatment consists of control of local tumour by radiotherapy (6000 rads), and control of metastasis by chemotherapy. Chemotherapy consists of Vincristine, Cyclophosphamide, and Adriamycin in cycles, repeated every 3-4 weeks for about 12-18 cycles.

**Prognosis:** It is very poor. Bone to bone secondaries are very common. With the availability of potent chemotherapeutic drugs, 5-year survival (which was only 10 per cent), has now improved to 30-40 per cent.

**MULTIPLE MYELOMA**

It is a malignant neoplasm derived from plasma cells.
<table>
<thead>
<tr>
<th>Tumour</th>
<th>Age (Yrs)</th>
<th>Common sites</th>
<th>Location</th>
<th>Clinical features</th>
<th>X-ray picture</th>
<th>Differential diagnosis</th>
<th>Pathology</th>
<th>Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Osteosarcoma</td>
<td>15-25</td>
<td>Lower end of femur, upper end of tibia</td>
<td>Metaphysis</td>
<td>Pain ++</td>
<td>Sun ray appearance, Codman’s triangle, tumour new bone +</td>
<td>Ewing’s tumour</td>
<td>Tumour cells with osteoid or bone formation</td>
<td>Local ablation+ Chemotherapy</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Swelling ++</td>
<td></td>
<td></td>
<td>Alkaline phosphatase increased in 50% cases</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Duration is wks - mths</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ewing’s tumour*</td>
<td>5-15</td>
<td>Femur, tibia flat bones*, multi-centric**</td>
<td>Diaphysis</td>
<td>Pain ++</td>
<td>Onion-peel appearance</td>
<td>Osteosarcoma, Osteomyelitis</td>
<td>Sheaths of round cells</td>
<td>Radiotherapy + Chemotherapy</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Swelling+</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>often fever+</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Duration wks-mths</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Osteoclastoma</td>
<td>20-40</td>
<td>Lower femur upper tibia, Lower radius</td>
<td>Epiphysis region</td>
<td>Pain +</td>
<td>Soap bubble appearance, No tumour new bone</td>
<td>Aneurysmal bone cyst, Fibrous dysplasia</td>
<td>Multi-nucleate giant cells in fibrous stroma</td>
<td>Excision of tumour + Reconstruction</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Swelling +</td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td>Duration - mths</td>
<td></td>
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</tr>
<tr>
<td>Chondrosarcoma</td>
<td>30-60</td>
<td>Flat bones, upper end of femur</td>
<td>Anywhere in the bone</td>
<td>Pain +</td>
<td>Mottled calcification within the tumour</td>
<td>Osteosarcoma</td>
<td>Chondroblasts, and cartilaginous matrix</td>
<td>Local ablation + Radiotherapy</td>
</tr>
<tr>
<td></td>
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<td></td>
<td></td>
<td>Swelling +++</td>
<td></td>
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<td></td>
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<td>Duration is mths-yrs</td>
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* Ewing’s tumour is the commonest malignant tumour of flat bones.

** Ewing’s tumour is the commonest malignant bone tumour which has multicentric origin.
**Pathology:** The neoplasm characteristically affects flat bones i.e., the pelvis, vertebrae, skull, and ribs. It may occur as a solitary lesion (plasmacytoma), multiple lesion (multiple myeloma), extra-medullary myelomatosis or diffuse myelomatosis. The lesions are mostly small and circumscribed. The bone is simply replaced by tumour tissue and there is no reactive new bone formation.

Grossly, the tumour is soft, grey and friable. Microscopically, it consists of sheets of closely packed cells. Typically, the tumour cells have an eccentric nucleus with clumped chromatin.

**Clinical features:** The tumour affects adults above 40 years of age. Men are affected more often than women. Usual presentation is that of multiple site involvement. Common presenting complaint is increasingly severe pain in the lumbar and thoracic spine. *Pathological fractures*, especially of the vertebrae and ribs may result in acute symptoms. The patient is weak, and will have loss of weight. *Neurological symptoms* may result if the tumour presses on the spinal cord or the nerves in the spinal canal. There is local tenderness over the affected bones. There may be no swelling or deformity unless a pathological fracture occurs.

**Investigations:** Following investigations prove helpful:

- **Radiological examination:** Characteristic radiological features are as follows (Fig-28.6):
  - Multiple *punched out lesions* in the skull and other flat bones.
  - Pathological *wedge collapse* of the vertebra, usually more than one, commonly in the thoracic spine. The pedicles are usually spared.
  - Diffuse, severe rarefaction of bones.
  - Erosions of the borders of the ribs.

- **Other investigations** carried out to support the diagnosis of multiple myeloma are as follows:
  - **Blood:** Low haemoglobin, high ESR (usually very high), increased total protein, A/G ratio reversed, increased serum calcium, normal alkaline phosphatase.
  - **Urine:** Bence Jones proteins are found in 30 per cent of cases.
  - **Serum electrophoresis:** Abnormal spike in the region of gamma globulin (myeloma spike) is present in 90 per cent of cases.
  - **Sternal puncture:** Myeloma cells may be seen.
  - **Bone biopsy** from the iliac crest, or a CT guided needle biopsy from the vertebral lesion may show features suggestive of multiple myeloma.
  - **Bone scan:** This may be required in cases presenting as solitary bone lesion, where lesions at other sites may be detected on a bone scan.
  - **Open biopsy:** An open biopsy from the lesion may sometimes be required to confirm the diagnosis.

**Treatment:** It consists of control of the tumour by chemotherapy, and splintage to the diseased part by PoP, brace etc. Radiotherapy plays a useful role in cases with neurological compression, localised painful lesions, fractures and soft tissue masses. Complications like pathological fractures must be prevented by splinting the affected part. Treatment of a pathological fracture can be done by conservative or operative methods as discussed in Chapter 1.

![Fig-28.6 X-rays showing features of multiple myeloma](https://kat.cr/user/Blink99/)
Chemotherapy: Melphalan is the drug of choice. It is given in combination with Vincristine, Prednisolone, and sometimes Cyclophosphamide. The cycles are repeated every 3-4 weeks for 6-12 cycles.

SOME UNCOMMON MALIGNANT TUMOURS

CHONDROSARCOMA
This is a malignant bone tumour arising from cartilage cells. It may arise in hitherto normal bone (primary chondrosarcoma), or in a pre-existing cartilaginous tumour such as enchondroma (secondary chondrosarcoma). It may arise in any bone but is common in flat bones such as scapula, pelvis and ribs.

Diagnosis: It occurs commonly in adults between 30-60 years of age, and is rare in children. The tumour has a wide spectrum of aggressiveness; from low grade malignant to highly malignant. Metastasis occurs through the blood vessels, commonly to the lungs. Presenting symptoms are pain and swelling, often of long duration. X-ray shows erosion of the cortex and bone destruction. The tumour matrix may have mottled calcification, typical of a cartilaginous tumour (Fig-28.7). Diagnosis is confirmed by a biopsy.

Treatment depends upon the behaviour of the tumour. Amputation is necessary for most tumours. In some low grade tumours, after proper assessment, wide resection of the tumour is done (limb saving surgery). Role of chemotherapy and radiotherapy is doubtful.

SYNOVIAL SARCOMA
This is a malignant tumour, histologically a combination of synovial cells and fibroblasts. It occurs most commonly around the knee. It may not necessarily originate from the synovial membrane. More often than not, it is extra-articular in origin. The tumour spreads via the blood vessels, lymphatics, and along the soft tissue planes.

Treatment is by amputation, as for other bone malignancies. Prognosis is poor.

RETICULUM CELL SARCOMA
This is a tumour arising from the marrow reticulum cells. It has a clinical and pathological resemblance to Ewing’s tumour, but different behaviour. It has a more favourable prognosis. Long bones are commonly affected, but it also occurs in flat bones. The most common age group affected is 20-50 years, and males are affected more commonly. Overall, it has a slow rate of growth and metastasises late. Pathological fractures occur commonly.

Radiological findings: Small multiple irregular areas of destruction can be seen in the medulla with hardly any reactive new bone, either within the tumour or in the form of periosteal reaction. This is typically described as moth-eaten appearance.

Treatment: This is a highly radiosensitive tumour. Amputation with radiotherapy gives 5 years survival upto 50 per cent.

METASTASIS IN BONE
Metastatic tumours in the bone are commoner than the primary bone tumours. The tumours most commonly metastasising to bone are carcinoma of the lung in the male and carcinoma of the breast in the female. Other malignancies metastasising to the bone are carcinoma of prostate, carcinoma of thyroid etc.

CLINICAL FEATURES
A patient with secondaries in the bone may present in the following ways:

a) It may be a patient with known primary malignancy, who presents with symptoms suggestive of secondaries in the bone. These
symptoms are: (i) bone pain – in the spine (commonest site), ribs or extremities; and (ii) pathological fracture – commonly in the spine.

b) It may be a patient, not a known case of primary malignancy, who presents with: (i) bone pain, which on subsequent investigations is found to be due to a destructive lesion in the bone; or (ii) a pathological fracture through an area of bone weakened by such a lesion. On further investigations the lesion is found to be a secondary from somewhere else.

Malignancies which are known to present first time with secondaries (with silent primary) are carcinoma of thyroid, renal cell carcinoma, carcinoma of the bladder etc.

INVESTIGATIONS

A case of secondaries in the bone can be investigated as follows:

a) In a case with known primary, a complaint of bone pain may be due to metastatic lesion of the bone. On plain X-rays, 20-25 per cent or more of metastatic deposits are missed. Hence, in a case where bone secondaries are suspected, a bone scan should be performed (Fig-28.8). This also helps in evaluating the extent of spread of metastasis in bones. PET scan is the most recent imaging modality for early detection of metastasis.

b) In a case presenting first time with bony secondaries, a systematic investigation programme is required to detect the primary. In spite of the best efforts, it is not possible to detect the primary in 10 per cent of cases. The following investigations may be carried out:

- **Radiological examination:** Majority of bone secondaries are osteolytic, but a few are osteoblastic. Carcinoma of the prostate in males and carcinoma of the breast in females are the commonest tumours to give rise to sclerotic secondaries in the bone. Vertebral bodies are affected most frequently. Other common sites are the ribs, pelvis, humerus and femur. Secondaries in bone are uncommon distal to the elbow and knee.

- **Blood:** A high ESR, and an elevated serum calcium are indications of bony secondaries in a suspected case. Other tests may be positive, depending upon the nature of the primary e.g., elevated serum acid phosphatase in prostatic malignancy.

- **Other investigations:** These depend upon the site suspected on clinical examination. In a secondary without a known primary, useful investigations are an abdominal ultrasound, Ba studies, IVP, thyroid scan etc.

**Treatment:** It consists of symptomatic relief of pain, prevention of any pathological fracture, and control of secondaries by chemotherapy or radiotherapy, depending upon the nature of the primary tumour. Role of surgery is limited, mostly to the management of pathological fractures.

**TUMOUR-LIKE CONDITIONS OF THE BONE**

**OSTEOCHONDROMA**

This is the commonest benign “tumour” of the bone. It is not a true neoplasm since its growth stops with cessation of growth at the epiphyseal plate. It is a result of an aberration at the growth plate, where a few cells from the plate grow centrifugally as a separate lump of bone. Though the tumour originates at the growth plate, it gets ‘left behind’ as the bone grows in length, and thus comes to lie at the metaphysis. The stalk and part of the head of the tumour are made up of mature bone, but the tip is covered with cartilage.

**Clinical presentation:** The patient, usually around adolescence, presents with a painless swelling around a joint, usually around the knee. There may be similar swellings in other parts of the body in case of multiple exostosis (see page 317).
**Examination:** The swelling has all the features of a benign bony swelling. Usual location is metaphyseal, but often it comes to lie as far as the diaphysis. It may be a sessile or pedunculated swelling. There may be signs suggestive of complications secondary to the swelling. These are: (i) pain due to bursitis at the tip of the swelling or due to fracture of the exostosis; (ii) signs due to compression of the neurovascular bundle of the limb; and (iii) limitation of joint movements due to mechanical block by the swelling. Occasionally, the tumour undergoes malignant transformation (chondrosarcoma). A rapid increase in the size of the tumour and appearance of pain in a hitherto painless swelling may be suggestive of malignant transformation. Diagnosis is made on X-ray where one can see a bony growth made up of mature cortical bone and marrow (Fig-28.9). The cartilage cap is not visible on the X-ray.

**Treatment:** When necessary, the tumour should be excised. The excision includes the periosteum over the exostosis; since leaving it may result in leaving a few cartilage cells, which will grow again and cause recurrence of the swelling.

**ENCHONDROMA**

This is a benign tumour consisting of a lobulated mass of cartilage encapsulated by fibrous tissue. The intercellular matrix may undergo mucoid degeneration. Frequently the fibrous septae dividing the lobules are calcified. The tumour is seen commonly between the ages of 20-30 years. Small bones of the hands and feet are commonly affected. The presenting complaint is a long standing swelling from one or more phalanges or metacarpals, without much pain. The swelling increases in size very slowly, often totally replacing the bone. An X-ray shows expanding lytic lesions in one or more bones (Fig-28.10). Overlying cortices are thinned out. The tumour matrix has stippled calcification.

**Treatment:** An unsightly appearance is generally the indication for treatment. The lesion is curetted thoroughly, and the cavity, if it is big, is filled with bone grafts. Prognosis is good. Although, chondromas in small bones are not known to undergo malignant change, those in the long bones may change into chondrosarcoma.

**Uncommon presentations of Enchondromas (Ollier’s diseases):**

This is a non-hereditary disorder seen in childhood. In this, masses of unossified cartilage persist within the metaphysis of some long bones, usually multiple. Growth at the adjacent epiphyseal plates may be affected, leading to shortening and deformities.

**Maffucci syndrome:** This is a hereditary disorder where multiple enchondromas and cavernous haemangiomas occur together.

**SIMPLE BONE CYST**

This is the only true cyst of the bone, different from other lesions, which though appear clear ‘cyst-like’ on X-ray, are actually osteolytic, some-times solid lesions. Its aetiology is not known. Pathologically, it is a cavity in the bone lined by thin membrane, and contains serous or sero-sanguinous yellow coloured fluid.
Bone Tumours

Diagnosis: It occurs in children and adolescents. The ends of the long bones are the favourite sites; the commonest being the upper-end of the humerus. The cyst itself may not produce many symptoms, and attention is brought to it by a pathological fracture through it. X-rays show a well-defined, lobulated, radiolucent zone in the metaphysis or diaphysis of a bone (Fig-28.11a). Maximum width of the lesion is less than the width of the epiphyseal plate. A lesion close to the epiphyseal plate is considered ‘active’, as against the one away from it – say in the diaphysis.

The other common cyst of the bone with which this lesion often needs to be differentiated is the aneurysmal bone cyst (Fig-28.11b). It also needs to be differentiated from other causes of a solitary cystic lesion in a bone as discussed in Table–28.2.

Treatment: The cyst is known to undergo spontaneous healing, particularly after a fracture. One or two injections of methylprednisolone into the cyst results in healing. Some cases need curettage and bone grafting.

ANEURYSMAL BONE CYST

This is a benign bone lesion occurring in wide age group, and affects almost any bone. It consists of a blood-filled space enclosed in a shell, ballooning up the overlying cortex – hence its name.

Diagnosis: It is common between 10-40 years of age. Common sites are the long bones, usually at their ends. A gradually increasing swelling is the predominant presentation. There is little pain. Often it presents with a pathological fracture. Typical radiological fractures are as follows (Fig-28.11b):

- Eccentric well-defined radiolucent area.
- Expansion of the overlying cortex.
- Trabeculation within the substance of the tumour.

Treatment is by curettage and bone grafting. Recurrence occurs in 25 per cent cases. Some surgeons prefer to excise the lesion en bloc and fill the gap with bone grafts.

FIBROUS DYSPLASIA

This is a disorder in which the normal bone is replaced by fibrous tissue – hence its name. The mass of fibrous tissue thus formed grows inside the bone and erodes the cortices of the bone from within. A thin layer of sub-periosteal bone forms around the mass, so that the bone appears expanded.

CLINICAL FEATURES

It may affect only one bone (monostotic), or many bones (polyostotic). Often in the polyostotic variety, the bones of a single limb are affected. A polyostotic fibrous dysplasia in girls may have precocious puberty and cutaneous pigmentation (Albright’s syndrome). The disease commonly occurs in children and adolescents. Pain, deformity and pathological fracture are the common presenting symptoms. The bones commonly affected are the upper-ends of femur and tibia, and ribs.

Radiologically, the affected bone shows translucent to ground-glass appearance. The lesion is usually
multiloculated, expanding the cortex of the bone (Fig-28.12). Serum alkaline phosphatase is often elevated. Diagnosis is confirmed by biopsy.

Treatment is curettage and bone grafting.

Further Reading

What have we learnt?

- Primary malignant bone tumours occur in children.
- Knee is the commonest site of primary bone tumours.
- GCT is a locally aggressive tumour, sometimes metastasising.
- Secondaries in the bone is a common cause of pathological fractures.
- Limb salvage, rather than amputation is the aim of modern bone tumour surgery.

Additional information: From the entrance exams point of view

<table>
<thead>
<tr>
<th>Tumour</th>
<th>Site</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chondroblastoma</td>
<td>Epiphyseal (most common tumour in this region before puberty)</td>
</tr>
<tr>
<td>Osteoclastoma</td>
<td>Epiphyseal (most common tumour in this region after puberty)</td>
</tr>
<tr>
<td>Chondrosarcoma, osteochondroma, bone cyst, enchondroma, osteosarcoma, osteoelastoma</td>
<td>Metaphyseal</td>
</tr>
<tr>
<td>Ewing’s tumour, lymphoma, multiple myeloma, adamantinoma, osteoid osteoma</td>
<td>Diaphyseal</td>
</tr>
</tbody>
</table>

Most Common sites of individual bone tumours

<table>
<thead>
<tr>
<th>Tumour</th>
<th>Site</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solitary bone cyst</td>
<td>Upper end of humerus</td>
</tr>
<tr>
<td>Aneurysmal bone cyst</td>
<td>Lower limb metaphysis</td>
</tr>
<tr>
<td>Osteochondroma</td>
<td>Distal femur</td>
</tr>
<tr>
<td>Osteoid osteoma</td>
<td>Femur</td>
</tr>
<tr>
<td>Osteoblastoma</td>
<td>Vertebral</td>
</tr>
<tr>
<td>Osteoma</td>
<td>Skull, facial bones</td>
</tr>
<tr>
<td>Enchondroma</td>
<td>Short bones of hand</td>
</tr>
<tr>
<td>Chordoma</td>
<td>Sacrum</td>
</tr>
<tr>
<td>Adamantinoma</td>
<td>Tibia</td>
</tr>
<tr>
<td>Ameloblastoma</td>
<td>Mandible</td>
</tr>
<tr>
<td>Osteoclastoma</td>
<td>Lower end femur</td>
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Contd...
<table>
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<th>Tumour type and appearance</th>
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</thead>
<tbody>
<tr>
<td><strong>Tumour</strong></td>
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<tr>
<td>Fibrous dysplasia</td>
</tr>
<tr>
<td>Multiple myeloma</td>
</tr>
<tr>
<td>Osteosarcoma</td>
</tr>
<tr>
<td>Ewing's sarcoma</td>
</tr>
<tr>
<td>Chondrosarcoma</td>
</tr>
<tr>
<td>Secondary tumours</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Tumour</td>
</tr>
<tr>
<td>Fibrous dysplasia</td>
</tr>
<tr>
<td>Chondrogenic tumours</td>
</tr>
<tr>
<td>Osteogenic tumours</td>
</tr>
<tr>
<td>Unicameral bone cyst</td>
</tr>
</tbody>
</table>
RELEVANT ANATOMY

The intervertebral disc consists of three distinct components — the cartilage end-plates, nucleus pulposus and annulus fibrosus. The cartilage plates are thin layers of hyaline cartilage between adjacent vertebral bodies and the disc proper (Fig-29.1). The disc receives its nutrition from the vertebral bodies via these end-plates, by diffusion.

The *nucleus pulposus* is a gelatinous material which lies a little posterior to the central axis of the vertebrae. It is enclosed in *annulus fibrosus*, a structure composed of concentric rings of fibro-cartilaginous tissue. The nucleus pulposus is normally under considerable pressure and is restrained by the crucible-like annulus. The posterior longitudinal ligament is a strap-like ligament at the back of the vertebral bodies and discs.

PATHOLOGY

The term ‘prolapsed disc’ means the protrusion or extrusion of the nucleus pulposus through a rent in the annulus fibrosus. It is not a one time phenomenon; rather it is a sequence of changes in the disc, which ultimately lead to its prolapse. These changes consist of the following:

a) **Nucleus degeneration**: Degenerative changes occur in the disc before displacement of the nuclear material. These changes are: (i) softening of the nucleus and its fragmentation; and (ii) weaken-
ing and disintegration of the posterior part of
the annulus (Fig-29.2a).

b) **Nucleus displacement**: The nucleus is under
positive pressure at all times. When the
annulus becomes weak, either because a small
area of its entire thickness has disintegrated
spontaneously or because of injury, the nucleus
tends to bulge through the defect (Fig-29.2b).
This is called *disc protrusion*. This tendency is
greatly increased if the nucleus is degenerated
and fragmented. Finally, the nucleus comes
out of the annulus and lies under the posterior
longitudinal ligament; though it has not lost
contact with the parent disc. This is called *disc
extrusion* (Fig-29.2c). The extruded nucleus pulposus becomes flattened,
fibrosed and finally undergoes calcification.
At the same time, new bone formation occurs
at the points where the posterior longitudinal
ligament has been stripped from the vertebral
body and spur formation occurs.

The *site of exit* of the nucleus is usually postero-
lateral (Fig-29.3) on one or the other side.
Occasionally, it can be central (posterior-midline)
disc prolapse. The type of nuclear protrusion may
be: a protrusion, an extrusion or a sequestration.
A dissecting extrusion, (an extrusion with disc
material between the body of the vertebra and
posterior longitudinal ligament, stripping the latter
off the body), may occur. The *commonest level of disc
prolapse* is between L₄-L₅ in the lumbar spine and
C₅-C₆ in the cervical spine. In the lumbar spine, it
is uncommon above L₃-L₄ level.

**Secondary changes associated with disc prolapse**: As a
consequence of disc prolapse, changes occur in the
structures occupying the spinal canal and in the
intervertebral joints. These are as follows:

a) **Changes in structures occupying spinal canal**:
   - Commonly, the unilateral protrusion is in
     contact with the spinal theca and *compresses
     one or more roots* in their extra-thecal course.
     Usually, a single root is affected. Sometimes,
     two roots on the same or opposite sides are
     affected. The nerve root affected is usually
     the one which leaves the spinal canal below
     the *next* vertebra. This is because the root
     at the level of the prolapsed disc leaves
     the canal in the upper-half of the foramen
     (Fig-29.1). Thus, the nerve root affected in
     a disc prolapse between L₄-L₅ vertebrae is
L₄, although it is the L₅ root which exits the canal at this level.

- **Pressure effects on the intra-thecal roots of the cauda equina** may occur by a sudden large disc protrusion in the spinal canal and may present as cauda equina syndrome. This is uncommon.

b) **Changes in the intervertebral joints**: With the loss of a part of the nucleus pulposus and its subsequent fibrosis, the height of the disc is reduced. This affects the articulation of the posterior facet joints. The incongruity of the facet articulation leads to degenerative arthritis.

**DIAGNOSIS**

The diagnosis is mainly clinical. Investigations like CT scan and MRI scan may be done to confirm the diagnosis, especially if surgery is being considered.

**CLINICAL FEATURES**

The patient is usually an adult between 20-40 years of age, with a sedentary lifestyle. The commonest presenting symptom is low back pain with or without the pain radiating down the back of the leg (sciatica). A preceding history of trauma is present in some cases. In a few cases, there is a history of exertion such as having lifted something heavy or pushed something immediately preceding a sudden onset backache. The following symptoms are common:

- **Low backache**: The onset of backache may be acute or chronic. An acute backache is severe with the spine held rigid by muscle spasm, and any movement at the spine painful. The patient may be able to go about with difficulty. In extreme cases, he is completely incapacitated, any attempted movement producing severe pain and spasm. In chronic backache, the pain is dull and diffuse, usually made worse by exertion, forward bending, sitting or standing in one position for a long time. It is relieved by rest.

- **Sciatic pain**: This is usually associated with low back pain, but may be the sole presenting symptom. The pain radiates to the gluteal region, the back of the thigh and leg. The pattern of radiation depends upon the root compressed. In S₁ root compression, the pain radiates to the posterolateral calf and heel. In L₅ root compression the pain radiates to the anterolateral aspect of the leg and ankle. In a disc prolapse at a higher level (L₂-L₃ etc.), the pain may radiate to the front of the thigh. Often the radiation may begin on walking, and is relieved on rest (neurological claudication).

- **Neurological symptoms**: Sometimes, the patient complains of paraesthesias, most often described as ‘pins and needles’ corresponding to the dermatome of the affected nerve root. There may be numbness in the leg or foot and weakness of the muscles. In cases with large disc material compressing the theca and roots, a cauda equina syndrome results, where the patient has irregular LMN type paralysis in the lower limbs, bilateral absent ankle jerks, with hypoesthesia in the region of L₅ to S₄ dermatomes and urinary and bowel incontinence.

**EXAMINATION**

The back and limbs are examined with the patient undressed. The following observations are made:

- **Posture**: The patient stands with a rigid, flattened lumbar spine. The whole trunk is shifted forwards on the hips (Fig-29.4). The trunk is tilted to one side (sciatic tilt or scoliosis). The sideways tilt tends to exaggerate on attempted bending forwards.

- **Movements**: The patient is unable to bend forwards; any such attempt initiates severe muscle spasm in the paraspinous muscles.

- **Tenderness**: There is diffuse tenderness in the lumbo-sacral region. A localised tenderness in the midline or lateral to the spinous process is found in some cases.
Table–29.1: Neurological deficit in disc prolapse

<table>
<thead>
<tr>
<th>Level</th>
<th>Nerve root affected</th>
<th>Motor weakness</th>
<th>Sensory loss</th>
<th>Reflexes</th>
</tr>
</thead>
<tbody>
<tr>
<td>L₅-S₁</td>
<td>S₁ root</td>
<td>Weakness of plantar-flexors of the foot</td>
<td>Over lateral side of the foot</td>
<td>Ankle jerk sluggish or absent</td>
</tr>
<tr>
<td>L₄-L₅</td>
<td>L₃ root</td>
<td>Weakness of EHL* and dorsiflexors of the foot</td>
<td>Over dorsum of the foot and lateral side of the leg</td>
<td>Ankle jerk normal</td>
</tr>
<tr>
<td>L₃-L₄</td>
<td>L₄ root</td>
<td>Weakness of extensors of the knee</td>
<td>Over great toe and medial side of the leg</td>
<td>Knee jerk sluggish or absent</td>
</tr>
</tbody>
</table>

*EHL: extensor hallucis longus

- **Straight leg raising test (SLRT):** This test indicates nerve root compression (details in Annexure-III). A positive SLRT at 40° or less is suggestive of root compression. More important is a positive contralateral SLRT.
- **Lasegue test:** This is a modification of SLRT where first the hip is lifted to 90° with the knee bent. The knee is then gradually extended by the examiner. If nerve stretch is present, it will not be possible to do so and the patient will experience pain in the back of the thigh or leg.
- **Neurological examination:** A careful neurological examination would reveal a motor weakness, sensory loss or loss of reflex corresponding to the affected nerve root. Of special importance is the examination of the muscles of the foot supplied by L₄, L₅, and S₁ roots, as these are the roots affected more commonly. The extensor hallucis longus muscle is exclusively supplied by L₅ root and its weakness is easily detected by asking the patient to dorsiflex the big toe against resistance. Sensory loss may merely be the blunting of sensation or hypoaesthesia in the dermatome of the affected root. Table–29.1 gives the neurological findings as a result of compression of different roots.

**INVESTIGATIONS**

**Plain X-ray:** It does not show any positive signs in a case of acute disc prolapse. X-rays are done basically to rule out bony pathology such as infection etc. In a case of chronic disc prolapse, the affected disc space may be narrowed and there may be lipping of the vertebral margins posteriorly.

**Myelography:** With the availability of non-invasive imaging techniques like the CT scan, the usefulness of myelography has become limited. It is performed in cases where precise localisation of the neurological signs is not possible. It is also used in cases where facilities for a CT scan are not available. Following myelographic features suggest disc prolapse:
- Complete or incomplete block to the flow of dye at the level of a disc
- An indentation of the dye column

![Myelogram showing disc prolapse](https://kat.cr/user/Blink99/)

- **Root cut off sign:** Normally, the dye fills up the nerve root sheath. In cases where a lateral disc prolapse is pressing on the nerve root, the sheath may not be filled. It appears on the X-ray, as an abrupt blunting of the dye column filling the root sheath (Fig-29.5).

![CT scan showing a large central disc prolapse](https://kat.cr/user/Blink99/)
mimic a disc prolapse. These include ankylosing spondylitis, vascular insufficiency, extra-dural tumour, spinal tuberculosis etc. (Ref. page 185).

TREATMENT

PRINCIPLES OF TREATMENT

Aim of treatment is to achieve remission of symptoms, mostly possible by conservative means. Cases who do not respond to conservative treatment for 3-6 weeks, and those presenting with cauda equina syndrome may require operative intervention.

CONSERVATIVE TREATMENT

This consists of the following:
- **Rest:** It is most important in the treatment of a prolapsed disc. Rest on a hard bed is necessary for not more than 2-4 days.
- **Drugs:** These consist mainly of analgesics and muscle relaxants.
- **Physiotherapy:** This consists of hot fomentation, gentle arching exercises, etc.
- **Others:** These consist of lumbar traction, transcutaneous electrical nerve stimulation (TENS) etc.

OPERATIVE TREATMENT

Indications for operative treatment are: (i) failure of conservative treatment; (ii) cauda equina syndrome; and (iii) severe sciatic tilt. The disc is removed by the following techniques (Fig-29.8):
- **Fenestration:** The ligamentum flavum bridging the two adjacent laminae is excised and the spinal canal at the affected level exposed.
- **Laminotomy:** In addition to fenestration, a hole is made in the lamina for wider exposure.

DIFFERENTIAL DIAGNOSIS

A prolapsed disc is a common cause of low backache, especially the backache associated with sciatic pain. One must be extremely cautious and avoid misdiagnosing other diseases that may

CT scan: Normally, in an axial cut section, the posterior border of a disc appears concave. In a case where there is disc prolapse, it will appear flat or convex. There will be loss of pre-thecal fat shadow normally seen between the posterior margin of the disc and theca. The herniated disc material can be seen within the spinal canal, pressing on the nerve roots or theca (Fig-29.6).

MRI Scan: This is the investigation of choice. It shows the prolapsed disc, theca, nerve roots etc. very clearly (Fig-29.7).

Electromyography (EMG): Findings of denervation, localised to the distribution of a particular nerve root, helps in localising the offending disc in cases with multiple disc prolapse. This test is rarely required.
Prolapsed Intervertebral Disc

- **Hemi-laminectomy**: The whole of the lamina on one side is removed.
- **Laminectomy**: The laminae on both sides, with the spinous process, are removed. Such a wide exposure is required for a big, central disc producing cauda equina syndrome.

**CHEMONUCLEOSIS**

In this technique, an enzyme (chymopapain) with the property of dissolving fibrous and cartilaginous tissue, is injected into the disc, under X-ray control. This leads to dissolution and fibrosis of the disc and thus relief of symptoms. It can be done through a few puncture wounds.

**PERCUTANEOUS DISCECTOMY**

This is a more recent technique where the disc is removed by using an endoscope. Fine endoscopic instruments or laser probes are inserted percutaneously through small stab wounds. Though a minimally invasive technique, its indications are limited, and one requires adequate instrumentation and training.

**CERVICAL DISC PROLAPSE**

Prolapse of the intervertebral disc in the cervical spine is much less common than it is in the lumbar spine. The disc between C₅-C₆ is the one affected most frequently. Postero-lateral protrusion is the commonest. A typical patient presents with a vague history of injury to the neck, often a jerk or a twisting strain. Symptoms may begin hours after the episode of injury. The neck becomes stiff and the pain radiates down the shoulder to the outer aspect of the limb, up to the thumb. Paraesthesias may be felt in the hand. On examination, it may be possible to localise the neurological deficit to a particular nerve root, usually C₅. In some cases, there may be signs of cord compression from the front (UMN signs). X-rays do not show any abnormality. MRI scan is the imaging modality of choice but should be done if operative intervention is contemplated.

**TREATMENT**

There is a strong tendency to spontaneous recovery. Cases may present with signs of cord compression or root compression in the upper limb. Such cases may require surgery. The disc is exposed from the front and the material removed.

**Further Reading**


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**What have we learnt?**

- Disc prolapse is common at L₅-S₁.
- SLRT, particularly contralateral positive SLRT, is highly suggestive of disc prolapse.
- Treatment depends upon the stage of the disease.
Approach to a Patient with Back Pain

Table 30.1: Causes of low back pain

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<th><strong>Causes</strong></th>
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<td>Ankylosing spondylitis</td>
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<td>– Osteoid osteoma</td>
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<td>– Eosinophilic granuloma</td>
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<td>Malignant</td>
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<td>– Primary: Multiple myeloma, Lymphoma</td>
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<td>– Secondaries from other sites</td>
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<td><strong>Metabolic causes</strong></td>
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<td><strong>Miscellaneous causes</strong></td>
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<td>Functional back pain</td>
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<td>Postural back pain</td>
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<tr>
<td>– Protuberant abdomen</td>
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<tr>
<td>– Occupational bad posture</td>
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<td>– Habitual bad posture</td>
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LOW BACK PAIN

Back pain is an extremely common human phenomenon, a price mankind has to pay for their upright posture. According to one study, almost 80 per cent of persons in modern industrial society will experience back pain at some time during their life. Fortunately, in 70 per cent of these, it subsides within a month. But, in as many as 70 per cent of these (in whom pain had subsided), the pain recurs.

CAUSES

The specific aetiology of most back pains is not clear. Table 30.1 gives some common causes of back pain. Postural and traumatic back pains are among the commonest. Back pain could be a feature of an extra-spinal disease like a genitourinary or gynaecological disease. The following findings in the clinical examination are helpful in reaching a diagnosis.

HISTORY

Age: Some diseases are commoner at a particular age. Back pain is uncommon in children, but if present, it is often due to some organic disease. This is different from adults, in whom psychological factors play an important role in producing back pain. In adolescents, postural and traumatic back pain are commoner. In adults, ankylosing spondylitis and disc prolapse are common. In elderly persons, degenerative arthritis, osteo-
porosis and metastatic bone disease are usually the cause.

**Sex:** Back pain is commoner in women who have had several pregnancies. Lack of exercise leading to poor muscle tone, and nutritional osteomalacia are contributory factors in these patients. Some women put on a lot of weight during pregnancy, and later develop mechanical back pain.

**Occupation:** A history regarding the patient’s occupation may provide valuable clues to risk factors responsible for back pain. These are often not apparent to the patient, and could be a part of his ‘routine’. People in sedentary jobs are more vulnerable to back pain than those whose work involves varied activities. Back pain is common in surgeons, dentists, miners, truck drivers etc.

**Past history:** A past history of having suffered from a spinal disease such as a traumatic or inflammatory disease may point to that as the possible cause of back pain.

**Features of pain:** The following features are to be noted:

- **Location:** Pain may be located in the lower, middle or upper back. Disc prolapse and degenerative spondylitis occur in the lower lumbar spine; infection and trauma occur in the dorso-lumbar spine.
- **Onset:** Often, there is a history of significant trauma immediately preceding an episode of back pain, and may indicate a traumatic pathology such as a fracture, ligament sprain, muscle strain etc. A precipitating history is present in about 40 per cent cases of disc prolapse. The trauma may not particularly be a significant one. It may be subtle, resulting from a routine activity such as twisting to pull something out of a drawer. Careful questioning regarding leisure activities and exercise is important because inconsistency in activity levels during work and leisure time can precipitate back pain.

  - **Localisation of pain:** Pain arising from a tendon or muscle injury is localised, whereas that originating from deeper structure is diffuse. Often, pain referred to a dermatome of the lower limb, with associated neurological signs pertaining to a particular root, points to nerve root entrapment.

- **Progress of the pain** (Fig-30.1): In traumatic conditions, or in acute disc prolapse, pain is maximum at the onset, and then gradually subsides over days or weeks. Back pain due to disc prolapse often has periods of remissions and exacerbations. An arthritic or spondylitic pain is more constant, and is aggravated by activity. Pain due to infection or tumour takes a progressive course, with nothing causing relief.

- **Relieving and aggravating factors:** Most back pains are worsened by activity and relieved by rest. Pain due to ankylosing spondylitis, and seronegative spond-arthritis (SSA) is typically worse after rest, and improves with activity. Severe back pain at night that responds to aspirin may indicate a benign tumour. Pain initiated on walking or standing and relieved by rest, is a feature of spinal stenosis. An increase in pain during menstruation may indicate a gynaecological pathology.

**Associated symptoms:** The following associated symptoms may point to the cause of back pain:

- **Stiffness:** It is associated with most painful backs, but it is a prominent symptom in pain due to ankylosing spondylitis, more so early in the morning. There may be an associated limitation of chest expansion.

- **Pain in other joints:** In some rheumatic diseases, back pain may be the presenting feature, but on detailed questioning one may get a history of pain and swelling of other joints.

- **Neurological symptoms:** Symptoms like paraesthesias, numbness or weakness may point to a lesion of the nervous tissue, or a lesion in close proximity to it (e.g., a disc prolapse).
• **Extra-skeletal symptoms**: A history suggestive of abdominal complaints, urogenital complaints, or gynaecological complaints may indicate an extra-skeletal cause of back pain.

• **Mental status** of the patient must be judged to rule out any psychological cause of back pain (hysteria, malingering, etc.). A patient suffering from an organic disease may have a significant underlying psychological disturbance also.

**PHYSICAL EXAMINATION**

The patient should be stripped except undergarments, and examined in the standing and lying down positions:

**Standing position**: The following observations are made in the standing position:

• **Position**: Normally a person stands erect with the centre of the occiput in the line with the natal cleft (Fig-30.2), the two shoulders are at the same level, the lumbar hollows are symmetrical and the pelvis is ‘square’. In a case with back pain, look for scoliosis, kyphosis, lordosis, pelvic tilt and forward flexion of the torso on the lower limbs.

• **Spasm**: Muscle spasm may be present in acute back pain and can be discerned by the prominence of the para-vertebral muscles at rest, which stand out on slightest movement.

• **Tenderness**: Localised tenderness may indicate ligament or muscle tear. There may be trigger points or tender nodules in cases of fibrositis (see page 304). Pain originating from the sacro-iliac joint may have tenderness localised to the posterior superior iliac spine.

• **Swelling**: A cold abscess may be present, indicating tuberculosis as the cause.

**Lying down position**: In the supine position the following observations are made:

• **Range of movement**: There is limitation of movement in organic diseases of the spine. One must carefully differentiate spinal movements from the patient’s ability to bend at the hips (Fig-30.3).

**Fig-30.2 Posture and back pain**

Fig-30.3 Bending forwards

**Fig-30.2 Posture and back pain**

**Fig-30.3 Bending forwards**

• **Straight leg raising test (SLRT)**: This is a test to detect nerve root compression (Annexure-III).

• **Neurological examination**: Sensation, motor power and reflexes of the lower limb are examined. This helps in localising the site of spine pathology.

• **Peripheral pulses**: The peripheral pulses should be palpated to detect a vascular cause of low back
pain, which may be due to vascular claudication. The skin temperature in the affected leg may be lower.

- **Adjacent joints**: Often, the pain originates from the hip joints or the sacro-iliac joints, hence these should be examined routinely.
- **Abdominal, rectal or per vaginal examination**: may be done wherever necessary. Chest expansion should be measured in young adults with back pain.

**INVESTIGATIONS**

The diagnosis of back pain is essentially clinical. There is no use getting X-rays done in acute back pain less than 3 weeks duration, as it does not affect the treatment. On the contrary, X-ray examination is a must for back pain lasting more than 3 weeks; it is almost an extension of the clinical examination. There are a number of other investigations like CT scan, MRI scan, bone scan, blood investigations etc. One has to be very thoughtful in ordering these investigations. Order only when you think it is going to change your line of action, or if the clinical diagnosis is doubtful.

- **Radiological examination**: Routine X-rays of the lumbo-sacral spine (AP and lateral) and pelvis (AP) should be done in all cases. These are useful in diagnosing metabolic, inflammatory and neoplastic conditions. Though, X-rays are usually normal in non-specific back pain, these provide a base line. X-rays should be done after preparation of the bowel with laxatives and charcoal tablets.

- **CT scan**: has replaced more invasive techniques like myelography etc. It shows most bony and soft tissue problems around the spine and spinal canal.

- **MRI scan**: is an expensive investigation, now available in big cities. It delineates soft tissues extremely well, and may be needed in some cases.

- **Blood investigations**: These should be carried out if one suspects malignancy, metabolic disorders, or chronic infection (please refer to their respective Chapters for details).

- **Electromyography**: If nerve root compression is a possibility, electromyography (EMG) may be appropriate (please refer to page 69).

- **Bone scan**: It may be helpful if a benign or malignant bone tumour is suspected on clinical examination but is not seen on plain X-rays.

**TREATMENT**

**Principles of treatment**: For specific pathologies, treatment is discussed in respective chapters. Most back pains falling in the ‘non-specific’ category have a set programme of treatment, mostly conservative. It consists of rest, drugs, hot packs, spinal exercises, traction, corset and education regarding the prevention of back pain.

- **Rest**: In the acute phase, absolute bed rest on a hard bed (a mattress is allowed) is advised. Bed rest for more than 2-3 weeks is of no use; rather, a gradual mobilisation using aids like brace is preferred.

- **Drugs**: Mainly analgesic—anti-inflammatory drugs are required. In cases with a stiff spine, muscle relaxants are advised.

- **Physiotherapy**: This consists of heat therapy (hot packs, short-wave diathermy, ultrasonic wave etc.). Gradually, a spinal exercises programme is started.

- **Traction**: It is given to a patient with back pain with lot of muscle spasm. It also sometimes help in ‘forcing’ the patient to rest in the bed.

- **Use of corset**: This is used as a temporary measure in treating acute back pain, in back pain due to lumbar spondylosis, etc.

- **Education**: Patients must be taught what they can do to alleviate the pain and to avoid injury or re-injury to the back. This includes education to avoid straining the back in activities of daily living such as sitting, standing, lifting weight etc. ‘Back Schools’ are formalised approach to this education.

**MAJOR CAUSES OF LOW BACK PAIN**

**CONGENITAL DISORDERS**

- **Spina bifida** (see page 230 also): This and other minor congenital anomalies of the spine are present in about half the population, but are not necessarily the cause of back pain. Therefore, other pathological conditions should be ruled out before diagnosing this as the cause of symptoms. Treatment is as for non-specific back pain.

- **Transitional vertebrae**: A transitional vertebra is the one at the junction of two segments of the spine, so that the characteristics of both segments is present in one vertebra. It is common in the lumbo-sacral region, either as lumbarisation (S₁ becoming L₅) or sacralisation (L₃ fused with the sacrum, either completely or partially).
TRAUMATIC DISORDERS

Back strain (acute or chronic): The terms back strain and back sprain are often used interchangeably. Most often this arises from a ‘trauma’ sustained in daily routine activities rather than from a definite injury. People prone to back strain are athletes, tall and thin people, those in a job requiring standing for long hours and those working in bad postures. Sedentary workers and women after pregnancy are also frequent candidates for back strain. Acute ligament sprain may occur while lifting a heavy weight, sudden straightening from bent position, pushing etc. Treatment is ‘non-specific’ as discussed earlier.

Compression fractures: These fractures occur commonly in the thoraco-lumbar region (see page 267). Treatment depends upon the severity of compression. It is important to be suspicious of any underlying pathology. Diseases such as early secondary deposits in an elderly, may produce a fracture spontaneously, in one or multiple vertebrae.

INFLAMMATORY DISORDERS

Tuberculosis: Spinal tuberculosis is a common cause of persistent back pain, especially in undernourished people living in unhygienic conditions. Early diagnosis and treatment is crucial for complete recovery (details on page 185).

Ankylosing spondylitis: This should be suspected in a young male presenting with back pain and stiffness. Symptoms are worst in the morning and are relieved on walking about. Spinal movements may be markedly limited along with limitation of chest expansion.

DEGENERATIVE DISORDERS

Osteoarthritis: See page 295.

Prolapsed disc: See page 252.

Spinal stenosis: Narrowing of the spinal canal may occur in the whole of the lumbar spine (e.g., achondroplasia), or more often, in a segment of the spine (commonly in the lumbo-sacral region). Stenosis may be in all parts of the canal or only in the lateral part; the latter is called as root canal stenosis. It may give rise to pressure or tension on the nerves of the cauda equina or lumbar nerve roots. Typically, the patient complains of pain radiating down the lower limbs on walking some distance, and is relieved on taking rest for a few minutes (neurological claudication). Diagnosis is confirmed by a CT scan or MRI. Treatment is by decompression of the spinal canal or root canal, as the case may be.

TUMOURS

Both benign and malignant tumours occur in the spine and the spinal canal. Tumours of the spinal canal, usually benign, are classified as extradural or intradural; the latter can be either intra-medullary or extra-medullary. These tumours are usually diagnosed on myelogram or CT scan. Tumours of the spine are mostly malignant, usually secondaries from some other primary tumours (details on page 246). Some commoner tumours of the spine are as discussed below.

Benign tumours: These are uncommon. Osteoid osteoma is the commonest benign tumour of the spine. It causes severe back pain, especially at night. Typically the pain is relieved by aspirin. The tumour, usually the size of a pea, is found in the pedicle or lamina. Haemangioma also occurs in the vertebral body. Meningioma is a common intradural, extra-medullary tumour which presents with back pain or radiating pain.

Malignant tumours: Multiple myeloma is the commonest primary malignancy of the spine. Metastatic deposits are extremely common in the spine because of its rich venous connections, especially with the vertebral venous plexus. Pain often precedes X-ray evidence of a metastatic deposit. By the time a deposit is visible on X-ray, the tumour has replaced about 30 per cent of the bony content of the vertebra. A bone scan can detect the lesion earlier.

OTHER CAUSES

Metabolic disorders: Osteoporosis and osteomalacia are common causes of back pain (see page 307).

Spondylolysis and spondylolisthesis discussed on page 285.

Facet arthropathy and subtle arthritis of the facet joints can result from a degenerative disease and mal-development of the facets (facet tropism).

APPROACH TO A PATIENT WITH BACK PAIN

The source of back pain is difficult to find because of variable factors. The aim is to identify the pathology that needs immediate treatment, such
as an infection, neoplasm, disc prolapse etc. All other back pains are treated as ‘non-specific back pain’ with more or less common treatment programme. While the patient is on this treatment programme, he is reviewed at regular intervals for any additional signs suggesting an organic illness. First establish whether the problem is acute (3 to 6 months) or chronic (longer than 6 months). If it is an acute pain, whether it is related to a definite episode of trauma or is spontaneous in onset. The causes are accordingly worked out (Flow chart-30.1). In cases with chronic back pain, it is helpful to judge whether it is mechanical or inflammatory by asking the patient whether rest
Flow chart-30.1 gives an outline of how to approach a patient with low back pain.

**SCIATICA**

Sciatica is a symptom and not a diagnosis. It means a pain radiating down the back of the thigh and calf. Degenerative arthritis and disc prolapse are the common causes. Some other causes are given in Table–30.2. Broadly, sciatica can either be because of inflammation of the sciatic nerve or because of compression of one of the roots constituting the sciatic nerve.

_Further Reading_


<table>
<thead>
<tr>
<th>Table–30.2: Causes of sciatica</th>
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<tr>
<td><strong>Inflammatory</strong></td>
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<tr>
<td>• Sciatic neuritis</td>
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<td>• Arachnoiditis</td>
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<tr>
<td><strong>Nerve root compression</strong></td>
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<tr>
<td>• Compression in the vertebral canal by disc, tumour, tuberculosis</td>
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<tr>
<td>• Compression in the intervertebral foramen due to root canal stenosis because of OA, spondylolisthesis, facet arthropathy or tumours</td>
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<tr>
<td>• Compression in the buttock or pelvis by abscess, tumour, haematoma</td>
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brings relief or makes the pain worse. Accordingly, further signs and symptoms help in diagnosis.

**What have we learnt?**

- 70 per cent of acute back pains recover with rest.
- There are two types of back pain: (a) Inflammatory, which are worst in the morning (after rest); and (b) Mechanical, which come up after exertion.
- Treatment depends upon the cause.
Fractures and dislocations of the spine are serious injuries because they may be associated with damage to the spinal cord or cauda equina. Thoraco-lumbar segment is the commonest site of injury; lower cervical being the next common.

About 20 per cent of all spinal injuries result in a neurological deficit in the form of paraplegia in thoraco-lumbar spine injuries or quadriplegia in cervical spine injuries. Often, the patient does not recover from the deficit, resulting in prolonged invalidism or death.

RELEVANT ANATOMY

STRUCTURE
The vertebral column consists of 33 vertebrae (7 cervical, 12 dorsal, 5 lumbar, 5 sacral and 4 coccygeal) joined together by ligaments and muscles. Each vertebra consists of an anterior body and a posterior neural arch (Fig-31.1). Each vertebral body has a central part of cancellous bone and a peripheral cortex of compact bone. The margins of the upper and the lower surfaces of the vertebral body are thickened to form vertebral...
The neural arch is constituted by pedicles, laminae, spinous process and articulating facets. Between any two vertebrae is a strong ‘cushion’—the intervertebral disc. It consists of two portions, a central nucleus pulposus and a peripheral annulus fibrosus. The nucleus pulposus is a remnant of the notochord and is made up of muco-gelatinous material. The annulus fibrosus is made up of fibrous tissue and surrounds the nucleus pulposus.

**ARTICULATION**

The entire vertebral column has similar articulation (except atlanto-axial joint). The vertebral bodies are primarily joined by intervertebral discs. Anteriorly, the vertebral bodies are connected to one another by a long, strap-like, anterior longitudinal ligament and posteriorly by a similar posterior longitudinal ligament.

The neural arches of adjacent vertebrae articulate through facet joints. These are synovial joints with a thick capsule. The adjacent laminae are joined together by a thick elastic ligament, the ligamentum flavum. Interspinous ligaments connect the adjacent spinous processes. The supraspinous ligament connects the tips of the adjacent spinous processes. Inter-transverse ligaments connect the adjacent transverse processes. These ligaments are together often termed the posterior ligament complex.

The direction and size of the articular facets forming the facet joints is different in different parts of the spine (Fig-31.2). In the cervical spine, they are short and more horizontally placed, becoming stouter and more vertical lower down the vertebral column. The facets of the lumbar spine are stout and vertically placed, hence pure dislocation (without associated fracture) does not occur in this region.

**BIOMECHANICS OF INJURY**

**MODE OF INJURY**

A fall from height, e.g., a fall from a tree, is the commonest mode of sustaining a spinal injury in developing countries. In developed countries, road traffic accidents account for the maximum number. Other modes are: fall of a heavy object on the back e.g., fall of a rock onto the back of a miner, sports injuries etc.

**STABLE AND UNSTABLE INJURIES**

For purpose of treatment, it is crucial to assess the stability of an injured spine. A stable injury is one where further displacement between two vertebral bodies does not occur because of the intact ‘mechanical linkages’. An unstable injury is one where further displacement can occur because of serious disruption of the structures responsible for stability. Often, it is difficult to decide with some surety whether the spine is stable; in all such cases it is safer to treat them as unstable injuries.

Recent biomechanic studies show that from the viewpoint of stability, the spine can be divided into three columns: anterior, middle and posterior (Fig-31.3). The anterior column consists of the anterior longitudinal ligament and the anterior part of annulus fibrosus along with the anterior half of the vertebral body. The middle column consists of the posterior longitudinal ligament and the posterior part of the annulus fibrosus along with the posterior half of the vertebral body. The posterior...
column consists of the posterior bony arches along with the posterior ligament complex.

In different spinal injuries, the integrity of one or more of these columns may be disrupted, resulting in threat to the stability of the spine. When only one column is disrupted (e.g., a wedge compression fracture of the vertebra) the spine is stable. When two columns are disrupted (e.g., a burst fracture of the body of the vertebra) the spine is considered unstable. When all the three columns are disrupted, the spine is always unstable (e.g., dislocation of one vertebra over other).

CLASSIFICATION
Spinal injuries are best classified on the basis of mechanism of injury into the following types:

- Flexion injury
- Flexion-rotation injury
- Vertical compression injury
- Extension injury
- Flexion-distraction injury
- Direct injury
- Indirect injury due to violent muscle contraction

FLEXION INJURY
This is the commonest spinal injury.

Examples: (i) heavy blow across the shoulder by a heavy object; (ii) fall from height on the heels or the buttocks (Fig-31.4).

Results: In the cervical spine, a flexion force can result in: (i) a sprain of the ligaments and muscles of the back of the neck; (ii) compression fracture of the vertebral body, C₅ to C₇; and (iii) dislocation of one vertebra over another (commonest C₅ over C₆). In the dorso-lumbar spine, this force can result in the wedge compression of a vertebra (L₁ commonest, followed by L₂ and D₁₂). It is a stable injury if compression of the vertebra is less than 50 per cent of its posterior height.

FLEXION-ROTATION INJURY
This is the worst type of spinal injury because it leaves a highly unstable spine, and is associated with a high incidence of neurological damage.

Examples: (i) heavy blow onto one shoulder causing the trunk to be in flexion and rotation to the opposite side; (Fig-31.5) (ii) a blow or fall on postero-lateral aspect of the head.

Results: In the cervical spine this force can result in: (i) dislocation of the facet joints on one or both sides; and (ii) fracture-dislocation of the cervical vertebra. In the dorso-lumbar spine, this force can result in a fracture-dislocation of the spine. Here one vertebra is twisted off in front of the one below it. While dislocating, the upper vertebra takes a slice of the body of the lower vertebra with it. There is extensive damage to the neural arch and posterior ligament complex. It is a highly unstable injury.

VERTICAL COMPRESSION INJURY
It is a common spinal injury.

Examples: (i) a blow on the top of the head by some object falling on the head; (ii) a fall from height in erect position (Fig-31.6).

Results: In the cervical spine, this force results in a burst fracture i.e., the vertebral body is crushed
This injury results in a chip fracture of the anterior rim of a vertebra. Sometimes, these injuries may be unstable.

**FLEXION-DISTRACTION INJURY**

This is a recently described spinal injury, being recognised in western countries where use of a seat belt is compulsory while driving a car.

**Example:** With the sudden stopping of a car, the upper part of the body is forced forward by inertia, while the lower part is tied to the seat by the seat belt. The flexion force thus generated has a component of ‘distraction’ with it (Fig-31.8).

**Results:** It commonly results in a horizontal fracture extending into the posterior elements and involving a part of the body. It is termed a ‘Chance fracture’. It is an unstable injury.

**EXTENSION INJURY**

This injury is commonly seen in the cervical spine.

**Examples:** (i) motor vehicle accident – the forehead striking against the windscreen forcing the neck into hyperextension; (ii) shallow water diving – the head hitting the ground, extending the neck (Fig-31.7).

**DIRECT INJURY**

This is a rare type of spinal injury.

**Examples:** (i) bullet injury; (ii) a *lathi* blow hitting the spinous processes of the cervical vertebrae.

**Results:** Any part of the vertebra may be smashed by a bullet, but, a *lathi* blow generally causes a fracture of the spinous processes only.

**VIOLENT MUSCLE CONTRACTION**

This is a rare injury.

**Example:** Sudden violent contraction of the psoas.

**Results:** It results in fractures of the transverse processes of multiple lumbar vertebrae. It may...
be associated with a huge retro-peritoneal haematoma.

CLINICAL FEATURES

Presenting complaints: A patient with a spinal injury may present in the following ways:

- Pain in the back following a severe violence to the spine: The history is often so classic that one can predict the type of injury likely to have been sustained. At times the pain is slight, and one may not even suspect a spinal injury. Sometimes, a mild compression fracture of a vertebra may occur from a little jerk in the osteoporotic spine of an elderly person.

- Neurological deficit: Sometimes, a patient is brought to the hospital with complaints of inability to move the limbs and loss of sensation. Mostly there is a history of violence to the spine immediately preceding the onset of these complaints. Sometimes, the paralysis may ensue late, or may extend proximally due to traumatic intra-spinal haemorrhage.

EXAMINATION

A patient with suspected spinal injury should be treated as if it were certain unless proved otherwise on further clinical examination and investigation. Utmost care is required during examination and moving such a patient. Examination consists of the following:

- General examination: A quick general examination should be carried out to evaluate any hypovolaemic shock and associated injuries to the head, chest or abdomen.

- Neurological examination: It is carried out before examining the spine per se. By doing so, it will be possible to find the expected segment of vertebral damage. The level of motor paralysis, loss of sensation and the absence of reflexes are a guide to the neurological level of injury. It is easy to calculate the expected vertebral level from the neurological level (Table–23.4 on page 185).

- Examination of the spine: In a patient with a suspected spinal injury, utmost care must be observed during examination of the spinal column. If such care is not observed, in an unstable spine, movement at the fracture site may cause damage to the spinal cord. The patient should be tilted by an assistant just enough to permit the surgeon’s hand to be introduced under the injured segment. One may be able to feel the prominence of one or more of the spinous processes, tenderness, crepitus or haematoma at the site of injury.

INVESTIGATIONS

Good antero-posterior and lateral X-rays centering on the involved segment provide reasonable information about the injury. Sometimes, special imaging techniques are required e.g., CT scan, MRI etc.

Plain X-rays: This is helpful in: (i) confirmation of diagnosis; (ii) assessment of mechanism of injury; and (iii) assessment of the stability of the spine. Following features may be noted on plain X-rays (Fig-31.9).

- Change in the general alignment of the spine i.e., antero-posterior bending (kyphosis) or sideways bending (scoliosis).
- Reduction in the height of a vertebra.
- Antero-posterior or sideways displacement of one vertebra over another.
- Fracture of a vertebral body.
- Fracture of the posterior elements i.e., pedicle, lamina, transverse process etc.

Occasionally, plain X-rays may appear normal in the presence of a highly unstable spinal injury. This is commonly seen in ‘whiplash’ injury to the cervical spine where all the three columns of the spine are disrupted in a sudden hyperflexion followed by sudden hyperextension of the neck e.g., after the...
sudden stopping of a car. Sometimes, a dislocation of the cervical spine may be spontaneously reduced so that there are only minimal findings on X-ray. Following are some of the radiological features suggestive of an unstable injury:
- Wedging of the body with the anterior height of the vertebra reduced more than half of the posterior height.
- A fracture-dislocation on X-ray.
- Rotational displacement of the spine.
- Injury to the facet joints, pedicle or lamina.
- An increase in the space between the adjacent spinous processes as seen on a lateral X-ray.

**CT scan:** This can be done, where facility for MRI is not available. One can see the damaged structures more clearly, and make note of any bony fragments in the canal.

Essential features of different types of spinal injuries is given in Table 31.1.

### Treatment

The treatment of spinal injuries can be divided into three phases, as in other injuries:

- **Phase I** Emergency care at the scene of accident or in emergency department.
- **Phase II** Definitive care in emergency department, or in the ward.
- **Phase III** Rehabilitation.

### Table 31.1: Essential features of different types of spinal injuries

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>Type of injury</th>
<th>Common site</th>
<th>Column failure</th>
<th>X-ray features</th>
<th>Stability</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flexion injury</td>
<td>Compression fracture</td>
<td>L₁ &gt; L₂ &gt; D₁₂, C₅ &gt; C₁</td>
<td>Only anterior column failure</td>
<td>Diminished anterior height of vertebra, posterior part remains intact</td>
<td>Stable</td>
</tr>
<tr>
<td>Flexion-rotation injury</td>
<td>Fracture-dislocation</td>
<td>L₁, D₁₂, C₅ &gt; C₁</td>
<td>All the three columns failure</td>
<td>Direct evidence (subluxation/dislocation)&lt;br&gt;Posterior arch fracture&lt;br&gt;Broken facets&lt;br&gt;Vertebral body offset anteriorly&lt;br&gt;Indirect evidence&lt;br&gt;Broken ribs&lt;br&gt;Broken transverse processes&lt;br&gt;Increased disc height&lt;br&gt;Increased interspinous distance</td>
<td>Unstable</td>
</tr>
<tr>
<td>Vertical compression injury</td>
<td>Burst fracture</td>
<td>C₅, C₆</td>
<td>Anterior &amp; middle columns failure</td>
<td>Diminished anterior and posterior heights of vertebra&lt;br&gt;CT scan may be of help in demonstrating compromise of the spinal canal by bony fragments</td>
<td>May be unstable</td>
</tr>
<tr>
<td>Extension injury</td>
<td>Avulsion fracture of anterior lip of vertebra</td>
<td>C₅, C₆, lumbar spine</td>
<td>Only anterior column failure</td>
<td>Small chip from margin of vertebra, CT scan of no help</td>
<td>Stable</td>
</tr>
<tr>
<td>Flexion-distraction injury</td>
<td>Chance fracture</td>
<td>Dorsal spinel</td>
<td>Middle &amp; posterior column failure</td>
<td>Horizontal fracture line through posterior arch and posterior part of body of the vertebra</td>
<td>May be unstable</td>
</tr>
<tr>
<td>Direct injury</td>
<td>Fracture of spinous or transverse processes</td>
<td>Any region</td>
<td>Any/All columns failure</td>
<td>Variable</td>
<td>Variable</td>
</tr>
</tbody>
</table>

Injured bones and soft tissues, it shows very well the anatomy of the cord.

**MRI:** It is the best modality of imaging an injured spine. In addition to showing better the details of
**PHASE I - EMERGENCY CARE**

**At the site of accident:** An acute pain in the back following an injury is to be considered a spinal injury unless proved otherwise. Also, all suspected spinal injuries are to be considered unstable unless their stability is confirmed on subsequent investigation. Based on this, a patient with a spinal injury has to be given utmost care right at the site of accident; the basic principle being to avoid any movement at the injured segment.

While moving a person with a suspected cervical spine injury, one person should hold the neck in traction by keeping the head pulled. The rest of the body is supported at the shoulder, pelvis and legs by three other people. Whenever required, the whole body is to be moved in one piece so that no movement occurs at the spine. The same precaution is observed in a case with suspected dorso-lumbar injury.

**In the emergency department:** The patient should not be moved from the trolley on which he is first received until stability of the spine is confirmed. In cases with cervical spine injury, two sand bags should be used on either side of the neck in order to avoid any movement of the neck. A quick general examination of the patient is carried out in order to detect any other associated injuries to the chest, abdomen, pelvis, limbs etc. A thorough neurological examination of the limbs is performed. The spine is examined for any tenderness, crepitus, haematoma etc. X-ray examination, as desired, is requisitioned.

**Medical management of spinal cord injury:** If the patient presents within 8 hours of injury, IV methylprednisolone is administered as a bolus dose followed by maintenance dose. Naloxone, thyrotropin-releasing hormone and GM1 gangliosides have been used.

**PHASE II - DEFINITIVE CARE**

Definitive care of a patient with spinal injury depends upon the stability of the spine and the presence of a neurological deficit. The aim of treatment is: (i) to avoid any deterioration of the neurological status; (ii) to achieve stability of the spine by conservative or operative methods; and (iii) to rehabilitate the paralysed patient to the best possible extent. Treatment of the various type of spinal injuries, as practiced most widely is as discussed below:

**Treatment of cervical spine injuries:** Cervical spine injuries are often associated with head injury, the effect of which may mask the spinal lesion. Therefore, it is necessary to get an X-ray of the cervical spine in any serious case of head injury.

Aim of treatment is to achieve proper alignment of vertebrae, and maintain it in that position till the vertebral column stabilises. Operative stabilization of the fractured spine has become the treatment of choice, as it enhances rehabilitation. Where facilities are not available, reduction and stabilisation can be done by non-operative methods as discussed below:

**Reduction** is achieved by skull traction applied through skull calipers – Crutchfield tongs (Fig-31.10). A weight of up to 10 kg is applied and check X-rays taken every 12 hours. Also a close watch is kept on the patient's neurological status, because it is possible to damage the spinal cord or the medulla by injudicious traction. When it is confirmed on X-rays that reduction has been achieved, light traction is continued for 6 weeks. This is followed by immobilisation of the neck in a moulded PoP cast or a plastic collar. In about 3-4 months, a bony bridge forms between the subluxed vertebrae, and the spine stabilises. The collar can then be discarded.

**Operation:** This may be particularly required for: (i) irreducible subluxation because of 'locking' of the articular processes or (ii) persistent instability. The operation consists of inter-body fusion (anterior fusion) or fusion of the spinous processes and laminae (posterior fusion). Internal fixation may be required.
Even with an apparently good looking alignment on the X-ray, the spine may be highly unstable.

**Fig-31.11 Cervical collar**

**Common cervical spine injuries:**

- **Wedge compression fracture** of the vertebral body: This results from a flexion force. The posterior elements are usually intact so that the injury is stable.

  *Treatment:* Reduction is not required. The neck is kept immobilised with the help of skull traction/sling traction. Once pain and muscle spasm subside, the neck is supported in a cervical collar, PoP cast or a brace (Fig-31.11). Exercises of the neck are started after 8-12 weeks.

- **Burst fracture** of the vertebral body: This results from a vertical compression force. The posterior elements are usually intact but because of the severity of crushing of the vertebra, fracture is considered unstable. It may be associated with a neurological deficit if a broken fragment from the body gets displaced inside the spinal canal.

  *Treatment:* Where there is no neurological deficit, the injury can be treated on the same lines as for wedge compression fractures mentioned above. Management of a patient with neurological involvement is discussed later.

- **Subluxation or dislocation** of the cervical spine:

  A flexion rotation force or a severe flexion force may result in the forward displacement of one vertebra over the other (commonly C₃ over C₄). The displacement may be partial or complete. Sometimes, the displacement may be spontaneously reduced*, leaving a well aligned spine but significantly devoid of supporting ligament; these are unstable injuries. For proper assessment,
A pure dislocation is more often associated with a neurological deficit. The displacement is commonly anterior. Treatment consists of skull traction, followed by immobilisation in a Minerva jacket. In due course, the fracture unites and a bridge of bone joins $C_1$ to $C_2$ anteriorly, thereby stabilising the spine.

- **Clay shoveller’s fracture**: This is a fracture of the spinous process of $D_1$ vertebra. It is caused by muscular action as occurs in shovelling by labourers, hence its name.
- **Displacement of intervertebral disc**: A violent flexion-compression force can sometimes result in sudden prolapse of the nucleus pulposus of a
cervical disc into the vertebral canal resulting in quadriplegia. An early decompression may give good results.

A practical plan of treatment of cervical spine injuries is as shown in Flow chart-31.1.

**Treatment of thoracic and lumbar spine injuries:**
Definitive treatment of a thoracic spine injury depends upon the presence of neurological deficit and on the type of vertebral injury i.e., whether it is stable or unstable. In general, conservative treatment is sufficient for stable injuries. Recently, some centres have adopted a more aggressive approach i.e., treatment by operative methods. Though they have reported good results, to use these methods one needs facilities of a high standard.

**Stable injuries:** Most of these need a period of bed rest and analgesics followed by mobilisation. Initial mobilisation may be by some external support, like a brace etc., but gradually these are discarded and an active programme of rehabilitation continued till full functions are achieved. During the period of bed rest, one must take special care of possible complications such as bed sores, chest infection, urinary tract infection etc.

**Unstable injuries:** These are either associated with a neurological deficit or are likely to develop it during treatment. Open reduction and surgical stabilisation gives the best choice of recovery but

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Flow chart-31.2 Treatment plan of dorso-lumbar spine injury
conventionally, these cases have been treated non-operatively with: (i) bed rest for 6 weeks; (ii) bracing till spine stabilises; and (iii) care of the back.

**Operative intervention:** This is particularly required under the following circumstances:

a) Partial neurological deficit with CT or MRI proven compromise of the spinal canal.
b) Worsening of the neurological deficit.
c) Multiple injured patient.

**Operative methods:** Whenever necessary the following operative methods are performed (Fig-31.14):

- Harrington instrumentation – bilateral
- Luque instrumentation
- Hartshill rectangle fixation
- Pedicle screw fixation
- Moss Miami system

Flow chart-31.2 shows a practical plan of treatment of injuries to dorso-lumbar spine.

**Further Reading**

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**What have we learnt?**

- Spinal injuries are a complex combination of bony and neural injuries.
- Meticulous care, right from the scene of injury, till final rehabilitation is necessary to prevent neurological damage.
- Stability of the injured spine is the most important parameter in deciding whether to go for operative or non-operative treatment.
- The trend is towards operative stabilisation of the spinal injuries, and their treatment in specialised centres.

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**Additional information: From the entrance exams point of view**

- Motorcyclist’s fracture is a ring fracture of the skull base.
- Most common cause of spinal cord injury in India (developing countries) is fall from height whereas in developed countries, it is road traffic accidents.
- Dislocation without fracture can be seen in the cervical spine.
- Vertebroplasty: Polymethylmethacrylate is injected into fractured (compressed) vertebral bodies to decrease pain and strengthen the bone. This does not restore the height of the vertebrae or prevent deformity.
- Baloon kyphoplasty: A small balloon is inflated in the compressed vertebral body to restore its height and alignment.
- Both procedures are absolutely contraindicated in infection, untreated coagulopathy and healed osteoporotic fractures.
Only a small proportion of cases of spinal injuries are complicated by injury to the neural structures within the vertebral column. In the cervical spine, it may lead to paralysis of all four limbs (quadriplegia). In thoracic and thoraco-lumbar spine, it may result in paralysis of the trunk and both lower limbs (paraplegia). The terms quadriplegia and paraplegia are sometimes used for incomplete paralysis of all four limbs or the lower limbs respectively.

The commonest spinal injury to be associated with paraplegia is a fracture-dislocation (flexion-rotation injury) of the dorso-lumbar spine. Quadriplegia most commonly results from fracture-dislocation (flexion-rotation injuries) at the C5-C6 junction. Only severely displaced lumbar spine injuries below L1 level, produce cauda equina type of paralysis.

**PATHOLOGY**

The displaced vertebra may either damage the cord (very unlikely), the cord along with the nerve roots lying by its side or the roots alone. Pathologically, damage to neural structures may be a cord concussion, cord transection or root transection (Fig-32.1).

**Cord concussion:** In this type, the disturbance is one of functional loss without a demonstrable anatomical lesion. Motor paralysis (flaccid), sensory loss and visceral paralysis occur below the level of the affected cord segment. Recovery begins within 8 hours, and eventually the patient recovers fully.

**Cord transection:** In this type, the cord and its surrounding tissues are transected. The injury is anatomical and irreparable. Initially, the motor paralysis is flaccid because the cord below the level of injury is in a state of ‘spinal shock’. After some time, however, the cord recovers from shock and acts as an independent structure, without any control from the higher centres. In this state, though the cord manifests reflex activity at spinal level, there is no voluntary control over body parts below the level of injury. There is total loss of sensation and autonomic functions below the level of injury.

The appearance of signs suggestive of reflex cord activity i.e., bulbo-cavernosus reflex, anal reflex and plantar reflex, without recovery of motor power or sensations is an indicator of cord transection. These reflexes usually appear within 24 hours of the injury. In a few days or weeks, the flaccid paralysis (due to spinal shock) becomes spastic, with exaggerated tendon reflexes and clonus. Involuntary flexor spasms at different joints and spasticity leads to contractures. Sensation and autonomic functions never return.

![Fig-32.1 Pathology of neural injury. (a) Only roots affected, (b) Only the cord affected, (c) Roots + cord affected](image-url)
**Root transection:** Spinal nerve roots may be damaged alone in injuries of the lumbar spine, or in addition to cord injury, in injuries of the dorso-lumbar spine. Neurological damage in nerve root injury is similar to that in cord transection except that in the former residual motor paralysis remains permanently flaccid and regeneration is theoretically possible. A discrepancy between the neurological and skeletal levels may occur in spinal injuries below D₁₀ level because the roots descending from the segments higher than the affected cord level may also be transected, thereby producing a higher neurological level than expected.

**Incomplete lesions:** Occasionally, the neurological lesion may be incomplete i.e., affecting only a portion of the cord. In these cases, there is evidence of neurological sparing distal to the injury (perianal sensation sparing is common). Such sparing is an indication of a favourable prognosis. Incomplete lesions may be of the following types.

a) **Central cord lesion:** This is the commonest incomplete lesion. There is initial flaccid weakness followed by a lower motor neurone type of paralysis of the upper limbs and upper motor neurone (spastic) paralysis of the lower limbs, with preservation of bladder control and perianal sensations (sacral sparing).

b) **Anterior cord lesion:** There is complete paralysis and anaesthesia but deep pressure and position sense are retained in the lower limbs (dorsal column sparing).

c) **Posterior cord lesion:** It is a very rare lesion. Only deep pressure and proprioception are lost.

d) **Cord hemi-section (Brown-Sequard syndrome):** There is ipsilateral paralysis and contralateral loss of pain sensation.

**NEUROLOGICAL DEFICIT AND SPINAL INJURIES**

**Cervical spine:** In these injuries, the segmental level of the cord transection nearly always corresponds to the level of bony damage. A high cervical cord transection (above C₅) is fatal because all the respiratory muscles (thoracic and diaphragmatic) are paralysed. Transection at the C₅ segment results in paralysis of the muscles of the upper limbs, thorax, trunk, and lower limbs, with loss of sensation and visceral functions. With transection at level below the C₅ segment, some muscles of the upper limbs are spared, resulting in characteristic deformities, depending upon the level.

**Thoracic lesion (between T₁ and T₁₀):** In cord transection from T₁ to T₁₀, trunk and lower limb muscles are paralysed. At the tenth thoracic vertebra, the corresponding cord segment is L₁, so in injuries at this level, only the lower limbs are affected.

**Dorso-lumbar lesions (between D₇ and L₁):** Between 11th dorsal and 1st lumbar vertebrae lie all the lumbar and sacral segments along with their nerve roots. Hence, injuries at this level cause cord transection with or without involvement of nerve roots. This is the cause of difference in neurological deficit in fractures and fracture-dislocations with apparently similar X-ray appearances. In injuries of the cord with nerve root transection, paralysis in the lower limbs is mixed (UMN+LMN type). It is important to differentiate it from a lesion of cord transection with root escape, as the latter has a better prognosis.

**Lesions below L₁:** This area of the canal has only bunch of nerve roots, which subsequently emerge at successive levels of the lumbo-sacral spine. Thus, injury in this area results in root damage, resulting in flaccid paralysis, sensory loss and autonomic disturbances in the distribution of the affected roots.

**CLINICAL EXAMINATION**

A neurological deficit following trauma to the spine is difficult to miss. More important is to perform a thorough neurological examination to evaluate the following: (i) the level of neurological deficit; (ii) any evidence of an incomplete lesion; and (iii) any indication of complete cord transection.

**INVESTIGATIONS**

**Radiological examination:** Often there is no correlation between the severity of the injury on the X-rays and the degree of neurological deficit.

**CT and MRI scan:** This may be indicated in cases with incomplete paralysis, particularly if it is increasing. It is also indicated in cases where no bony lesions are visible on plain X-rays. MRI has become the imaging modality of choice for these cases.

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* The root, being made up of myelinated fibres, behaves like any other nerve as far as recovery is concerned. But, because the distance between the level of injury and the neuromuscular junction is big, motor recovery is only a theoretical possibility.
TREATMENT

A patient with traumatic paraplegia, wherever possible, should be admitted to specialised units, where necessary facilities for management of these cases are available. In developing countries, these cases are still managed in general hospitals. Treatment can be discussed in 3 phases:

**Phase I**
Emergency care at the scene of accident and in the emergency department

**Phase II**
Definitive care on in-patient basis

**Phase III**
Rehabilitation

**PHASE I – EMERGENCY CARE**
The care in phase I is along the lines already discussed in ‘treatment of spinal injuries’ on page 270.

**PHASE II – DEFINITIVE CARE**
Care in phase II consists of: (i) clinical assessment of the neurological deficit; (ii) radiological and special investigations to understand the type of vertebral lesion, and to detect the possibility of persistent cord compression by a bone fragment in the vertebral canal; and (iii) care of paraplegic in the ward. (i) and (ii) are discussed in Chapter 31; (iii) is being discussed here.

**Ward care of a paraplegic:** Ward care of a traumatic paraplegic or quadriplegic consists of: (i) management of the fracture; (ii) nursing care; (iii) care of the bladder and bowel; and (iv) physiotherapy.

**Management of the fracture:** Treatment of the fracture or fracture-dislocation per se is the same as that for spinal injury at that level without neurological lesion. This is as discussed on page 271. Role of operative treatment is controversial. It consists of stabilisation of the spine by internally fixing it. This ensures better nursing care of the patient but offers no security about the recovery of neurological function. The generally accepted indications for surgery in developing countries, with limited expertise, can be considered as follows:

a) *Incomplete paralysis,* particularly if it is increasing, and a CT scan shows fragments of bone encroaching upon the spinal canal.

b) *Patient with multiple injuries,* in whom it is desirable to stabilise the spine for overall optimum care of the patient.

**Nursing care:** Specialised nursing care has dramatically changed the prognosis of a traumatic paraplegic. It can be considered under the following heads:

a) *Positioning in bed:* The patient is nursed flat on a hard bed with a mattress. The limbs are positioned with pillows so that contractures do not develop; also pressure points are adequately padded (Fig-32.2).

b) *Care of the back:* Frequent turning in bed is vital so that the patient lies for equal periods on his
back and on either side. The bed is kept dry and free of wrinkles. Special beds are available which provide an ease of turning the patient periodically (Stryker frame), and constantly changing pressure-point (water-bed, alpha-bed).

c) **Personal hygiene:** All personal hygiene of the patient from top to toe, is to be looked after. This includes combing hair, cleaning teeth, mouth wash, care of the skin and nails etc.

**Care of the bladder:** Intermittent catheterisation is the **best** but for convenience an indwelling catheter is used. Catheter is changed once a week, and the patient is kept on prophylactic antiseptic drugs. A urine culture is done once every two weeks. As the patient becomes haemodynamically stable, catheter is periodically clamped so that the bladder capacity is maintained.

In most cases of cord transection, satisfactory automatic emptying is established within one to three months of the injury (**automatic bladder**). In a case, where the sacral segments are irrecoverably damaged, as in a cauda equina lesion, reflex emptying does not occur. In such cases, micturition will have to be started or aided by other mechanisms like abdominal straining or manual compression etc. (**autonomous bladder**).

**Care of the bowel:** The patient develops bowel incontinence and constipation. The latter may result in periodic bloating up of the abdomen. A frequent soap water enema or manual evacuation of the bowel may be required.

**Physiotherapy:** Aim of physiotherapy in the initial few weeks is to maintain mobility of the paralysed limbs by moving all the joints through the full range gently, several times a day. Later, in cases where partial recovery occurs, exercises specifically for building up the muscle groups are taught.

**PHASE III – REHABILITATION**

In most cases with traumatic paraplegia and quadriplegia, the deficit is permanent. With concentrated efforts at rehabilitation, a majority of these cases can be made reasonably independent and enabled to lead a useful life within the constraints of their disability. Rehabilitation can be considered under the following headings: (i) physical rehabilitation; (ii) psychological and social rehabilitation; and (iii) economic rehabilitation.

**Physical rehabilitation:** It consists of making the patient as independent in his activities of daily living (ADL) as possible. The patient may be given special appliances like calipers, wheelchair etc. for this.

**Psychological and Social rehabilitation:** Keeping the morale of a paraplegic high is a great challenge. The doctor, nursing staff, family, friends and social organisations have a great role to play in this.

**Economic rehabilitation:** This is an important aspect of rehabilitation of a paraplegic. As soon as the patient is able to do a worthwhile job, efforts should be made to procure some form of renumerative employment for him.

In developed countries, these patients are managed in special **spinal injury centres**. There are now more and more surgeons in these centres who believe in the operative treatment of most cases of paraplegia and quadriplegia. According to them, stabilisation of the spine after reduction of the displacement gives the patient: (a) best chance of relieving compression on the cord, if at all; and, (b) helps in better nursing care of the patient.

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**What have we learnt?**

- Traumatic paraplegia is one of the most common spinal injury.
- Recovery depends upon the nature of neural damage, and whether it is a complete or incomplete cord damage.
- Nursing care during recovery phase is crucial.
- Prolonged rehabilitation is required.
Scoliosis

Scoliosis is the sideways curvature of the spine.

**Classification**

It is of two types: non-structural (transient) and structural (permanent). In structural scoliosis, the vertebrae, in addition to sideways tilt, are rotated along their long axis; in non-structural scoliosis they are not.

**Non-structural scoliosis:** This is a mobile or transient scoliosis. It has three subtypes, as discussed below:

- **Postural scoliosis:** It is the commonest overall type, often seen in adolescent girls. The curve is mild and convex, usually to the left. The main diagnostic feature is that the curve straightens completely when the patient bends forwards.
- **Compensatory scoliosis:** In this type, the scoliosis is a compensatory phenomenon, occurring in order to compensate for the tilt of the pelvis (e.g., in a hip disease or for a short leg). The scoliosis disappears when the patient is examined in a sitting position (in case the leg is short) or when the causative factor is removed.
- **Sciatic scoliosis:** This is as a result of unilateral painful spasm of the paraspinal muscles, as may occur in a case of prolapsed intervertebral disc.

**Structural scoliosis:** It is a scoliosis with a component of permanent deformity. The following are the different subtypes:

- **Idiopathic:** It is the commonest type of structural scoliosis. It may begin during infancy, childhood or adolescence. *Infantile scoliosis* begins in the first year of life, and is different from the other in that, it can be a *resolving or progressive* type. Scoliosis beginning later in life progresses at a variable rate, and leads to an ugly deformity. The deformity is most obvious in thoracic scoliosis because of the formation of a rib hump. In the lumbar region, even a moderate curve goes unnoticed because it gets masked by the compensatory curvature of the adjacent part of the spine. Idiopathic curves progress until the cessation of skeletal growth.
- **Congenital scoliosis:** This type is always associated with some form of radiologically demonstrable...
anomaly of the vertebral bodies (Fig-33.1). These are: (i) hemivertebrae (only one-half of the vertebra grows); (ii) block vertebrae (two vertebral bodies fused); or (iii) an unsegmented bar (a bar of bone joining two adjacent vertebrae on one side, thereby preventing growth on that side). These curves grow, often at a very fast rate. Sometimes, there are associated anomalies in the growth of the neural structures, leading to a neurological deficit in the lower limbs.

- Paralytic scoliosis: An unbalanced paralysis of the trunk muscles results in paralytic scoliosis of the spine. Poliomyelitis is the commonest cause in developing countries. Other common causes are cerebral palsy and muscular dystrophies.

- Other pathologies: There are other causes of structural scoliosis such as neurofibromatosis which produces a sharp kyphoscoliosis.

**PATHOLOGY**

The main pathology is lateral curvature of a part of the spine. This is called the primary curve. The spine above or below the primary curve undergoes compensatory curvature in the opposite direction. These are called the compensatory or secondary curves (Fig-33.2a). The lateral curvature is associated with rotation of the vertebrae. In curves of the thoracic spine, rotation of the vertebrae leads to prominence of the rib cage on the convex side, giving rise to a rib hump (Fig-33.2b).

Any part of the thoraco-lumbar spine may be affected. The pattern of the curve and its natural evolution are fairly constant for each site. The following types are recognised: (i) dorsal scoliosis; (ii) dorso-lumbar scoliosis; and (iii) lumbar scoliosis.

**DIAGNOSIS**

**Clinical features:** In most cases, visible deformity is the only symptom. Pain is occasionally a feature in adults with a long-standing deformity. In exceptional cases of severe long-standing scoliosis, sharp angulation of the spinal cord over the apex of the curve may result in interference with cord functions, leading to a neurological deficit.

**Radiological features:** For proper assessment of scoliosis, a full antero-posterior X-ray of the spine in supine and erect positions, plus a lateral view are necessary. Severity of the curve is measured by Cobb’s angle – an angle between the line passing through the margins of the vertebrae at the ends of the curve (Fig-33.3a). Radiological assessment regarding the likelihood of progress of the curve can be made by looking at the iliac apophysis (Fig-33.3b). It fuses with the iliac bone at maturity and indicates the completion of growth, and thus no possibility of the curve worsening. This is called Reisser’s sign.
Rotation of a vertebra can be appreciated by looking at the position of the spinous processes and pedicles on AP view. Normally, a spinous process is in the centre of the vertebral body. In a case where there is a rotation of a vertebra, the spinous process is shifted to one side (Fig-33.3c). Also, there will be asymmetry in the position of the pedicles on the two sides.

In congenital scoliosis, one may find wedging, hemivertebrae, an unsegmented bone bar between the vertebrae, fused ribs etc. In scoliosis associated with neurofibromatosis an erosion of the vertebral bodies may be seen. Intervertebral foramina may be widened in a dumbell-shaped neurofibroma producing scoliosis.

TREATMENT

Principles of treatment: Aim of treatment is to assess the prognosis of the curve in terms of the visible deformity it is likely to produce. This depends upon: (i) the type of the curve; (ii) age at onset; and (iii) the site of the curve. Congenital curves progress at variable rates depending upon the type of vertebral malformation, but overall they grow faster than idiopathic curves. Neurofibromatotic curves progress faster. In general, younger the patient, the worse the prognosis. Thoracic curves produce the worst deformities.

As soon as it is realised that a curve is likely to progress and result in an ugly deformity, the affected part of the spine is fused. The basic guiding principle is that a straight, stiff spine is better than a curved, flexible one. Treatment of postural curves is non-operative. Proper training and exercises form the mainstay of treatment. Structural curves of less than 30°, and well-balanced double-curves can also be successfully treated by non-operative methods. The following are the indications for surgical intervention:

- Congenital scoliosis, where the radiological signs suggest the possibility of fast progression of the curve, especially those in the thoracic spine.
- Curves showing deterioration radiologically, and are in the region where they are likely to produce ugly deformities at pubertal growth spurt.
- Scoliosis associated with backache.

For all other curves, the patient is started on a non-operative regimen consisting of exercises and a brace. The progress of the curve is monitored clinically and radiologically every 6 months. Following are the non-operative and operative methods of treatment:

Non-operative methods: These consist of exercises to tone up the spinal muscles and give support to the spine. Following supports are commonly used:

- Milwaukee brace: This is named after the city of Milwaukee where it was designed (Fig-33.4).
- Boston brace: It is cosmetically more acceptable.
- Reisser's turn-buckle cast: This is a body cast with a turn-buckle in between. Tightening of the turn-
Scoliosis and Other Spinal Deformities

commonly in tuberculosis where usually two or more vertebrae are affected.

CAUSES
The following are the common causes of diffuse kyphosis:

a) **Postural:** This is the *commonest* type, seen in tall individuals, especially in some tall women, because of their tendency to stand with a forward stoop. It occurs in the upper dorsal spine, and can be corrected by postural training and physiotherapy.

b) **Compensatory:** If there is an exaggerated lumbar lordosis due to some disease, the thoracic spine develops compensatory kyphosis.

c) **Scheurmann’s disease:** It is a common type. There is a gentle round kyphosis in the lower thoracic spine. It is due to osteochondritis affecting the ring-epiphyses of the vertebral bodies. On X-rays, the vertebral bodies appear wedge-shaped, narrower in front. There may be a dull constant pain during early stages of the disease, but later, only kyphosis remains. Conservative treatment is adequate for most patients with pain as the complaint. If the deformity is severe, especially if it is compromising the activities in any way, surgical intervention may be required.

d) **Ankylosing spondylitis:** The disease produces a stiff and kyphotic spine. It begins in young men as low backache, which gradually spreads to affect the whole spine. Chest expansion is reduced because of the limitation of movements at the costo-vertebral joints. In a few cases, hips and shoulders are also affected.

**SPONDYLOLISTHESIS**
Spondylolisthesis is forward displacement of a vertebra over the one below it (Fig-33.5a). It commonly occurs between $L_5-S_1$ and between $L_4-L_5$. Occasionally, the displacement is backwards (retrolisthesis).

**PATHOLOGY**
Normally, forward displacement of a vertebral body is prevented primarily by the engagement of its articular processes with that of the vertebra below it. The attachments of the intervertebral disc and ligaments between vertebrae also check this displacement, but to a small extent. Thus, any defect in this ‘check’ mechanism leads to

Followed by a diagram and references for further reading.
spondylolisthesis. Accordingly, spondylolisthesis has been divided into the following types:

a) **Isthmic**: This is the *commonest* type overall. The lesion is in the pars interarticularis*. Three subtypes are recognised:
   - Lytic: Fatigue fracture of the pars interarticularis
   - Intact but elongated pars interarticularis
   - Acute fracture of the pars interarticularis.

The defect allows the separation of the two halves of the vertebra. The anterior half (i.e., the body with the pedicles and superior articular facet) along with the whole of the spinal column above it, slips forwards over the vertebra below. The posterior half of the affected vertebra (i.e., laminae and inferior articular facets), remain with the lower vertebrae (Fig-33.5b).

b) **Dysplastic**: In this, the *least common* type, there is a congenital abnormality in the development of the vertebrae, so that one vertebra slips over the other.

c) **Degenerative**: This is seen fairly commonly in elderly people. The posterior facet joints becomes unstable because of osteoarthritis, and subluxate. Vertebral displacement is occasionally backwards rather than forwards (retrolisthesis). Displacement is usually not severe, and neurological disturbance is unusual.

d) **Pathological**: This type results from a generalised or localised bone disease weakening the articulation between the vertebrae.

e) **Traumatic**: This is a very rare type, where one vertebra slips over other following an injury.

**DIAGNOSIS**

**Clinical features**: The isthmic type of spondylolisthesis presents in adolescents and young adults. The degenerative type occurs in old age. The presenting symptom is usually backache, with or without sciatica. Symptoms become worse on standing or walking. Sometimes, there may be neurological symptoms in the lower limbs. In a large number of cases, the abnormality is symptomless, and is detected on a routine X-ray taken during screening for a health checkup.

**On examination**, there is often a visible or palpable ‘step’ above the sacral crest due to the forward displacement of the spinal column. There may be increased lumbar lordosis. There may be evidence of stretching of the sciatic nerve, as found by the straight leg raising test (SLRT).

**X-ray examination**: Anterior displacement of one vertebra over other can be seen on a lateral view of the spine (Fig-33.6). The displacement can be graded into four categories depending upon the severity of slip. Grade I spondylolisthesis means vertebral displacement up to 25 per cent of the antero-posterior width of the lower vertebral body, whereas grade IV means the complete forward displacement of the affected vertebra. An *oblique*...
Scoliosis and Other Spinal Deformities

**TREATMENT**

**Principles of treatment:** For a mild symptomless spondylolisthesis, no treatment is required. When symptoms are mild, they are adequately relieved by conservative methods, such as a brace and spinal exercises. When symptoms are moderately severe or more, especially if these hamper the activity of the patient, an operation may be required.

**Methods of treatment:** These consist of conservative and operative methods.

*Conservative methods* consist of rest and external support to the affected segment followed by flexion exercises. The patient is advised to change his job to a physically less demanding one.

*Operative methods* consist of decompression of the compressed nerves if any, followed by fusion of the affected segments of the spine. This is commonly achieved by fusion between the transverse processes of adjacent vertebrae (inter-transverse fusion). Use of internal fixation devices like pedicular screws and rods has helped in early mobilisation of the patient.

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**What have we learnt?**

- Scoliosis is sideways curvature of spine, different from kyphosis and lordosis, which are antero-posterior curves.
- Non-structural scoliosis is transient, whereas structural is permanent.
- Treatment depends upon severity of the scoliotic curve, its location and how it is likely to affect cosmesis and functions.
Arthritis and Related Diseases

**DEFINITIONS**

**Arthritis** is an inflammation of a joint. It is characterised by pain, swelling and limitation of joint movement. The cause may be purely a local pathology such as pyogenic arthritis, or a more generalised illness such as rheumatoid arthritis.

**Arthralgia** is a term used for pain in a joint, without any associated signs of inflammation.

**CLASSIFICATION**

From the clinical viewpoint, arthritis can be divided into two types: (i) monoarthritis; and (ii) polyarthritis. Some common causes of the two types are given in Table–34.1.

**Table–34.1: Types of arthritis**

<table>
<thead>
<tr>
<th>Monoarthritis</th>
<th>Polyarthritis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyogenic arthritis</td>
<td>Rheumatoid arthritis</td>
</tr>
<tr>
<td>Tubercular arthritis</td>
<td>Rheumatic fever</td>
</tr>
<tr>
<td>Haemophilic arthritis</td>
<td>Juvenile chronic polyarthritis</td>
</tr>
<tr>
<td>Secondary osteoarthritis</td>
<td>Primary osteoarthritis</td>
</tr>
<tr>
<td>Gout - sometimes</td>
<td>Seronegative spondarthritis</td>
</tr>
</tbody>
</table>

**RHEUMATOID ARTHRITIS**

Rheumatoid arthritis is a chronic non-suppurative inflammation of the synovial joints diagnosed as per the criteria laid down by American Rheumatism Association in 1987 (Table–34.2).

**AETIOPATHOLOGY**

**Aetiology:** The exact aetiology is not known. Following factors have been thought to play a role in causation of the disease:

**Table–34.2: New diagnostic criteria for rheumatoid arthritis (1987)**

- Morning stiffness
- Swelling of three or more specified joints
- Swelling of joint(s) in the hands and wrist
- Symmetrical swellings
- Rheumatoid nodule
- Rheumatoid factor positive
- X-ray changes – erosion or unequivocal peri-articular osteopenia

If four or more of these are present, it is rheumatoid arthritis

- A genetic predisposition is strongly suspected because of certain histocompatibility markers associated with it (HLA-drw4/HLA-DR1).
- Agents such as mycoplasma, clostridium and some viruses (EB virus) have been implicated in its aetiology.

It is now believed that rheumatoid arthritis results from exposure of a genetically predisposed individual to some infectious agent. This leads to autoimmunity and formation of immune complexes with IgM antibodies in the serum. These immune complexes are deposited in the synovial membrane and initiate a self-perpetuating chronic granulomatous inflammation of the synovial membrane.
Pathology: Initially the synovium becomes oedematous, filled with fibrin exudates and cellular infiltrates. There is an increase in synovial fluid. As the inflammation persists, the synovium gets hypertrophied and surrounds the periphery of the articular cartilage to form a pannus. The articular cartilage loses its smooth shiny appearance. The pannus extends over the cartilage from the periphery and burrows into the subchondral bone. With further progress of the disease, the cartilage becomes worn off and the bone surfaces become raw. The joints gets deformed, initially because of severe muscle spasm associated with pain, but later due to fibrosis of the capsule and other soft tissue structures.

In some cases, adhesions develop between opposing layers of pannus, leading to fibrous ankylosis, and later bony ankylosis. In an advanced disease, the joint capsule gets distended by the hypertrophied synovium and synovial fluid, and the ligaments supporting the joint are stretched, resulting in subluxation of the joint. Osteoporosis develops in the bones adjacent to the diseased joint. Peri-articular tissues, notably tendons and muscles become oedematous and infiltrated with cells, and may rupture spontaneously.

The course of the disease varies from patient to patient. In some, it is no more than a mild arthritis which totally recovers; in others it may be a severe, chronic debilitating disease, ultimately ending up in deformities. A typical case has a history of spontaneous remissions and exacerbations. Some of the factors known to precipitate an attack are physical exertion, psychological stress, infections and occasionally, trauma.

Stages of rheumatoid arthritis: From clinical viewpoint rheumatoid arthritis can be divided into three stages:

1. Potentially reversible soft tissue proliferations: In this stage, the disease is limited to the synovium. There occurs synovial hypertrophy and effusion. No destructive changes can be seen on X-rays.

2. Controllable but irreversible soft tissue destruction and early cartilage erosions: X-rays show a reduction in the joint space, but outline of the articular surfaces is maintained.

3. Irreversible soft tissue and bony changes: The pannus ultimately destroys the articular cartilage and erodes the subchondral bone. The joint becomes ankylosed usually in a deformed position (fibrous ankylosis). It may be subluxated or dislocated.

Associated changes: In rheumatoid arthritis there is sometimes evidence of diffuse vasculitis. The most serious lesions occur in the arterial tree; which may be mild non-necrotising arteritis, or severe and fulminant arteritis akin to polyarteritis nodosa. The latter is fatal.

DIAGNOSIS

Clinical features: It occurs between the age of 20 to 50 years. Women are affected about 3 times more commonly than men. Following presentations are common:

a) An acute, symmetrical polyarthritis: Pain and stiffness in multiple joints (at least four), particularly in the morning, mark the beginning of the disease. This may be followed by frank symptoms of articular inflammation. The joints affected most commonly are the metacarpophalangeal joints, particularly that of the index finger. Other joints affected commonly are as given in Table–34.3.

b) Others: The onset may be with fever, the cause of which cannot be established (PUO), especially in children. Sometimes, visceral manifestations of the disease such as pneumonitis, rheumatoid nodules etc. may antedate the joint complaints.

On examination, one finds swollen boggy joints as a result of intra-articular effusion, synovial hypertrophy and oedema of the peri-articular structures. The joints may be deformed (Table–34.4).
Table–34.4: Deformities in Rheumatoid arthritis

<table>
<thead>
<tr>
<th>Joint</th>
<th>Deformities</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hand</td>
<td>Ulnar drift of the hand</td>
</tr>
<tr>
<td></td>
<td>Boutonniere deformity</td>
</tr>
<tr>
<td></td>
<td>Swan neck deformity</td>
</tr>
<tr>
<td>Elbow</td>
<td>Flexion deformity</td>
</tr>
<tr>
<td>Knee</td>
<td>Early - flexion deformity</td>
</tr>
<tr>
<td></td>
<td>Late - triple* subluxation</td>
</tr>
<tr>
<td>Ankle</td>
<td>Equinus deformity</td>
</tr>
<tr>
<td>Foot</td>
<td>Hallux valgus, Hammer toe, etc.</td>
</tr>
</tbody>
</table>

* Flexion, posterior subluxation and external rotation.

Joints of the hand show typical deformities as shown in Fig-34.1. There may be severe muscle spasm. Range of motion of the joints may be limited. In later stages, the joints may be subluxated or dislocated. There may be fever, rash and signs suggestive of systemic vasculitis. The rash in rheumatoid arthritis is typically non-pruritic and maculo-papular on the face, trunk and extremities.

ExTRA-ARTICULAR MANIFESTATIONS OF RHEUMATOID ARTHRITIS: Although, rheumatoid arthritis is primarily a chronic polyarthritis, extra-articular manifestations are very common, and sometimes govern the prognosis of a case. These are given in Table–34.5.

Investigations: Following investigations are useful:

1) Radiological examination: This consists of X-rays of both hands and of the affected joints. Following features may be present (Fig-34.2):
   - Reduced joint space
   - Erosion of articular margins
   - Subchondral cysts

2) Blood: It shows the following changes:
   - Elevated ESR
   - Low haemoglobin value
   - Rheumatoid factor (RF): This is an auto antibody directed against the Fc fragment of immunoglobulin G (IgG). RF can belong to

Fig–34.1 Deformities in rheumatoid arthritis

Fig–34.2 X-rays of both hands, AP view, showing juxta-articular rarefaction
any class of immunoglobulins i.e., IgG-RF, IgM-RF, IgA-RF, or IgE-RF, but commonly done tests detect only the IgM type of RF. It can be detected in the serum of the patient by the following tests:

- **Latex fixation test:** This is an agglutination test where the antibodies are coated to latex particles. Positivity in titres more than 1/20 is significant. Sensitivity is 80 per cent.

- **Rose-Waaler test:** In this agglutination test sheep’s red blood cells are used as a carrier. Sensitivity is 60 per cent.

- **All patients with positive rheumatoid factor do not have rheumatoid arthritis. Conversely, all patients with rheumatoid arthritis do not have a positive rheumatoid factor. It is the constellation of signs and symptoms, titre in which the RF is positive, presence or absence of other positive tests, etc, which determine whether the patient has rheumatoid arthritis or not.**

- **Synovial fluid examination**—See Table–22.2 on page 177.

- **Synovial biopsy:** This can be obtained arthroscopically or by open methods.

**DIFFERENTIAL DIAGNOSIS**

Rheumatoid arthritis must be differentiated from the following diseases:

a) **Systemic lupus erythematosus (SLE):** In SLE, the joint involvement is not symmetrical; nor are ankylosis and erosions common. Absence of anti-nuclear antibody factor (ANF) is in favour of rheumatoid arthritis, although its presence does not confirm SLE. It is present in 25 per cent cases of rheumatoid arthritis, though in low titres.

b) **Osteoarthritis:** This occurs in older patients. There is complete lack of the systemic features of rheumatoid arthritis such as fever, weight loss, fatigue etc. Distal inter-phalangeal joints are often involved. Duration of morning stiffness, joint swelling, ESR etc., are less compared to rheumatoid arthritis.

c) **Psoriatic arthropathy:** Characteristic skin and nail lesions may be present. Distal inter-phalangeal joints are usually involved. Rheumatoid factor is negative.

**TREATMENT**

**Principles of treatment:** Aims of treatment are as follows:

a) **Induction of remission and its maintenance:** Disease activity is brought under control by drugs.

b) **Preservation of joint functions and prevention of deformities** during the activity of the disease and thereafter, by physiotherapy and splinting.

c) **Repair of joint damage** which already exists, if it will relieve pain or facilitate functions. It sometimes requires surgical intervention e.g., synovectomy.

**Methods of treatment:** Above mentioned goals can be achieved by medical and orthopaedic treatment.

**Medical treatment:** Medical treatment essentially consists of anti-rheumatic drugs. These consist of: (i) non-steroidal anti-inflammatory drugs (NSAIDs); (ii) disease modifying anti-rheumatic drugs (DMARDs); and (iii) steroids. For details please refer to a Medicine textbook.

**Orthopaedic treatment:** Orthopaedic treatment aims at prevention of deformity, preservation of joint functions and rehabilitation. It falls essentially into non-operative and operative methods of treatment.

**Non-Operative Methods:** These consist of the following:

- **Physiotherapy:** This consists of: (i) splintage of the joints in proper position during the acute phase: (ii) heat therapy – wax bath, hot water fomentation for symptomatic relief; (iii) joint mobilisation exercises to maintain joint functions; and (iv) muscle building exercises to gain strength.

- **Occupational therapy:** Role of occupational therapy is to help the patient cope with his occupational requirements in the most comfortable way, by modifying them.

- **Rehabilitation:** Role of rehabilitation is to improve the functions of the patient with the help of devices like braces, walking aids etc.

**Operative Methods:** Surgical treatment of rheumatoid arthritis can be divided into: (i) preventive surgery; (ii) palliative surgery; (iii) reconstructive surgery; and (iv) salvage surgery.
• **Preventive surgery**: This is done to prevent damage to the joint and nearby tendons by the inflamed, hypertrophied synovium. It consists of synovectomy of the wrist, knee and MP joints.

• **Palliative surgery**: This is done in situations where general condition of the patient does not permit corrective surgery, but some relief can be provided by limited surgical procedures such as bone block operations, tendon lengthening etc.

• **Reconstructive surgery**: This has revolutionised the rehabilitation of patients with deformed and painful joints. It includes tendon transfers, interposition arthroplasties and total joint replacement. With improvement in surgical techniques and better design of artificial joints, it is now possible to replace practically any joint of the body. The joints where total replacement is most popular are the hip, knee and metacarpophalangeal joints.

**Plan of treatment**: Management depends upon the stage of the disease, as discussed below:

1. **Potentially reversible** soft tissue proliferation, where drug therapy constitutes the mainstay of treatment.
2. **Controllable** but irreversible soft tissue destruction and early cartilage erosion, where a combination of drug therapy and orthopaedic treatment is required.
3. **Advanced stage** of joint destruction with subluxation or dislocation, where primarily surgical treatment is necessary. Drugs alone are of no use at this stage.

Plan of treatment in these three stages is as given in Table–34.6.

**Prognosis**: Following factors decide the outcome of a patient diagnosed to have rheumatoid arthritis.

**Natural history of the disease**: It is well known that rheumatoid arthritis is a disease with variable natural history. It may be fulminant i.e., damaging joints quickly and producing deformities in spite of best care, or more usually a disease with persistent course punctuated with remissions and exacerbations. It is not possible to predict the precise nature of the disease in a particular patient.

• **Sex and age at onset**: Women of child bearing age with predominant upper extremity involvement have a progressively severe disease. Males, with sparing of upper extremity, where onset of disease is under the age of 30 years, show less severe disease.

• **Type of onset**: It is generally believed that insidious onset disease progresses to have more severe disease.

• **Anaemia**: Anaemia is associated with progressive rheumatoid arthritis. Also, it is believed that unresponsiveness of anaemia to oral iron therapy is a bad prognostic indicator.

• **ESR and C-reactive protein**: High levels are associated with more erosive arthritis.

• **Rheumatoid factor**: A positive rheumatoid factor is associated with more progressive disease. High titres of rheumatoid factor, appearing early in the disease, carry a bad prognosis.

• **Radiological erosions**: Presence of erosions within 2 years of onset of the disease, is a bad prognostic indicator.

• **Histopathological changes**: A case with synovial proliferation, with increased number of synovial cells with DR antigen, carries bad prognosis.

**ANKYLOSING SPONDYLITIS**

*(Marie strumpell disease)*

Ankylosing spondylitis is a chronic disease characterised by a progressive inflammatory...
stiffening of the joints, with a predilection for the joints of the axial skeleton, especially the sacroiliac joints.

AETIOPATHOLOGY
The exact aetiology is not known. A strong association has been found between a genetic marker—HLA-B27 and this disease. Whereas, the incidence of HLA-B27 is less than 1 per cent in general population, it is present in more than 85 per cent of patients with ankylosing spondylitis.

Pathology: Sacro-iliac joints are usually the first to get involved; followed by the spine from the lumbar region upwards. The hip, the knee and the manubrio-sternal joints are also involved frequently. Initially synovitis occurs; followed later, by cartilage destruction and bony erosion. Resultant fibrosis ultimately leads to fibrous, followed by bony ankylosis. Ossification also occurs in the anterior longitudinal ligament and other ligaments of the spine. After bony fusion occurs, the pain may subside, leaving the spine permanently stiff (burnt out disease).

CLINICAL FEATURES
Presenting complaints: This is a disease of young adults, more common in males (M : F=10 : 1). The following clinical presentations may be seen:
a) Classic presentation: The patient is a young adult 15-30 years old male, presenting with a gradual onset of pain and stiffness of the lower back. Initially, the stiffness may be noticed only after a period of rest, and improves with movement. Pain tends to be worst at night or early morning, awakening the patient from sleep. He gets better only after he walks about or does some exercises. There may be pain in the heel, pubic symphysis, manubrium sterni and costo-sternal joints. In later stages, kyphotic deformity of spine and deformity of the hips may be prominent features.
b) Unusual presentations: Patient may occasionally present with involvement of peripheral joints such as the shoulders, hips and knees. Smaller joints are rarely involved. Sometimes, a patient with ankylosing spondylitis may present with chronic inflammatory bowel disease; the joint symptoms follow.

On examination it is found that the patient walks with a straight stiff back. There may be a diffuse kyphosis. Following clinical signs may be present:
- Stiff spine: There may be a loss of lumbar lordosis. Lumbar spine flexion may be limited.
- Tests for detecting sacro-iliac involvement: Following tests may be positive in a case with sacro-iliac joint involvement:
  - Tenderness, localised to the posterior superior iliac spine or deep in the gluteal region.
  - Sacro-iliac compression: Direct side to side compression of the pelvis may cause pain at the sacro-iliac joints.
  - Gaenslen’s test: The hip and the knee joints of the opposite side are flexed to fix the pelvis, and the hip joint of the side under test is hyperextended over the edge of the table. This will exert a rotational strain over the sacro-iliac joint and give rise to pain (Fig-34.3a).
  - Straight leg raising test: The patient is asked to lift the leg up with the knee extended.

![Fig-34.3 Tests for sacro-iliac joint affections](https://kat.cr/user/Blink99/)
This will cause pain at the affected sacroiliac joint.

- **Pump-handle test:** With the patient lying supine, the examiner flexes his hip and knee completely, and forces the affected knee across the chest, so as to bring it close to the opposite shoulder (Fig-34.3b). This will cause pain on the affected side.

- **Tests for cervical spine involvement:** In advanced stages, the cervical spine gets completely stiff. The Fle’che test may detect an early involvement of the cervical spine.

- **Fle’che test:** The patient stands with his heel and back against the wall and tries to touch the wall with the back of his head without raising the chin. Inability to touch the head to the wall suggests cervical spine involvement.

- **Thoracic spine involvement:** Maximum chest expansion, from full expiration to full inspiration is measured at the level of the nipples. A chest expansion less than 5 cm indicates involvement of the costo-vertebral joints.

**Extra-articular manifestations:** In addition to articular symptoms, a patient with ankylosing spondylitis may have the following extra-articular manifestations:

a) **Ocular:** About 25 per cent patients with ankylosing spondylitis develop at least one attack of acute iritis sometimes during the natural history of the disease. Many patients suffer from recurrent episodes, which may result in scarring and depigmentation of the iris.

b) **Cardiovascular:** Patients with ankylosing spondylitis, especially those with a long standing illness, develop cardiovascular manifestations in the form of aortic incompetence, cardiomegaly, conduction defects, pericarditis etc.

c) **Neurological:** Patients may develop spontaneous dislocation and subluxation of the atlanto-axial joint or fractures of the cervical spine with trivial trauma, and may present with signs and symptoms of spinal cord compression.

d) **Pulmonary:** Involvement of the costo-vertebral joints lead to painless restriction of the thoracic cage. This can be detected clinically by diminished chest expansion, or by performing pulmonary function tests (PFT). There may also occur bilateral apical lobe fibrosis with cavitation, which remarkably simulates tuberculosis on X-ray.

e) **Systemic:** Generalised osteoporosis occurs commonly. Occasionally, a patient may develop amyloidosis.

**INVESTIGATIONS**

**Radiological examination** (Fig-34.4): In a suspected case, X-rays of the pelvis (AP), and dorso-lumbar spine (AP and lateral) are required. Oblique views of sacro-iliac joints may be required in early stages to appreciate their involvement. Following changes may be seen on X-ray of the pelvis:

- Haziness of the sacro-iliac joints
- Irregular subchondral erosions in SI joints

Fig-34.4 X-rays showing changes in ankylosing spondylitis. (a) X-ray of the pelvis, AP view, showing bilateral SI joint and hip involvement (b) X-ray of the lumbar spine, Lateral view, showing calcification of the ligaments

- Sclerosis of the articulating surfaces of SI joints
- Widening of the sacro-iliac joint space
- Bony ankylosis of the sacro-iliac joints
- Calcification of the sacro-iliac ligament and sacro-tuberous ligaments
- Evidence of enthesopathy – calcification at the attachment of the muscles, tendons and ligaments, particularly around the pelvis and around the heel.

X-ray of the lumbar spine may show the following:

- **Squaring** of vertebrae: The normal anterior concavity of the vertebral body is lost because
Table–34.7 Differential diagnosis of Ankylosing spondylitis

<table>
<thead>
<tr>
<th>Disease</th>
<th>Clinical</th>
<th>Investigations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stiffness</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TB spine</td>
<td>Occurs at any age or sex</td>
<td>Signs of TB spine on X-rays</td>
</tr>
<tr>
<td></td>
<td>Localised tenderness present</td>
<td>Primary lesion in the chest</td>
</tr>
<tr>
<td></td>
<td>Cold abscess present</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Constitutional symptoms present</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Family or past history of TB present</td>
<td></td>
</tr>
<tr>
<td>Fluorosis</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Any age or sex</td>
<td>Posterior longitudinal ligament calcified, seen on X-rays</td>
</tr>
<tr>
<td></td>
<td>Dental mottling present</td>
<td>Interosseous membrane calcification seen on X-rays</td>
</tr>
<tr>
<td></td>
<td>Chest expansion normal</td>
<td>Serum and urine fluoride studies – high levels</td>
</tr>
<tr>
<td>Back pain</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lumbo-sacral strain</td>
<td>Non-specific, Localised tenderness present</td>
<td>X-rays normal</td>
</tr>
<tr>
<td>Disc prolapse</td>
<td>SLRT positive</td>
<td>No radiological changes</td>
</tr>
<tr>
<td>Osteoarthritis</td>
<td>Seen in elderly patients</td>
<td>Osteophytes present</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ESR normal</td>
</tr>
<tr>
<td>SI joint diseases</td>
<td></td>
<td></td>
</tr>
<tr>
<td>TB of SI joint</td>
<td>Any age affected</td>
<td>Lytic lesions with sclerosis seen on X-ray, Cold abscess may be seen on CT, ESR high</td>
</tr>
<tr>
<td></td>
<td>Generally unilateral</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cold abscess present</td>
<td></td>
</tr>
<tr>
<td>Osteitis condensans iliac</td>
<td>Bilateral, non-specific sclerosis of ilium, Seen in parous women</td>
<td>ESR normal</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sclerosis of subchondral bone on the iliac side of the SI joint only</td>
</tr>
</tbody>
</table>

- Loss of the lumbar lordosis.
- Bridging ‘osteophytes’ (*syndesmophytes*).
- *Bamboo spine* appearance.

In the peripheral joints, X-ray changes are similar to those seen in rheumatoid arthritis, except that there is formation of large osteophytes and peri-articular calcification. Bony ankylosis occurs commonly.

**Other investigations:** These are the following:
- ESR: elevated
- Hb: mild anaemia
- HLA-B27: positive (to be tested in doubtful cases)

**DIFFERENTIAL DIAGNOSIS**

In early stages, ankylosing spondylitis may be confused with other disorders, as given in Table-34.7.

**OTHER RHEUMATOLOGICAL DISEASES**

**Gout**

Disturbed purine metabolism leading to excessive accumulation of uric acid in the blood — an inherited disorder; or impaired excretion of uric acid by the kidneys. The result is accumulation of *sodium biurate* crystals in some soft tissues. Tissues of predilection are cartilage, tendon, bursa

Patient, usually beyond 40 years of age, presents as (i) arthritis – MP joint of the big toe being a *favourite site*, onset is acute, pain is severe; (ii) bursitis – commonly of the olecranon bursa; or (iii) tophi formation deposit of uric acid salt in the soft tissue

Confirmation of diagnosis – urate crystals in the aspirate from a joint or bursa, high serum uric acid levels

Treatment – NSAIDs, uricosuric drugs, uric acid inhibitors.

**Pseudogout**

*sodium pyrophosphate* crystal deposition

Symptoms like those of gout

*Meniscus calcification* may be seen on X-rays of the knee

Treatment - NSAIDs.
Psoriatic arthropathy

Presentation is like rheumatoid arthritis – a polyarthritis, *distal IP joints of hands involved* (unlike rheumatoid arthritis, where these are spared)
Classic skin lesions help in diagnosis
Treatment is by steroids.

Alkaptonuric arthritis

(Ocronosis)

An inherited defect in enzyme system involved in metabolism of phenylalanine and tyrosine. As a result homogentisic acid is excreted in patient’s urine. As a long term result, it accumulates in the cartilage and other connecting tissues.
Joint symptoms occur after 40 years of age. Spine and shoulder joint are commonly affected. There may be evidence of pigment deposit in the sclera. Homogentisic acid is present in the urine, and results in the colour of the urine turning dark brown on standing (due to oxidation of homogentisic acid on exposure to air)
X-ray – *disc space calcification*, peri-articular calcification in large joints
Treatment same as that for osteoarthritis.

Haemophilic arthritis

Occurs due to a number of bleeding disorders
Occurs in males
Joints affected commonly are knee, elbow and ankle
May present as acute or chronic haemarthrosis. There are other manifestations of bleeding disorders
X-ray – non-specific signs including bone resorption, cyst formation, osteoporosis, *widening of intercondylar notch in the knee.*
Treatment – rest during acute stage along with factor VIII supplementation or other deficient factor replacement. In the chronic stage, physiotherapy, bracing etc. are required. Deformities may be corrected by conservative or operative methods.

Neuropathic arthropathy

(Charcot's joint)

These are changes seen in a neuropathic joint, where repeated strain on a joint due to loss of sensations leads to severe degeneration
Clinically, the joint manifests as painless effusion, deformity or instability
The X-ray changes are those of severe osteoarthritis but without much clinical findings like pain, muscle spasm etc.
Treatment is difficult. Bracing is usually advised for some joints. Fusion of the joint may be required.

### TREATMENT

*No specific therapy* is available. Aim is to control the pain and maintain maximum degree of joint mobility. This can readily be achieved by life long pursuit of a structured exercise programme. In some cases surgical intervention is required.

**Conservative methods:** These consist of: (i) *drugs*—NSAIDs are given for pain relief; Indomethacin in effective in most cases; long acting preparations are preferred; (ii) *physiotherapy* – this consists of proper posture guidance, heat therapy and mobilisation exercises; (iii) *radiotherapy* – in some resistant cases; and (iv) *yoga therapy.*

**Operative methods:** Role of operative treatment is in correction of kyphotic deformities of the spine by spinal osteotomy, and joint replacement for cases with hip or knee joint ankylosis.

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**What have we learnt?**

- There are two types of arthritis: inflammatory and degenerative.
- Rheumatoid arthritis is a chronic polyarthritis of inflammatory nature, typically affecting peripheral joints.
- Orthopaedic management of rheumatoid arthritis is aimed at prevention of deformity, correction of deformity and joint replacement.
- Ankylosing spondylitis occurs in young men. Treatment is aimed at physiotherapy.
PATHOLOGY

Osteoarthritis is a degenerative condition primarily affecting the articular cartilage. The first change observed is an increase in water content and depletion of the proteoglycans from the cartilage matrix. Repeated weight bearing on such a cartilage leads to its fibrillation. The cartilage gets abraded by the grinding mechanism at work at the points of contact between the apposing articular surfaces, until eventually the underlying bone is exposed. With further ‘rubbing’, the subchondral bone becomes hard and glossy (eburnated). Meanwhile, the bone at the margins of the joint hypertrophies to form a rim of projecting spurs known as osteophytes. A similar mechanism results in the formation of subchondral cysts and sclerosis.

The loose flakes of cartilage incite synovial inflammation and thickening of the capsule, leading to deformity and stiffness of the joint. Often one compartment of a joint is affected more than the other. For example, in the knee joint, the medial compartment is affected more than the lateral, leading to a varus deformity (genu varum).

CLINICAL FEATURES

The disease occurs in elderly people, mostly in the major joints of the lower limb, frequently bilaterally. There is a geographical variation in the joints involved, depending probably upon the daily previous disease producing a damaged articular surface; (iv) internal derangement of the knee, such as a loose body; (v) mal-alignment (bow legs etc.); and (vi) obesity and excessive weight.
activities of a population. The hip joint is commonly affected in a population with western living habits, while the knee is involved more commonly in a population with Asian living habits i.e., the habit of squatting and sitting cross legged.

Pain is the earliest symptom. It occurs intermittently in the beginning, but becomes constant over months or years. Initially, it is dull pain and comes on starting an activity after a period of rest; but later it becomes worse and cramp-like, and comes after activity. A coarse crepitus may be complained of by some patients. Swelling of the joint is usually a late feature, and is due to the effusion caused by inflammation of the synovial tissues. Stiffness is initially due to pain and muscle spasm; but later, capsular contracture and incongruity of the joint surface contribute to it. Other symptoms are: a feeling of 'instability' of the joint, and 'locking' resulting from loose bodies and frayed menisci.

**EXAMINATION**

Following findings may be present:
- Tenderness on the joint line
- Crepitus on moving the joint
- Irregular and enlarged-looking joint due to formation of peripheral osteophytes
- Deformity – varus of the knee, flexion-adduction-external rotation of the hip
- Effusion – rare and transient
- Terminal limitation of joint movement
- Subluxation detected on ligament testing
- Wasting of quadriceps femoris muscle

**INVESTIGATIONS**

**Radiological examination:** The diagnosis of osteoarthritis is mainly radiological (Fig-35.1). The following are some of the radiological features:
- Narrowing of joint space, often limited to a part of the joint e.g., may be limited to medial compartment of tibio-femoral joint of the knee.
- Subchondral sclerosis – dense bone under the articular surface
- Subchondral cysts
- Osteophyte formation
- Loose bodies
- Deformity of the joint

**Other investigations** are made primarily to detect an underlying cause. These consist of the following:
- Serological tests and ESR to rule out rheumatoid arthritis
- Serum uric acid to rule out gout
- Arthroscopy, if a loose body or frayed meniscus is suspected

**TREATMENT**

**Principles of treatment:** Once the disease starts, it progresses gradually, and there is no way to stop it. Hence efforts are directed, wherever possible, to the following:
- To delay the occurrence of the disease, if the disease has not begun yet.
- To stall progress of the disease and relieve symptoms, if the disease is in early stages.
- To rehabilitate the patient, with or without surgery, if his disabilities can be partially or completely alleviated.

**Methods of treatment:** To achieve the above objectives, the following therapeutic measures may be undertaken:
- **Drugs:** Analgesics are used mainly to suppress pain. A trial of different drugs is carried out to find a suitable drug for a particular patient. Long-acting formulations are preferred.
- **Chondroprotective agents:** Agents such as Glucosamine and Chondroitin sulphate have been introduced, claiming to be the agents which result in repair of the damaged cartilage. Their role as disease modifying agents has yet not been established, but these could be tried in some early cases.
c) **Viscosupplementation:** Sodium Hylarunon has been introduced. It is injected in the joint 3-5 times at weekly interval. It is supposed to improve cartilage functions, and is claimed to be chondroprotective.

d) **Supportive therapy:** This is a useful and harmless method of treatment and often gives gratifying results. It consists of the following:

- **Weight reduction**, in an obese patient.
- **Avoidance of stress** and strain to the affected joint in day-to-day activities. For example, a patient with OA of the knee is advised to avoid standing or running whenever possible. Sitting cross legged and squatting is harmful for OA of the knee.
- **Local heat** provides relief of pain and stiffness.
- **Exercises** for building up the muscles controlling the joint help in providing stability to the joint.
- The local application of *counter-irritants* and liniments sometimes provide dramatic relief.

e) **Surgical treatment:** In selected cases, surgery can provide significant relief. Following are some of the surgical procedures performed for OA:

- **Osteotomy:** Osteotomy near a joint has been known to bring about relief in symptoms, especially in arthritic joints with deformities. A high tibial osteotomy for OA of the knee with genu varum (Fig-35.2a), and inter-trochanteric osteotomy for OA of the hip have been shown to be useful for pain relief.

- **Joint replacement:** For cases crippled with advanced damage to the joint, total joint replacement operation (Fig-35.2b) has provided remarkable rehabilitation. These are now commonly performed for the hip and knee. An artificial joint serves for about 10-15 years.

- **Joint debridement:** This operation is not so popular now. In this, the affected joint is opened, degenerated cartilage smoothened, and osteophytes and the hypertrophied synovium excised. The results are unpredictable.

- **Arthroscopic procedures:** Arthroscopic removal of loose bodies, degenerated meniscal tears and other such procedures have become popular because of their less invasive nature. In arthroscopic chondroplasty, the degenerated, fibrillated cartilage is excised using a power-driven shaver under arthroscopic vision. Results are unpredictable.

### CERVICAL SPONDYLOSIS

This is a degenerative condition of the cervical spine found almost universally in persons over 50 years of age. It occurs early in persons pursuing ‘white collar jobs’ or those susceptible to neck strain because of keeping the neck constantly in one position while reading, writing etc.

**PATHOLOGY**

The pathology begins in the intervertebral discs. Degeneration of disc results in reduction of disc space and peripheral osteophyte formation. The posterior intervertebral joints get secondarily involved and generate pain in the neck. The osteophytes impinging on the nerve roots give rise to radicular pain in the upper limb. Exceptionally, the osteophytes may press on the spinal cord, giving rise to signs of cord compression. Cervical spondylosis occurs most commonly in the lowest three cervical intervertebral joints (the commonest is at C₅-C₆).

**CLINICAL FEATURES**

Complaints are often vague. Following are the common presentations:

- **Pain and stiffness:** This is the commonest presenting symptom, initially intermittent but
later persistent. Occipital headache may occur if the upper-half of the cervical spine is affected.

- **Radiating pain:** Patient may present with pain radiating to the shoulder or downwards on the outer aspect of the forearm and hand. There may be paraesthesia in the region of a nerve root, commonly over the base of the thumb (along the C₆ nerve root). Muscle weakness is uncommon.
- **Giddiness:** Patient may present with an episode of giddiness because of vertebro-basilar syndrome.

**EXAMINATION**

There is loss of normal cervical lordosis and limitation in neck movements. There may be tenderness over the lower cervical spine or in the muscles of the para-vertebral region (myalgia). The upper limb may have signs suggestive of nerve root compression – usually that of C₆ root involvement. Motor weakness is uncommon. The lower limbs must be examined for signs of early cord compression (e.g. a positive Babinski reflex etc.).

**RADIOLOGICAL FINDINGS**

X-rays of the cervical spine (AP and lateral) are sufficient in most cases. Following radiological features may be present (Fig-35.3):

- Narrowing of intervertebral disc spaces (most commonly between C₅-C₆).
- Osteophytes at the vertebral margins, anteriorly and posteriorly.
- Narrowing of the intervertebral foramen in cases presenting with radicular symptoms, may be best seen on oblique views.

**DIFFERENTIAL DIAGNOSIS**

The diseases to be considered in differential diagnosis of cervical spondylosis are: (i) other causes of neck pain such as infection, tumours and cervical disc prolapse; and (ii) other causes of upper limb pain like Pancoast tumour, cervical rib, spinal cord tumours, carpal tunnel syndrome etc.

**TREATMENT**

**Principles of treatment:** The symptoms of cervical spondylosis undergo spontaneous remissions and exacerbations. Treatment is aimed at assisting the natural resolution of the temporarily inflamed soft tissues. During the period of remission, the prevention of any further attacks is of utmost importance, and is done by advising the patient regarding the following:

a) **Proper neck posture:** Patient must avoid situations where he has to keep his neck in one position for a long time. Only a thin pillow should be used at night.

b) **Neck muscle exercises:** These help in improving the neck posture.

During an episode of acute exacerbation, the following treatment is required:

- Analgesics
- Hot fomentation
- Rest to the neck in a cervical collar
- Traction to the neck if there is stiffness
- Anti-emetics, if there is giddiness

In an exceptional case, where the spinal cord is compressed by osteophytes, surgical decompression may be necessary.

**LUMBAR SPONDYLOSIS**

This is a degenerative disorder of the lumbar spine characterised clinically by an insidious onset of pain and stiffness and radiologically by osteophyte formation.

**CAUSE**

Bad posture and chronic back strain is the commonest cause. Other causes are, previous injury to the spine, previous disease of the spine, birth defects and old intervertebral disc prolapse.
PATHOLOGY
Primarily, degeneration begins in the intervertebral joints. This is followed by a reduction in the disc space and marginal osteophyte formation. Degenerative changes develop in the posterior facet joints. Osteophytes around the intervertebral foramen may encroach upon the nerve root canal, and thus interfere with the functioning of the emerging nerve.

DIAGNOSIS
Clinical features: Symptoms begin as low backache, initially worst during activity, but later present almost all the time. There may be a feeling of ‘a catch’ while getting up from a sitting position, which improves as one walks a few steps. Pain may radiate down the limb up to the calf (sciatica) because of irritation of one of the nerve root. There may be complaints of transient numbness and paraesthesia in the dermatome of a nerve root, commonly on the lateral side of leg or foot (L₅, S₁ roots) respectively.

EXAMINATION
The spinal movements are limited terminally, but there is little muscle spasm. The straight leg raising test (SLRT) may be positive if the nerve root compression is present.

RADIOLOGICAL FINDINGS
Good AP and lateral views of the lumbo-sacral spine (Fig-35.4) should be done after preparing the bowel with a mild laxative and gas adsorbent like charcoal tablets. It is particularly difficult in obese patients, the ones usually suffering from this disease. Following signs may be present:

- Reduction of disc space
- Osteophyte formation
- Narrowing of joint space of the facet joints
- Subluxation of one vertebra over another

TREATMENT
Principles of treatment: Like cervical spondylosis, lumbar spondylosis also undergoes spontaneous remissions and exacerbations. Treatment is essentially similar to cervical spondylosis. In the acute stage, bed rest, hot fomentation and analgesics are advised. As the symptoms subside, spinal exercises are advised. In some resistant cases, a lumbar corset may have to be used at all times. Spinal fusion may occasionally be necessary.

What have we learnt?
- Osteoarthritis is a degenerative, progressive disorder. It commonly affects the knee and the back.
- Treatment is preventive, if predisposing factors are known.
- Once progressed, physiotherapy and surgery remain the only options.
### Additional information: From the entrance exams point of view

<table>
<thead>
<tr>
<th>Characteristic findings</th>
<th><strong>Osteoarthritis</strong></th>
<th><strong>Rheumatoid arthritis</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Osteophytes and new bone formation</td>
<td>Juxta-articular osteoporosis, osteophytes and new bone formation usually absent</td>
</tr>
<tr>
<td>Joints Involved commonly</td>
<td>Distal interphalyngeal joints of hand (Heberden’s nodes), Proximal interphalyngeal joints of hand (Bouchard’s nodes), 1st carpometacarpal joint, hip, knee</td>
<td>Wrist joint, metacarpophalyngeal joints, proximal interphalyngeal joints, knee, and hip</td>
</tr>
<tr>
<td>Joints not involved</td>
<td>Metacarpophalyngeal joints and wrist joint</td>
<td>Lumbar spine, distal interphalyngeal joint</td>
</tr>
</tbody>
</table>

- **Type A** synovial cells are phagocytes that engulf joint debris
- **Type B** synovial cells secrete synovial fluid
Bursitis

Inflammation may occur in a normally situated bursa or in an adventitious bursa. It may arise from mechanical irritation or from bacterial infection. Accordingly, there are two types of bursitis:

**Irritative bursitis:** This is the commoner of the two types. It is caused by excessive pressure or friction, occasionally due to a gouty deposit. Inflammation of the bursa results in the effusion of a clear fluid within the bursal sac. With prolonged inflammation, the sac gets thickened and may cause pressure erosion on the adjacent bone. Some commonly seen bursites are given in Table–36.1.

<table>
<thead>
<tr>
<th>Table-36.1: Common bursites</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prepatellar bursitis</td>
</tr>
<tr>
<td>Infrapatellar bursitis</td>
</tr>
<tr>
<td>Olecranon bursitis</td>
</tr>
<tr>
<td>Ischial bursitis</td>
</tr>
<tr>
<td>On lateral malleolus</td>
</tr>
<tr>
<td>On great toe</td>
</tr>
</tbody>
</table>

*Treatment:* Most cases respond to analgesics, rest to the part and removal of the causative factor i.e., friction or pressure. In some resistant cases, the sac is infiltrated with hydrocortisone. Very rarely, excision of the bursa is required.

**Infective bursitis:** Uncommonly, a bursa may get infected by a pyogenic or tubercular infection.

It occurs commonly in trochanteric bursa or prepatellar bursa. Treatment is by surgical drainage and antibacterial drugs.

Tenosynovitis

Inflammation of the thin synovial lining of a tendon sheath (Fig-36.1) is termed tenosynovitis. It may arise from mechanical irritation or from bacterial infection.

**Irritative tenosynovitis** is commonly seen in the tendons of the hand and results in pain and swelling. Treatment is by rest, analgesics and ultrasonic therapy. Some cases need local hydrocortisone infiltration.

**Infective tenosynovitis** is an infection of the synovial lining of the tendon by pyogenic or tubercular bacteria. Pyogenic infection is common
in the flexor tendons of the hand. Tubercular tenosynovitis of the sheaths of the flexor tendons of the forearm at the level of the wrist occurs commonly (compound palmar ganglion).

**DUPUYTREN’S CONTRACTURE**  
*(Contracture of the palmar aponeurosis)*

This is a condition characterised by a flexion deformity of one or more fingers due to a thickening and shortening of the palmar aponeurosis. The cause is unknown, but a hereditary predisposition has been established. There is an increased incidence of the disorder among cirrhotic patients and in epileptics on sodium hydantoïn.

**PATHOANATOMY**

Normally, the palmar aponeurosis is a thin but tough membrane, lying immediately beneath the skin of the palm. Proximally, it is in continuation with the palmaris longus tendon. Distally, it divides into slips, one for each finger. The slip blends with the fibrous flexor sheaths covering the flexor tendon of the finger, and extends up to the middle phalanx (Fig-36.2a). In Dupuytren’s contracture, the aponeurosis or a part of it becomes thickened and slowly contracts, drawing the fingers into flexion at the metacarpophalangeal and proximal inter-phalangeal joints (Fig-36.2b). The ring finger is the one affected most commonly. The contracture is generally limited to the medial three fingers. Sometimes, it may be associated with a thickening of plantar fascia or that of the penile fascia (Peyronie’s disease).

**CLINICAL FEATURES**

In early stages, thickening of the palmar aponeurosis is felt at the bases of ring and little fingers. Later, a flexion deformity of the fingers develops. Dupuytren’s contracture can be differentiated from a similar deformity due to contracture of the flexor tendons; in the former only the MP and PIP joints are flexed, unlike the latter where the DIP joints are also flexed.

**TREATMENT**

An elderly patient with mild contracture does not need any treatment. If the deformity is significant and hampers the activity of the patient, excision of the palmar aponeurosis (subtotal excision) may be required.

**TENNIS ELBOW**  
*(Lateral epicondylitis)*

This is a condition characterised by pain and tenderness at the lateral epicondyle of the humerus due to non-specific inflammation at the origin of the extensor muscles of the forearm. Although, it is sometimes seen in tennis players, other activities such as squeezing clothes, carrying a suitcase etc. are frequently responsible.

**CLINICAL FEATURES**

One finds tenderness, precisely localised to the lateral epicondyle of the humerus. Pain is aggravated by putting the extensor tendons to a stretch; for example, by palmar-flexing the wrist and fingers with the forearm pronated. Elbow movements are normal. X-ray does not reveal any abnormality.

**TREATMENT**

The patient is initially treated with analgesics–anti-inflammatory drugs for a week or so. If there is no response, a local injection of hydrocortisone at the point of maximum tenderness generally brings relief.

**GOLFER’S ELBOW**  
*(Medial epicondylitis)*

This is a condition similar to tennis elbow where the inflammation is at the origin of the flexor tendons at the medial epicondyle of the humerus. Treatment is on the lines of a tennis elbow.
**de QUERVAIN’S TENOSYNOVITIS**

This is a condition characterised by pain and swelling over the radial styloid process. It results from inflammation of the common sheath of abductor pollicis longus and extensor pollicis brevis tendons (Fig-36.3). On examination, the tenderness is localised to the radial styloid process. Pain is aggravated by adducting the thumb across the palm and forcing ulnar deviation and on asking the patient to perform radial deviation against resistance (Finkelstein’s test). There may be a palpable thickening of the sheath.

**TREATMENT**

In early stage, rest to the wrist in a crepe bandage or a slab, analgesics and ultrasonic radiation may bring relief. In some cases, a local infiltration of hydrocortisone is required. A chronic case may need slitting and excision of a part of the tendon sheath.

**TRIGGER FINGER/THUMB**

This is a condition resulting from the constriction of the fibrous digital sheath, so that free gliding of the contained flexor tendon does not occur.

**CLINICAL FEATURES**

Initially, the only symptom is pain at the base of the affected finger, especially on trying to passively extend the finger. As the sheath further thickens, the contained tendon gets swollen proximal to it (Fig-36.4). The swollen segment of the tendon does not enter the sheath when an attempt is made to straighten the finger from the flexed position. This is called ‘locking of finger’. This locking can be overcome either by a strong effort in which case the finger extends with a snap-like trigger of a pistol or by extending the finger passively with other hand.

**TREATMENT**

In early stages, local ultrasonic therapy provides relief. In a long standing problem, a local injection of hydrocortisone relieves the pain. In some cases, splitting of the tight tendon sheath may be required.

**GANGLION**

It is the commonest cystic swelling on the dorsum of the wrist. It results from mucoid degeneration of the tendon sheath or the joint capsule. Ordinarily, there are no symptoms other than the swelling itself. Sometimes, a mild discomfort or pain is experienced. The cyst may sometimes be so tense as to resemble a solid tumour of the tendon sheath. Often the cyst is multi-loculated. Aspiration of the cyst is performed and an injection of hylase given. If the cyst recurs, excision may be required.

**CARPAL TUNNEL SYNDROME**

This is a syndrome characterised by the compression of the median nerve as it passes beneath the flexor retinaculum (Fig-36.5).

**CAUSES**

Any space occupying lesion of the carpal tunnel may be responsible. Some of the common causes are given in Table–36.2.
PLANTAR FASCITIS

This is a common cause of pain in the heel. It occurs as a result of inflammation of the plantar aponeurosis at its attachment on the tuberosity of the calcaneum (Fig-36.6). The pain is worst early in the morning, and often improves with activity. On examination, there is marked tenderness over the medial aspect of the calcaneal tuberosity, at the site of attachment of the plantar fascia.

Fig-36.6 Causes of heel pain
1. Diseases of calcaneum
2. Plantar fascitis
3. Fat pad inflammation
4. Retro-calcaneal bursitis
5. Achilles tendinitis
6. Diseases of subtalar joint

X-rays sometimes show a sharp bone spur projecting forwards from the tuberosity of the calcaneum. Its significance is doubtful since it is also found in some cases without heel pain. Analgesics, the use of a heel pad and local induction of steroids brings relief in most cases.

FIBROSITIS

This is a non-specific condition where there is pain in certain muscles, with tenderness when they are gripped. One can palpate small, firm nodules, mostly over the trapezius and spinal muscles. These nodules are supposed to be trigger points. There are no other objective signs. The patient responds to ultrasonic therapy or local steroid infiltration.

PAINFUL ARC SYNDROME

This is a clinical syndrome in which there is pain in the shoulder and upper arm during the mid-range of gleno-humeral abduction (Fig-36.7). Following are the common causes:

- Minor tears of the supraspinatus tendon
- Supraspinatus tendinitis

CLINICAL FEATURES

The patient is generally a middle aged woman complaining of tingling, numbness or discomfort in the thumb and radial one and a half fingers i.e., in the median nerve distribution. Tingling is more prominent during sleep. There is a feeling of clumsiness in carrying out fine movements. On examination, features of low median nerve compression are found (see page 66). Nerve conduction studies show delayed or absent conduction of impulses in the median nerve across the wrist. Treatment is by dividing the flexor retinaculum, and thus decompressing the nerve.

‘FROZEN’ SHOULDER
(Periarthritis shoulder)

This is a disease of unknown aetiology where the gleno-humeral joint becomes painful and stiff because of the loss of resilience of the joint capsule, possibly with adhesions between its folds. Often, there is a history of preceding trauma. The disease is commoner in diabetics.

CLINICAL FEATURES

It produces pain and stiffness of the shoulder. In early stages, the pain is worst at night, and the stiffness limited to abduction and internal rotation of the shoulder. Later, the pain is present at all times and all the movements of the shoulder are severely limited.

TREATMENT

This is a self-limiting disease lasting for 6-9 months, after which in most cases, the inflammation subsides, leaving a stiff but painless shoulder. Treatment is by analgesics, hot fomentation and physiotherapy. An intra-articular injection of hydrocortisone may speed up the recovery. Stiffness can be prevented by continuous shoulder mobilising exercises.

Table–36.2: Causes of carpal tunnel syndrome

<table>
<thead>
<tr>
<th>Cause</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Idiopathic</td>
<td>The commonest cause</td>
</tr>
<tr>
<td>Inflammatory causes</td>
<td>Rheumatoid arthritis</td>
</tr>
<tr>
<td>Post-traumatic causes</td>
<td>Wrist osteoarthritis</td>
</tr>
<tr>
<td>Endocrine causes</td>
<td>Bone thickening after a Colles’ fracture</td>
</tr>
<tr>
<td></td>
<td>Myxoedema, Acromegaly</td>
</tr>
</tbody>
</table>

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- Minor tears of the supraspinatus tendon
- Supraspinatus tendinitis
• Calcification of supraspinatus tendon
• Subacromial bursitis
• Fracture of the greater tuberosity

In all these conditions, the space between the upper-end of the humerus and the acromion gets compromised, so that during mid-abduction the tendon of the rotator-cuff gets nipped between the greater tuberosity and acromion. X-ray of the shoulder may show calcific deposit, or a fracture of the greater tuberosity or acromion. Treatment consists of ultrasonics to the tender point and anti-inflammatory drugs. Some cases need an injection hydrocortisone in the subacromial space or excision of the anterior, often prominent part of the acromion.

**MERALGIA PARAESTHETICA**

This is a feeling of tingling, burning, and numbness in the skin supplied by the lateral cutaneous nerve of the thigh as it gets entrapped in the fascia just medial to the anterior superior iliac spine. Treatment is non-specific with analgesics, local hydrocortisone etc. Sometimes, surgical decompression of the nerve may be required.

**FIBROMYALGIA**

This is a widespread disease characterized by multiple tender points, affecting both sides of the body – both above and below the waist, lasting more than 3 months. It is known to be associated with irritable bowel, headache, dysmenorrhea and chronic fatigue syndrome. It is an entity, distinct from fibrositis which is a localised disorder. Aetiology is not known, but it is proposed to be a part of fatigue syndrome. Diseases like hypothyroidism, SLE, hyperparathyroidism and osteomalacia need to be ruled out by careful investigation. Treatment is by patient counselling, avoidance of aggravating factors, physical therapy and antidepressants.

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**What have we learnt?**

- Bursae are deflated balloons around areas which are subjected to friction. They get inflamed and produce pain.
- Dupuytren’s contracture is contracture of palmer fascia.
- Tennis elbow and golfer’s elbow are epicondylitis.
- de Quervain’s disease is tenosynovitis of tendons around the wrist.
- Frozen shoulder is idiopathic capsular inflammation of shoulder, usually responds to physiotherapy.
- Painful arc syndrome occurs because of impingement of rotator-cuff.
## Additional information: From the entrance exams point of view

- Types of superficial heat therapy: Hot bath, chemical pack, paraffin wax bath, infrared lamp, moist air cabinet.
- Types of deep heat therapy: Short wave diathermy, microwave therapy, ultrasound therapy.
- ‘O’ Brien’s test done for a tight iliotibial band.
- Dupuytren’s contracture: Surgery required if proximal interphalyngeal joint contracture >15° and metacarpophalyngeal joint contraction >30°.
- Level of tendon sheath constriction in trigger finger is metacarpophalyngeal joint (A1 pulley).
- Bursa pes anserinus is between the tendons of the sartorius, gracilis, semitendinosus and the tibial collateral ligament.
- Thoracic outlet syndrome is best diagnosed via clinical examination.
- Athletic pubalgia is due to strain of the rectus abdominus muscle.
CONSTITUTION OF BONE

Osseous tissue is made up of organic and inorganic material along with water. Relative proportions of these constituents are as shown in Flow chart-37.1.

Inorganic constituents of bone: It constitutes 65 per cent of the dry weight of bones. The bulk of this is calcium and phosphate, which in an adult is primarily crystalline (hydroxyapatite crystals). Besides calcium, other minor mineral constituents are Magnesium, Sodium, Potassium etc. Although, the bone contains a large amount of calcium, only about 65 per cent of it is in an exchangeable form.

Organic constituents of bone: On a dry weight basis, the organic matrix constitutes 35 per cent of the total weight of human bones. Of this, 95 per cent is collagen; other constituents being polysaccharides (mucoproteins or glycoproteins) and lipids (including phospholipids).

BONE AND CALCIUM

Bone serves as a storehouse for 99 per cent of the body’s calcium. Changes in the calcium ion activity in the extracellular fluid affects multiple biological processes. Hence, special regulatory mechanisms are required to provide an overall control of this

Table 37.1: Effects of PTH, vitamin D and calcitonin on kidney, GIT and bone

<table>
<thead>
<tr>
<th>Agent</th>
<th>Effect on serum levels</th>
<th>Kidney</th>
<th>GIT</th>
<th>Bone</th>
</tr>
</thead>
<tbody>
<tr>
<td>PTH</td>
<td>↑ Ca, ↓ P</td>
<td>Direct action ↑ renal phosphate excretion (phosphaturia)</td>
<td>—</td>
<td>↑ mobilisation of Ca from bone</td>
</tr>
<tr>
<td></td>
<td></td>
<td>↑ resorption of Ca</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vitamin D</td>
<td>↑ Ca, ↓ P</td>
<td>—</td>
<td>Direct action ↑ absorption of Ca and P from gut</td>
<td>↑ mobilisation of Ca from bone</td>
</tr>
<tr>
<td>Calcitonin</td>
<td>↑ Ca, ↓ P</td>
<td>↓ urinary</td>
<td>— from bone</td>
<td>—</td>
</tr>
</tbody>
</table>

https://kat.cr/user/Blink99/
activity. If for any reason, the serum level of calcium falls below its normal value, the body can react in three specific ways: (i) it may increase intestinal absorption; (ii) it may decrease urinary excretion; or (iii) it may increase the release of calcium from bone. The factors responsible for monitoring these activities are parathyroid hormone (PTH), vitamin D and calcitonin as given in Table–37.1.

In cases with acute lowering of the serum calcium level, such as that induced by the administration of calcium complexing substances, the intestinal tract and kidney cannot act swiftly enough to restore the level to normal. In this instance, the first supply of calcium comes from the lacunar and canalicular surfaces of the bone. This effect is believed to be an equilibrium exchange. It accounts for only a small amount of mineral and lasts for only a few minutes. After this source is depleted, and if the serum level of calcium is still low, parathyroid hormone secretion is stimulated as a direct consequence of the lowered serum level of calcium. This hormone provides a second source of calcium from the bone, from the zone surrounding the osteocytes. This is called osteocytic osteolysis. This happens first in cancellous bone and finally in cortical bone. Increase in the number and activity of osteoclasts results in resorption of large amount of bone. This type of resorption is called osteoclastic resorption.

There are four types of metabolic bone diseases (Fig-37.1).

a) **Osteopenic diseases:** These diseases are characterised by a generalised decrease in bone mass (i.e., loss of bone matrix), though whatever bone is there, is normally mineralised (e.g., osteoporosis).

b) **Osteosclerotic diseases:** These are diseases characterised by an increase in bone mass (e.g., fluorosis).

c) **Osteomalacic diseases:** These are diseases characterised by an increase in the ratio of the organic fraction to the mineralised fraction i.e., the available organic matter is under-mineralised (e.g., osteomalacia).

d) **Mixed diseases:** These are diseases that are a combination of osteopenia and osteomalacia (e.g., hyperparathyroidism).

**OSTEOPOROSIS**

Osteoporosis is by far the commonest metabolic bone disease. It is characterised by a diffuse reduction in the bone density due to a decrease in the bone mass. It occurs when the rate of bone resorption exceeds the rate of bone formation.

**CAUSES**

Several aetiological factors may be operative in a given patient. Commonest factor in males is senility and in females is menopause. Table–37.2 gives some of the causes of generalised osteoporosis.

### Table–37.2: Causes of generalised osteoporosis

- **Senility**
- **Post-immobilisation e.g. a bed-ridden patient**
- **Post-menopausal**
- **Protein deficiency**
  - Inadequate intake – old age, illness
  - Malnutrition
  - Mal-absorption
  - Excess protein loss (3rd degree burns, CRF etc.)
- **Endocrinal**
  - Cushing’s disease
  - Cushing’s syndrome
  - Hyperthyroid state
- **Drug induced**
  - Long term steroid therapy
  - Phenobarbitone therapy

**CLINICAL FEATURES**

Osteoporosis is an asymptomatic disorder unless complications (predominantly fractures) occur. Loss of bone mass leads to loss of strength so that a trivial trauma is sufficient to cause a fracture. Dorso-lumbar spine is the most frequent site. Pain from these fractures is usually the reason for a person to consult a physician. Other fractures whose aetiology has been linked to underlying osteoporosis are Colles’ fracture and fracture of the neck of femur.
Metabolic Bone Diseases

Densitometry: This is a method to quantify osteoporosis. In this method absorption of photons (emitted from gamma emitting isotopes) by the bone calcium is measured. Two types of bone densitometry are available – ultrasound based and X-ray based. DEXA scan is an X-ray based bone densitometry, and is the gold standard in the quantification of bone mass.

Neutron activation analysis: In this method, calcium in the bone is activated by neutron bombing, and its activity measured.

Bone biopsy.

TREATMENT

Since the aetiology of osteoporosis is multifactorial and the diagnosis usually delayed, treatment becomes difficult. There are no set treatment methods as yet. The principle objectives of treatment are alleviation of pain and prevention of fractures. Treatment can be divided into medical and orthopaedic.

Medical treatment: This consists of the following:

- High protein diet: Many elderly patients suffer from malnutrition. Increasing their protein intake may increase the formation of organic matrix of the bone.
- Calcium supplementation: Its role is doubtful, but may be helpful in cases with deficiency of calcium in their diet.
- Androgens: These hormones have an anabolic effect on the protein matrix of bone, and in some instances ameliorate symptoms.
- Estrogens: Estrogens have been shown to halt the progressive loss of bone mass in post-menopausal osteoporosis.
- Vitamin D: This is given, in addition to the above, to increase calcium absorption from the gut.
- Fluoride: The use of fluoride is still under study. It is supposed to make the crystallinity of the bone greater; thereby making bone resorption slower.
- Alendronate: These are used in once a day dose, empty stomach. Oesophagitis is a troubling complication.
- Calcitonin: Parenteral administration of calcitonin helps in building up the bone mass and also acts as an analgesic.

On examination, the findings are subtle and can be missed. A slight loss of height and increased kyphosis due to compression of the anterior part of the vertebral bodies is seen in most cases.

RADIOLOGICAL FEATURES

Radiological evidence of decreased bone mass is more reliable, but about 30 per cent of the bone mass must be lost before it becomes apparent on X-rays (Fig-37.2). Following features may be noticed on X-rays:

- Loss of vertical height of a vertebra due to collapse.
- Cod fish appearance: The disc bulges into the adjacent vertebral bodies so that the disc becomes biconvex.
- Ground glass appearance of the bones, conspicuous in bones like the pelvis.
- Singh’s index: Singh et al. graded osteoporosis into 6 grades based on the trabecular pattern of the femoral neck trabeculae. Details are outside the scope of this book.
- Metacarpal index and vertebral index are other methods of quantification of osteoporosis.

OTHER INVESTIGATIONS

These include the following, some of them more recent:

- Biochemistry: Serum calcium, phosphates and alkaline phosphatase are within normal limits. Total plasma proteins and plasma albumin may be low.
- **Teriparatide**: Anabolic agent increasing osteoblastic new bone formation.
- **Denosumab, Strontium**: Antiresorptive agents.

**Orthopaedic treatment**: This consists of the following:
- **Exercises**: Weight bearing is a major stimulus to bone formation. Increased guarded activity would therefore be of benefit to the patient.
- **Bracing**: Prophylactic bracing of the spine by using an ASH brace or Taylor brace may be useful in prevention of pathological fractures in a severely osteoporotic spine.

### Rickets and Osteomalacia

Rickets and osteomalacia are the diseases where the organic matrix of the bone fails to calcify properly, leaving large osteoid seams. Manifestations of the two diseases are different only with respect to the stage in life at which they occur. Rickets occurs in the growing bones of children; osteomalacia in the bones of adults. Both conditions are primarily due to a deficiency of vitamin D or a disturbance in its metabolism secondary to renal disease.

**Vitamin D and its metabolism**: The *endogenous* form of vitamin D i.e., cholecalciferol, is found in the skin as a product of cholesterol metabolism in a process requiring ultraviolet radiation. The *exogenous* form of the vitamin is usually D<sub>3</sub>. The two most important nutritionally useful forms of vitamin D are D<sub>2</sub> (ergocalciferol) and D<sub>3</sub> (cholecalciferol).

**Steps in activation**: The basic forms, vitamin D<sub>2</sub> and D<sub>3</sub> are inactive until hydroxylated. The first step of hydroxylation (25 hydroxylation) occurs in the liver and the second step (1 hydroxylation) occurs in the kidney. 1,25 dihydroxylated form is the active form and stimulates the intestinal absorption of calcium and also acts on the bone.

**Control**: The most sensitive of the physiological actions of 1-25 dihydroxy vitamin D is to increase intestinal absorption of calcium. The action of vitamin D metabolites in bone tissue is controversial. To calcify the bone matrix properly, small amounts of the metabolites are necessary along with sufficient calcium.

### Rickets

Rickets is a disease of the growing skeleton. It is characterised by failure of normal mineralisation, seen prominently at the growth plates, resulting in softening of the bones and development of deformities.

**CAUSES**

There are two types of rickets i.e., Type I and Type II (Table–37.3). In Type I, there is either a deficiency of vitamin D or a defect in its metabolism. In Type II, the rickets occurs due to a deficiency of phosphates in the extra-cellular fluid because of defective tubular resorption or diminished phosphate intake. Nutritional deficiency continues to be the *commonest* cause of rickets in developing countries because of poor socio-economic conditions.

### CLINICAL FEATURES

Nutritional rickets occurs in children about 1 year old. It may occur in older children with mal-absorption syndrome. Following are the clinical features:

- **Craniotabes**: This is the manifestation of rickets seen in young infants. Pressure over the soft membranous bones of the skull gives the feeling of a ping pong ball being compressed and released.
- **Bossing of the skull**: Bossing of the frontal and parietal bones becomes evident after the age of 6 months.

---

**Table-37.3: Types of rickets**

<table>
<thead>
<tr>
<th>Type 1</th>
<th>Due to a deficiency of vitamin D</th>
</tr>
</thead>
<tbody>
<tr>
<td>a)</td>
<td>Diminished intake e.g., malnutrition</td>
</tr>
<tr>
<td></td>
<td>Diminished absorption e.g.,</td>
</tr>
<tr>
<td></td>
<td>– mal-absorption syndrome</td>
</tr>
<tr>
<td></td>
<td>– gastric abnormalities</td>
</tr>
<tr>
<td></td>
<td>– biliary diseases</td>
</tr>
<tr>
<td></td>
<td>Lack of exposure to sunlight</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Type II</th>
<th>Due to disturbance in vitamin D metabolism</th>
</tr>
</thead>
<tbody>
<tr>
<td>a)</td>
<td>Defective absorption of phosphates through renal tubules</td>
</tr>
<tr>
<td></td>
<td>Hypophosphataemic rickets (x-linked dominant)</td>
</tr>
<tr>
<td></td>
<td>Fanconi syndrome</td>
</tr>
<tr>
<td></td>
<td>Renal tubular acidosis</td>
</tr>
<tr>
<td></td>
<td>Oncogenic rickets</td>
</tr>
<tr>
<td>b)</td>
<td>Diminished intake or absorption of phosphates</td>
</tr>
</tbody>
</table>

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• **Broadening of the ends of long bones**, most prominently around wrists and knees. It is seen around 6-9 months of age.
• **Delayed teeth eruption** is noticed in infants.
• **Harrison’s sulcus**: A horizontal depression, along the lower part of the chest, corresponding to the insertion of diaphragm.
• **Pigeon chest**: The sternum is prominent.
• **Rachitic rosary**: The costo-chondral junctions on the anterior chest wall become prominent, giving rise to appearance of a rosary.
• **Muscular hypotonia**: The child’s abdomen becomes protruberant (pot belly) because of marked muscular hypotonia. Visceroptosis and lumbar lordosis occurs.
• **Deformities**: Deformities of the long bones resulting in knock knees or bow legs is a common presentation of rickets, once the child starts walking.

**RADIOLOGICAL FEATURES**
Early radiological changes are observed in the lower ends of the radius and ulna. X-rays of both wrists and knees – antero-posterior views are used for screening a patient suspected of rickets. Following radiological signs may be seen (Fig-37.3):

• **Delayed appearance of epiphyses**.
• **Widening of the epiphyseal plates**: Normal width of the epiphyseal plate is 2-4 mm. In rickets it is increased because of excessive accumulation of uncalcified osteoid at the growth plate.
• **Cupping of the metaphysis**: Normally, the metaphysis meets the epiphyseal plate as a smooth line of sclerosis (zone of provisional calcification). In rachitic bones, this line is absent and the metaphyseal end appears irregular. The cartilage cells accumulating at the growth plate create a depression in the soft metaphyseal end, giving rise to a cup shaped appearance (Fig-37.4).

![Fig-37.3 X-ray of the wrists, AP view, showing changes in rickets](https://kat.cr/user/Blink99/)

![Fig-37.4 Radiological features of rickets](https://kat.cr/user/Blink99/)

• **Splaying of the metaphysis**: The end of the metaphysis is splayed because of the pressure by the cartilage cells accumulating at the growth plate.
• **Rarefaction** of the diaphyseal cortex occurs late.
• **Bone deformities**: Knock knees, bow legs and coxa vara are common deformities in older children.

**OTHER INVESTIGATIONS**
Serum calcium is usually normal or low, serum phosphate is low, but serum alkaline phosphatase is high.

**TREATMENT**
It consists of medical and orthopaedic treatment.

**Medical treatment**: Administration of vitamin D 6,00,000 units as a single oral dose induces rapid healing. If the line of healing (a line of sclerosis on the metaphyseal side of the growth plate) is not seen on X-rays within 3-4 weeks of therapy, same dose may be repeated. In cases where the child responds to vitamin D therapy, a maintenance dose of 400 I.U. of vitamin D is given per day. If there is no response even after the second dose, a diagnosis of **refractory rickets** is made. Such patients are evaluated in detail by multi speciality team of nephrologist, endocrinologist and physician.

**Orthopaedic treatment**: It is required for the correction of deformities by conservative or operative methods.

a) **Conservative methods**: Mild deformities correct spontaneously, as rickets heals. Some surgeons use specially designed splints (mermaid
splints) or orthopaedic shoes for correction of knee deformities.

b) **Operative methods:** Moderate or severe deformities often require surgery. This can be performed any time after 6 months of starting the medical treatment. Corrective osteotomies, depending upon the nature of deformities, are performed.

### OSTEOMALACIA

Osteomalacia, which means softening of bones, is the adult counterpart of rickets. It is primarily due to deficiency of vitamin D. This results in failure to replace the turnover of calcium and phosphorus in the organic matrix of bone. Hence, the bone content is demineralised and the bony substance is replaced by soft osteoid tissue.

### AETIOLOGY

It is common in women who live in ‘purdah,’ and lack exposure to sunlight. Other causes are: dietary deficiency of vitamin D, under-nutrition during pregnancy, mal-absorption syndrome, after partial gastrectomy, etc.

### CLINICAL FEATURES

In its early stages, symptoms and signs are non-specific and the diagnosis is often missed. Following presentations may be seen:

- **Bone pains:** Skeletal discomfort ranging from backache to diffuse bone pains may occur. Bone tenderness is common.
- **Muscular weakness:** The patient feels very weak. He may have difficulty in climbing up and down the stairs. A waddling gait is not unusual. Tetany may manifest as carpopedal spasm and facial twitching.
- **Spontaneous fractures** occur usually in spine, and may result in kyphosis.

### INVESTIGATIONS

Following investigations may be carried out:

- **Radiological examination:** Plain X-rays appear to be of ‘poor quality’ i.e., not sharp and well-defined. Following findings may be present (Fig-37.5):
  - Diffuse rarefaction of bones.
  - Looser’s zone (pseudo-fractures): These are radiolucent zones occurring at sites of stress. Common sites are the pubic rami, axillary border of scapula, ribs and the medial cortex of the neck of the femur. These are caused by rapid resorption and slow mineralisation and may be surrounded by a collar of callus.
  - Triradiate pelvis in females.
  - Protrusio-acetabuli i.e., the acetabulum protruding into the pelvis.
  - **Bone biopsy:** A bone biopsy from the iliac crest usually confirms the diagnosis. The characteristic histological finding is excessive uncalcified osteoid.
  - **Serum:** The serum calcium level is low, the phosphates are low and alkaline phosphatase high.

### TREATMENT

When osteomalacia is due to defective intake, vitamin D supplementation therapy, as for rickets, brings dramatic results. Vitamin D in daily maintenance doses of 400 I.U. is sufficient. If there is mal-absorption, higher dose or intramuscular dose may be needed. In patients with renal disease, alfa-calcidol (an activated form of vitamin D) may be used. Calcium supplementation should also be given. In addition, the underlying cause is treated.

### HYPERPARATHYROIDISM

Clinical bone disease occurs in less than half of the patients with hyperparathyroidism. For reasons unknown, hyperparathyroidism tends to present either with bone disease or with renal stones, but not both.

Before we discuss this topic further, here is a brief account of the action of parathyroid hormone on bone.

**Parathyroid hormone and bone:** The parathyroid hormone acts directly on the bone to release calcium into the extra-cellular fluid by stimulating
osteoclastic resorption. It activates the adenyl cyclase so that the formation of cyclic AMP is increased, which in turn increases the synthesis of specific lysosomal enzymes. These enzymes break down the organic matrix of bone and release calcium.

**CLINICAL FEATURES**

The disease can affect either sex, but is more common in women. The majority of cases occur from the third to fifth decades of life. Following are the more common presenting complaints:

- **Bone pains**: This is the most common initial feature. There is tenderness on palpating the bones, especially in the lower limbs and back. Pain is usually associated with general weakness, pallor and hypotonia.

- **Pathological fracture**: Fractures occur with trivial injuries and unite in a deformed position. Common sites of fractures are dorso-lumbar spine, neck of the femur and pubic rami.

- **Brown’s tumour**: This is an expansile bone lesion, a collection of osteoclasts. It commonly affects the maxilla or mandible, though any bone may be affected.

- **Anorexia, nausea, vomiting and abdominal cramps** are common presenting complaints.

- **Occasionally, renal colics** with haematuria, because of renal calculus, may occur.

**RADIOLOGICAL INVESTIGATIONS**

*X-ray examination* consists of the lateral view of the skull, dorso-lumbar spine (AP/lateral views), both hands (AP view), pelvis (AP view) and X-ray of the region with symptoms. Following signs may be present (Fig-37.6):

- Irregular, diffuse rarefaction of the bones.

- **Salt pepper appearance**: The skull bones show a well-marked stippling, but the opaque areas are small pin head size.

- **Loss of lamina dura**: A tooth socket is made up of thin cortical bone seen as a white line surrounding the teeth. This is called the lamina dura. It gets absorbed in hyperparathyroidism.

- **Sub-periosteal resorption** of the phalanges is a diagnostic feature of hyperparathyroidism (generalised variety). Resorption may also occur at lateral end of the clavicle.

- Spine shows central collapse of the vertebral body and biconvex discs.

- Pelvis and other bones show coarse striations with clear cyst-like spaces.

- **Brown’s tumour** is an expansile lytic lesion, which appears like a bone tumour, generally affecting the maxilla/mandible.

- Extra-osseous radiological features such as renal calculi etc. may be present.

**OTHER INVESTIGATIONS**

- **Serum** calcium is high, phosphates low and alkaline phosphatase high.

- **Urinary** excretion of calcium is low and that of phosphates high, as found on 24 hour urine analysis.

- Investigations for finding the underlying cause of hyperparathyroidism i.e., CT scan of the neck for parathyroids, and for the evaluation of other organs for ectopic secretion of parathormone.

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**Fig-37.6 X-rays showing changes in hyperparathyroidism**

https://kat.cr/user/Blink99/
**TREATMENT**

It consists of: (i) removal of the basic cause; (ii) orthopaedic treatment; and (iii) urologic treatment.

a) *Treatment* of the basic cause is by surgical excision of the hormone secreting tissue.

b) *Orthopaedic treatment* is directed towards adequate protection of the softened bones from all deforming stress and strain. Once the disease has been arrested and recalcification of bones occurs, the established deformities may be corrected surgically.

c) *Urology treatment*: This is directed towards the removal of calculi and maintenance of renal functions.

**FLUOROSIS**

Fluorosis is a disease where excessive deposition of calcium occurs in bone and soft tissues. It results from excessive ingestion of fluorides in drinking water. The fluoride content of normal water is less than 1 PPM. In India and parts of south east Asia, large areas have been reported with high water fluoride content. Fluorosis is an endemic disease and a public health problem in some states of India i.e., Punjab, Andhra Pradesh, Tamil Nadu etc.

**CLINICAL FEATURES**

The symptoms and signs may pertain to skeletal system or teeth.

a) *Skeletal fluorosis*: The patient complains of pain in the back and joints. There may be associated stiffness of the spine and paraesthesia in the limbs. In advanced stages, the patient presents with spastic paraparesis and anaemia. Paraparesis occurs because of cord compression resulting from calcification of the posterior longitudinal ligament and subsequent pressure on the cord.

b) *Dental fluorosis*: This is the earliest sign of fluorosis. The earliest to occur is mottling of the enamel, best seen in the incisors of the upper jaw. Later the teeth get eroded and fall off.

**RADIOLOGICAL INVESTIGATIONS**

X-rays of the spine, pelvis and forearms are required in a suspected case. Following features may be seen (Fig-37.7):

- Spine: Increased density, calcification of the posterior longitudinal ligament.
- Pelvis: Increased density, calcification of the ischio-pubic and sacro-iliac ligaments.
- Forearm and leg: Interosseous membrane calcification.

**OTHER INVESTIGATIONS**

- Elevated serum fluoride levels.
- Elevated fluoride levels in urine and drinking water.
- A biopsy shows high fluoride levels in bones.

**DIFFERENTIAL DIAGNOSIS**

Fluorosis is an osteosclerotic disease, and must be differentiated from other causes of osteosclerosis given in Table–37.4.

**TREATMENT**

Prevention is the most important aspect of treatment of this difficult disease. Defluorination
of the water is carried out as a public health programme. Patients improve symptomatically once the defluorinated water is used for some time.

DISTURBANCES OF ORGANIC CONSTITUENTS

LATHYRISM
Ingestion of certain agents called lathyrogens causes profound alterations in the collagen of connective tissues and bones. The bones of such individuals are soft, their arteries become weak and develop aneurysms. Treatment is removal of causative factor.

SCURVY
This disease is caused by deficiency of vitamin C (ascorbic acid). The result is decreased production and poor quality collagen. In adults, it presents with swollen gums, gingivitis and abnormal bleeding tendencies typically producing *perifollicular* haemorrhages over the lower part of the thighs. Petechial haemorrhages and spontaneous bruises may occur anywhere in the body, but usually first in the lower extremities. In infantile scurvy, important features are lassitude, anaemia, painful limbs due to sub-periosteal haematoma, and scurbutic rosary (bead like thickening of the ribs due to calcified sub-periosteal haematomas). Treatment is by supplementation with vitamin C.

POLYSACCHARIDOSIS
The principle polysaccharide of bone is a mucopolysaccharide — chondroitin-4-sulfate (chondroitin sulfate A). In certain diseases called as mucopolysaccharidosis, there is increased excretion of polysaccharides in the urine. Loss of these polysaccharides from bone and cartilage results in specific skeletal deformities.

What have we learnt?
- 65% of weight of the bone is inorganic.
- Osteoporosis is deficiency in matrix, whereas osteomalacia is deficiency of mineralisation of bone.
- Rickets and osteomalacia are diseases of deficiency in bone mineralisation.
- Hyperparathyroidism results in mobilisation of calcium from bone.
- Fluorosis is excessive deposition of calcium in bone and soft tissues.

Additional information: From the entrance exams point of view
- Rugger jersey spine is seen in renal osteodystrophy and is due to hyperparathyroidism.
- Milkman fractures are pseudofractures in adults.
GENERALISED BONE DISORDERS

ACHONDROPLASIA
This is a condition caused by the failure of normal ossification of bones, mainly the long bones, resulting in dwarfism (Fig-38.1). Since growth of the trunk is only marginally affected, the dwarfism is disproportionate; the limbs being out of proportion with the trunk. Shortening is especially marked in the proximal segments of the limbs.

OSTEOGENESIS IMPERFECTA
(Fragilitas ossium, Vrolik’s disease)
This is a condition characterised by tendency for frequent fractures because of weak and brittle bones. It results from defective collagen synthesis, and thus affects other collagen containing soft tissues such as the skin, sclera, teeth, ligaments etc., as well. The disease is commonly inherited as an autosomal dominant disorder, but a severe variant is known to occur where the inheritance is autosomal recessive. Typically, the patient has a tendency for frequent fractures, usually with minimal trauma. Associated features are blue sclera, joint laxity and otosclerosis in adulthood. Patients with the severe type do not survive beyond a few years, but those with the milder disease live their full life interrupted by frequent fractures. The fractures unite normally but deformities secondary to malunion or joint contractures may occur. The tendency to fracture often reduces with age. Some of the common causes of tendency for easy fracture are given in Table–38.2.

Table–38.1: Common causes of dwarfism

- Achondroplasia
- Dyschondroplasia
- Diaphysial aclasis
- Multiple epiphyseal dysplasia
- Cretinism
- Malnutrition
- Morquio’s, Hurler’s, Hunter’s diseases

The disease is of autosomal dominant inheritance, but many cases arise from a fresh gene mutation. Intelligence is normal. Typically, the patient has a large skull with a bulging vault and forehead, a flat nose, short limbs, short and stubby fingers and increased lumbar lordosis. These patients lead a near normal life, except for a few who develop spinal canal stenosis. Common causes of dwarfism are given in Table–38.1.
Table–38.2: Causes of tendency for easy fractures
- Osteogenesis imperfecta
- Osteoporosis
- Osteopetrosis
- Osteomalacia
- Polio limb

DIAPHYSIAL ACLASIS
(Multiple exostosis)
This is a condition characterised by multiple, cartilage-capped bony outgrowths from the metaphysis of long bones, and some from flat bones. It is inherited as an autosomal dominant disorder. The basic defect is that of remodelling. The columns of cartilage at the epiphyseal plate grow rapidly and sideways due to poor remodelling forces. X-rays typically show a ‘trumpet-shaped’ metaphysis and bony projections from it (Fig-38.2). The problem is of dwarfism, pressure effects of the exostosis, deformities, and a tendency of the exostosis to undergo malignant change. Since, it is impractical to excise all the exostosis, the one causing symptoms is excised.

OSTEOPETROSIS
(Marble bone disease, Albers-Schonberg disease)
This is a disorder characterised by dense but brittle bones (marble bones). In a less severe, autosomal dominant variety the patient has a tendency to fracture. In a severe, congenital, autosomal recessive variety, the child may have severe anaemia, jaw osteomyelitis and cranial nerve palsies. Most of the patients of the latter type do not survive for long.

PAGET’S DISEASE
(Osteitis deformans)
This is a condition characterised by a progressive tendency for one or more bones to bend, get thickened and spongy. Tibia is the bone affected most commonly. The cause is not known, but it is understood to be a defect in the osteoclast functions, so that irregular bone resorption and increased bone turnover occurs. The bone is soft and vascular in the initial stages, but becomes dense and hard later. The disease begins after 40 years of age. Presenting complaints are dull pain, and bowing and thickening of the affected bone. X-rays show multiple confluent lytic areas with interspread new bone formation. Bone scan shows an increased uptake. Serum alkaline phosphatase is elevated. Usual complications are pathological fracture and malignant change. Treatment is by calcitonin or diphosphonate.

NEUROFIBROMATOSIS
A generalised variety of neurofibromatosis (von Recklinghausen’s disease) may have skeletal disturbances like scoliosis, solitary bone lesions, pseudarthrosis of the tibia, compressive myelopathy, local gigantism, limb length inequality etc. In addition, other soft tissue manifestations of the disease may be present. For details, please refer to a Surgery textbook.

HISTIOCYTOSIS X
This comprises of a group of diseases characterised by proliferation of histiocytes in the bones. Three clinical entities are recognised:

a) Eosinophilic granuloma: This is a solitary bone lesion, commonly seen in the femur, tibia, spine and ribs. The patient usually presents with a dull pain or a pathological fracture. Treatment is curettage and bone grafting.

b) Hand-Schuller-Christian disease: It is a variation of (a), where the lesions are found at multiple sites.

c) Litterer-Siwe disease: This is the most severe form. There is involvement of multiple bones. It begins in childhood and progresses rapidly to death.

OSTEOCHONDRITIS
These are a group of miscellaneous affections of the growing epiphyses in children and adolescents. Typically, a bony nucleus of the epiphysis affected by osteochondritis becomes temporarily softened;
and while in the softened stage it is liable to deformation by pressure. Perthes’ disease, the osteochondritis of the epiphysis of the head of the femur is the commonest. Osteochondritis is sometimes classified into: (i) crushing type or osteochondrosis; (ii) osteochondritis dissecans; and (iii) traction osteochondritis or traction apophysitis. Table–38.3 gives names of some of the common osteochondrites.

Table–38.3: Common osteochondrites

<table>
<thead>
<tr>
<th>Name</th>
<th>Site affected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Perthes’ disease</td>
<td>Femoral head</td>
</tr>
<tr>
<td>Panner’s disease</td>
<td>Capitulum</td>
</tr>
<tr>
<td>Kienbock’s disease</td>
<td>Lunate bone</td>
</tr>
<tr>
<td>Osgood Shlatter’s disease</td>
<td>Tibial tubercle</td>
</tr>
<tr>
<td>Sever’s disease</td>
<td>Calcaneal tuberosity</td>
</tr>
<tr>
<td>Kohler’s disease</td>
<td>Navicular bone</td>
</tr>
<tr>
<td>Freiberg’s disease</td>
<td>Metatarsal head</td>
</tr>
<tr>
<td>Scheurmann’s disease</td>
<td>Ring epiphysis of vertebrae</td>
</tr>
<tr>
<td>Calve’s disease</td>
<td>Central bony nucleus of vertebral body</td>
</tr>
<tr>
<td>Iselin’s disease</td>
<td>5th metatarsal</td>
</tr>
</tbody>
</table>

PERTHES’ DISEASE
(Coxa plana, Pseudocoxalgia)

This is an osteochondritis of the epiphysis of the femoral head. In this disease, the femoral head becomes partly or wholly avascular and deformed. The cause is not definitely known, but it is supposed to be due to recurrent episodes of ischaemia of the head in the susceptible age group, probably precipitated by episodes of synovitis. Pathologically, the disease progresses in three ill-defined stages: (i) stage of synovitis; (ii) stage of trabecular necrosis; and (iii) stage of healing.

The disease occurs commonly in boys in the age group of 5-10 years. The child presents with pain in the hip, often radiating to the knee. There may be limping or hip stiffness. On examination, findings may be minimal – sometimes the only findings being a limitation of abduction and internal rotation and shortening. Radiological examination reveals collapse and sclerosis of the epiphysis of the femoral head. Hip joint space is increased. In fact, the contrast between the paucity of symptoms and signs in the presence of gross X-ray changes is striking (Fig-38.3). Bone scan may show a decreased uptake by the head of the femur. Four groups have been described by Catterall (1972), depending upon the extent of involvement of the head. He also describes the adverse prognostic signs (head at risk signs).

Treatment: Preventing the head from mis-shapening while the bone is in softening phase, is the primary aim of the treatment. The head is required to be kept inside the acetabulum while the revascularisation takes place (head containment). This may be achieved by conservative methods (plaster, splint etc) or by operation (containment osteotomy).

AVASCULAR NECROSIS

Avascular necrosis (AVN) of the bone due to loss of a vascularity of a part of the bone occurs commonly after a fracture or dislocation (e.g., AVN of head of the femur in a fracture of the femoral neck). Sometimes, a part of the bone undergoes avascular necrosis spontaneously. The head of the femur is a common site of AVN. It occurs in adults between the ages of 20-40 years. Some of the causes of avascular necrosis of the femoral head are given in Table–38.4.

Table–38.4: Causes of avascular necrosis of femoral head

- Idiopathic – commonest
- Alcoholism
- Steroid therapy
- Sickle cell disease
- Patient on renal dialysis
- Patient on anti-cancer drugs
- Post-partum necrosis
- Goucher’s disease
- Caisson’s disease

Diagnosis: The disease is often bilateral (75%). Patient complains of pain in the groin or in front of the thigh. Pain is present at all times, but increases

Fig-38.3 X-rays showing changes of Perthes’ disease of the hip. (Both the hips should be included to be able to appreciate early changes)
on exertion. X-rays may be normal in early stage; if suspicion is strong, a bone scan may be done. In later stage, an osteolytic lesion can be seen in supero-lateral part of the head. There may be diffuse osteosclerosis of the head, but the shape of the head may be maintained. In advanced stage, the head collapses (Fig-38.4). Eventually, changes of secondary osteoarthritis become apparent. MRI scan is the best modality for early diagnosis of avascular necrosis.

**Treatment:** In early stages, diagnosis is often missed as there are no X-ray findings. Core decompression and fibular grafting is done if the head of the femur has not got deformed. In later stages, once the head gets deformed, muscle-pedicle bone grafting using tensor facia lata has been shown to produce good results (Baksi). In some cases with involvement of a part of the head, an intertrochanteric osteotomy has been described. In cases with changes of advanced osteoarthritis, total hip replacement becomes necessary.

**SOME OTHER DEVELOPMENTAL ABNORMALITIES OF ORTHOPAEDIC INTEREST**

<table>
<thead>
<tr>
<th>1. Ollier’s disease</th>
<th>Not familial</th>
</tr>
</thead>
<tbody>
<tr>
<td>(multiple enchondromas)</td>
<td>Masses of cartilage in the metaphysis remain unossified</td>
</tr>
<tr>
<td>Dyschondroplasia</td>
<td>Defective ossification</td>
</tr>
<tr>
<td>2. Melorheostosis</td>
<td>Candle bone disease</td>
</tr>
<tr>
<td>3. Osteopathia striata</td>
<td>Striped bones disease</td>
</tr>
<tr>
<td>4. Osteopoikilosis</td>
<td>Spotted bones disease</td>
</tr>
<tr>
<td>5. Morquio’s disease</td>
<td>Familial (autosomal recessive) disease</td>
</tr>
<tr>
<td></td>
<td>Gives rise to dwarfism affecting both, limbs and trunk</td>
</tr>
<tr>
<td></td>
<td>Mental development normal</td>
</tr>
<tr>
<td></td>
<td>Corneal opacity sometimes present</td>
</tr>
<tr>
<td></td>
<td>X-ray – typical ‘tonguing’ of lumbar vertebrae</td>
</tr>
<tr>
<td></td>
<td>Keratan sulphate in urine</td>
</tr>
<tr>
<td>6. Hunter’s disease</td>
<td>Familial x-linked disease</td>
</tr>
<tr>
<td></td>
<td>Defect is the excretion of large amount of Keratan sulphate in urine</td>
</tr>
<tr>
<td></td>
<td>Dwarf with dorso lumbar kyphosis, knock knees, flat feet</td>
</tr>
<tr>
<td></td>
<td>Mental deficiency may occur</td>
</tr>
<tr>
<td></td>
<td>No corneal opacity</td>
</tr>
<tr>
<td>7. Hurler’s disease</td>
<td>Familial (autosomal recessive) disease</td>
</tr>
<tr>
<td>(Gargoylism)</td>
<td>Gives rise to dwarfism of both, limbs and trunk</td>
</tr>
<tr>
<td></td>
<td>Defect is an error in development of fibroblasts</td>
</tr>
<tr>
<td></td>
<td>There is excretion of dermatan sulphate and heparitan sulphate in urine</td>
</tr>
<tr>
<td></td>
<td>Typical facial appearance</td>
</tr>
<tr>
<td></td>
<td>Mental development abnormal</td>
</tr>
<tr>
<td></td>
<td>Corneal opacity present</td>
</tr>
<tr>
<td></td>
<td>X-ray typical ‘beak’ in 2nd lumbar vertebra</td>
</tr>
<tr>
<td>8. Engelmann’s disease</td>
<td>Familial (autosomal recessive) disease</td>
</tr>
<tr>
<td></td>
<td>Symmetrical, fusiform enlargement and sclerosis of shafts of the long bones in children. Femur affected commonly.</td>
</tr>
<tr>
<td></td>
<td>Epiphysis is spared</td>
</tr>
<tr>
<td>9. Caffey’s disease</td>
<td>Non-familial disease</td>
</tr>
<tr>
<td>(infantile cortical hyperostosis)</td>
<td>Starts early in life (before the 5th month)</td>
</tr>
<tr>
<td></td>
<td>There is a formation of sub-periosteal bone on the shafts of long bones, and on the mandible</td>
</tr>
<tr>
<td></td>
<td>Self limiting course, resolves by 3 years of age</td>
</tr>
<tr>
<td></td>
<td>Tibia more common than ulna in familial form</td>
</tr>
</tbody>
</table>

**Contd...**
| 10. | Albright's syndrome | • Polyostotic fibrous dysplasia and precocious puberty |
| 11. | Arthrogryposis multiplex congenita (AMC) | • Defective development of muscles |
|      |                           | • Stiff, deformed joints |
|      |                           | • Multiple joint dislocations with ‘shapeless extremities’ |
|      |                           | • May present as clubfoot |
| 12. | Myositis ossificans progressiva | • Ectopic ossification, often beginning in trunk |
| 13. | Multiple epiphyseal dysplasia | • Least rare type |
|      |                           | • Affects all the epiphyses, resulting in stunted growth, deformities (varum, valgum etc.) |
|      |                           | • Epiphysis looks ill defined, irregular on X-rays |
| 14. | Spondylo-epiphyseal dysplasia | • Spine is also involved in addition to limb epiphyses |
| 15. | Metaphyseal dysplasia (Pyle’s disease) | • Autosomal recessive |
|      |                           | • A modelling defect results in ‘Erlenmeyer flask’ deformity of the distal femur and proximal tibia |
| 16. | Blount’s disease | • The growth of the medial-half of the proximal tibial epiphysis is retarded, resulting in severe tibia vara deformity in childhood, common in West Indies |
| 17. | Cleido-cranial dysostosis | • Faulty development of membranous bones |
|      |                           | • Clavicles are absent |
|      |                           | • Skull sutures remain open |
|      |                           | • Coxa vara |
|      |                           | • Wide foramen magnum |
| 18. | Nail patella syndrome | • Familial disorder |
|      |                           | • Hypoplastic nails and absence of patella |
| 19. | Marfan’s syndrome (Arachnodactily) | • Spider fingers |
|      |                           | • Associated atrial regurgitation |
|      |                           | • Occular lens dislocation |
| 20. | Apert syndrome | • Tower shaped head |
|      |                           | • Syndactyly |

**What have we learnt?**
- Achondroplasia produces disproportionate dwarfism.
- Osteogenesis imperfecta is the cause of frequent fractures.
- Diaphyseal aclasis is an autosomal dominant disorder.
- Avascular necrosis commonly affects head of the femur. It is common in patients with sickle cell disease.

**Additional information: From the entrance exams point of view**
- Investigation of choice to diagnose early Perthe’s disease is MRI.
- Vertebra plana seen in eosinophilic granuloma, Ewing’s sarcoma, TB, Calve’s disease, leukemia and metastasis.
- Trident hand is seen in achondroplasia.
- Osteogenesis imperfecta is due to abnormal type I procollagen in the body.
- Muscles most commonly absent congenitally are pectoralis major and minor.
- Osteochondritis dessicans most commonly affects lateral part of medial femoral condyle.
TORTICOLLIS (Wry neck)

This is a deformity of the neck where the head and neck are turned and twisted to one side. It may be permanent, temporary, or spasmodic. Spasmodic torticollis is the commonest. Table-39.1 gives some of the common causes of torticollis. Most often, torticollis is secondary to pain and reflex muscle spasm and recovers once the inflammatory process subsides. Congenital torticollis, a common cause of permanent torticollis, is of orthopaedic interest.

Table-39.1: Causes of torticollis

<table>
<thead>
<tr>
<th>Type</th>
<th>Cause</th>
</tr>
</thead>
<tbody>
<tr>
<td>Congenital</td>
<td>Sterno-mastoid tumour</td>
</tr>
<tr>
<td>Infection</td>
<td>Tonsillitis</td>
</tr>
<tr>
<td></td>
<td>Atlanto-axial infections</td>
</tr>
<tr>
<td></td>
<td>Labyrinthitis</td>
</tr>
<tr>
<td>Reflex spasm</td>
<td>Acute disc prolapse (cervical)</td>
</tr>
<tr>
<td>Neurogenic</td>
<td>Spasmodic condition</td>
</tr>
<tr>
<td></td>
<td>Paralytic condition</td>
</tr>
<tr>
<td>Ocular</td>
<td>Compensation for squint</td>
</tr>
<tr>
<td>Others</td>
<td>Rheumatoid arthritis</td>
</tr>
<tr>
<td></td>
<td>Spasmodic torticollis</td>
</tr>
</tbody>
</table>

CONGENITAL TORTICOLLIS
(Infantile torticollis, Sterno-mastoid tumour)

The sterno-mastoid muscle on one side of the neck is fibrosed and fails to elongate as the child grows, and thus results in a progressive deformity. The cause of fibrosis is not known, but it is possibly a result of ischaemic necrosis of the sterno-mastoid muscle at birth. Evidence in favour of this theory is the presence of a lump in the sterno-mastoid muscle in the first few weeks of life, probably a swollen ischaemic muscle. This is termed *sterno-mastoid tumour*. The lump disappears spontaneously within a few months, leaving a fibrosed muscle. Torticollis occurs more commonly in children with breech presentation.

**Diagnosis:** The child usually presents at 3-4 years of age, often as late as puberty. The head is tilted to one side so that the chin faces to the opposite side (Fig-39.1). The sterno-mastoid is prominent on the side the head tilts, and becomes more prominent on trying to passively correct the head tilt. In cases presenting in the first few weeks of life, a lump may be felt in the sterno-mastoid muscle. Facial asymmetry develops in cases who present later in life. Radiological examination is normal, and is carried out to rule out an underlying bone defect such as scoliosis.

**Treatment:** In a child presenting with a sterno-mastoid tumour, progress to torticollis can be prevented by passive stretching and splinting. The same may also be sufficient for mild deformities in...
younger children. For severe deformities, especially in older children, release of the contracted sternomastoid muscle is required. It is usually released from its lower attachment, but sometimes both attachments need to be released. Following surgery, the neck is maintained in the corrected position in a Callot’s cast.

CERVICAL RIB

This is an additional rib which arises from the 7th cervical vertebra. It is usually attached to the first rib close to the insertion of the scalenus anterior muscle, and is present in less than 0.5 per cent of the population. It may be a complete rib, but more often it is present posteriorly for a short distance only; the anterior part being just a fibrous band. The cervical rib is usually unilateral and is more common on the right side.

CLINICAL FEATURES

In 90 per cent of cases, there are no symptoms; an extra rib is detected on an X-ray made for some other purpose. In others, it produces symptoms after the age of 30 years, probably because with declining youth the shoulders sag, increasing the angulation of the neurovascular structures of the upper limb as they come out of the neck. It is more often symptomatic in females. A patient may present with the following symptoms:

a) **Neurological symptoms:** Tingling and numbness along the distribution of the lowest part of the brachial plexus (T1 dermatome), along the medial border of the forearm and hand, is the commonest complaint. There may be weakness and wasting of the hand muscles and clumsiness in the use of the hand.

b) **Vascular symptoms:** These are uncommon. Compression of the subclavian artery may result in an aneurysm distal to constriction. This is a potential source of tiny emboli to the hand and may cause gangrene of the finger tips. There may be a history of pain in the upper limb on using the arm or elevating the hand (claudication).

c) **Local symptoms:** Occasionally, the patient presents with a tender supraclavicular lump (the anterior end of the cervical rib) which, on palpation, is bony hard and fixed.

RADIOLOGICAL EXAMINATION

X-ray examination may show a well-formed rib articulating posteriorly with transverse process of C7 vertebra. It is attached anteriorly to middle of the 1st rib. More often there is no fully developed cervical rib but merely an enlargement of the transverse process of the seventh cervical vertebra (Fig-39.2).

DIFFERENTIAL DIAGNOSIS

A patient with cervical rib is to be differentiated from those presenting with pain radiating down the upper limb due to other causes. Some of these causes are as follows:

a) **Carpal tunnel syndrome:** The symptoms are in the median nerve distribution. Nocturnal pain is characteristic.

b) **Cervical spine lesions:** In cases with cervical disc prolapse and spondylosis, pain radiates to the outer side of the arm and forearm. Associated limitation of neck movement and characteristic X-ray appearance may help in diagnosis.

c) **Spinal cord lesions:** Syringomyelia or other spinal cord lesions may cause wasting of the hand, but other neurological features help in reaching a diagnosis.

d) **Ulnar neuritis** may mimic this lesion but can be differentiated on clinical examination or by electrodiagnostic studies.

TREATMENT

Conservative treatment is usually rewarding. It consists of ‘shrugging the shoulder’ exercises to
build up the muscles, and avoidance of carrying heavy objects like shopping bag, bucket full of water, suitcase etc. Occasionally, surgical excision of the first rib may be required to relieve compression on the neurovascular bundle of the upper limb.

**OBSERVATION HIP**
*(Transient synovitis)*

This is a non-specific synovitis of the hip seen in children 4-8 years of age. It results in a painful stiffness of the hip which subsides after 2-3 weeks of rest and analgesics. X-ray examination and the ESR are normal. It is termed ‘observation hip’ because it must be ‘observed’ and differentiated from the following conditions:

a) **Early infective arthritis**: Some cases of early tuberculosis or septic arthritis may have features similar to observation hip. A high ESR, systemic symptoms, and persistent signs may necessitate a biopsy; especially in countries where tuberculosis is common.

b) **Chronic synovitis**: A mono-articular rheumatoid arthritis may resemble an ‘observation hip’.

c) **Perthes' disease**: In its early stages, before X-ray findings appear, Perthes’ disease may resemble a transient synovitis, but further follow up shows characteristic X-ray changes of the former.

**Treatment**: It consists of bed rest and analgesics. Recovery occurs within a few weeks.

**COXA VARA**

Coxa vara is a term used to describe a reduced angle between the neck and shaft of the femur. It may be congenital or acquired.

**INFANTILE COXA VARA**

This is coxa vara resulting from some unknown growth anomaly at the upper femoral epiphysis. It is noticed as a painless limp in a child who has just started walking. In severe cases, shortening of the leg may be obvious. On examination, abduction and internal rotation of the hip are limited and the leg is short. X-rays will show a reduction in neck-shaft angle (Fig-39.3). The epiphyseal plate may be too vertical. There may be a separate triangle of bone in the inferior portion of the metaphysis, called Fairbank’s triangle (Fig-39.4). Treatment is by a subtrochanteric corrective osteotomy.

**SLIPPED CAPITAL FEMORAL EPIPHYSIS**

In this condition, the upper femoral epiphysis may get displaced at the growth plate, usually postero-medially, resulting in coxa vara. The slip occurs gradually in majority of cases, but in some it occurs suddenly.

**CAUSES**

Aetiology is not known but it is thought to be a result of trauma in the presence of some not yet understood underlying abnormality. It occurs more commonly in unduly fat and sexually underdeveloped; or tall, thin sexually normal children.

**CLINICAL FEATURES**

Following are the salient clinical features:

- **Age**: It occurs at puberty (between 12-14 years).
**Sex:** It is commoner in boys.

**Side:** It occurs on both sides in 30 per cent of cases.

**There is a definite history of trauma in some cases.**

**It is commoner in patients with endocrine abnormalities.**

**Presenting symptoms:** Pain in the groin, often radiating to the thigh and the knee is the common presenting complaint. Often in the initial stages, the symptoms are considered due to a ‘sprain’, and are disregarded. They soon disappear only to recur. Limp occurs early and is more constant.

**Examination:** The leg is found to be externally rotated and 1-2 cm short. Limitation of hip movements is characteristic – there is limited abduction and internal rotation, with a corresponding increase in adduction and external rotation. When the hip is flexed, the knee goes towards the ipsilateral axilla (Fig-39.5). Muscle bulk may be reduced. Trendelenburg’s sign may be positive.

**TREATMENT**

It is based on the following considerations:

a) **Treatment of an acute slip:** This is by closed reduction and pinning, as for a fracture of the neck of the femur.

b) **Treatment of a gradual slip:** This depends upon the severity of the slip present. If it is less than 1/3 the diameter of the femoral neck, the epiphysis is fixed internally in situ. If the slip is more than 1/3, a corrective osteotomy is performed at the inter-trochanteric region.

c) **Treatment of the unaffected side** in unilateral cases: Since the incidence of bilateral involvement is 30 per cent, prophylactic pinning of the unaffected side in a case with unilateral slip is justified.

**DEFORMITIES OF THE KNEE**

**KNOCK KNEES (Genu Valgum)**

This is a condition where the knees are abnormally approximated and the ankles abnormally divergent (Fig-39.7a).

**Causes:** The commonest type is idiopathic, almost invariably bilateral. The deformity basically results from the unequal growth of two sides of the growth plate of the lower femoral epiphysis or upper tibial epiphysis (Table–39.2).

**Clinical features:** Physiological genu valgum appears at the age of 2-3 years and nearly always corrects by the age of 6. It may be associated with flat feet. The degree of deformity is estimated by measuring the inter-malleolar distance, with the child lying supine, with the knees in contact. In genu valgum secondary to a disease such as rickets, there will be findings suggestive of the primary disease.
Treatment: Spontaneous recovery occurs in most idiopathic cases. A medial shoe raise (3/16 inch) is sometimes prescribed. It has no proven scientific rationale but does help in satisfying anxious parents. If the inter-malleolar distance is 10 cm or more by the age 4, the child may need an operation. A supracondylar closed wedge osteotomy is performed.

BOW LEGS (Genu varum)
This is a condition where the knees are abnormally divergent (bow like) and the ankles abnormally approximated (Fig-39.7b).

Causes: Idiopathic is the commonest type. In others, causes similar to those for genu valgum can be identified, except that the defective growth is on the medial side of the epiphyseal plate. Blount’s disease is a special type of genu varum where the posteromedial part of the proximal tibial epiphysis fails to grow during the first 3 years of life.

Clinical features: An ugly deformity is the main complaint. Severity of deformity can be estimated by measuring the distance between the two knees with the ankles held together. If the distance is more than 8 cm, further investigations for an underlying cause are required.

Treatment: Idiopathic type usually corrects spontaneously. Shoes with an outer raise (3/16 inch) are usually prescribed. If bowing persists beyond childhood, surgical correction may be required.

GENU RECURVATUM (Fig-39.7c)
This means hyperextension at the knee joint. It may be congenital or acquired. Polio is the commonest cause of acquired genu recurvatum. Others causes are: (i) diseases known to produce lax ligaments (Marfan’s syndrome, Charcot’s arthropathy); (ii) epiphyseal growth defects; and (iii) malunited fractures.

Treatment: It is difficult. Generally, support with braces is required. In some cases, upper tibial corrective osteotomy may be required.

Table 39.2: Causes of genu valgum

<table>
<thead>
<tr>
<th>Causes</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Idiopathic</td>
<td>The commonest type.</td>
</tr>
<tr>
<td>Post-traumatic</td>
<td>Fractures of the lateral femoral or tibial condyles.</td>
</tr>
<tr>
<td></td>
<td>Damage to the lateral side of the lower femoral or upper tibial epiphyses or epiphyseal plates.</td>
</tr>
<tr>
<td>Post-infectious</td>
<td>Damage to the lateral side of the lower femoral or upper tibial epiphyses or epiphyseal plates by infection.</td>
</tr>
<tr>
<td>Neoplastic causes</td>
<td>A tumour causing a growth disparity at the epiphyseal plate e.g., chondroblastoma.</td>
</tr>
<tr>
<td>Bone softening</td>
<td>Rickets and osteomalacia.</td>
</tr>
<tr>
<td></td>
<td>Bone dysplasias.</td>
</tr>
<tr>
<td></td>
<td>Rheumatoid arthritis.</td>
</tr>
<tr>
<td>Stretching of joints</td>
<td>Charcot’s disease.</td>
</tr>
<tr>
<td></td>
<td>Paralytic disease.</td>
</tr>
<tr>
<td>Cartilage thinning</td>
<td>OA of the lateral compartment of the knee.</td>
</tr>
</tbody>
</table>

Fig-39.7 Deformities of the knee

https://kat.cr/user/Blink99/
POPLITEAL CYST
This follows a synovial rupture or its herniation in the popliteal region. It may be osteoarthritic (Morrant-Baker’s cyst) or secondary to rheumatoid arthritis. The lump is in the midline and fluctuant, but is not tender. It may shrink following knee aspiration if it is connected to the knee, or may leak or rupture so that the fluid tracks down the calf. Arthroscopic excision is the treatment of choice for symptomatic cases not responding to conservative treatment.

LOOSE BODIES IN JOINTS
This is a common problem, seen most frequently in the knee joint. A fractured osteophyte, becoming loose in an osteoarthritic, is the commonest cause. Other causes are knee osteochondritis, osteochondral fractures, synovial chondromatosis etc. In synovial chondromatosis, the number of bodies is more than 50-60. The complaint of a patient with a loose body in the joint is sudden locking of the joint. Often he can feel the loose body within the joint. Most loose bodies are radiopaque and can be seen on plain X-rays. In some, an arthroscopic examination may be required. Treatment is removal of the body arthroscopically or by opening the joint.

FLAT FOOT
This is a foot with less developed longitudinal arches.

Relevant anatomy: A normal foot has longitudinal and transverse arches. The longitudinal arch consists of medial and lateral components resting on a common pillar posteriorly – the tuberosity of the calcaneum (Fig-39.8). The talus is the keystone of the arches. It receives the body weight and transmits it to the arches below. Through the arches, the weight is transmitted to the ground via the tuberosity of the calcaneum and the heads of first and fifth metatarsals.

The integrity of the arches is maintained by the plantar ligaments, the plantar aponeurosis, the extrinsic and intrinsic muscles and the structure of the bones of the foot. Ligaments are the most important of these structures; especially the spring ligament, long plantar ligament, short plantar ligament, interosseous ligaments and plantar aponeurosis. Of the muscles, tibialis posterior and peroneus longus are more important.

Causes: Idiopathic flat foot is the commonest. There are factors related to the anatomical development of the foot which predispose to formation of a flat foot. Some common causes of flat foot are given in Table–39.3.

CONGENITAL FLAT FOOT (vertical talus)
The feet of all newborns appear flat because the postural tone of the intrinsic muscles has not yet developed; but in some, the foot is not only flat but also its undersurface is convex (rocker-bottom foot). Such a foot may be in severe valgus. This is due to a congenital anomaly where the talus lies in a vertical position rather than the normal horizontal (Fig-39.9). Diagnosis can usually be confirmed by taking an X-ray of the foot (lateral view), on which one can see the head of the talus facing vertically downwards. The navicular, along with rest of the foot, rests on the dorsal surface of the talus. Treatment is difficult. In mild cases, the footwear is modified to provide an arch support at mid foot. In severe cases, corrective surgery is required.

<table>
<thead>
<tr>
<th>Table–39.3: Causes of flat foot</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Biomechanical causes</strong></td>
</tr>
<tr>
<td><strong>Congenital</strong></td>
</tr>
<tr>
<td>• Infantile or physiological</td>
</tr>
<tr>
<td>• Congenital vertical talus</td>
</tr>
<tr>
<td><strong>Acquired</strong></td>
</tr>
<tr>
<td>• Occupational</td>
</tr>
<tr>
<td>• Obesity</td>
</tr>
<tr>
<td>• Postural</td>
</tr>
<tr>
<td>• Secondary to anatomical defect elsewhere</td>
</tr>
<tr>
<td>• External rotation of the limb</td>
</tr>
<tr>
<td>• Genu valgum</td>
</tr>
<tr>
<td>• Equinus deformity of the ankle</td>
</tr>
<tr>
<td>• Varus deformity of the foot</td>
</tr>
<tr>
<td><strong>Others</strong></td>
</tr>
<tr>
<td>• Paralytic</td>
</tr>
<tr>
<td>• Flaccid flat foot</td>
</tr>
<tr>
<td>• Spasmodic</td>
</tr>
<tr>
<td>• Due to peroneal spasm</td>
</tr>
<tr>
<td>• Arthritic</td>
</tr>
<tr>
<td>• Rheumatoid arthritis</td>
</tr>
<tr>
<td>• Traumatic</td>
</tr>
<tr>
<td>• Fracture calcaneum</td>
</tr>
</tbody>
</table>
INFANTILE FLAT FOOT
This is the most common type. The child is brought usually soon after he starts walking with the complaints that he walks on flat feet. There is sometimes a tendency for frequent falls. Arches develop as the child grows, and no special treatment is required. In some children, the feet may remain flat but there are no symptoms. Either of the parents usually have flat feet. Such individuals lead a normal life except that they are prone to developing foot strain and are unfit for joining professions requiring high levels of physical fitness (e.g., army). Foot exercises are taught and arch support is given in the shoes. In late adulthood, pain in the foot and stiffness after physical exertion are common complaints.

ACQUIRED FLAT FOOT
These are static flat feet, where there has been a structural change in the foot e.g., flail foot in a fat person, post-traumatic flat foot following a fracture of the calcaneum, flat foot secondary to genu valgum etc. The other type is spasmodic flat foot where there is a spasm of the peronei muscles due to some painful condition of the foot such as rheumatoid arthritis, tuberculosis, inter-tarsal bar etc.

DEFORMITIES OF THE TOES

HALLUX VALGUS
It is the lateral deviation of the great toe at the metatarso-phalangeal joint. Causes are many, but it is commonly due to rheumatoid arthritis, wearing pointed shoes with high heels, idiopathic etc. Usually there are no symptoms. If symptoms are present, surgical correction may be necessary. Common operations performed are: (i) osteotomy of neck of the first metatarsal (Mitchell's osteotomy); (ii) excision of the metatarsal head (Mayo's operation); and (iii) excision of the base of the proximal phalanx (Keller's operation).

HALLUX RIGIDUS
This means a stiff big toe due to OA of the metatarso-phalangeal joint of the great toe. It is usually a result of old trauma, arthritis etc.

HAMMER TOE
It is a fixed flexion deformity of an inter-phalangeal joint of the toe, usually with callosity over the prominent proximal joint.

What have we learnt?
- Fibrosis of sterno-mastoid muscle is the cause of torticollis.
- Cervical rib arises from 7th cervical vertebra.
- In infantile coxa vara, neck-shaft angle is reduced.
- Slipped capital femoral epiphysis is a disease of adolescence, commonly associated with endocrine disorders.
- Genu valgus, Genu varum, flat foot are developmental disorders which often get corrected with growth.

Additional information: From the entrance exams point of view
- Steel’s metaphyseal blanch sign and Scham’s loss of dense triangular appaearance of infero-medial articular neck is seen in SCFE.
- Phocomelia is a defect in the development of long bones.
- Madelung’s deformity is seen in the wrist.
- Congenital pseudoarthrosis of the tibia and fibula and musculoskeletal deformities are seen in neurofibromatosis type II. They are treated by internal fixation and bone grafting.
Amputations, Prosthetics and Orthotics

AMPUTATIONS

Amputation is a procedure where a part of the limb is removed through one or more bones. It should be distinguished from disarticulation where a part is removed through a joint. For simplifying this discussion, the term ‘amputation’ is applied to both these procedures. Amputation of lower limb is more commonly performed than that of upper limb; however, partial amputation of fingers or hand is common in developing countries, mainly as a sequelae of farm and machine injuries.

INDICATIONS

Overall, injury is the commonest cause of amputation in developing countries. The injury may be sustained in traffic accidents, in agriculture fields during harvesting season, in riots etc. Upper limb amputations occur commonly by kutti chopper or thresher machines. Train accidents, at a level railway crossing, unaware of a coming train, is a common cause of lower limb amputation. Some common indications for amputation are given in Table–40.1.

Indications for amputation vary in different age groups. In the elderly (50-75 years), peripheral vascular disease with or without diabetes is the main cause. In younger adults (25-30 years), amputation is most often secondary to injury or its sequelae. In children, limbs may be deficient since birth. Amongst the acquired causes, injury and malignancy top the list.

TYPES

Guillotine or Open Amputation

This is where the skin is not closed over the amputation stump, usually when the wound is not healthy. The operation is followed, after some period, by one of the following procedures for constructing a satisfactory stump:

- **Secondary closure**: Closure of skin flaps after a few days.
- **Plastic repair**: Soft tissues are repaired without cutting the bone and skin flaps are closed.
- **Revision of the stump**: Terminal granulation tissue and scar tissue, as well as a moderate amount of bone is removed and the stump reconstructed.

Table–40.1: Indications for amputation

<table>
<thead>
<tr>
<th>Indications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Injury</td>
</tr>
<tr>
<td>Peripheral vascular disease, including diabetes</td>
</tr>
<tr>
<td>Infections e.g., gas gangrene</td>
</tr>
<tr>
<td>Tumours</td>
</tr>
<tr>
<td>Nerve injuries</td>
</tr>
<tr>
<td>Congenital anomalies</td>
</tr>
</tbody>
</table>
• **Re-amputation**: This is amputation at a higher level, as if an amputation is being performed for the first time.

### Closed Amputation

This is where the skin is closed primarily (e.g., most elective amputations).

### SURGICAL PRINCIPLES – FOR CLOSED TYPE

Amputation surgery is a very important step in the rehabilitation of an amputee, and must be approached as a plastic and reconstructive procedure. Following are some of the basic principles to be followed meticulously:

a) **Tourniquet**: Use of a tourniquet is highly desirable except in case of an ischaemic limb.

b) **Ex-sanguination**: Usually a limb should be squeezed (ex-sanguinated) by wrapping it with a stretchable bandage (Esmarch bandage) before a tourniquet is inflated. It is contraindicated in cases of infection and malignancy for fear of spread of the same proximally.

c) **Level of amputation**: With modern techniques of fitting artificial limbs, strict levels adhered to in the past are no longer tenable. Principles guiding the level of amputations are as follows:

- **The disease**: Extent and nature of the disease or trauma, for which amputation is being done, is an important consideration. One tends to be conservative with dry-gangrene (vascular) and trauma, but liberal with acute life threatening infections and malignancies.

- **Anatomical principles**: A joint must be saved as far as possible. These days, it is possible to fit artificial limbs to stumps shorter than ‘ideal’ length, as long as the stump is well healed, non-tender and properly constructed.

- **Suitability for the efficient functioning of the artificial limb**: Sometimes, length is compromised for efficient functioning of an artificial limb to be fitted on a stump. For example, a long stump of an above-knee amputee may hamper with optimal prosthetic fitting.

### Classification of amputation on the basis of its level

This is given in Table 40.2.

**Table 40.2: Nomenclature of amputation by levels**

<table>
<thead>
<tr>
<th>Name</th>
<th>Part of the limb removed</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Upper limb</strong></td>
<td></td>
</tr>
<tr>
<td>Forequarter</td>
<td>Scapula + lateral 2/3 of clavicle + whole of the upper limb</td>
</tr>
<tr>
<td>amputation</td>
<td></td>
</tr>
<tr>
<td>Shoulder disarticulation</td>
<td>Removal through the gleno-humeral joint</td>
</tr>
<tr>
<td>Above elbow</td>
<td>Through the arm</td>
</tr>
<tr>
<td>amputation</td>
<td></td>
</tr>
<tr>
<td>Elbow disarticulation</td>
<td>Through the elbow</td>
</tr>
<tr>
<td>Below elbow</td>
<td>Through the forearm bones</td>
</tr>
<tr>
<td>amputation</td>
<td></td>
</tr>
<tr>
<td>Wrist disarticulation</td>
<td>Through the radio-carpal joint</td>
</tr>
<tr>
<td>Ray amputation</td>
<td>Removal of a finger with respective metacarpal from carpo-metacarpal joint</td>
</tr>
<tr>
<td>Krukenburg’s</td>
<td>Making ‘forceps’ with two forearm bones</td>
</tr>
<tr>
<td>amputation</td>
<td></td>
</tr>
<tr>
<td><strong>Lower limb</strong></td>
<td></td>
</tr>
<tr>
<td>Hindquarter</td>
<td>Whole of the lower limb with one side of the ilium removed</td>
</tr>
<tr>
<td>amputation</td>
<td></td>
</tr>
<tr>
<td>Hip disarticulation</td>
<td>Through the hip</td>
</tr>
<tr>
<td>Above knee</td>
<td>Through the femur</td>
</tr>
<tr>
<td>amputation</td>
<td></td>
</tr>
<tr>
<td>Knee disarticulation</td>
<td>Through the knee</td>
</tr>
<tr>
<td>Below knee</td>
<td>Through the tibia-fibula</td>
</tr>
<tr>
<td>amputation</td>
<td></td>
</tr>
<tr>
<td>Syme’s amputation</td>
<td>Through the ankle joint</td>
</tr>
<tr>
<td>Chopart’s amputation</td>
<td>Through talo-navicular joint</td>
</tr>
<tr>
<td>Lisfranc’s</td>
<td>Through inter-tarsal joints</td>
</tr>
<tr>
<td>amputation</td>
<td></td>
</tr>
</tbody>
</table>

### Skin flaps:

The skin over the stump should be mobile and normally sensitive, but atypical skin flaps are preferable to amputation at a more proximal level.

#### Muscles:

Muscles should be cut distal to the level of bone. Following methods of muscle sutures have been found advantageous:

- **Myoplasty** i.e., the opposite group of muscles are sutured to each other.
- **Myodesis** i.e., the muscles are sutured to the end of the stump.

These are contraindicated in peripheral vascular diseases.

#### Nerves:

Nerves are gently pulled distally into the wound, and divided with a sharp knife so that the cut end retracts well proximal to the level of bone section.
Large nerves such as the sciatic nerve contain relatively large vessels and should be ligated before they are divided.

**Major blood vessels** should be isolated and \textit{doubly} ligated using non-absorbable sutures. The tourniquet should be released before skin closure and meticulous haemostasis should be secured.

**Bone level** is decided as discussed earlier. Excessive periosteal stripping proximally may lead to the formation of ‘ring sequestrum’ from the end of the bone. Bony prominences which are not well padded by soft tissues should be resected. Sharp edges of the cut bone should be made smooth.

**Drain:** A corrugated rubber drain should be used for 48-72 hours post-operatively.

**After treatment:** Treatment, from the time amputation is completed till the definitive prosthesis fitted, is important if a strong and maximally functioning stump is desired. Following care is needed:

- **Dressing:** There are two types of dressings used after amputation surgery: (i) conventional or soft dressing; and (ii) rigid dressing. The latter has been found to be advantageous for wound healing and early prosthetic fitting.

**Soft Dressing:** This is conventional dressing using gauge, cotton and bandage.

**Rigid Dressing:** In this type of dressing, after a conventional dressing, a well moulded PoP cast is applied on to the stump at the conclusion of surgery. This helps in enhancing wound healing and maturation of the stump. In addition, the patient can be fitted with a temporary artificial limb with a prosthetic foot (pilon) for almost immediate mobilisation.

- **Positioning and elevation of the stump:** This is required to prevent contracture and promote healing.

- **Exercises:** Stump exercises are necessary for maintaining range of motion of the joint proximal to the stump and for building up strength of the muscles controlling the stump.

- **Wrapping** the stump helps in its healing, shrinkage and maturation. This can be done with a crepe bandage.

- **Prosthetic fitting and gait training:** This is started usually 3 months after the amputation.

**COMPLICATIONS**

1. **Haematoma:** Inadequate haemostasis, loosening of the ligature and inadequate wound drainage are the common causes. Haematoma results in delayed wound healing and infection. It should be aspirated and a pressure bandage given.

2. **Infection:** The cause generally is an underlying peripheral vascular disease, diabetes or a haematoma. Wound breakdown and occasionally spread of infection proximally may necessitate amputation at a higher level. A wound should not be closed whenever the surgeon is in doubt about the vascularity of the muscles or the skin at the cut end. Any discharge from the wound should be treated promptly.

3. **Skin flap necrosis:** A minor or major skin flap necrosis indicates insufficient circulation of the skin flap. It can be avoided by taking care at the time of designing skin flaps that as much subcutaneous tissues remain with the skin flap as possible. Small areas of flap necrosis may heal with dressings but for larger areas, redesigning of the flaps may be required.

4. **Deformities of the joints:** These results from improper positioning of the amputation stump, leading to contractures. A mild or moderate contracture is treated by appropriate positioning and gentle passive-stretching exercises. Severe deformity may need surgical correction.

5. **Neuroma:** A neuroma always forms at the end of a cut nerve. In case a neuroma is bound down to the scar because of adhesions, it becomes painful. Painful neuroma can usually be prevented by dividing the nerves sharply at a proximal level and allowing it to retract well proximal to the end of the stump, to lie in normal soft tissues. If it does form, it is to be excised at a more proximal level.

6. **Phantom sensation:** All individuals with acquired amputations experience some form of phantom sensation, a sensation as if the amputated part is still present. This sensation is most prominent in the period immediately following amputation, and gradually diminishes with time. Phantom pain is the awareness of pain in the amputated limb. Treatment is difficult.
SPECIAL FEATURES OF AMPUTATIONS IN CHILDREN

Amputations in children have the following special features:

• Children may have amputation since birth.
• A disarticulation is preferred to an amputation through the shaft of a long bone at a more proximal level. This is because disarticulation preserves the epiphysis distally, and therefore growth of the stump continues at the normal rate.
• As the child grows, terminal overgrowth of the bone occurs and needs frequent revisions.
• A child needs frequent changes in the size of the artificial limb.
• Children tolerate artificial limbs much better and get used to wearing it more quickly.

PROSTHESES IN ORTHOPAEDIC PRACTICE

Prosthetics is a unit of rehabilitation medicine dealing with the replacement of whole or a part of a missing extremity with an artificial device. The device so manufactured is called a prosthesis.

Uses of prostheses: A prosthesis may be used to replace a body part externally (e.g., an artificial limb) or internally (e.g., an artificial hip joint). During the past two decades considerable progress has occurred in prosthetics and rehabilitation of an amputee. Improved materials, new designs, and better evaluation and fitting techniques have resulted in prostheses that are lighter and stronger, and provide improved function, cosmesis and comfort. By and large, prosthetic replacement of the lower limb offers excellent restoration of function, and the cosmetic appearance is satisfactory. However, providing prosthesis for the upper limb is more difficult. It is almost impossible for a mechanical device to reproduce the versatility, dexterity and appearance of the natural hand. One of the most important aspect of a rehabilitation programme for a patient with amputation is to orient the patient realistically as to what the prosthesis can and cannot do.

A prosthesis can be: (i) cosmetic – to provide normal appearance or (ii) functional – to provide function of the missing part. The prosthesis does not have sensation, proprioception or muscle power. The power is provided to a prosthesis by forces arising from movement of the residual or other side limb.

These are called body powered prostheses; in others an external source of power, usually rechargeable batteries is used.

In general, more distal the amputation, more functional the individual is with the use of a prosthesis. Poor candidates for functional prosthetic fitting are the following:

a) A lower limb amputee with ischaemic limb, with an open or poorly healed wound.
b) An above-knee amputee with 45 degrees flexion contracture at the hip.
c) A below-elbow amputee with a flail elbow and shoulder.
d) Bilateral above-knee amputee with short stumps.

Parts of a prosthesis: The prosthesis consists of a socket, designed to be in close contact with the stump; a suspension to hold the socket to the stump; a prosthetic extension with substitute joints; and a terminal device (Fig-40.1). The sockets are shaped according to the shape of the stump. These could be end bearing sockets – where end of the stump bears the weight, or total contact socket – where the weight is distributed evenly throughout the surface of the socket. The socket is the fundamental component to which the remaining components are attached. Most sockets are double-walled. A plaster cast moulding of the stump is used to fabricate the socket for optimal fit, function and comfort.

* Solid Ankle Cushioned Heel.
Traditionally, the terminal device of a lower limb prosthesis is a prosthetic foot, called SACH foot. It is a simple device that has a wooden core surrounded by a solid rubber foot. This permits a combination of stiffness with pliability. The cushioned heel absorbs the impact of heel strike. In India, at Jaipur, SACH foot has been modified in a number of ways to make it suitable for barefoot walking. Essentially, these modifications are: (i) appearance of the foot is that of a normal foot; and (ii) it allows movement at forefoot and midfoot, making walking on uneven surfaces easier. Similarly, for an above-knee amputee, a prosthesis has been developed at AIIMS, New Delhi, which permits squatting and sitting cross-legged. Upper limb prostheses are named by the level of amputation. Some of the commonly used prostheses are given in Table–40.3.

Table–40.3: Commonly used prostheses

<table>
<thead>
<tr>
<th>Prosthesis Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Above-knee amputation</td>
<td>Quadrilateral socket prosthesis</td>
</tr>
<tr>
<td>Below-knee amputation</td>
<td>PTB (Patellar Tendon Bearing) prosthesis</td>
</tr>
<tr>
<td>Syme’s amputation</td>
<td>Canadian Syme’s prosthesis</td>
</tr>
<tr>
<td>Partial foot amputation</td>
<td>Shoe fillers</td>
</tr>
</tbody>
</table>

Recently, electrically operated prostheses have been developed. These have opened up a new world of freedom and function for persons with amputation, but these are very expensive. There have been a number of advances in designing of prosthesis. With the help of computers, the socket can be so designed keeping in mind particular areas over which pressure could be relieved.

**ORTHOSIS IN ORTHOPAEDIC PRACTICE**

Orthotics is the unit of rehabilitation which deals with improving function of the body by the application of a device which aids the body part. The device so manufactured is called an orthosis.

**NOMENCLATURE OF ORTHOSES**

Until recently, the terms braces, calipers, splints, and corsets, used to name and describe orthoses were not uniform. Now a logical, easy to use system of standard terminology has been developed. This system uses the first letter of the name of each joint which the orthosis crosses in correct sequence, with the letter O (for orthosis) attached at the end. Some of the commonly used orthoses are given below:

- AFO  Ankle Foot Orthosis
  (previously called below-knee caliper)
- KAFO  Knee-Ankle-Foot Orthosis
  (previously called above-knee caliper)
- HKAFO  Hip-Knee-Ankle-Foot Orthosis
  (previously called above-knee caliper with pelvic band)
- KO  Knee Orthosis (previously called knee brace)
- CO  Cervical Orthosis
  (previously called cervical collar)
- WHO  Wrist Hand Orthosis
  (previously called cock up splint)
- CTLSO  Cervico-Thoraco-Lumbo-Sacral Orthosis
  (previously called body brace)
- FO  Foot Orthosis
  (previously called surgical shoes)

Orthoses can be divided into static and dynamic types. Static orthoses are used: (i) to support an arthritic joint or a fractured bone; (ii) to prevent joint contractures in a paralytic limb; and (iii) for serial splinting of a joint to correct contracture. Dynamic orthoses are used to apply forces to a joint which is damaged by arthritis or when the muscles that normally control the joint are weak.

**USES OF ORTHOSES**

Orthoses are used for the following functions:

- To immobilise a joint or body part e.g., a painful joint
- To prevent a deformity e.g., in a polio limb
- To correct a deformity e.g., in Volkmann’s contracture
- To assist movement e.g., in a polio limb
- To relieve weight bearing e.g., in an un-united fracture
- To provide support e.g., to a fractured spine

Some common clinical conditions requiring orthoses are cervical spondylosis or whiplash injury (common cervical collar or cervical orthosis),

Table–40.4: Surgical shoes

<table>
<thead>
<tr>
<th>Shoe Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shoe with Thomas heel</td>
<td>Flat foot</td>
</tr>
<tr>
<td>(C and E heel*)</td>
<td></td>
</tr>
<tr>
<td>Shoe with arch support</td>
<td>Flat foot</td>
</tr>
<tr>
<td>CTEV shoes</td>
<td>Clubfoot</td>
</tr>
<tr>
<td>Shoe with heel pad</td>
<td>Plantar fascitis</td>
</tr>
<tr>
<td>Shoe with metatarsal pad</td>
<td>Corns</td>
</tr>
<tr>
<td>Shoe with metatarsal bar</td>
<td>Metatarsalgia</td>
</tr>
<tr>
<td>Shoe with medial raise</td>
<td>Genus valgum</td>
</tr>
<tr>
<td>Shoe with lateral raise</td>
<td>Genus varus</td>
</tr>
<tr>
<td>Shoe with universal raise</td>
<td>Short leg</td>
</tr>
</tbody>
</table>

* Crooked & Elongated heel
wrist drop (WHO), foot drop (AFO), poliomyelitis (orthosis depending upon muscle power), rheumatoid arthritis, and spinal injury (Fig-40.2). Some surgical shoe modifications made for different orthopaedic conditions are given in Table–40.4.

In recent years, quality of orthosis has improved with availability of better material and designing facilities. For polio, traditional calipers can now be replaced by aesthetic plastic inserts which can go inside the shoes and can be worn under clothes (Fig-40.2c). More and more orthoses are made available in ‘ready to use’ designs; these can be adjusted to fit individual patients. Also, custom made components of orthosis are available. These can be assembled and a caliper/orthosis made, thus saving time. In newer designs, adjustment of height of the orthosis is possible in growing children.

Contributed by: Dr. Sanjay Wadhwa, Professor, Department of Physical Medicine All India Institute of Medical Sciences, New Delhi.

What have we learnt?
- Amputations are named according to their level.
- Well constructed stump and well fitted prosthesis are key to good functions.
- Modern nomenclature of orthoses is based on the joints the orthosis is supposed to control.
- Shoe modifications help settle foot disorders.
Sports medicine has become a fast growing sub-speciality of orthopaedics. Initially, it was to do with the knee injuries in competitive athletes, but now it has expanded to include the overall care of an athlete at every level. The speciality consists of care of the injured athlete, his pulmonary and cardiovascular build up, training techniques, nutrition etc. Hence, it has become a speciality with multi-disciplinary approach involving trainers, physical therapists, cardiologists, pulmonologists, orthopaedic surgeons and general practitioners.

Arthroscopy is a technique of surgery on the joints in which tip of a thin (4 mm diameter) telescope called arthroscope is introduced into a joint, and the inside of the joint examined (Fig-41.1). This is called diagnostic arthroscopy. Once the diagnosis is made, necessary correction can be done, there and then, by introducing micro-instruments through another small skin puncture. This is called arthroscopic surgery. Today, most operations on the joints, particularly on the knee and shoulder, can be carried out arthroscopically. This technique has revolutionised the treatment of joint disorders.

**ADVANTAGES OF ARTHROSCOPIC SURGERY**

- **Minimally invasive technique:** The operation is performed through small punctures, without cutting open the joint. There is almost no blood loss.
- **Day-care surgery:** The surgery is performed on day-care basis, which means that the patient is admitted on the morning of the operation and sent home the same evening.
- **Little immobilisation required:** The only immobilisation of the knee is in the form of a small dressing for 48 hours. It allows the knee to be bent. It is possible for the patient to be up and about in the house within 48 hours. Very little or no physiotherapy is required.
- **Barely visible scars:** Since the whole operation is performed through multiple small punctures, the scars are barely visible.
- **Possible under local anaesthesia:** In selected cases, it is possible to perform the operation under local anaesthesia. The patient can literally walk into the operation theatre and walk out of it.
- **Better assessment of the joint:** Arthroscopy is the best modality for diagnosing a joint pathology. Even MRI, which is a close next to arthroscopy gives only limited information. MRI, being a sensitive investigation, can sometime pick up lesions which may not be
clinically significant (false positive), and also may miss lesions which are better picked up by actually seeing them and probing them (false negative).

- **Dynamic assessment of the joint possible:** Since it is possible to move the joint while arthroscopy is being performed, one can actually see how the structures inside the joint appear when the joint is moved. A new group of abnormalities in the joint have come to light due to the possibility of dynamic assessment. For example, an abnormal tracking of the patella (patella not moving concentrically in the trochlear notch) may be seen very convincingly arthroscopically.

- **New diagnostic possibilities:** A number of new diagnostic possibilities have come to knowledge since the availability of arthroscope. A whole new group of conditions in the knee called Plicas have been understood to be associated with patient’s symptoms. Similarly, some lesions such as SLAP* lesions, which cause shoulder pain, can be diagnosed only arthroscopically.

- **Research possibility:** Being a minimally invasive procedure, arthroscopy offers the possibility of studying the changes in the intra-articular structures e.g., changes in an implanted artificial ligament and its process of acceptance by the body.

## INDICATIONS FOR ARTHROSCOPIC SURGERY

Arthroscopy may be done to confirm a diagnosis in case it has not been possible to do so otherwise. In most cases, a provisional diagnosis is made before proceeding with arthroscopic surgery. Once the diagnosis is confirmed arthroscopically, necessary corrective measures are taken. Some of the common procedures which can be successfully performed arthroscopically are as shown in Table–41.1.

### EQUIPMENT

Arthroscopic surgery is an equipment dependent surgery. Most of the equipment is imported and expensive. The following equipment is necessary.

(a) **To visualise inside the joint:** Arthroscope, light source, fibre-optic cable, video camera and TV monitor.

(b) **To perform basic operations:** Hand instruments such as a probe, cutters, graspers, scissors, knives etc.

(c) **To perform complex operations:** Instruments such as motorized shaver, underwater cutting cautery etc. Some special instruments are required for particular operations such as ACL, PCL reconstructions.

An arthroscope is a 4 mm telescope having a 30° forward oblique angle (Fig-41.2). This obliquity helps in increasing the field of vision. Smaller size arthroscope is used for smaller joints.

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* Superior Labrum Anterior-Posterior
PROCEDURE

One needs to develop special psychomotor skills to be able to perform arthroscopic surgery. Following are the commonly scoped joints.

KNEE ARTHROSCOPY

Procedure on the knee is done with the patient under spinal or general anaesthesia. A tourniquet is applied on the thigh. The knee is cleaned and draped as would be done for any other major knee operation. The arthroscope and instruments are introduced through small cuts called portals, as shown in Fig-41.3. The commonest portal is antero-lateral portal located just lateral to the patellar tendon, at the level of the joint. This is the one through which the arthroscope is introduced. A small video camera is attached to the arthroscope, and the inside of the knee can be seen on the TV monitor. The arthroscope can be moved to different parts of the joint, and all the structures inside the joint are thoroughly examined. A second portal is used for introducing probe or other instruments. The portal used commonly for this purpose is made on the medial side of the patellar tendon (antero-medial portal). The crux of performing arthroscopic surgery is the ability to bring the tip of the instruments in front of the telescope (triangulation).

SHOULDER ARTHROSCOPY

It is very useful in making a correct diagnosis in shoulder problems. The usual approach to arthroscopic shoulder examination is via a posterior portal. This is located 2 cm below and medial to the postero-lateral angle of the acromian. Other instruments such as a probe, are passed from anterior portals, all of which are lateral to the coracoid process. In order to ensure clear visibility, and since a tourniquet cannot be used, clarity is maintained by inflating the joint with saline and maintaining it under pressure with the help of a fluid pump.

LIMITATIONS OF ARTHROSCOPIC SURGERY

CASE SUITABILITY

Arthroscopic surgery is not a panacea for each and every joint disorder. It has no role where the disease is too early and can be managed with medicines and physiotherapy. Sometimes the damage is beyond arthroscopic repair. In the knee, arthroscopic procedures have failed to produce significant relief in advanced stages of osteoarthritis. Also, a stiff knee with quadriceps scarring and adhesions cannot be managed only arthroscopically, and open surgery is required.

In the shoulder, arthroscopy is not effective if the exact cause of pain has not been diagnosed before surgery. Arthroscopy has limited role in treatment of shoulder osteoarthritis, massive rotator-cuff tears and multidirectional instability. The fascinating aspect of arthroscopic surgery is that what its limitation is today, may not remain so in future as advances in technology makes it possible.

LEARNING CURVE

Arthroscopy has a steep learning curve. One has to work within a confined space, and manoeuvering the scope as well as instruments is difficult. Rough movements can cause damage to the intra-articular structures and breakage of the rather delicate instruments.
EQUIPMENT
The equipment used for arthroscopy is expensive. The instruments being delicate, need continuous care and replenishment. A number of procedure specific instruments are necessary. One has to keep a big inventory of instruments and implants. There is no role of make-shift (Jugaad) in arthroscopic surgery.

What have we learnt?
• Arthroscopy is a fast developing field of orthopaedics with a steep learning curve.
• Most operations on joints can be performed by keyhole surgery. It is particularly useful for knee and shoulder.

Additional information: From the entrance exams point of view
Microfracturing is done for osteochondral defects.
Joint replacement is a procedure whereby one or both the components forming a joint are replaced with artificial components (called prosthesis). The prostheses are made up of special metal alloy or special high density polyethylene. A lot of research has gone into choice of the material, designing of the prosthesis and technique of their implantation. But, even till today, no artificial joint is as good as God given joint. Following are some of the commonly performed joint replacement procedures.

**HEMIARTHROPLASTY**  
*(Partial joint replacement)*

This means replacing only one side of a joint. For instance the head of the femur is replaced with an artificial component while the acetabulum is left as it is. Hemiarthroplasty is indicated in situations where only one half of the joint is affected, e.g., fracture neck of the femur in the elderly. A variety of prostheses are used – it could be a single piece (*monopolar*) or two piece (*bipolar*) prosthesis (Fig-42.1). In the latter, motion occurs between the two parts of the prosthesis itself. The prosthesis could be modular, where the prosthesis could be assembled on the table from a choice of combination of stem and head sizes. The prosthesis could be cemented (bonded to the host bone by bone cement), or uncemented (a press-fit design where natural bonding occurs between the host bone and the prosthesis). The operative technique consists of exposing the hip, dislocating the hip, resecting the ends, preparing the medullary canal for receiving the prosthesis, implanting the prosthesis in the
canal, reducing the hip and closing the wound. Post-operative rehabilitation is very important. A similar hemiarthroplasty operation is also done in the shoulder where the damaged head of the humerus is replaced with a prosthesis.

**TOTAL JOINT REPLACEMENT**

This means that both the components of the joint are replaced – e.g., the head as well as the acetabulum are replaced in a total hip replacement operation. This procedure is often required in patients suffering from arthritic afflictions of the joint. The procedure was first developed by Sir John Charnley in 1960. It has proved to be a successful operation giving 15-20 years of good function. Success of this operation depends upon the skill of the surgeon, his understanding of the basic biomechanics and the functional status of the joint before surgery.

These are expensive operations because good quality artificial joints are imported. Just for an idea, the cost of the artificial joint itself is approximately Rs. 30,000–100,000 (variable). Good quality Indian joints have become available and give satisfactory results in the hands of those using them. Apart from the joint, training of the surgeon, standard of the operation theatre and post-operative care constitute essential ingredients to making this operation successful.

Total joint replacement operations started with hip replacement, quickly went on to the knee, the shoulder, the elbow etc. Today, almost all joints of the body have been replaced with varying degree of success. Two most popular replacement operations are the hip and the knee replacement.

**TOTAL HIP REPLACEMENT**

This is an operation where both, the acetabulum and the head of the femur are replaced with artificial components. For the acetabulum, a cup made of high density polyethylene is used, and for the head a specially designed prosthesis made of metal alloy (cobalt-chromium alloy) is used. Both components are fixed in place with or without bone cement (Fig-42.2).

**Indications:** An overall indication of total hip replacement is incapacitating arthritis of the hip, severely affecting patient’s functions. It could result from a variety of reasons such as rheumatoid arthritis, osteoarthritis etc. Before considering a hip for replacement, full non-operative treatment should have been tried. Also should have been taken into consideration, other less invasive joint preserving procedures such as osteotomy, joint debridement and hemiarthroplasty. An arthrodesis may be a more suitable option in some cases.

Choice must be made between cemented and uncemented joint replacement. In general, cemented arthroplasty is used in elderly people with expected life of 10-15 years and uncemented in younger people.

**Complications:** It is a highly demanding operation. The following complications can occur:

a) **Deep venous thrombosis (DVT):** This occurs due to inadvertent manipulation of the thigh during surgery, venous stasis in the limb due to immobility, and some inherent factors in the patient which put him at a higher risk for developing DVT. Treatment consists of prevention of DVT by pharmacologic agents such as heparin and its newer derivatives, and by mechanical means such as continuous exercises of the leg, compression garments, elevation of the leg etc.

b) **Nerve palsies:** These are relatively infrequent. Sciatic nerve is the most commonly affected, particularly in procedures requiring complex hip reconstruction.

c) **Vascular injury:** This is uncommon, but can occur mainly due to technical reasons.

d) **Fracture:** These may occur during the process of implantation of the prothesis, mainly on the femoral side, or later due to stress concentration. The latter usually occurs just
distal to the tip of the femoral stem. Treatment depends upon the site and type of fracture, and it does prolong the rehabilitation.

e) **Dislocation**: The rate of dislocation of an artificial hip joint is between 1-8 per cent. It is primarily due to malpositioning of the limb during the early post-operative period, malposition of the replaced components, and later, loosening of the components.

f) **Infection**: This is the most serious of all complications. Prevention is the best way.

g) **Heterotrophic bone formation**: New bone formation around the components occurs in some cases such as ankylosing spondylitis, and results in decreased range of joint movements.

**TOTAL KNEE REPLACEMENT**

This is a relatively newer operation. In true sense, the term total knee replacement is a misnomer, since unlike the hip replacement where a part of the head and neck are actually removed and replaced with similar shaped artificial components, in the knee only the damaged articular surface is sliced off to prepare the bone ends to take the artificial components which 'cap' the ends of the bones. In a way, this could be more appropriately called a knee resurfacing operation (Fig-42.3).

**Indications**: Like in the hip, painful disabling arthritis is the main indication of doing a total knee arthroplasty. It is contraindicated if there is a focus of sepsis, extensor mechanism is insufficient or if the joint is neuropathic. Relative contraindications are: a younger patient (less than 50 years), obesity and those in physically demanding profession where results may not be as good.

**The Implant and the Procedure**: The artificial knee joint consists of the following parts (Fig-42.4):

a) A U-shaped femoral component to ‘cap’ the prepared lower end of the femur.

b) A tibial base plate to cover the cut flat surface of the upper end of the tibia. Either both cruciates or only anterior cruciate is excised.

c) A plastic tray inserted between the above two metallic components.

d) A patellar button made of polyethylene to replace the damaged surface of the patella.

The procedure consists of a series of steps based on specially designed jigs. These jigs are used in a step by step manner. The whole idea is to prepare the ends of tibia and femur to take the artificial components. The important goal of the procedure is to achieve optimal alignment of the leg and soft tissue balance between ligaments around the knee. This provides crucial stability to the artificial joint (Fig-42.5). The most recent advance in knee replacement surgery is use of computer navigation during surgery. This ensures accuracy.

It is fair to expect 10-15 years of excellent functions after a properly executed total knee replacement. The success of this operation depends upon proper selection of the patient, technically perfect execution of the procedure and sincere rehabilitation effort.

**Complications**: Following complications can occur:

1. **Infection**: Infection could be minor in the form of wound breakdown, or a major infection necessitating another operation to clean up...
the joint. Sometimes the infection may not be controlled, and removal of the prosthesis and fusion of the joint may become necessary.

2. **Deep Venous Thrombosis (DVT):** It occurs as a result of immobility. Treatment is on lines as discussed in hip section.

3. **Nerve palsy:** Common peroneal nerve palsy sometimes occurs in cases requiring dissection on the lateral side of the knee. Spontaneous recovery occurs in most cases.

4. **Fractures:** Fractures may occur while performing the operation, particularly in osteoporotic bones of a bedridden rheumatoid patient. Fractures may occur late through the bones near the prosthesis due to stress concentration in that area.

5. **Extensor mechanism complications:** Handling of extensor mechanism is required during the course of the operation. These may occur due to avulsion of the patellar tendon, inadvertent cutting of the tendon etc.

6. **Knee stiffness:** The patient may not be able to regain range of motion due to heterotropic bone formation or intra-articular adhesions.

### PARTIAL KNEE REPLACEMENT (Unicondylar Replacement)

This is a newer operation, done for a knee where only a part is damaged (partial damage). Here the knee is opened using a small incision, a cap is put on top of the damaged part without removing any ligaments, muscles, etc. In selected cases, this works as well as the more invasive total knee replacement. It is indicated in strictly partially damaged knee.

### TOTAL SHOULDER REPLACEMENT

This operation has limited indications because osteoarthritis of the shoulder is an uncommon condition. Most patients with stiff, painful shoulder due to other causes manage to live with it. Like in the hip, here also, the cup (glenoid) and the ball (head humerus) are replaced.

### TOTAL ELBOW REPLACEMENT

This is indicated in stiff and painful elbows due to rheumatoid arthritis and as a fall out of elbow injuries. The techniques have now got established to ensure good functions for 10-15 years.

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**What have we learnt?**

- Joint replacement surgery is well established with practically all joints of the body having been replaced.
- Hip, knee and shoulder replacement are common.
- Joint replacement can be partial or total depending upon whether one or both articulating surfaces are replaced.

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**Additional information: From the entrance exams point of view**

- Metal on metal joints are contraindicated in women of child bearing age.
- Most common cause of death after total hip replacement is pulmonary thromboembolism.
- Site to harvest first order, primary and cancellous bone graft is pelvis and iliac crest. It is also the best site to harvest the same.
Clinical Methods

GENERAL

The art of clinical methods can be mastered only in the ward. It essentially consists of the following:

Developing a rapport with the patient: The patient should be made comfortable in a chair or on a couch. Initial general talking will make the patient feel at home, and give the examiner an idea of the mental status of the patient. Patients usually feel at ease with one of their relatives with them. Patients have a concept that doctors are extremely busy people, and forget half the things when they face a doctor. It is the duty of a doctor to present himself as a well-composed, well-dressed, full of concern, not-in-a-hurry person.

Establishing communication with the patient so that you can understand what he says and means. Patients have their own concepts about diseases and their causes. Do not get carried away by what they say. Ascertain what they mean by intelligent cross questioning.

History taking: This consists of two parts. The first part is the presenting complaint i.e., what complaint has brought the patient to the hospital. The second part is the history of present illness i.e., the sequence of events starting from the onset of the problem till the time of presentation. It is best to let the patient say whatever he has to, or whatever he feels about his illness. You can always extract relevant information by moderating history.

Examination: This consists of examining the patient to look for salient features which may be in support of or against the diagnosis. It is best to arrive at some differential diagnoses on the basis of the history, before beginning the examination. Therefore, the aims of clinical history and examination are as follows:

a) To arrive at a diagnosis i.e., to find out the cause of the problem.
b) To find out whether the basic disease has produced any complication. For example, a patient with chronic osteomyelitis may have developed shortening of the bone due to effect of the disease on the growth plate.
c) To determine what way has the disease or its complication, if any, affected the functions of the patient. For example, in the case of a patient with affection of the lower limb there may be decreased ability to walk – there may be limp, or support may be needed for walking.

The last part of the work-up — the functional disability, is the most important in orthopaedics. It represents the way the disease has affected the functions of the patient; and it is this that concerns us and the patient the most. Upon it depends the treatment planning.

HISTORY TAKING

History taking is not merely a record of what the patient says, but it is an art of understanding and collecting information regarding what happened to the patient, what could have caused it, what way the patient has been affected by it, what treatment has the patient taken for it, and ultimately, what functional level of activity does the patient possess. For extracting relevant information, background knowledge of different diseases, their presentation, their complications etc. are necessary. The following discussion is only to form concepts in clinical orthopaedics. There is no ‘always’ in medicine. At places the reader may find over-emphasis; these are only meant to highlight some concepts.

Broadly, the history tells us about the disease aetiology (i.e., whether it is infection, tumour etc.), and the examination about the site of involvement (i.e., whether it is the bone or joint or the tendon etc., the tissue affected).

GENERAL INFORMATION

First note the name, age, sex, address and occupation of the patient. Some of this information may be helpful in thinking about the possible diagnosis, as discussed below:

Table-1: Diseases which occur at a particular age

<table>
<thead>
<tr>
<th>Disease</th>
<th>Age</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polio</td>
<td>1-2 yrs.</td>
</tr>
<tr>
<td>Rickets (Nutritional)</td>
<td>1-2 yrs.</td>
</tr>
<tr>
<td>Perthes’ disease</td>
<td>5-10 yrs.</td>
</tr>
<tr>
<td>Slipped capital epiphysis</td>
<td>12-16 yrs.</td>
</tr>
<tr>
<td>Acute Osteomyelitis/Arthritis</td>
<td>&lt;15 yrs.</td>
</tr>
<tr>
<td>Bone malignancies</td>
<td>10-20 yrs.</td>
</tr>
<tr>
<td>PIVD</td>
<td>20-40 yrs.</td>
</tr>
<tr>
<td>Rheumatoid arthritis</td>
<td>20-40 yrs.</td>
</tr>
</tbody>
</table>

Age: There are fractures which occur more commonly in children, others occur more often in adults or in the elderly. Hence by knowing the age, one can think of possible injuries which could occur at that age. Patients with congenital malformations such as CDH present early in life. Infections and bone tumours are common in children. Degenerative diseases occur at an older age. Some diseases occur in a particular age group and age consideration becomes very important in the diagnosis of these diseases (Table-1).

Sex: Some diseases are more common in males; some others in females. Table-2 lists some of these diseases. In general, all type of injuries are nearly as frequent in males as in females.

Table-2: Sex predisposition in Orthopaedic diseases

<table>
<thead>
<tr>
<th>Disease</th>
<th>Sex</th>
</tr>
</thead>
<tbody>
<tr>
<td>CDH</td>
<td>Females</td>
</tr>
<tr>
<td>Slipped epiphysis</td>
<td>Males</td>
</tr>
<tr>
<td>Rheumatoid arthritis</td>
<td>Females</td>
</tr>
<tr>
<td>Ankylosing spondylitis</td>
<td>Females</td>
</tr>
<tr>
<td>Osteomalacia</td>
<td>Females</td>
</tr>
</tbody>
</table>

Occupation: What the patient does has a lot of relevance in orthopaedics in two ways: (a) a number of complaints can be traced back to the kind of occupation. For example a patient, who is required to bend forward and lift heavy weight in the course of his job, may develop back strain; (b) in cases where cure is not possible, physical requirements of the patient become the basis for deciding the treatment. For example, a
little limp due to instability, at the cost of gaining movements at the hip, may be acceptable for a housewife. The same may be severely disabling for a heavy manual worker, who may prefer a stiff but stable hip. Similarly the living style of a patient (e.g., the habit of sitting on the floor), may become an important consideration in planning the treatment.

**PRESENTING COMPLAINTS**

History taking begins with asking the patient what exactly bothers him (i.e., what is his complaint?), and for how long. It takes a little while to be able to understand what and for how long has the problem been. One should let the patient say what he has to say, rather than obstructing his flow of thoughts and trying to fit them into the ‘sequence of questions’ you have learnt in the ‘book’. Often, the patient’s story is required to be guided by some clarification and direct questions. The following are some of the common complaints of an orthopaedic patient.

- Pain
- Difficulty in using the limb (usually upper limb)
- Inability to walk (patient is brought in a wheel chair, trolley, in the lap)
- Limp
- Deformity of a limb
- Swelling
- Stiffness
- Weakness
- Discharging sinus
- Altered sensation

There are usually more than one presenting complaints. If so, note the sequence in which they appeared.

**HISTORY OF PRESENTING ILLNESS**

One must give the patient time to settle down. A general greeting, or a nonspecific talk will make the patient at ease. One should let the patient narrate the ‘story’ of his illness. The following points need to be brought out from patient’s account:

**Onset of Symptoms**: Broadly, orthopaedic diseases can be divided into two groups – trauma related and nontrauma related. Hence, the first question to be asked is whether or not there was a trauma preceding the onset of symptoms. A number of patients may falsely implicate an unrelated episode of trauma as the cause of their disease. A detailed inquiry into the nature of the injury, the period between the injury and onset of the symptoms etc. can help in deciding whether injury did play a role in causing the disease or not. The following leading questions help in this assessment.

a) When did the injury occur in relation to the onset of symptoms i.e., immediately preceding, or a few days or weeks before*.

b) How did the injury occur? i.e., to assess whether severity of the trauma was sufficient to cause whatever the patient complains of, and also to know the exact mode of the injury.

c) Did the patient have symptoms (such as pain, swelling etc.) immediately following the trauma or did they occur after a few days or weeks?

d) Was the patient able to carry out his activities despite the injury or in the case of a child, did the child continue to play after the ‘injury’? Obviously, if this was so, the episode of trauma is unlikely to be related to the symptoms.

e) Was the patient given any treatment? Did he get any X-ray done at that time? These suggest that the injury was serious enough.

In case one is sure that the disease is not related to trauma inquire into the type of onset of the symptoms – whether acute, subacute or chronic.

**Progress of the disease**: This consists of finding out how the symptoms progressed over a period of time. The questions one must ask are: Whether it is a progressively worsening disease? Is it a disease with remissions and exacerbations? Is it a disease which came rather suddenly and subsided over a period of time? etc. etc. Any treatment carried out during this period and its effect should also be noted. At the end, one should make an assessment of the current status of the patient, his functional activity, severity of pain, etc. (Table–3).

**Table–3: Sample history**

After a history taking session, a student should be able to arrange the sequence of events in this way:

The patient was all right till .......... when he noticed .......... There was no* history of trauma related to the onset. The symptom appeared slowly**. Gradually the patient noticed additional symptoms such as (........ ). He consulted .......... and was prescribed .......... There was some*** relief with that treatment. In the meanwhile, the symptoms worsened.**** The patient could not do .......... things. Now the patient can not do .......... * or yes. ** or suddenly *** or no relief. ****or improved/ did not change.

**COMMON COMPLAINTS OF AN ORTHOPAEDIC PATIENT**

The following is an account of some of the common complaints of an orthopaedic patient and the way they are analysed.

**Pain**: This is the commonest complaint. The pain may be at the site of the disease or it may be a pain referred from some other part. The following details about the pain need to be elicited.

a) What is the exact site of the pain? Try to be as specific as possible. It helps to ask the patient to point to the site of pain.

b) Does the pain radiate to some other area? It is common in limbs to have pain originating in one part and radiating to another part (Table–4).

**Table–4: Radiation of pain**

<table>
<thead>
<tr>
<th>Site from</th>
<th>Radiation to</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neck</td>
<td>shoulder, arm</td>
</tr>
<tr>
<td>Shoulder</td>
<td>arm</td>
</tr>
<tr>
<td>Elbow</td>
<td>forearm and hand</td>
</tr>
<tr>
<td>Thoracic spine</td>
<td>girdle pain</td>
</tr>
<tr>
<td>Lumbar Spine</td>
<td>loin</td>
</tr>
<tr>
<td>Lumbo-sacral spine</td>
<td>gluteal region</td>
</tr>
<tr>
<td>SI joint</td>
<td>back of thigh and knee</td>
</tr>
<tr>
<td>Hip</td>
<td>front of thigh and knee</td>
</tr>
<tr>
<td>Thigh</td>
<td>knee</td>
</tr>
<tr>
<td>Knee</td>
<td>shin of tibia</td>
</tr>
</tbody>
</table>

c) Is the pain present at all times? A pain due to neoplasia is present at all times; it may fluctuate, but is persistent. A pain due to trauma is maximum within 4 to 6 hours of injury and then starts subsiding. A pain of inflammatory origin builds up rather suddenly and then subsides. Remissions and exacerbations are seen in pain due to chronic inflammatory diseases such as rheumatoid arthritis. A sudden appearance of pain in a rather painless
d) What aggravates or relieves the pain? A pain of mechanical origin becomes worse with activity, and improves with rest. On the other hand, a pain of chronic inflammation like osteoarthritis and rheumatoid arthritis comes up after a period of rest, and improves with activity.

e) What term can best describe the pain? This sometimes helps in localising the cause of the pain. A dull ache usually arises from a deeper structure; a shooting pain may indicate a neurogenic pain or that due to acute inflammation.

f) Are there any other symptoms associated with the pain? In most painful conditions of inflammatory, neoplastic or traumatic origin, pain is associated with the swelling, though in some cases it may not be clinically detectable. A referred pain or a pain of neurogenic origin may not have any local symptoms.

**Difficulty in using the limb:** This is usually a result of the pain. Sometimes, stiffness of joints deformity or muscle weakness may be responsible for the difficulty in using the limb.

**Inability to walk:** The cause of this is the same as above. It is important to know at what rate has the disease progressed to cause whatever limitation of walking i.e. whether it has been sudden, over days, over weeks etc.

**Limp:** This is a common early symptom in a patient with lower limb disease. Limp is of two types – painful or painless. Causes of limp are as given in Table–5.

**Table–5: Causes of Limp**

- **Painful limp**
  - Any traumatic condition of the limb
  - Any inflammatory condition of the limb e.g., TB hip
  - Osteoarthritis hip

- **Painless limp**
  - Polio affecting lower limb
  - Coxa vara deformity of the hip
  - CDH
  - Deformity of a joint or bone
  - Fused hip, knee or ankle

**Weakness:** Weakness of a limb is due to loss of muscle power. This could be secondary to disuse atrophy of the muscle or due to some neurological condition. The cause of neurological weakness may be affection of the brain, (e.g., a stroke), spinal cord (e.g., poliomyelitis), nerve (e.g. neuropathy), neuro-muscular junction (e.g., myasthenia) or muscle (e.g., myopathy). If there is no associated sensory loss, the cause may be either myopathy, neuropathy of a motor nerve, polio, or other motor neurone diseases. The onset of weakness may be sudden as in injury; or insidious as in myopathy, leprosy etc. Weakness is progressive in neuropathy and myopathy, but it improves with time (in the first few months) in polio.

**Discharging sinus:** A sinus discharging pus over a period of time, not healing with usual treatment, may indicate deeper infection. This could be an underlying bone infection. History of discharge of a piece of bone (sequestrum) from the sinus is a sure evidence of bone involvement. Other causes of a persistent discharging sinus are as listed in Table–7.

**Table–7: Causes of persistent discharging sinus**

- Generalised disease like diabetes
- Resistant bacteria
- Fungal infection
- Osteomyelitis
- Foreign body
- Epithelialisation of the sinus
- Scar tissue around the sinus
- Malignant change in the sinus

* Sometimes, a neoplastic swelling may reduce in size due to tumour degeneration.
PAST ILLNESS
Some illnesses in the past may give rise to symptoms years after apparent ‘healing’ of the disease. Some of these are as follows:
- An old injury: Osteoarthritis, presenting with pain and stiffness, is common many years after a joint is damaged due to injury or infection.
- In an old infection, recurrence may occur years after apparent healing of the infection.
- An old tubercular lesion anywhere in the body, may present as TB in the bone or joint.

PERSONAL HISTORY
The occupation of the patient, his living style, the kind of physical activity he is required to do, etc., have a bearing on his treatment.

FAMILY HISTORY
This may be relevant in a genetically transmitted disorder and in tuberculosis.

EXAMINATION
Before beginning the examination of a patient, the doctor must ensure the following:
- Patient is comfortably lying on a couch, or sitting on a chair.
- The part to be examined is exposed, and also the opposite limb, in the case of examination of a limb. This provides an opportunity of comparing the involved limb with the opposite, normal one.
- Things required to examine a patient are available. These are as follows:
  - An inch tape
  - Patellar hammer
  - Cotton wool, pins, a tuning fork
  - Skin marking pen
  - Goniometer to measure angles

GENERAL EXAMINATION: A general review of the different systems of the body, as is done in any other case, is performed.

REGIONAL EXAMINATION: This differs from region to region, and will be discussed subsequently.

GAIT ANALYSIS: Evaluation of gait constitutes an important part of orthopaedic examination for the following reasons:
- It gives a clue to the cause of gait abnormality and hence the diagnosis.
- It gives an idea of the extent of disability caused by the abnormal gait so that the treatment could be aimed at correction of the gait.

Gait can be evaluated by observing a person walk in slow motion. Normal gait has a definite pattern. It is made up of a number of gait cycles (Fig-1). One gait cycle constitutes the period from heel strike of a leg to its next heel strike. Gait cycle can be divided into two phases:
- Stance phase
- Swing phase

Stance phase: This is the part of the gait cycle when the foot is on the ground. It starts with heel strike and ends with toe off. It constitutes 60 percent of the gait cycle and consists of essentially three events:
  - Heel strike – when heel strikes the ground
  - Mid stance – when the whole foot is flat on the ground, and
  - Push off – when the body is propelled by taking a push from the foot; first the heel goes off the ground, and finally the toes.

Swing phase: This is the part of the gait cycle when the foot is off the ground. It starts with toe off and finishes when the foot is ready to strike the ground again. It constitutes 40 percent of the gait cycle, and consists of essentially the following events:
  - Acceleration: Once the foot is off the ground, the leg moves forward with the help of hip flexors.
  - Mid swing: This is the mid part of the swinging leg.
  - Deceleration: The swinging leg is slowed down to get the foot ready for heel strike.

Normal gait: In normal walking, each leg goes through a stance phase and a swing phase alternately. The rhythmic repetition of such cycles provides grace to the gait. Normal gait is mechanically efficient, and therefore, only minimal energy is consumed while walking. In case the rhythm of the gait is disturbed due to any reason, one lands up using extra energy for walking, and thus gets easily tired.

Abnormal gait: There are number of reasons for abnormality of gait. Usually there are a combination of factors. Some of the typical abnormal gait which are of value in making a diagnosis are as shown in Table–8.

EXAMINATION OF THE HIP
The hip joint is special in the following ways:
- It is a joint thickly covered with soft tissues, thus making it difficult to elicit signs.
Table-8: Abnormal gaits

<table>
<thead>
<tr>
<th>Gait</th>
<th>Pattern</th>
<th>Cause</th>
</tr>
</thead>
<tbody>
<tr>
<td>Antalgic or painful gait</td>
<td>Time taken on the affected leg is reduced. Body weight is shifted quickly to the normal leg.</td>
<td>Painful condition of the leg</td>
</tr>
<tr>
<td>Stiff hip gait</td>
<td>Lifts the pelvis, and swings it forward with leg in one piece</td>
<td>TB hip, Rheumatoid hip, Ankylosing Spondylitis</td>
</tr>
<tr>
<td>Stiff knee gait</td>
<td>The leg is circumducted and brought forward in order to get clearance.</td>
<td>TB knee, Painful stiff knee</td>
</tr>
<tr>
<td>Short limb gait</td>
<td>Becomes apparent only if the limb is shorter than 2 inches. The body on the affected side moves up and down every time the weight is borne on the affected leg.</td>
<td>Congenital short femur, Shortening secondary to fracture</td>
</tr>
<tr>
<td>Trendelenburg gait or Gluteus medius gait</td>
<td>The body swings to the affected side every time weight is borne on that side</td>
<td>Dislocated hip, CDH Congential coxa vara Fracture neck of femur Gluteus medius paralysis</td>
</tr>
<tr>
<td>Gluteus maximus lurch</td>
<td>The body swings backwards every time weight is borne</td>
<td>Gluteus maximus paralysis in polio</td>
</tr>
<tr>
<td>Quadriceps lurch</td>
<td>The person walks by hyperextending, and thereby locking the knee</td>
<td>Quadriceps paralysis</td>
</tr>
<tr>
<td>Hand-knee gait</td>
<td>The person walks with hand on the knee to prevent the knee from buckling in a quadriceps deficient knee with flexion deformity.</td>
<td>Polio</td>
</tr>
<tr>
<td>High stepping gait or Foot drop gait</td>
<td>Due to drop of the foot, the leg is lifted more in order to get clearance. First to touch the ground is the forefoot, and not the heel.</td>
<td>Common peroneal nerve palsy, Sciatic nerve palsy</td>
</tr>
<tr>
<td>Scissor gait</td>
<td>Legs are crossed in front of each other while walking due to spasm of the adductors of the hip</td>
<td>Cerebral Palsy</td>
</tr>
</tbody>
</table>

b) There are a number of diseases exclusive to the hip, for example, Perthes’ disease, slipped femoral epiphysis.

c) The compensatory mechanisms mask the deformities at the hip e.g., the flexion deformity is masked by forward tilting of the pelvis.

d) It is near the private parts, hence proper exposure and cooperation of the patient becomes difficult.

**HISTORY TAKING**

**Presenting Complaints:** As the hip is a deep joint, the patient often cannot localize the site of his problem. Rather, he complains of what he finds difficult to do. Common complaints of a patient with hip disease are as follows:

**Pain in the groin,** in the front of the thigh or sometimes in the knee. Pain in the groin can be referred pain from upper lumbar spine.

**Inability to squat:** This is due to stiffness of the hip. The stiffness may be due to painful spasm of the muscles around the hip or because of the adhesions within or around the hip.

**Limb:** This may be painless as in CDH or coxa vara, or painful as in early arthritis.

*A pain from the hip is often referred to the knee.

**Inability to walk:** This may be due to a painful condition or due to mechanical failure in the region of the hip (e.g., fracture neck of the femur, polio etc.)

**Swelling:** A swelling arising from the hip comes to notice very late, except when it is from the greater trochanter or pubic bone.

**Deformity:** Deformity of the hip may be the presenting symptom. The patient walks with a bend at the hip. The cause of the deformity could be the hip joint per se or the structures around the hip (e.g., psoas spasm due to inflammatory lesion in the vicinity of the psoas).

**HISTORY OF PRESENTING COMPLAINTS**

**Pain:** When pain is the major presenting complaint, the following details need to be elicited.

- Where is the pain? In the groin, in front of the thigh, outer side of the hip, back of the hip**.
- Does the pain radiate? Pain from the hip radiates to the knee, but not beyond. If the pain radiates beyond, its origin could be from the spine.
- Duration of the pain: Short duration pains are due to trauma, acute infections, acute arthritis etc. Long duration

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Any compensatory mechanism – increased lumbar lordosis to compensate for the flexion deformity, and pelvic tilt (as noted by position of the ASIs on the two sides) to compensate for the abduction or adduction deformities.

Gross shortening can be observed when the patient, trying to keep the leg on the ground produces plantar-flexion at the ipsilateral ankle or by keeps the opposite normal knee flexed.

Wasting of muscles on the affected side: This is an index of disuse atrophy of the muscles, and indicates long duration of the illness. Note especially the gluteal muscles and the quadriceps.

Any swelling: Note especially in the gluteal region, in the region of greater trochanter and in the groin. The greater trochanter may appear more prominent due to its proximal migration in some hip diseases.

Any active sinus or a scar of a healed sinus or previous operation: Scar of a healed sinus is puckered as against that of a superficial skin infection.

**Trendelenburg’s test:** This is a test to establish the stability of the hip. A hip is stable if the abduction mechanism of the hip is effective in preventing the pelvis from dipping on the opposite side, when weight is borne on the limb. The test and its explanation are as follows:

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**Examination**

**Exposure:** Proper exposure is essential for examination of the hip. The part of the body below the mid-thorax should be exposed, except for the area of the private parts, which should be covered with a small cloth. In Indian culture, especially in a female patient, such an exposure may not be socially acceptable. It is a must to have a female attendant/nurse while examining the hip of a female patient. While examining a patient with hip disease, the examination couch should be away from the wall. This makes it possible to go to both sides of the body to examine the respective hip. It also allows space for abduction of both the hips.

**Gait:** Observe the gait of the patient. The following are some of the common gait patterns in hip diseases. A combination of these may be present.

- **Antalgic gait:** In a painful hip disease, the patient can hardly bear weight on the affected side. So, he quickly takes the weight off the affected limb to the normal limb. Hence, he keeps the affected limb on the ground for a shorter time than the normal side.

- **Trendelenburg gait:** In a hip disease, where the hip joint is not stable, (i.e., the abductor mechanism of the hip is not effective) in order to avoid falling, the torso of the patient tilts to the affected side. In case of a bilateral unstable hip, the swing may be bilateral – the so called waddling gait (e.g., in bilateral CDH).

- **Short limb gait:** If the affected limb has become short due to some disease, when the patient walks, the whole affected side of the body dips down in order to make it possible for the patient to bring the foot to the ground. It is the ‘up and down’ movement of the half of the body, which is characteristic of a short limb gait as against the ‘sideways lurching’ seen in a Trendelenburg gait.

- **Circumduction gait:** When the hip is ‘fixed’ in abduction, there occurs apparent lengthening of the limb. In order to walk in such a situation, the patient has to take the affected ‘long limb’, in a round about fashion, and thus take the step forward.

- **In flexion deformity:** With mild flexion deformity of the hip, the patient manages to walk ‘straightly’ by compensatory lumbar lordosis. If the deformity is more than 30°, the patient can no longer compensate, and is required to stoop forward at the hip to be able to walk. This also happens in patients with ankylosing spondylitis, where compensatory lumbar lordosis is not possible due to stiffness of the spine.

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**EXAMINATION WITH THE PATIENT STANDING**

The patient should be examined first in standing position. The examiner observes him from front, from the side and from the back. The following points are noted:

- **Any obvious deformity** – flexion, abduction, adduction or rotation deformity at the hip.
muscles of the hip (mainly gluteus medius). This abductor mechanism can be compared to a lever (Fig-2. box). The fulcrum of the lever is the centre of the hip; the load is the weight of the body trying to tilt the pelvis down. This load is counter-acted by the abductor muscle force which acts through the lever-arm (the neck of the femur). Any failure in the effectiveness of the abductor mechanism causes lifting of the ASIS on the opposite (normal) side. This could occur if: (a) there is no fulcrum – e.g., dislocation of the hip, destruction of the head; (b) ineffective lever-arm (the neck of the femur) – e.g., fracture of the neck of the femur; (c) ineffective contraction of abductor muscles – e.g., weakness of the muscles due to polio or abductor muscles acting ineffectively through a short lever-arm (as in coxa vara).

**EXAMINATION WITH THE PATIENT LYING ON THE COUCH**

**Inspection:** Ask the patient to lie as straight as he can and observe the following:

**Position of the ASIS on the affected side,** if it can be distinctly seen. Normally, with the patient lying straight, both the ASISs should be square (i.e., at the same level). If the ASIS on the affected side is more proximal, an adduction deformity may be present. The reverse of this may occur in abduction deformity.

**Lumbar lordosis:** An exaggerated lumbar lordosis may be a result of tilt of the pelvis to compensate for the flexion deformity.

The patient may be keeping the hip flexed. There may be a rotational deformity of the hip as noticed by in or out turning of the patella.

**Palpation:** Following points are noted on palpation.

**Temperature** especially of the groin and over the swelling, if any.

**Tenderness** especially in the groin, over the greater trochanter. Tenderness in the gluteal region is usually due to sciatic pain arising from the spine.

**Abnormal swelling:** Any abnormal swelling is examined with regard to its site of origin, size, shape, surface, consistency, tender or not, margins, fixity to the bone and other structures.

**Thickening of the greater trochanter:** Greater trochanter is the most lateral, bony structure around the hip. It is often difficult to feel it in an obese person. The way is to palpate the shaft of the femur, and move the hand up. The most prominent bony structure at the proximal end of the thigh is the greater trochanter. It can be confirmed by moving the thigh – it should move with the thigh. Another bony prominence which can often be mistaken as ASIS is the ischial tuberosity, but can be differentiated as the latter does not move with the movement of the thigh. A dislocated head or a myositic mass around the femur can be confused as the trochanter. The trochanter is thickened in diseases involving the trochanter as in – malunited inter-trochanteric fracture, fibrous dysplasia and trochanteric bursitis.

**Proximal migration of the greater trochanter:** In diseases of the hip where the head of the femur is dislocated or damaged, or if there is a fracture of the neck of the femur, the greater trochanter is proximal than on the opposite side. This can be roughly judged by keeping the thumb at the ASISs and feeling the greater trochanters with middle finger so as to appreciate the distance between the two on two sides (Fig-3). The other method of finding this out is by drawing a Bryant’s triangle as discussed subsequently:

**Swelling:** Whether the swelling is in relation to the pelvis or the femur can be found by observing whether it moves with the femur. A dislocated head may be palpable in the gluteal region (in posterior dislocation of the hip) or in the groin (in anterior dislocation of the hip). A swelling in relation to the trochanters similarly moves on moving the thigh.

![Fig-3 Proximal migration of the greater trochanter. A is ASIS. B is greater trochanter. C is an imaginary point, and gives an idea about position of the trochanter](image)

**Deformity and Range of Movements:** Hip deformities are often not apparent because of compensatory mechanisms. It is customary to look for the deformity and test for range of motion simultaneously. In a normal person the position of complete extension is taken as zero position. In cases with deformities, the arc of movement from the deformed position of the hip to whatever further movement is possible, is noted. The following are the methods of finding out the extent of different deformities of the hip.

**Flexion deformity:** This is the commonest deformity of the hip, probably because the flexors of the hip are stronger than the extensors. When there is spasm of the muscles, the stronger flexors pull the hip in flexion. The test to evaluate the degree of the flexion deformity is called Thomas’ test, as discussed below:

- **Thomas’ Test:** Aim of the test is to remove the compensatory lumbar lordosis so that the flexion deformity becomes obvious and can be measured. The patient is asked to lie supine on a hard surface, with legs straight. He may be able to do so despite the flexion deformity by producing excessive lumbar lordosis. The same can be appreciated by the examiner passing his hand behind the patient’s lumbar spine. Now, the sound hip of the patient is flexed gradually. After the hip flexion is complete, the pelvis begins to tilt (Fig-4). This obliterates the lumbar lordosis, as can be felt by the hand under the lumbar spine. As this happens, the affected hip will automatically come to be in the deformed position (flexion position). The angle between the affected thigh and the bed is the degree of flexion deformity. One must be careful not to overflex...
the normal hip, as this results in excess tilting of the pelvis anteriorly, thereby falsely exaggerating the flexion deformity.

Problems of Thomas’ test: These are as follows:
- It is difficult to perform in a female patient as proper exposure is not always possible.
- It is difficult to perform in fat patients as in them lordosis cannot be appreciated.
- In a painful hip, the patient may be hurt during the test and thus may become uncooperative.
- It is difficult to perform this test if both the hips are affected or if the ipsilateral knee is stiff and deformed. In the case of bilateral hip deformity, better method is to put the patient prone at the edge of the couch in such a way that the body is on the couch with the legs hanging out (Fig-5). The lumbar spine is seen straight (no lordosis) and flexion deformity at the hip becomes obvious. With the palm of the hand stabilising the lumbar spine, the hip is extended gently, till the lordosis starts showing up. The angle between the body and the thigh indicates the flexion deformity at the hip.

**Range of Flexion:** Once the flexion deformity is measured the patient is asked to hold the normal knee flexed, the examiner keeping his one hand under the lumbar spine. The affected hip is now gently flexed further, beyond the position of the deformity. The arc of motion (from deformed position to the position of possible flexion) constitutes the range of motion of the hip. Normally, it is possible to flex the hip so much that the front of thigh touches the abdomen. In cases, where the hip flexion is limited, the pelvis will start tilting as the hip is forced beyond the limit of flexion. This becomes apparent as the hand under the lumbar spine can feel the movement at the spine. Hence we write that the range of hip flexion is from $20^\circ - 120^\circ$ ($150^\circ$). The figure in the bracket shows the ROM on the normal side. It is important to keep one hand over the ASIS so as to detect tilting of the pelvis while performing this test. It is possible to ‘flex’ a completely fused hip by $30^\circ - 40^\circ$, the movement actually occurring at the spine.

**Abduction Deformity:** A patient with abduction deformity compensates, and may appear ‘straight’ by tilting the pelvis (Fig-6). In abduction deformity, the pelvis on the affected side tilts down (hence the ASIS is lower). The opposite of this occurs in adduction deformity (i.e., ASIS on the affected side goes up). By removing the compensatory affect of the pelvic tilt one can make the deformity obvious, and measure it. This is done in the following way:

- **Test for detecting adduction and abduction deformities:** Let the patient lie as straight as he can with both the legs parallel to each other. In doing so, in case an abduction or adduction deformity is present, the patient will tilt the pelvis depending upon the deformity and conceal it. The examiner first palpates the ASISs on the two sides. This is done by moving his thumb from the groin laterally, and the first bony prominence detected is the ASIS. These are marked. Possibilities are that: (a) both the ASISs are at the same level (pelvis is square) which means that there is no abduction or adduction deformity; (b) ASIS on the affected side is higher (more proximal) than that on the normal side, which means that there is adduction deformity,
compensated by the pelvic tilt; (c) ASIS on the affected side is lower than that on the normal side, which means that there is abduction deformity compensated by the pelvic tilt. Once it is known that the pelvis is not square, we know which deformity is present. The next step is to square the pelvis to be able to measure the deformity. This is done as follows:

Depending upon which deformity is present, the only thing one has to do is to produce that very deformity. As this is being done, the ASIS on the affected side will move up or down as the case may be, and the pelvis will be squared (Fig-7). This is checked by feeling the two ASISs and joining them with a measuring tape. The angle between the long axis of the body and that of the leg is the degree of abduction-adduction deformity.

**Range of Adduction and Abduction:** Once the adduction-abduction deformity is measured, the next step is to see how much further adduction-abduction movement is possible. One must remember that if a hip has an adduction deformity, no abduction movement will be possible and vice versa. The only movement which may be present is movement in the direction of the deformity. This is again measured as arc of movement from deformity position to whatever further movement is possible. For example, it could be 20° of abduction deformity with further abduction from 20° to 50°. It is noted as ‘abduction deformity 20°, with ROM 20° – 50° (60°)’. The figure in the bracket is the range of abduction on the normal side. The precaution required while measuring the range of abduction and adduction movement is that the pelvis should not be allowed to move while this is being done. This is checked by keeping one hand over the opposite ASIS while moving the hip, and detecting any movement of the ASIS, (and hence that of the pelvis).

- Problem of the test: Sometimes, squaring is not possible due to fixed pelvic tilt, as may occur in a patient with lumbar scoliosis. It may also not be possible to square a pelvis with old injury where the normal anatomy is disturbed.

**Rotation Deformity:** Gross rotational deformities may be noticed by looking at the patella or the foot. Normally, the patella faces 5° to 10° outward. If it faces inwards compared to the opposite side, internal rotation deformity is present and vice-versa for the external rotation deformity. Minimal internal or external rotation deformities become more noticeable when one observes from the foot end of the patient’s bed. Comparing the two sides is important for this. Rotational deformities cannot be compensated or concealed.

**Range of Rotations:** Range of motion of rotation can be measured with the hip in extension or in flexion. With the hip extended, the leg is held by the thigh and the knee. The leg is gently turned inward and outward. This gives an idea whether there is any gross limitation of rotations. Precise measurements can be made by testing rotations with the hips in flexion. This is done on one leg at a time. The leg is held at the knee with one hand and at the ankle with the other hand (Fig-8a). The hip and knee are flexed to 90°. The rotation movement is produced at the hip by moving the leg as a lever. The arc made by the leg shows the amount of internal or external rotation. This can be compared with the same on the opposite side. Range of rotations can be tested on the two sides simultaneously (Fig-8b). This gives an instant idea of limitation of rotation on the affected side.

**Range of movement in other positions:** Range of hip movements is tested in other positions as discussed:

- **Abduction-in-flexion:** This is a good, quick method of comparing abduction movement on the two sides. The hips are flexed to 45° with the knees and ankles together. Both the knees are now ‘opened apart’ so as to allow the outer side of the knees to touch the couch. A limitation of abduction becomes obvious as the knee on the affected side remains at a higher level (Fig-9).

**Limb Length Measurement:** Shortening of the limb is common in hip diseases. Some of the shortening is compensated by the patient by:

- tilting the pelvis down on that side;
- plantar...
flexing the foot; and (c) flexing the knee on the normal side. While examining for shortening, it is important to note: (a) whether shortening is present; (b) if yes, whether it is true or apparent shortening; (c) if it is true shortening, whether it is from the hip (supra-trochanteric) or from some other part of the limb. It is customary to measure the apparent length (the length of the limb with compensatory mechanisms allowed) and true length (the actual length of the limb after removing the compensatory mechanism). Accordingly, after comparing the lengths on the two sides, apparent and true shortenings are calculated. It is the apparent shortening which concerns the patient i.e., the shortening which remains even after compensation by the body. True shortening is of significance to the clinician for diagnosis, as it is the shortening produced by the disease due to actual destruction or shortening of the bone.

There may be a situation where all bones and joints are all right, but the limb is ‘short’. This will be due to deformity at the hip, and will be called apparent shortening. There will be no true shortening in this situation. On the other hand, there may be true shortening of the bones, but the body, by compensating this shortening, may make the limb appear equal. Hence, there will be no shortening effectively (no apparent shortening), although true shortening is present, and can be detected by unmasking the compensatory mechanism.

- **Measurement of apparent length:** This is simpler to measure. The patient lies supine on the couch, as straight as he can. Both the legs should be parallel and in alignment with the body. Measurement is taken from any fixed point in the midline of the trunk (e.g., Xiphisternum, suprasternal notch etc.) upto the prominent tip of the medial malleolus. No attempt is made to correct any deformity while measuring the apparent length.

- **Measurement of true length:** The patient lies supine. The first step is to check whether the pelvis is square. If yes, the length is measured from ASIS to the tip of the medial malleolus. If the pelvis is not square, the same is done first (as discussed on page 351). As the pelvis is square, the hip deformity will show up. The limb length, from ASIS to tip of the medial malleolus is measured in the deformed position of the limb. When the normal limb is being measured for comparison, it is necessary that it be placed in the position as that of the affected limb. Hence, before measuring the normal limb, the pelvis must be squared, and the limb should be in a position, identical to that of the affected limb.

- **Leg length measurement in standing position:** In a hip without deformity, a quick and accurate method of measuring true shortening is as follows: The patient is asked to stand against a wall, facing the examiner. The pelvis may be tilted due to shortening of the limb. The examiner puts wooden blocks under the foot on the shorter side, one after another, till the ASISs on the two sides are level (Fig 10). The thickness of the blocks is measured. This indicates the amount of true shortening. Similarly, if the affected limb is longer, insert wooden blocks under the foot on the normal side, till the pelvis is square. The height of the wooden blocks indicates true lengthening of the affected limb. CT scanogram is the radiological method of accurately measuring the limb length.

- **Supra-trochanteric shortening:** Any disparity in length (true length) of the limb has to be further examined to find out as to which segment of the limb is short i.e., whether the leg is short, the thigh is short or the shortening is above the trochanter. The last one is called supra-trochanteric shortening and is important in the diagnosis of hip diseases.

- **Measurement of supra-trochanteric shortening:** A quick assessment of supra-trochanteric shortening can be made by feeling the greater trochanters in relation to respective ASISs. The patient lies supine. The examiner places his hands on both the hips as shown in the Fig 3. page 349. The thumbs are placed on ASISs, the tips of the middle fingers over the tips of the trochanters and tip of the index finger over an imaginary point at the intersection of two perpendiculats – one dropped from ASIS over the bed and the other from tip of the greater trochanter on to the first one. This gives a rough idea about proximal migration of the greater trochanter mostly* due to supra-trochanteric shortening. Supra-trochanteric shortening can be accurately measured by drawing Bryant’s triangle (Fig 11).

* Sometimes, the trochanter is pointed as in coxa vara.
Bryant’s Triangle: The patient lies supine with the pelvis square and the limbs in identical position. The tips of the greater trochanters and ASISs on both the sides are marked. A perpendicular is dropped from each ASIS on to the bed. From the tip of the greater trochanter, another perpendicular is dropped on to the first one. The tips of the greater trochanters are joined to the ASISs on the respective sides. This forms a triangle ABC. Each side of the triangle is compared with its counterpart on the normal side. The side BC of the triangle measures supra-trochanteric shortening. This may be due to: (a) dislocation of the hip; (b) central fracture-dislocation of the hip; (c) destruction of the head or acetabulum or both; (d) fracture of the neck of the femur; (e) coxa vara deformity of the hip; and (f) malunited inter-trochanteric fracture.

Some other tests have been described to roughly assess the position of the greater trochanter, but as these are difficult to perform and are not accurate, these are no longer used. Some of these are as follows:

- **Nelaton’s line:** With the hip in 90° of flexion, a line joining ASIS and ischial tuberosity passes through the tip of the greater trochanter on that side. Therefore, in cases with supra-trochanteric shortening, the trochanter will be proximal to this line.

- **Shoemaker’s line:** With the patient lying supine, the line joining ASIS and tip of the greater trochanter is extended on the side of the abdomen on both sides. Normally, these lines meet in the midline, above the umbilicus. In case one of the greater trochanters has migrated proximally, the lines will meet on the opposite side of the abdomen, and below the umbilicus.

- **Chiene’s lines:** With patient lying supine, lines are drawn joining the two ASISs and the two greater trochanters. Normally, these make two parallel lines. In case one of the trochanter has moved proximally, the lines will converge on that side.

- **Morris’ bistrochanteric test:** This is used for detecting inward migration of the greater trochanter, as may occur in a central fracture-dislocation of the hip. It is no longer used.

Telescopy: It is to test stability of a hip. The patient lies supine on the couch, with the affected side towards the examiner. Keep one hand (the right hand for examination of the left hip) to stabilise the pelvis using the thenar eminence over the ASIS and the fingers of this hand on the greater trochanter (Fig-12). The knee and the hip are flexed to 90°. With the other hand holding the knee, a gentle push and pull force is applied along the long axis of the thigh. An up and down movement of the greater trochanter can be felt by the fingers in case the hip is unstable. A positive telescopy means that the greater trochanter can be gripped between the chest wall and the arm to be able to apply push and pull force. It is easy to perform this test in young children with CDH, in which it is a very useful test.

**OTHER EXAMINATION**

Examination of the ipsilateral knee, the contralateral hip, the spine and neurovascular status of the limb must always be done in a case with hip disease. A per rectal examination may be required if it is a suspected case of TB hip, central lymph nodes should be done. Examination of the inguinal lymph nodes should be done. For any intra-abdominal cause for the deformity of the hip (e.g., a psoas abscess) may be done. Examination of the inguinal lymph nodes should be done.

**DIFFERENTIAL DIAGNOSIS**

The most important sign, the key to the diagnosis of a hip disease is movements of the hip. If movements are markedly restricted in all directions (ankylosis), the disease could be a severely damaging arthritis such as septic arthritis, tuberculosis, rheumatoid arthritis etc. If the hip movements are well preserved but there is pain and terminal limitation of movements, a secondary OA of the hip is more likely. Some movements may be more limited than others if the head is deformed, as may occur in avascular necrosis, old Perthes’ disease etc. Limitation of movement in only one direction usually indicates an extra-articular cause. For example, a child with psoas spasm due to infective focus in the vicinity of the psoas may have flexion deformity of the hip (hence no extension possible), but other movements, especially rotations, will be normal. Similarly, in coxa vara deformity of the hip, abduction is limited but with increased abduction (actually it is merely a change in the arc of motion). Hip movements may be increased in all directions in a case of non-union of fracture of the femoral neck or in a case of old Tom-Smith arthritis. The other important sign of hip disease is stability of the hip as seen by performing telescopy or by Trendelenburg’s test. If positive, it narrows the possibilities of diagnosis to a few.
The third important sign is the amount of true supra-trochanteric shortening. Only a little true shortening occurs in most hip diseases. Greater amount of shortening occurs in a dislocated hip, a non-union of fracture of the femoral neck, Tom-Smith arthritis etc. Classic deformities at the hip may also help in diagnosis.

**Examination of a patient with hip disease**

**Inspection**

*Patient Standing*
- Gait
- Obvious deformity
- Compensating mechanism - lordosis, pelvic tilt
- Shortening
- Wasting of muscles
- Swelling all around
- Sinus, scar
- Trendelenburg’s test

*Patient lying*
- ASISs on two sides
- Lumbar lordosis

**Palpation**
- Temperature
- Tenderness
- Swelling - details of swelling
- Greater trochanter
  - Thickening
  - Proximal migration

**Measurement**
- Degree of deformities: Flexion, Add.–Abd., Rotation
- Range of movement: Flexion, Add.–Abd., Rotation
- Apparent shortening/lengthening
- True shortening/lengthening, and in which component of the leg is it?
- Bryant’s triangle

**Telescopy**

**Ipsilateral knee**

**Contralateral hip**

**Spine**

**Neurovascular structures of the limb**

**Examination of the abdomen, if needed**

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**EXAMINATION OF THE KNEE**

The knee joint is special in the following ways:

a) It is the major weight bearing joint of the body, hence its diseases are very disabling.

b) It is a superficial joint, hence more prone to injuries.

c) It is a joint whose stability is dependent primarily on the ligaments, and hence ligament injuries are common.

d) The joint has intra-articular structures like the menisci, a common source of knee symptoms.

e) The joint has a large synovial space, hence it is commonly involved in the diseases affecting the synovium.

A number of orthopaedic diseases such as osteomyelitis and sarcoma occur around the knee.

The knee is therefore, affected in a wide variety of orthopaedic conditions. Broadly, these can be divided into trauma-related and non-traumatic. While examining a patient with knee complaints, one must think of the conditions affecting the knee joint per se (e.g., arthritis); those affecting the bones constituting the joint (tumours around the knee); and diseases elsewhere, which may present with pain in the knee (e.g., a referred pain from a disease of the hip, presenting as pain in the knee).

**HISTORY-TAKING**

**Presenting Complaints:** Following are the usual presenting complaints:
- Pain in the knee
- Swelling
- Deformity
- Stiffness
- Mechanical symptoms such as a give-way, something getting stuck, a catch etc.

**History of Present Illness:** A detailed account of the presenting complaints, looking at since when is it present; how it started; how the other symptoms have added on; how activity makes a difference, if any; how any treatment has made a difference; natural progress of the symptoms (whether intermittent, gradually progressive, gradually subsiding, etc.) will constitute the contents of the history of presenting complaints. Most deformities related to arthritis are painful. Painless deformities may occur in paralytic diseases (e.g., polio, CP) or if the joint is completely destroyed and fused. Deformity in arthritis is the flexion deformity; varus (bowing of legs) and valgus (knock knees) deformities may be present. Recurvatum (hyperextension) deformity may occur in polio or due to a fracture in the region of the knee.

**EXAMINATION**

The patient should be examined in the lying down position: first in supine position and then in prone position. Always compare the affected knee with the opposite, normal knee.

**Exposure:** The whole limb on the affected as well as unaffected side should be exposed. It is difficult to examine the knee when the thigh is half covered by tightly rolled up trousers or pyjama.

**Gait:** Observe the gait of the patient. A deformity of the knee will be obvious. Recurvatum deformity can be best appreciated when the patient walks. A patient with weakness of the quadriceps muscles may walk with ‘hand-knee gait’ i.e., he supports his knee on the front with his hand when he takes weight on the leg, and thus ‘prevents the knee from buckling’.

**Inspection:** The following points are noted on inspection:

**Deformity and attitude:** Flexion deformity is the commonest. Initially, it occurs due to spasm of the hamstring muscles in any painful condition of the knee. Later, the capsule and other structures around the knee develop contracture, and the deformity becomes permanent. A slight flexion deformity (basically an inability to extend the knee completely) is often termed as ‘locking’. True locking means inability to extend the knee for terminal 15° to 20°, but, flexion from there is possible. This kind of block to extension, if due to meniscus tear, is more springy’. Locking due to hamstring spasm, osteoarthritis or loose body (pseudo-locking) is not springy. Locking due to loose body occurs in different positions of the knee, and gets locked and unlocked early. Flexion movement from the position of locking may not be free and complete in osteoarthritis.
In advanced stages of arthritis, the capsule and ligaments of the knee become lax. This leads to flexion, posterior subluxation and lateral rotation of the tibia, the so-called triple displacement. The leg may be abnormally abducted (valgus) or adducted (varus).

**Swelling:** An early swelling of the knee can be appreciated on inspection. Comparison of the two knees will show that hollows, normally present on each side of the patella have been filled up. The swelling may be diffuse – which indicates an intra-articular pathology; or localized to one part of the joint. In the latter case, depending upon the location, it could be: (a) an inflamed bursa; (b) a tumour arising from within or in the vicinity of the knee; or (c) a malunited fracture of one of the bones constituting the knee. Different bursae in relation to the knee, which may present as swelling are: (a) semi-membranosus bursa causes painless oval swelling at the postero-medial aspect of the knee; (b) infra-patellar bursa (Clergyman’s knee) lying deep to the ligamentum patellae; (c) pre-patellar bursa (Housemaid’s knee) lying in front of the patella; (d) Morrant-Baker’s cyst – a posterior herniation of the synovial membrane in the popliteal fossa. A swelling can be seen distending the suprapatellar pouch, giving rise to a horseshoe shaped swelling, and is suggestive of effusion into the knee. Thickening of the capsule and bones can only be appreciated on palpation, specially on comparing it with the opposite, normal side. An extra-articular swelling of diffuse nature (e.g., cellulitis), extends all over the knee, over the skin over the knee: muscles should also be noted.

**Skin over the knee:** It may be stretched and shiny in an inflammatory disease. There may be an active sinus or a scar of a healed sinus, indicating an infective pathology. A scar of an old injury may suggest a direct hit on the knee.

**Muscle wasting:** Wasting of the thigh muscles is indicative of significant knee pathology. It can be appreciated on inspection when both thighs are exposed, side by side. Wasting of the leg muscles should also be noted.

**Skin over the knee:** It may be stretched and shiny in an inflammatory disease. There may be an active sinus or a scar of a healed sinus, indicating an infective pathology. A scar of an old injury may suggest a direct hit on the knee.

**Palpation:** Palpation is carried out to find the following:

**Temperature** of the overlying skin.

**Swelling:** If there is swelling, its nature i.e., fluid, synovial thickening or bony swelling, should be made out as discussed below:

a) **Fluid within the joint:** Fluid within the joint can be detected by one of the following tests:

   - **Cross fluctuation test:** When there is adequate fluid in the joint, it fills up the suprapatellar pouch. With one hand over the pouch and the other on the sides of the patellar tendon, one can feel cross-fluctuation between the fluid in the suprapatellar pouch and that on the side of the patella.

   - **Patellar tap:** With the knee fully extended, the suprapatellar pouch is emptied by pressing it with one hand. The fluid comes to lie between the patella and femoral condyles, and thus ‘lifts’ the patella. Now one can, with a gentle tap on the patella, feel it hitting the femoral condyle and springing back. This sign may be negative even in the presence of fluid if: (a) either there is very large, tense effusion not allowing the patella to hit the femoral condyles: or (b) if it is too little to be able to lift patella enough. It will also be negative if there is a flexion deformity of the knee. Hence, it is not a very reliable test.

   The fluid within the joint could be effusion, blood or pus. What exactly it is, can be guessed from history of onset of the symptoms and associated symptoms. Haemarthrosis builds up quickly, effusion slowly. In case there is pus inside the joint, signs of inflammation may be prominent.

b) **Synovial thickening:** Hypertrophied synovium and thickened capsule is a feature of chronic arthritis. The thickening may be appreciated in the suprapatellar pouch where it feels like a boggy swelling. Minimal synovial thickening can be appreciated by rolling one’s fingers over the medial femoral condyle where one can feel a ‘chord like’ structure, suggestive of the thickened synovium and capsule.

c) **Bony thickening** can be appreciated by palpation of the swelling. The swelling may be all around due to osteophytes, or localised to one of the condyles — as may occur in a bone tumour. In order to appreciate an early bony swelling, one should feel the bones forming the knee on both sides, and appreciate any difference in thickness, smoothness etc.

**Tenderness:** The joint may be diffusely tender, as in cases of infective arthritis. Tenderness may be localised to a particular area. Joint line tenderness, on medial or lateral side occurs in meniscus tears or osteoarthritis. With the knowledge of surface anatomy, different parts of the bones, patellar tendon, medial and lateral collateral ligaments are pressed systematically with tip of the thumb, and tenderness correlated with the underlying structure.

**Muscle wasting:** Wasting of the muscles can be measured by measuring the girth of the thigh and that of the leg at fixed points from the pole of patella. Obtain the measurement on both sides and compare.

**Deformity:** Full extension of the knee is taken as zero degree, and from there how much it is bent constitutes the flexion deformity. Attempt at gently correcting the deformity may give a ‘springy’ feel (indicative of locking), or it may result in muscle spasm in a painful knee. The block may be bony as occurs in an osteoarthritic knee. Any varus or valgus deformity can be measured with the help of a goniometer. Posterior subluxation of the tibia becomes obvious when one looks at the knee from side.

**Range of Movement:** Active range of movement shows the capability of the patient to use his muscles within the constraints of pain. It may be diminished if the muscles are weak. Passive range of movements show how much the destruction of joint articulating surfaces, and resultant adhesions have occurred. Limitation of joint movement, both flexion and extension suggests intra-articular pathology. Sometimes, there may be extra-articular block to flexion (due to bony mass behind the knee) or due to tight quadriceps muscle holding the knee on the front (as occurs in quadriceps fibrosis). Normally, the range of flexion is enough to bring the heel in contact with the buttock, but comparison with the opposite normal knee is the best. A few degrees of hyperextension is possible in a normal knee.
Tests for integrity of the ligaments: There are four main ligaments in the knee. The medial collateral ligament (MCL), lateral collateral ligament (LCL), anterior cruciate ligament (ACL) and posterior cruciate ligament (PCL). Integrity of these can be tested by the following tests.

- **Medial and lateral collateral ligaments (Fig-13a):** With the patient lying supine, the leg is lifted and held in the axilla. The knee is kept in 20° to 30° of flexion*. Gentle adduction (to test LCL) and abduction (to test MCL) force is applied, as if one is trying to ‘force open’ the joint on one or the other side. The fingers over the joint line can appreciate ‘opening-up’ of the joint. Even if the joint does not open up but an attempt to do so produces pain at the ligament, it indicates a partial tear of the ligament being tested.

- **Anterior cruciate ligament:** This is the most frequently injured ligament of the knee. It can be tested by the following methods:
  - **Anterior drawer test (Fig-13b):** The patient lies supine. The knee is flexed to 90° with the foot flat on the couch. The examiner sits lightly on the foot to stabilise it. The upper end of the tibia is held in such a way that fingers are behind the knee, the thenar eminences over the tibial condyles and the tips of the thumbs, one on each femoral condyle. The fingers behind the knee check for relaxation of the hamstrings when this test is being performed. A gentle pull is applied on the upper end of tibia and forward movement of the tibia in relation to the femoral condyles appreciated. Normally, there is a glide of up to half a centimeter. Anything more than this is suggestive of ACL laxity.
  - **Lachmann test:** This test is considered better than anterior drawer test. In this test, the knee is kept in 15 to 20° of flexion. One hand supports the thigh just above the knee and the other grasps the upper end of the tibia (Fig-13c). The extent of anterior glide indicates integrity of anterior cruciate ligament. This test is difficult to perform in bulky, muscular individuals, as in them it is difficult to hold the thigh and tibia.

- **Posterior cruciate ligament:** This ligament is injured uncommonly. One can suspect such an injury by carefully observing backward sagging of the upper end of the tibia. It can be further confirmed by the following test:
  - **Posterior drawer test:** It is like anterior drawer test except, one has to make a note of how much is it possible to push the tibia backwards.

Tests for meniscus injury: These are as follows:

- **McMurray’s test (Fig-13d):** With the patient lying on a couch, the surgeon stands at the side of the injured limb. He grasps the foot firmly with one hand and the knee with the other. The knee joint is completely flexed. The foot is rotated externally and the leg abducted. The joint is now slowly extended keeping the leg externally rotated and abducted. As the torn cartilage gets caught during this manoeuvre, the patient will experience pain or a click may be heard and felt. The angle at which these symptoms occurs indicates the position of the tear. The more posterior the tear, the more flexed position of the knee is, when the sign becomes positive. A similar test with the foot internally rotated and leg adducted is carried out for lateral meniscus tears.

- **Apley’s grinding test:** The patient lies prone on the couch. The surgeon places one hand on the back of the thigh, and with the other hand flexes the knee is flexed to 90°. The surgeon now applies compression along the long axis of the tibia while rotating it on the femur (grinding movement). Pain during this movement indicates a meniscal tear. Pain on later rotation indicates a medial meniscal tear while that on medial rotation indicates a lateral meniscal tear.

* With the knee in flexion (20-30°), the main restraints to medio-lateral instability are the collateral ligaments.
Examination with the patient lying prone: In prone position, one looks for any tenderness over the muscle attachments (sprain), any swelling over site for semimembranosus bursa, any swelling in the popliteal fossa (due to Morrant-Baker’s cyst or lymph nodes etc.).

Examination of the neurovascular structures of the limb distal to the knee is carried out in all cases. The ipsilateral hip may be examined in case no significant abnormal findings are evident on examination of the knee, as the knee pain could be a pain referred from the hip. The opposite knee should also be examined, as often knee diseases are bilateral.

**DIFFERENTIAL DIAGNOSIS**

**Patient with knee swelling:** A non-traumatic knee swelling could be due to arthritis of the knee. If it involves only one joint (monoarthritis), the usual causes are tuberculosis, septic arthritis, villo-nodular synovitis, chronic traumatic synovitis and haemophilic arthritis (see relevant sections of the book for details). In children, it could be a presentation of juvenile chronic polyarthritis – monoarticular type. Bilateral knee symptoms may be due to osteoarthritis (in elderly people), rheumatoid arthritis (in younger, usually females), gout, osteo-chondritis, etc.

**Patient with deformity:** The cause of the flexion deformity could be arthritis affecting the joint, in which case there will be painful limitation of movements. If the deformity is due to ‘burnt out’ arthritis or due to polio, it is painless. A severe limitation of movement usually indicates an infective arthritis either tubercular or pyogenic. Valgus and varus deformities of the knee occur commonly. The causes of these are as discussed on page 324.

**Pain in the knee:** It is a very complex symptom. The causes can be divided broadly into traumatic (meniscus tear, ligament tears, fracture, etc.) or inflammatory (arthritis group). Lack of significant generalised signs and presence of specific signs go in favour of traumatic causes. Often, it is difficult to diagnose the cause, and such a case is broadly termed as “Internal Deviation of the Knee” or IDK.

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**EXAMINATION OF THE ELBOW**

The elbow joint is special in the following ways:

a) It is superficial joint
b) A number of important neurovascular structures lie in close proximity of the elbow, and are prone to damage in disorders of the elbow.

c) Bones around the elbow are commonly injured during childhood.

d) The elbow is very prone to stiffness.

**HISTORY TAKING**

**Presenting complaints:** Following are the usual presenting complaints:

**Pain:** This occurs commonly in the arthritis affecting the elbow. The elbow is one of the joint affected in a polyarticular disease, but uncommonly it could be involved alone e.g., in tuberculosis of the elbow. More commonly, the pain around the elbow is due to extra-articular diseases such as lateral epicondylitis (tennis elbow); medial epicondylitis (golfer’s elbow); olecranon bursitis (student’s elbow, etc.).

**Swelling:** Pain and swelling usually occur together. With limitation of movements of the joint, an arthritic condition is more likely. A swelling without much pain may point to a neoplasm in the elbow region. History of remissions is an important indicator of inflammatory pathology.

**Stiffness:** This is a common and disabling symptom. It hampers the utility of the hand by restricting its reach. It is usually as a result of painful arthritis and associated muscle spasm. In late conditions, intra-articular and extra-articular adhesions contribute to stiffness. Elbow joint is highly prone to develop post-traumatic stiffness due to myositis.

**Deformity:** Flexion deformity occurs in any arthritic condition, or as a result of post-traumatic stiffness. Varus or valgus deformities occur, usually following fractures around the elbow. Cubitus varus occurs commonly due to a malunited supracondylar fracture of the humerus. Cubitus valgus occurs in fracture of the lateral condyle of the humerus. Hyperextension deformity occasionally occurs in a supracondylar fracture malunited in extension.

**Past history:** In a case with an old elbow injury, details of injury and treatment received are important. One must ask for history of massage, in particular. This is often the cause of stiffness due to myositis ossificans.

**EXAMINATION**

**Exposure:** The whole upper limbs on both the sides should be exposed.

**Inspection:** Following points are to be noted:

**Deformity and attitude:** Flexion deformity is obvious on putting the affected limb next to the normal limb. Varus and valgus deformities become apparent only in full extension of the elbow, because it is only in full extension that the deformed part of the lower end of the humerus articulates with the forearm bones. Hyperextension deformity is usually mild and can be appreciated by looking from the side.
**Swelling:** Early swelling of the elbow joint may be noticed on looking at the elbow from behind with the patient sitting on a stool with his hands on the thigh (elbow in about 30° flexion). Fullness on the two sides of the triceps tendon indicate fluid in the joint. A swelling just proximal to the joint, or on one side of the joint may be due to a malunited fracture, callus formation, myositis ossificans or a tumour.

**Muscle wasting:** This can be appreciated on exposing the other arm. Wasting of the arm muscles and that of the shoulder muscles should be noticed.

**Skin over the elbow:** Any healed sinus, scars of operation may be present.

**Palpation:** Following points should be noticed:

- **Temperature:** This is increased in arthritic conditions or inflammatory conditions.
- **Tenderness:** Diffuse tenderness indicates arthritis. Localised tenderness may occur in tennis elbow (over lateral epicondyle), in golfer’s elbow (over medial epicondyle), and in students’ elbow (over the tip of the olecranon).
- **Muscle wasting:** The severity of muscle wasting can be measured at a fixed distance from a bony point, and compared with the opposite normal side.
- **Stability:** Medio-lateral stability of the elbow is ascertained by alternatively stressing the elbow.

**Three bony point relationship:** This is an important sign. It is helpful in diagnosing different traumatic conditions around the elbow. With the elbow in 90° flexion, the three bony points around the elbow i.e., the medial epicondyle, lateral epicondyle and tip of the olecranon form a near-isosceles triangle (page-81). The base of the triangle is formed by the line joining the two epicondyles and the apex by the tip of the olecranon. In a supracondylar fracture the relationship of three bony points is maintained (normal). In posterior dislocation of the elbow, the triangle is reversed. In intercondylar fractures the base of the triangle is broadened.

Sometimes, identification of the three bony points becomes difficult due to a number of other bony prominences – either from a malunited fracture, or due to myositis masses of bone. It is therefore, best to identity the three bony points as follows: Palpate the medial and lateral supracondylar ridges of the humerus, about 4 to 5 cm proximal to the elbow. The most prominent points, as one follows the ridges, are the epicondyles. For identifying the tip of the olecranon, one follows the subcutaneous border of the ulna proximally.

**Deformity:** Flexion deformity of the elbow is measured considering full extension as zero. Varus, valgus deformities can be measured as an angle between the long axis of the arm and that of the forearm, with the forearm supinated. Hyperextension deformity is measured with full extension as the zero reference point.

**Range of movement:** of flexion is measured from zero position or from the position of the deformity — up to as much flexion possible. It is noted as flexion 20° to 80° (150°), with the range in the bracket being the movement on the normal side. Any pain, muscle spasm, crepitus during movement is noted. The nature of limitation of flexion; soft in arthritis, and ‘bony block’ in malunion and myositis may be appreciated.

**EXAMINATION OF THE SHOULDER**

The shoulder joint is special in the following ways:

- **a)** It is a joint complex made up of mainly two joints — the gleno-humeral joint (shoulder joint proper), and the scapulo-thoracic joint.
- **b)** It is a very unstable joint because the ball (the head of the humerus) is bigger than the cup (the glenoid). The capsule is lax, and thus allows a large range of movement.
- **c)** The shoulder is prone to stiffness, primarily because the lax capsule has a tendency to develop contracture whenever immobilised.

**HISTORY TAKING**

**Presenting complaints:** The following are some of the common presenting complaints:

- **Pain**
- **Stiffness**
- **Instability**
- **Swelling**

**HISTORY OF PRESENTING COMPLAINTS**

**Pain:** A shoulder pain may be of traumatic origin, when it is sudden onset with a history of clear cut trauma. It could be a fracture in the region of the shoulder, a subluxation or dislocation of the shoulder, or tear of one of the soft tissues around the shoulder (e.g., rotator-cuff tear, deltoid contusion, etc). Pain without a history of antecedent trauma could be from the shoulder joint per se — as may
occur in periarthritis of the shoulder; or from structures around the shoulder—as may be due to rotator-cuff tendinitis, biceps tendinitis, acromio-clavicular (AC) joint arthritis, etc. An important cause of pain in the shoulder is referred pain. It could be from cervical spine disease, visceral pathologies such as angina, cholecystitis, etc.

Location of the pain may point toward its aetiology. Pain at the top of the shoulder is usually from the AC joint. The patient can, more or less point to the site of pain with a finger. Pain at the lateral side of the arm, in the region of the deltoid is usually from rotator-cuff disease or a disease from deep shoulder joint. The patient points to the pain with whole of his palm over the deltoid. Pain in the front of the shoulder and forearm is usually due to biceps tendinitis or subacromial bursitis.

Stiffness: It is a very disabling symptom, and makes it difficult for the patient to take his hand in different directions, particularly while changing clothes. The shoulder joint is very prone to get stiff. Stiffness could be due to pain and the associated muscle spasm—as occurs in acute painful conditions. It could be primarily stiffness with not much pain, as in chronic conditions such as periarthritis. Shoulder commonly gets stiff following trauma or immobilisation due to any reason. Stiffness in all directions, specially limitation of rotations points to intra-articular pathology (e.g., periarthritis); limitation in only one direction (e.g., limitation of mainly abduction) points to a localised, extra-articular cause (e.g., rotator-cuff tendinitis). In cases of visceral diseases presenting as pain in the shoulder, the range of movement of the shoulder is normal.

Instability: The patient presents with symptom that the shoulder 'comes out'. Less frequently the complaint is more vague—as such a sudden onset pain or the arm dropping ‘dead’ (dead arm syndrome). Symptoms occur while throwing something or doing some overhead activity. A careful history into the first episode is important. The history about what happened and how it was treated helps. An X-ray taken, if any, at the time of the first episode may leave no doubt whether the ‘instability’ is due to recurrent dislocation or not.

Examination

Exposure: The patient is examined sitting on a stool, so that it is possible to go around the shoulder. The trunk is exposed (except the brassiere in a female patient).

Inspection: The shoulder is inspected from front, from side, and from behind. Following findings are noted:

Contour of the shoulder: Normally, the shoulder is round—the roundness contributed by the head of the humerus and the bulky deltoid. The shoulder may appear flat if the head is not in place (i.e., dislocated) or destroyed; or if the deltoid has got wasted due to diseases such as polio, tubercular arthritis, etc. The shoulder may appear swollen due to effusion into the joint or due to subdeltoid bursitis. If it is due to effusion, the swelling extends all around; and also, fullness can be seen (and later felt) in the axilla. Swelling may also be due to old injury or a tumour in the region of the shoulder; in which case, the swelling will be localised to one side. The AC joint may be unusually prominent in cases with AC joint subluxation or arthritis.

Muscle wasting: This occurs in any chronic problem of the shoulder. It is more marked in the region of supraspinatus and infraspinatus when there is a rupture of the rotator-cuff. In long standing cases, the deltoid and the arm muscles may also be wasted.

Skin over the swelling may be inspected for stretching and engorged veins, or any healed or active sinuses, particularly in the axilla, in an infective pathology.

Attitude: In most affections of the shoulder, the arm is held by the side of the chest. Any deviation from normal can be noticed by comparing the two sides. An attitude of internal rotation may be present in a case with posterior dislocation of the shoulder. Sometimes the shoulder girdle appears elevated due to a high scapula (Sprengle’s shoulder).

Palpation: Following points are noted on palpation:

Temperature rise, if any, of the skin overlying the shoulder should be noted.

Tenderness: Different bones forming the shoulder are examined for tenderness. Start from the sterno-clavicular joint, shaft of clavicle, lateral end of the clavicle, AC joint, acromion, spine of the scapula and borders of the scapula. The base of the neck, rotator-cuff area just distal to the margin of the acromion process, biceps tendon, and the deltoid are also examined for tenderness. A diffuse tenderness is present in an arthritis of the shoulder. Localised tenderness may indicate a disease of the underlying structure.

Swelling: If there is a diffuse, fluctuant swelling, the cause could be fluid in the joint. It is best felt in the axilla. A cystic swelling beneath the acromion, without any fullness in the axilla occurs in subacromial bursitis. The swelling may be localised to lateral end of the clavicle (due to AC joint arthritis), subacromial area (subacromial bursitis), below the coracoid (in a dislocated shoulder), or any other place (due to a tumour).

Range of movement: The movements present at the shoulder joint are flexion, extension, abduction, adduction, internal and external rotations, and circumduction. Abduction and adduction movements occur in the plane of the scapula. Thus, in abduction, the arm is carried not only laterally but also forward. Flexion and extension occur in a plane perpendicular to that in which adduction—abduction occur. It is important that the movements at the shoulder joint (the gleno-humeral joint) are tested in isolation. This is done by stabilising the scapula. Movement of the scapula may be wrongly considered as that at the shoulder, by a novice. A ‘good’ range of movement may be possible even in the presence of a stiff shoulder as a result of movement of the scapula. Following are the methods of testing passive and active shoulder movements:

Passive movements: The patient sits on the stool. The examiner stands behind him, stabilises his scapula with one hand (Fig-14a), and holds his flexed elbow with the other. The arm is gradually abducted till the scapula starts moving (this can be made out by the hand stabilising the scapula). Normally, up to 100° of abduction is possible at the gleno-humeral joint. Abduction beyond 100° occurs at the shoulder girdle. Adduction can be carried out only up to neutral position because the arm very soon comes in contact with the chest wall. The arm is brought in flexion and extension. Normal range of flexion is 75° and that of extension is 45°. For testing rotations, with one hand the scapula is stabilised, with the other, the elbow is held flexed. The forearm acts as a
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pointer, showing how much range of internal and external rotation is present (Fig-14b). Normally, about 90° of internal and external rotation are present. External rotation of the two sides can be compared by doing the above manoeuvre on both sides simultaneously (Fig-14c).

**Active movements**: The importance of examining active range of movements lies in the fact that these may be limited in patients with normal passive movements. This occurs in paralytic diseases of the shoulder, and incomplete tear of the supraspinatus tendon. Active abduction may be limited due to pain caused by impingement in the subacromial space, commonly due to supraspinatus tendinitis. In diseases of the AC joint, the extreme of shoulder abduction may be limited due to pain.

**Measurement**: This involves measuring the length of the arm, a circumference of the arm (for muscle wasting). Length of the arm is measured from the angle of acromion process to tip of the lateral condyle of the humerus. The angle of the acromion is felt as follows: one feels the spine of the scapula and palpates laterally. The angular prominence felt is the angle of the acromion. Muscle bulk is measured on both the arms at a fixed distance from the point of acromion.

**Examination of a patient with shoulder disease**

**Inspection**
- Contour of the shoulder
- Muscle wasting
- Swelling
- Skin over the swelling/shoulder
- Attitude of the arm

**Palpation**
- Temperature
- Tenderness
- Swelling and its details

**Deformity**

**Measurements**
- Arm shortening
- Muscle wasting

**Range of movements – active and passive**

**Special signs**
- Painful arc
- Drop-arm sign
- Apprehension test

**SPECIAL SIGNS**

- **Painful arc**: This is a test to detect subacromial impingement of the rotator-cuff as a cause of shoulder pain. The patient is asked to gradually abduct his shoulder with the arm rotated internally. It will be noticed that the pain starts at around 40° to 50° of abduction, and disappears at about 120° abduction. This is because the rotator-cuff gets impinged between the head of the humerus and the acromion between this arc of abduction.

- **Drop-arm sign**: This is a sign suggestive of complete tear of the rotator-cuff. The examiner abducts the arm of the patient, while stabilising the scapula with the other hand. Once 90° of abduction is achieved, the patient is asked to hold the arm in the air as the examiner leaves the elbow. In case there is a complete tear of the rotator-cuff, the patient will not be able to hold the arm, and it will drop by the side of the trunk.

- **Apprehension sign**: This is a test to detect an unstable shoulder. The shoulder is abducted and externally rotated. As the examiner loads the shoulder along the long axis of the arm, the patient becomes apprehensive, and tries to resist any further movement by using his hand or by making the shoulder stiff by muscle spasm.

**EXAMINATION OF A PATIENT WITH OLD FRACTURE**

Examination of a patient with an old fracture is carried out with an aim to find out the following:

a) Whether the fracture has united or not: If it has united, whether the union has occurred in proper position or not. When a fracture has united but not in acceptable position, it is called as malunion. If the fracture has not united, it is judged whether it is on way to union (delayed union), or there are signs suggestive of non-union, as will be discussed subsequently.

b) What secondary effects has the fracture produced on the limb as a whole (e.g., joint stiffness, muscle wasting or myositis).

c) Whether there is any damage to the neurovascular structures of the affected limb, with the injury or due to treatment.

**HISTORY TAKING**

Usually, the patient gives a history of clear trauma. Often, there is an underlying disease in the bone to have lead to the fracture.
and subsequent problem in union. So it is wise to ask a direct question whether the patient was alright before the episode of injury. Details of the type of fracture (whether open or not); details of treatment especially whether the immobilisation was sufficient; and finally, what the patient is not able to do because of the fracture; should be brought out in the history.

**EXAMINATION**

**Exposure:** The patient should be seated comfortably, with the limb supported. The whole of the limb should be exposed, as also the opposite, normal limb.

**Inspection:** Important features to be noted are as follows:

- **Gross deformity and shortening:** A comparison with the opposite limb is important.
- **Swelling** may be due to malposition of the fracture fragments or due to callus formation.
- **Any scar,** suggestive of a compound fracture in the past. Wasting of the muscles and deformity of the joints may be present.

**Palpation:** Following features are noted on palpation:

- **Tenderness at the site of the fracture:** This is an important sign of an un-united fracture.
- **Palpation of bone ends:** This is to examine whether the alignment and apposition of the bone is alright. Any bony irregularity in the form of a gap, a sharp elevation or a bend indicates an improper position of the bone. This is a definite sign of old fracture.
- **Abnormal mobility at the fracture site:** This is a pathognomonic sign of non-union of a fracture. Mobility should be tested in both antero-posterior and medio-lateral planes. It is often difficult to appreciate minimal mobility in an obese person or if the fracture is close to a joint. Presence of a crepitus while looking for abnormal mobility and also any pain on stressing the fracture site are important signs of an un-united fracture.
- **Absence of transmitted movements:** This test is another way of judging whether the fracture has united or not. It is useful in fractures of the shaft of femur, tibia and humerus. One end of the bone is rotated with one hand, while with the other, the movement is felt at the other end. If there is no transmitted movement, the fracture is mobile.

**Limb length measurement:** It is important to keep the following in mind while measuring the limb length.

- a) Did the patient have any pre-existing limb length discrepancy?
- b) The normal limb must be placed in the same position as the affected limb.

Limb length is measured from any two prominent bony points of the affected bone. Shortening indicates the amount of overlapping at the fracture site.

**Examination of the joints** proximal and distal to the affected bone to detect any deformity, swelling, limitation of movements should be done. There may be an associated injury to the nearby joint.

**Examination of nerves and vessels** going across the fracture site is carried out by examining the part of the limb distal to the fracture.

**Any complication** of the fracture, such a dystrophy etc. are noted.

**DIFFERENTIAL DIAGNOSIS**

A fracture presenting late could be one of the following:

- **United:** No mobility, no pain on stressing, no deformity or shortening.
- **Malunited:** No mobility, no pain on stressing but with deformity and/or shortening.
- **Un-united:** Abnormal mobility with or without pain at the fracture site. If there is mobility without pain, it is called pseudarthrosis. In some cases, there may be no appreciable abnormal mobility, but only pain on stressing the fracture site. Some cases of non-union appear clinically united except that the patient cannot bear weight on the limb. It is difficult to differentiate these from a delayed union, and diagnosis is made only on X-rays.

**Examination of a patient with old fracture**

**Inspection**

- Deformity
- Shortening
- Swelling
- Wasting
- Scar

**Palpation**

- Temperature
- Tenderness
- Palpation of bone ends
- Abnormal mobility
- Absence of transmitted movements

**Measurement**

- Shortening
- Muscle wasting

**Range of movement of adjacent joints**

**Ipsilateral joints**

**Neurovascular bundle**

**EXAMINATION OF A PATIENT WITH BONY LESION**

**HISTORY TAKING**

Following are relevant in the history:

**Age:** Bone tumours occur at specific ages as shown in Table–28.7, page 244. Osteomyelitis is common in children, but may present any time in life.

**Sex:** Some tumours are more common in females, and others in males. Males develop osteomyelitis more commonly than females.

**Presenting complaints:** It may be only pain in early stages of malignant tumours, but pain and swelling may be present together. Benign tumours generally have little pain until a pathological fracture occurs through the tumour (e.g., through a bone cyst).

Following are the common presenting complaints of patients presenting with bony lesion:

- Pain
- Swelling
- Pain and swelling
• Inability to use the limb, due to weakness, pain or pathological fracture.

**HISTORY OF PRESENTING COMPLAINTS**

Onset of symptoms and their progress is important in considering differential diagnosis. Most tumours are insidious in onset, but often the patient gives a history of antecedent trauma at the onset. On detailed questioning, it can be ascertained whether the trauma was related; usually it is not. An insidious onset disease which comes rather suddenly, is usually inflammatory while an insidious onset progressive disease suggests a neoplasm or a chronic infection. Course of the disease is progressive in case of a neoplasm, howsoever slow it may be. On the other hand, an inflammatory swelling has remissions and exacerbations. The various symptoms of a patient presenting with a bony lesion should be evaluated in this light.

**Pain:** This is the commonest symptom. Onset of pain is insidious, but sometimes, a history of trauma (mostly insignificant or unrelated) is present. Pain is constant at all times in a neoplastic swelling.

**Swelling:** Benign tumours present with swelling and little or no pain. In some benign tumours like osteoid osteoma, pain is the main presenting symptom. Onset of the swelling is insidious. The swelling grows at a slow rate (over months or years) or remains static. Benign swellings such as an osteochondroma is related to growth of the patient, and stop growing once the child attains maturity. A change in the rate of growth of a pre-existing swelling is ominous – there may be a malignant change in the swelling. Similarly, appearance of pain in a painless swelling may indicate a malignant change, or a complication such as a pathological fracture.

A swelling from the bone expands usually in all directions. Some swellings from the bone grow eccentrically (e.g., GCT). Swelling arising from structures other than the bone are localised to one side of the limb. Swelling near a joint may produce limitation of joint movements; either by producing a mechanical block to motion, or due to the pain associated with motion. Swelling may produce pressure on the adjacent neurovascular bundle of the limb, and produce symptoms thereof. The latter does not occurs in benign swellings. If a swelling appears benign on the basis of the history, inquire about similar swellings elsewhere (e.g., diaphyseal aclasis).

**Pain and swelling:** This is a common complaint, mostly in malignant tumours. The appearance of pain first or swelling first, is of academic significance only. It has to be differentiated from pain and swelling of inflammatory origin. One key question is whether the pain and swelling ever subsided completely or significantly. This occurs in inflammatory disorders, and not in neoplastic disorders.

**Inability to use the limb:** Inability to walk is usually a complaint in tumours of the lower limb. A pathological fracture, often without any trauma at all, may occur in a tumour which is primarily an osteolytic lesion (e.g., GCT). The other reason for not being able to use the limb may be paralysis of the limb muscles due to a tumour pressing on some nerve or a tumour arising from a nerve.

**Associated complaints** such as fever may be present in some sarcomas. It is a common symptom in Ewing’s sarcoma, and can create confusion in differentiating it from an inflammatory swelling.

**EXAMINATION**

**Position:** A patient with suspected lower limb tumour should be made comfortable on a couch and the affected leg should be well-supported. A patient with upper limb swelling can be examined, sitting on a stool. A patient with swelling of the hand should be asked to rest his hands on the table.

**Exposure:** Exposure of the whole of the involved limb is essential. It should permit examination of the most proximal part of the limb (e.g., axillary lymph nodes in case of upper limb). It is wise to have the following questions in mind before proceeding for examination.

a) From which structure of the limb is the swelling arising: Is it from the bone, joint, muscle, fascia, nerve or vessel?

b) Whether the swelling is benign or malignant?

c) Whether the swelling has produced any secondary effects such as restriction of joint movement, pathological fracture etc.

d) Whether there is any evidence of regional (to lymph nodes) or distal metastasis (to lungs etc.)?

e) Whether the neurovascular status of the limb is okay?

Whatever sequence of examination is adopted, at the end, the examiner should be able to get answers to the above questions.

**Inspection:** Begin with something most striking. It could be swelling, a deformed joint, muscle wasting etc.

**Swelling:** Following features are noted on inspection:

- **Site:** Be precise about the site of the swelling, especially whether it involves the ends* of the bone or away from the ends. Is the appearance nodular? Is the swelling all around the limb or more on one side than the other? A swelling in all directions is usually malignant. A swelling on one side may be an eccentrically growing bone swelling such as GCT, osteochondroma etc.; or it could be a swelling arising from structures outside the bone.

- **Shape and size:** An approximate size; whether the swelling is diffuse or well-defined; whether it is spherical, fusiform or irregular should be noted.

- **Surface:** Whether the surface appears smooth or lobulated should be noted.

- **Skin over the swelling:** Whether the skin over the swelling has any signs suggestive of infection – these are discharging sinuses, redness and oedema of the skin. The skin becomes tense, glossy, and often red over a rapidly growing large tumour such as an osteosarcoma. The subcutaneous veins get engorged. There may be scar of the previous biopsy or an operation.

Any pressure effects on the limb, such as oedema of the distal limb, nerve palsy etc. should be noted.

**Any deformity of the joint:** Joints develop deformities as an after effect of osteomyelitis – either due to direct involvement of the joint (see page 75), or secondary to its effect on the growth plate. In tumour, deformity may occur due to its effect on the growth plate or due to painful contracture of the joint as a result of painful spasm of the muscles around the joint. Flexion deformity at the knee is common in tumours around

* A swelling at the end of the bone right next to the joint line means that it is originating from the epiphysis (as occurs in GCT)
the knee. Varus or valgus deformities occur at different joints due to irregularity at the growth plate, as may occur in an osteochondroma or osteomyelitis.

Wasting of the group of muscles around the swelling may be noted.

Any signs suggestive of involvement of the distal neurovascular structures e.g., loss of hair, shriveled up skin, ulcer, trophic changes in the nails etc. should be noted.

Palpation: Before examining the patient, ensure that he is comfortable, and has gained the confidence that you are not going to suddenly press or move the already painful part. Start examining from the least painful area to the most painful area. Keep talking to the patient to divert his attention, and thus allay his fears. One can never be too gentle in handling a patient! It is suggested to follow a defined order while palpating; otherwise, important findings may be missed. But, at the end, answers to questions mentioned above is sought – you may like to look for some specific signs; otherwise missed.

Local temperature: Local rise of temperature is best felt with the back of the fingers. A comparison with the other side or the nearby normal skin may be useful. A local rise of temperature is a characteristic finding of an inflammatory swelling, but the skin over a sarcoma may have a rise in temperature due to increased vascularity of the tumour.

Tenderness: It is best to ask the patient to point to the most tender area, and avoid palpating that area till the end (if at all necessary). Do not just keep palpating here and there with no aim. Look at the face of the patient. He will wince with pain if the area being palpated is tender. There is no need to ask the patient about ‘pain’. Tenderness is more marked in an inflammatory swelling than in a neoplastic swelling.

Ulcers and sinuses: To be able to say that the ulcer or sinus is related to the underlying bone, one must be able to demonstrate its fixity to the bone.

Details of the Swelling: Visual impression of the swelling is now corroborated. If it is a diffuse bony swelling, it could be due to osteomyelitis. A more localised swelling occurs in a neoplasm. Following points are considered in connection with any bony swelling:

- **Size of the swelling:** Measure in two directions (e.g., length and breadth), or simply, so many by so many cms.
- **Location:** For swelling arising near a joint, one has to make out whether it is from the joint itself, very near the joint (from epiphyseal region) or a little away from the joint (from metaphyseal region). For this, it is required to be able to define the joint line clearly.
- **Extent:** Look at the extent of the swelling – whether it is growing all around, or on one side; whether it is a pedunculated or sessile swelling.
- **Surface:** Note whether the swelling is smooth, or nodular. Malignant swellings are smooth, ill-defined as against the benign swellings which may be smooth or nodular, but well-defined.
- **Margins:** Palpate the margins of the swelling. Are these well defined, or is it that the swelling merges with the surrounding tissues rather imperceptibly?
- **Consistency:** The swelling may be bony hard, as in a case of benign bony swelling such as osteochondroma. The swelling may be firm as in most sarcomas. The consistency may be variable from soft to hard in a malignant growth. In swelling due to GCT, it may be possible to elicit ‘ping pong’ type of crepitis what is called egg shell crackling. This is due to ping pong ball-like springiness of the thin rim of bone surrounding the tumour. A soft swelling is usually due to fluid in a bursa, a cold abscess, or just a lump of fat.
- **Fixity to the surrounding structures:** First thing to decide is whether the swelling is fixed to the underlying bone. Grip the swelling carefully between your fingers and appreciate mobility in more than one directions. Beware of the feeling of ‘movement’ of the muscles over the swelling as that of the swelling. A swelling fixed to the bone is usually taken as arising from the bone or periosteum. Rarely, a swelling from outside the bone may be deep, and may appear to be ‘fixed’ to the bone.

If the swelling is from a bone, assess whether it has invaded the surrounding muscles, skin, nerve or vessels as discussed below. Fixity to the surrounding structures is an important sign to differentiate between a benign and a malignant swelling.

a) **Fixity to the muscles:** A swelling infiltrating into a muscle will restrict flexibility of that muscle, and hence, there will be checkrein type of limitation of the joint motion. Also, the power of the involved muscle will be reduced.

b) **Fixity to the skin:** Skin is ‘fixed’ early in a malignant growth from a subcutaneous bone such as the tibia. If this doesn’t happen, the growth is most likely benign. In other areas, where the bone is deep, it may take long before skin fixity occurs. The skin may sometime get stretched over a huge underlying tumour, and appear fixed to it.

c) **Fixity to the nerve or vessel** will cause signs of nerve palsy or vascular insufficiency distal to the tumour. A malignant tumour in the vicinity of a nerve will nearly always infiltrate the nerve if it has reached a reasonable size.

If it is clear that the swelling is not fixed to the bone (i.e., it is not arising from the bone), make out from which structure is it arising. First step is to decide whether it is deep to the muscles within the muscles or superficial to it. For this, the patient is asked to contract the concerned muscle against resistance. A tumour which is deep to the muscle becomes less prominent; a tumour superficial to the muscle becomes more prominent; and the one in the muscle remains same. Also, the tumour superficial to the muscle remains as mobile as it was before the muscle contracted; whereas the one within the muscle becomes ‘immobile’ due to fixity provided by the contracted muscle belly.

A swelling originating from a nerve is suspected if there is a major branch of a nerve in that area. Also, tapping such a swelling may produce paraesthesias in the region of sensory distribution of the nerve. A swelling in relation to the vessel may elicit pulsation, either transmitted or expansile depending upon the exact nature of the swelling.

**Other signs:** Presence of a thrill on palpation or a bruit on auscultation may indicate a highly vascular tumour or an arterio-venous malformation.

**Any deformity of the limb:** Abnormal mobility due to pathological fracture or any limb length discrepancy should be looked for.
Movement of the neighbouring joints: Regional lymphadenopathy and distal neurovascular status should always be examined.

A general examination of the patient, to look for secondaries in a suspected case of malignancy, should be done. Also, a general review of all systems is made to assess the overall health status of the person.

DIFFERENTIAL DIAGNOSIS
Diagnosis of bone tumours depends upon: (i) age of the patient; (ii) the bone affected; and (iii) the site (epiphysis, metaphysis or diaphysis). Characteristic clinical features of some of the common bone tumours are discussed below (for details consult relevant text).

Osteosarcoma
- Age group: 15 to 25 years, and after 45 years
- Bones: Around the knee, upper humerus
- Site: Metaphysis
- Others features: Usually short duration (3–6 months); pain and swelling present; signs of a malignant swelling such as diffuse margins, fixity to muscle and skin, dilated veins etc. present.

Ewing’s Sarcoma
- Age group: 10 to 15 years, occasionally up to 30 years
- Bones: Tibia, femur, also flat bones – ileum, scapula, ribs
- Site: Diaphysis
- Others features: Usually short duration (1–2 months); may present with fever, pain and swelling and thus confused with infection. Signs of a malignant swelling present.

Chondrosarcoma
- Age group: 20 to 50 years
- Bones: Upper femur, flat bones
- Site: Diaphysis or metaphysis
- Others features: Variable duration (few months to few years), usually slow growing. History of an underlying osteochondroma, usually well defined with a little pain.

Osteoclastoma (Giant cell tumour)
- Age group: 20 to 40 years (after fusion of epiphysis)
- Bones: Around the knee, lower end of radius
- Site : Epiphysis
- Others features: Variable duration of pain (3–6 months), often presents with sudden-onset pain due to pathological fracture. Usually well-capsulated, smooth, eccentric growth, not infiltrating the nearby tissues.

Osteochondroma: The commonest benign tumour (tumour-like swelling) of the bone.
- Age group: 10 to 20 years (during growth period)
- Bones: Around the knee or upper humerus (if a solitary osteochondroma). Around the knee, shoulder, and wrist (in multiple osteochondromas)
- Site: Metaphysis or diaphysis
- Other features: Long duration (months to year), slow growing, grows as long as the child grows. A well defined, painless benign swelling may produce mechanical block to movement of the adjacent joint. Deformity or distal neural deficit, may present with complications.

Enchondroma
- Age group: 15 to 30 years
- Bones: Small bones of the hand (phalanges, metacarpals etc., usually multiple
- Site: Diaphysis
- Others features: Long duration, benign swellings. Swelling is the main complaint.

Examination of a patient with bony lesion
Inspection
- Swelling
  - Site
  - Shape and Size
  - Surface
  - Skin over the swelling
  - Pressure effects
- Deformity of the adjacent joint
- Muscle wasting
- Signs suggestive of distal neurovascular involvement

Palpation
- Temperature
- Tenderness
- Ulcer, sinuses
- Details of the swelling
  - Size-measure it
  - Site
  - Extent
  - Surface
  - Margins
  - Consistency
  - Fixity to the surrounding structures

Other Signs
- Thrill, bruit over the swelling

Deformity of the limb
Neurovascular status of the limb

EXAMINATION OF THE SPINE
A patient with spine disorder presents either with pain usually in the cervical or lumbo-sacral region; or with a deformity. The deformity may be a kyphosis (stooping forward) or scoliosis (sideways bending). Sometimes, there may be no or minimal symptoms in the back, but are primarily in the limbs: upper limb pain in cervical disorders (brachalgia), and lower limb pain in lower limb disorders (sciatica).

At times, the presenting symptom of a patient with spine disorder is neurological deficit — quadriplegia, paraplegia or paraesthesias and weakness pertaining to one or more nerve roots.

HISTORY TAKING
Presenting complaints: Following are the common presenting complaints:

Pain in the neck or back.

Radiating pain in the upper limb, girdle pain along the trunk, or sciatic pain along the back or front of the leg.

Paraesthesia and weakness in a part of the limb due to involvement of one or more nerve roots.
More extensive weakness of limbs e.g., paraplegia or quadriplegia.

HISTORY OF PRESENTING ILLNESS

Pain: Pain is a common symptom. It is mostly non-specific but following are some characteristic pains indicating a specific diagnosis.

- Sharp, shooting pain down the limb, which is exaggerated by coughing or on minimal movements. This indicates a disc prolapse.
- Dull boring pain which increases on exertion and gets relieved on rest is due to osteoarthritis.
- Pain in a young male, associated with stiffness, more early in the morning, which wears off as the person gets involved in daily chores, could be seronegative spondarthritis (SSA).
- Backache associated with pain and numbness, radiating down the leg, especially on exertion and gets relieved on rest is indicative of spinal canal stenosis. Such a symptom is called neurological claudication.
- Back pain in the dorso-lumbar region in the young may be due to traumatic or infective pathology.

Neurological symptoms: Complaints such as weakness, numbness and paraesthesias are often associated with spinal disorders. Symptoms localised to one limb usually indicate disc pathology. Bilateral lower limb weakness and loss of sensation occurs usually in dorsal and dorso-lumbar spine diseases. A cauda equina syndrome presentation occurs in lumbar spine diseases. Neurological symptoms in TB spine and in tumours are gradual in onset; in disc prolapse these are rather sudden.

EXAMINATION

Exposure: A proper exposure of the whole spine is crucial. A female patient should be asked to change and wear a gown open from the back. A female attendant/nurse should be present when examining a female patient.

Position: A patient with cervical spine disease is examined sitting on a stool, so that the examiner can observe from front, side or back. A patient with lumbar spine or dorso-lumbar

spine disease is examined first standing, then lying supine and then lying prone.

Inspection: Following points are noted on inspection:

Gait: Observe the gait as the patient walks into the room. A side lurching gait may suggest a scoliosis. A patient with painful condition of the spine walks rather cautiously, with short steps and a stiff spine. A patient with acute disc prolapse has a forward stoop and sideways tilt of the torso on the pelvis.

Deformity: Normally, the neck has lordosis (forward curve), the dorsal spine is kyphotic and lumbar spine lordotic. The nape of the neck is in a straight line above the natal cleft. The position of the shoulder, scapular blades, lumbar hollows and iliac wings is symmetrical. Any deviation could be due to a disease.

A diffuse kyphosis occurs in ankylosing spondylitis, Schuermann’s disease, osteoporosis etc. A localised kyphosis may be very sharp due to collapse of one vertebra (a knuckle type) or localised to collapse of 2 to 3 vertebrae (gibbus
Examination of a patient with spine disease

Inspection
- Gait
- Posture
- Deformity
- Swelling
- Paravertebral muscle spasm

Palpation
- Tenderness
- Swelling
- Prominence of the spinous processes

Range of movements

Neurological testing of the legs
- Motor
- Sensory
- Reflexes

SI joint examination

Examination of abdomen, chest

- Laségue Test: This is a modification of SLRT where first the hip is lifted to 90° with the knee bent. The knee is then gradually extended by the examiner. If nerve stretch is present, it will not be possible to do so, and the patient will experience pain in the back of the thigh or leg.
- Motor power: These are examined in different muscle groups of the limb, especially that of EHL, ankle dorsiflexors in a case of disc prolapse.
- Sensory Loss: These are examined dermatome-wise, especially in L4, L5, S1 dermatomes.
- Reflexes: The deep and superficial reflexes, and Babinski reflex are examined.

Examination of the lower limb: Length of both the legs should be measured. Sometimes, a disparity is the cause of scoliosis. Both the hips should be examined, as there could be simultaneous involvement of the hips and spine; or the hip disease may be responsible for the spine deformity.

General examination: Following examination should be done in a case with spine disease:
- Look for cold abscesses away from the site of tuberculosis of the spine (see page 187).
- Chest should be examined to look for a tubercular focus there or to rule out an old chest disease as a cause of scoliosis.
- Examination of the breast, kidney, prostate, thyroid and abdomen is necessary if secondaries are being suspected in the spine.
Orthopaedic Terminology

**FRACTURES**

Fracture: A break in the continuity of bone
- **Avulsion**: bone piece pulled off by attached muscle or ligament
- **Burst**: vertebral body fracture where fragments burst out in different directions
- **Chip**: just a sliver of bone chipped off
- **Closed (Simple)**: the skin over the fracture intact
- **Comminuted**: fracture in multiple pieces
- **Complicated**: fracture associated with a complication such as a vascular injury
- **Compression**: vertebral body fracture where the body is compressed
- **Displaced**: fragments separated
- **Greensstick**: fracture in children where one cortex breaks and the other cortex bends
- **Impacted**: fracture where one fragment gets jammed with the other fragment
- **Open (Compound)**: the fracture communicates with outside through a rent in the skin and overlying soft tissues.
- **Pathological**: the broken bone had an underlying weakness
- **Segmental**: fracture at two levels in the same bone
- **Stress (Fatigue)**: fracture caused due to repeated stress at one point
- **Traumatic**: cause of the fracture is injury
- **Undisplaced**: not displaced, only a crack

**FRACTURES WITH EPONYMS**

- **Aviators**: fracture of the neck of the talus
- **Barton’s**: distal radius, intra-articular fracture
- **Bennett’s**: fracture of base of the 1st metacarpal, intra-articular
- **Boxers’**: fracture of neck of 5th metacarpal
- **Bumper**: comminuted fracture of lateral condyle of the tibia
- **Chauffer’s**: radial styloid fracture
- **Colles’**: distal radius, extra-articular fracture with dorsal tilt of the distal fragment
- **Cotton’s**: trimalleolar ankle fracture
- **Galleazzi**: fracture of distal 1/2 of the radius with dislocation of distal radio-ulnar joint
- **Hangman’s**: fracture pedicle-lamina of C2 vertebra
- **Jane’s**: fracture of the base of the 5th metatarsal
- **Malgaigne’s**: pelvic ring disruption with both pubic rami and sacro-iliac injury on the same side
- **Mallet**: avulsion of attachment of ext tendon from base of the distal phalanx
- **March**: stress fracture of shaft of 2nd metatarsal
- **Monteggia**: fracture of proximal 1/2 of the ulna with dislocation of head of the radius
- **Night stick**: isolated fracture shaft of the ulna
- **Pott’s**: bimalleolar ankle fracture
- **Rolando**: fracture of base of 1st metacarpal, extra-articular
- **Smith’s**: distal radius fracture, extra-articular with volar tilt of the distal fragment

**DISLOCATIONS**

Dislocation: Complete separation of joint surfaces
Subluxation: Incomplete separation of joint surfaces
- **Congenital**: present at birth
- **Acquired**: develop later in life
- **Habitual**: occurs every time the joint is moved
- **Pathological**: occurs due to some disease of the joint, e.g. sepsis.
- **Recurrent**: occurs again and again
- **Traumatic**: due to Injury

**DISLOCATIONS WITH EPONYMS**

- **Chopart’s**: dislocation through talo-navicular joints
- **Divergent**: elbow dislocation where ulna and radius dislocate in opposite directions
- **Lisfranc’s**: dislocation through inter tarsal joint
- **Lunate**: wrist injury where lunate bone comes out to lie in front of other carpal bones
- **Luxatio erecta**: inferior dislocation of shoulder
- **Otto pelvis**: gradual shift of the acetabulum into the pelvis (e.g. in osteomalacia)
- **Perilunate**: wrist injury where the lunate remains in its place and the other carpal bones dislocate around it dorsally
- **Spondylolisthesis**: movement of one vertebra over another (usually L4 over L5)
- **Sprain**: A break in the continuity of a ligament
- **Strain**: A break in muscle fibres

**SIGNS AND TESTS**

- **Adson’s test**: for thoracic outlet syndrome
- **Allen’s test**: for testing patency of radial and ulnar arteries
- **Alli’s test**: for CDH
- **Anvil test**: for testing tenderness of the spine
- **Apley’s grinding test**: for meniscus injury
- **Apprehension test**: for recurrent dislocation of the shoulder
- **Barlow’s test**: for CDH
- **Blue sclera**: Osteogenesis imperfecta
- **Bryant’s test**: for anterior dislocation of the shoulder
- **Callaways’ test**: for anterior dislocation of the shoulder
- **Chovstek’s sign**: for tetany
- **Claw hand**: for ulnar nerve injury
### SOME ABBREVIATIONS USED IN ORTHOPAEDICS

- **Abd**: Abduction
- **ACL**: Anterior cruciate ligament
- **Add**: Adduction
- **ADL**: Activities of daily living
- **AE**: Above elbow
- **AJ**: Ankle jerk
- **AK**: Above knee
- **AP**: Antero-posterior
- **ASIS**: Anterior superior iliac spine
- **B/L**: Bilateral
- **BB**: Both bones
- **BE**: Below elbow
- **BJ**: Biceps jerk
- **BK**: Below knee
- **Bx**: Biopsy
- **CDH**: Congenital dislocation of the hip
- **CP**: Cerebral palsy
- **CTEV**: Congenital talipes equino-varus
- **DIP**: Distal inter-phalangeal
- **DVT**: Deep vein thrombosis
- **EMG**: Electromyography
- **FDP**: Flexor digitorum profundus
- **FDS**: Flexor digitorum superficialis
- **FFD**: Fixed flexion deformity
- **FK**: Fixed flexion knee
- **FWD**: Fixed flexion wrist
- **HKA**: Hip-knee-ankle
- **HLD**: Hip-lumbar-dorsal
- **HLA**: Human leukocyte antigen
- **IDK**: Internal derangement of the knee
- **KJ**: Knee jerk
- **Lat.**: Lateral
- **LM**: Lateral meniscus
- **LS**: Lumbo-sacral
- **MM**: Medial meniscus
- **MP**: Metacarpo-phalangeal
- **MWD**: Micro wave diathermy
- **NCV**: Nerve conduction velocity
- **NWB**: Non-weight bearing
- **OA**: Osteoarthritis
- **ORIF**: Open reduction internal fixation
- **PCL**: Posterior cruciate ligament
- **PCL**: Proximal inter-phalangeal
- **PIP**: Proximal interphalangeal
- **PIVD**: Prolapsed intervertebral disc
- **PoP**: Plaster of Paris
- **PSS**: Peripheral systemic sclerosis
- **PTB**: Patellar tendon bearing
- **PWB**: Partial weight bearing
- **RA**: Rheumatoid arthritis
- **RoM**: Range of motion
- **SI**: Sacro-iliac
- **SLAP**: Superior labrum anterior posterior tear
- **SLE**: Systemic lupus erythematosus
- **SLRT**: Straight leg raising test
- **SOS**: If necessary
- **SSA**: Sero-negative spond-arthritis
- **SWD**: Short-wave diathermy
- **THR**: Total hip replacement
- **TJ**: Tendon jerk
- **TKR**: Total knee replacement
- **US**: Ultrasonic waves
- **WNL**: Within normal limit

### SOME ORTHOPAEDIC TERMS

- **Arthrocentesis**: aspiration of a joint
- **Arthrodesis**: fusing a joint
- **Arthrography**: imaging a joint with dye inside it
- **Arthrolysis**: releasing a stiff joint
- **Artroplasty**: creating a new joint
- **Arthroscopy**: looking into a joint with a telescope
- **Artrotomy**: opening up a joint
- **Closed reduction**: setting a fracture in position by manipulation
- **Epiphysiodysis**: knocking out an epiphyseal plate to stop its growth
- **Fenestration**: removing ligamentum flavum (from in-between the laminae)
- **Hemi-laminectomy**: removing half of the lamina
- **Laminectomy**: removing whole of the lamina
- **Laminotomy**: making a hole in the lamina
- **Neurectomy**: cutting a nerve (as in CP)
- **Neurolysis**: releasing a tight nerve
- **Neurorrhaphy**: repairing a nerve
- **Open reduction**: setting a fracture by operation
- **Osteoclasis**: rebreaking a uniting fracture (to obtain better reduction)
- **Osteogenesis**: new bone formation
- **Osteosynthesis**: reconstructing a fractured bone
• Osteotomy: making a cut in the bone
  • Derotation osteotomy for CDH
  • Dimon-Houston osteotomy for inter-trochanteric fracture
  • Dwyer’s osteotomy for CTEV
  • French osteotomy for cubitus varus deformity
  • High tibial osteotomy for OA knee with varus
  • McMurray’s osteotomy for fracture neck femur
  • Pauwel’s osteotomy for fracture neck femur
  • Pemerton osteotomy for CDH
  • Salter’s osteotomy for CDH
  • Sandwich osteotomy for slipped epiphysis
  • Spino osteotomy for ankylosing spondylitis
  • Wilson’s osteotomy for congenital coxa vara
• Tendon transfers: changing the direction or action of a tendon
  • Tenodesis: attaching a tendon to another tendon or bone
  • Tenolysis: releasing a tendon from adhesions
  • Tenotomy: cutting a tendon

IMPLANTS AND THEIR USES
• Austin-Moore prosthesis: for fracture neck of the femur
• Baksi’s prosthesis: for elbow replacement
• Buttress plate: for condylar fractures of the tibia
• Charnley prosthesis: for total hip replacement
• Condylar blade plate: for condylar fractures of the femur
• DHS: for inter-trochanteric fracture
• Ender’s nail: for fixing inter-trochanteric fracture
• GK nail: for femoral or tibial shaft fracture
• Gamma nail: for inter or sub-trochanteric fractures
• Harrington rod: for fixation of the spine
• Hartshill rectangle: for fixation of the spine
• Insall Burstein prosthesis: for total knee replacement
• Interlocking nail: for femoral or tibial shaft fractures
• Kirschner wire: for small bone fixation
• Kuntscher nail: for fracture shaft of the femur
• Luque rod: for fixation of the spine
• Moore’s pins: for fracture neck of the femur
• Neer’s prosthesis: for shoulder replacement
• Rush nail: for diaphyseal fractures of the long bone
• SP nail with McLaughlin’s plate: for inter-trochanteric fracture
• SP nail: for fracture neck of the femur
• Seidel nail: for fracture of the shaft of humerus
• Soutter’s prosthesis: for elbow replacement
• Steffi plate: for fixation of the spine
• Steinmann pin: for skeletal traction
• Swanson prosthesis: for finger joint replacement
• Talwalkar nails: for fracture of radius and ulna
• Thompson prosthesis: for fracture neck of the femur
• Keller’s operation: for hallux valgus correction
• Lambrinudi operation: for correcting equinus deformity of the foot
• Meyer’s operation: for fracture neck of the femur
• Putti-Plat procedure: for recurrent dislocation of the shoulder
• Soutter’s release: for flexion deformity of the hip in polio
• Steindler’s release: for cavus deformity of the foot
• Tension-band wiring: for fracture patella, olecranon
• Turco’s procedure: for CTEV
• Wilson’s release: for flexion deformity of the knee
• Yount’s release: for flexion deformity of the knee in polio

ANATOMICAL POSITIONS AND DIRECTIONS

PLANES
• Coronal: side-to-side, dividing into anterior and posterior portions
• Horizontal: transverse, dividing into superior and inferior portions
• Sagittal: antero-posterior, dividing into left and right portions

JOINT MOTION
• Abduction: movement of a part away from the body
• Adduction: movement of a part towards the body
• Apposition: being in close contact
• Eversion: turning the foot outward
• Extension: straightening a joint
• External rotation: outward rotation e.g., patella facing outward
• Flexion: bending a joint
• Internal rotation: inward rotation e.g., patella facing inward
• Inversion: turning the foot inward
• Pronation: twisting inward e.g., palm facing down
• Supination: twisting outward e.g., palm facing up

RADIOLOGICAL SIGNS

SPECIAL VIEWS
• Judet views: for acetabular fracture
• Mortice view: for ankle injuries
• Oblique view of the wrist: for fracture scaphoid
• Shenton’s line: hip X-ray in CDH
• Sunset view: for patello femoral dysplasia
• Von Rosen view: for CDH

ANGLES
• Bohler’s angle: fracture of the calcaneum
• Carrying angle: elbow
• Kite’s angle: Talo-navicular angle in CTEV
• Neck-shaft angle: of the femoral neck
• Pauwel’s angle: fracture neck of the femur

CLASSIC FEATURES
• Aneurysmal sign: TB spine (anterior type)
• Febella: sesamoid bone in the lateral head of gastronemius
• Onion-peel appearance: Ewing’s Sarcoma
• Patchy calcification: Chondrosarcoma
• Risser’s sign: Epiphysis of iliac bone
• **Sagging rope sign**: Perthes’ disease
• **Shepherd Crook deformity**: Fibrous dysplasia
• **Soap-bubble appearance**: Osteoclastoma
• **Spondylolisthesis**: slip of one vertebra over other
• **Spondylolysis**: break in posterior elements (at pars interarticularis)
• **Spondylosis**: degenerative spine disease
• **Sun-ray appearance**: Osteosarcoma
• **Tonguing of vertebra**: Morquio-Brails disease
• **Trethowan’s sign**: Slipped capital femoral epiphysis
• **Wormian bones**: Osteogenesis imperfecta

### GAITS

• **Antalgic gait**: occurs in painful condition of lower limb
• **Charlie Chaplin gait**: occurs in tibial torsion
• **Circumduction gait**: occurs in hemiplegia
• **Duck waddling gait**: occurs in bilateral CDH
• **High stepping gait**: occurs in foot drop
• **Sailor’s gait**: occurs in bilateral CDH
• **Scissoring gait**: occurs in CP
• **Stiff hip gait**: occurs in ankylosis of the hip
• **Trendelenburg gait**: occurs in an unstable hip due to CDH, gluteus medius weakness etc.

### CLASSIFICATIONS

• **Garden’s**: for fracture neck of the femur
• **Gustilo’s**: for open fractures
• **Lauge-Hansen**: for ankle injuries

• **Neer’s**: for upper end of humerus fractures
• **Pauwel’s**: for fracture neck of the femur
• **Salter and Harris**: for epiphyseal injuries

### MISCELLANEOUS

• **Bone grafting**: A technique where ‘spare’ bone is taken from some part and put where required.
• **Delayed union**: A fracture not uniting in expected time
• **Image intensifier**: A modified portable X-ray machine, where a much clearer X-ray image of a part can be seen on a TV screen. Radiation exposure is much less than a conventional X-ray exposure.
• **Malunion**: A fracture united in unacceptable alignment.
• **Nail**: A rod made of steel, usually hollow, used for internal fixation of fractures
• **Non-union**: Failure of a fracture to unite
• **Osteoarthritis**: Wear and tear arthritis
• **Osteophyte**: A bony spur at the margin of an osteoarthritic joint
• **Plate**: A thick strip of a metal (usually steel) with holes, used for internal fixation of fractures
• **Pseudarthrosis**: Painless, mobility at a fracture due to non-union (as if a ‘false’ joint has formed)
• **Spica**: is a plaster cast in which a limb and a part of the trunk are included (e.g., shoulder spica)
• **Valgus deformity**: The distal part goes outwards (e.g. knock knee – Genu valgus).
• **Varus deformity**: The distal part goes inwards (e.g. bow legs – Genu varum)
Orthopaedic Instruments and Implants

INSTRUMENTS

PERIOSTEUM ELEVATOR
The periosteum elevator is used to elevate the periosteum. Elevation of the periosteum is necessary in all operations on the bone because all the important structures such as vessels, nerves, tendons, etc. are outside the periosteum, and therefore, once the periosteum is elevated, the surgeon is in a safe plane. All the muscles of the extremity are attached to the periosteum, and are lifted off the bone with periosteum.

![Fig-1 Periosteum elevator (Farabeauf)](image)

The periosteum is not elevated in some operations such as excision of osteochondroma, where the periosteum is excised with the osteochondroma to avoid recurrence. Periosteum elevators are of different shapes and sizes depending upon their uses (Fig-1).

BONE LEVER
It is used to lever out a bone from the depth of a wound after the periosteum has been elevated (Fig-2). It is placed between the bone and the periosteum, and thus retracts the soft tissues.

![Fig-2 Different types of bone levers](image)

BONE NIBBLER
It is used for nibbling the bone (Fig-3a). It is available in various sizes and with different angle of the nose. Some of the common bone nibblers are: (i) straight nibbler – for general use; (ii) curved nibbler – for spinal surgery; and (iii) double action nibbler – straight or curved. The double-action nibblers are mechanically superior.

BONE CUTTER
It is used for cutting a bone into small pieces e.g., for cutting bone grafts (Fig-3b). It is also available with straight or curved ends, and with double-action type.

OSTEOTOME
It is used for osteotomy – cutting a bone. Its both edges are bevelled (Fig-4a). It is available in different widths of the blade. Some of the osteotomies commonly performed are: (i) McMurray’s osteotomy for fracture of the neck of the femur; (ii) corrective osteotomy for deformities such as genu varum (bow legs), genu valgum (knock knees), etc.

![Fig-4 (a) Osteotome (b) Bone chisel (note the cutting edges)](image)

BONE CHISEL
It is like an osteotome except that only one of its surfaces is bevelled (Fig-4b). It is used for removing a protruding bone or levelling a bone surface e.g., for levelling excessive callus, removing an osteochondroma, etc.

MALLET
It is used for hammering osteotome, chisel etc. (Fig-5).

BONE CURETTE
This is used for curetting a cavity in the bone or for removing fibrous tissue from fracture ends of an old fracture (Fig-6).
Curettage is performed for: (i) benign tumours such as enchondroma, giant cell tumour; and (ii) infections such as tubercular cavity of the bone, osteomyelitis, etc.

**Bone Gouge**
This is a concave bladed chisel used for cutting on round bone surfaces (Fig-7), or sometimes for making a round hole in the bone.

**Bone Awl**
This is a pointed thin instrument for making a hole in the bone (Fig-8). There is an eye at its tip to thread a wire through the bone e.g., for tendon attachment.

**Bone Holding Forceps**
There are different types of forceps for holding a bone (Fig-9). These are: (i) Lane’s forceps – for holding the femur, tibia, etc.; (ii) lion-toothed forceps; and (iii) self-retaining – AO type forceps.

**Plate-holding forceps:** Once the reduction is achieved, a plate of suitable size is placed over the fracture and held with the help of the following plate holding forceps: (i) Lowman’s clamp; and (ii) AO type self-retaining forceps (Fig-10).

**Traction Instruments**

**Kirschner wire:** This is thin, straight steel wire, of diameter ranging from 1 to 3 mm (Fig-11a). It is used (i) for internal fixation of small bones; (ii) for giving traction e.g., for applying traction through the olecranon; (iii) for fixing fractures in children; and (iv) for Ilizarov’s fixation system.

**Steinmann pin:** This is a stout, straight steel rod, of diameter ranging from 3 to 6 mm (Fig-11b). It is used for skeletal traction—common sites being upper end of tibia, supracondylar region of the femur and calcaneum.

**Bohler’s stirrup:** This is a device used for holding a Steinmann pin and applying traction (Fig-11c). The screws on the sides of the stirrup are used to hold the pin. It is possible to change the direction of traction without moving the pin inside the bone, thus avoiding loosening of the pin.

**K-wire stirrup with tensioner:** When skeletal traction is to be applied with the help of K-wire, the strength of the wire is increased by subjecting it to an axial tension by a tensioner (Fig-11d).

**Skull traction tongs:** These are tongs to apply skull traction in cases of cervical spine injury or disease (Fig-11e). Examples are Crutchfield tongs, Blackburn tongs, etc.

**Implants**

**Nails**
Nails are devices used for the intra-medullary fixation of fractures of long bones. Some of the nails used commonly are as follows:

- **Kuntscher’s nail:** This is used for internal fixation of fracture of the femoral shaft.
- **Smith Petersen nail (SP nail):** This is used for internal fixation of fracture of the femoral neck.
- **V Nail:** This is used for internal fixation of fracture of the tibial shaft.
• **Talwalkar nail**: This is used for fractures of forearm bones.
• **Rush nail**: This is used for some special situations in long bone fractures.
• **Ender’s nail**: This is used for internal fixation of intertrochanteric fractures of the femur.

**Kuntscher’s cloverleaf intra-medullary nail (K-nail)**: Kuntscher, a German surgeon devised the intramedullary nail for internal fixation of femoral fractures. The nail is a hollow tube with a slot on one side (Fig-12). It is cloverleaf shape in cross section. The fixation by K-nailing is based on the concept of three point fixation i.e., when a straight rod passes through the curved medullary cavity of the femur, it fixes the bone at three points — at either ends and at the isthmus (Fig-12a). The cloverleaf shape is designed to give good rotational stability to the fracture (Fig-12b). The nail has an ‘eye’ at its either end; in which the hook of the extractor is introduced while removing the nail (Fig-12c).

The size of a K-nail required for a particular case is found by determining the length and diameter of the nail required. The length is measured from the tip of the greater trochanter to the lateral joint line of the knee, and subtracting 2 cm from it. The diameter is determined on an X-ray, from the width of the medullary cavity at the isthmus.

The nail can be inserted by two techniques. In the first technique, the nail is inserted from the fracture site, and is hammered proximally till it comes out of the trochanter. The fracture is reduced and the nail driven back into the distal fragment. This is called retrograde nailing. In the second technique, the nail is introduced from the greater trochanter over a guide-wire passed from the fracture site. Once, the nail comes up to the fracture site, the guide-wire is removed, the fracture reduced under vision, and the nail driven home. About 2 cm nail is left protruding at the trochanter to facilitate removal usually a minimum two years after operation. For this, the hook of an extractor is engaged into the nail at the ‘eye’ and the nail pulled out by outward stroking of the extractor.

Some common complications of K-nailing are: (i) nail getting stuck; (ii) splintering of the cortex while hammering the nail; (iii) proximal migration of the nail, leading to bursitis over its protruding end; (iv) distal migration of the nail leading to stiffness of the knee; and (v) infection.

**Smith-Peterson nail (SP nail)**: Smith Peterson (Fig-12) cannulated triflanged nail is an implant used for internal fixation of a fracture of the neck of the femur. The advantages of its triflanged shape are that: (a) it prevents axial rotation...
of the fragments; and (b) it cuts only a little bone to provide good stability. The nail is cannulated because it is threaded over a guide-wire introduced at the correct site under X-ray control. It can be used along with a McLaughlin’s plate for the fixation of inter-trochanteric fractures (Fig-12).

**Dynamic Hip Screw (DHS):** This is a device used for the internal fixation of trochanteric fractures (Fig-12). It has two components – the lag screw and the barrel. The lag screw slides freely inside the barrel, so that if there is collapse at the fracture site, the screw does not cut out of the cortex; it telescopes into the barrel.

**PLATES AND SCREWS**

These are used for fixing two bony fragments. Different types of plates are available; these may be heavy duty broad and narrow plates or semi-tubular plate (Fig-13).

Screws may be used alone or in combination with a plate. Different types of screws used in orthopaedic practice are as shown in Fig-13. In the past, machine screws (self-tapping screws) were used, but now AO screws (non-tapping screws) are used. A non-tapping screw is better than a self tapping screw because in the latter, while tightening, heat is produced at the bone–screw interface causing necrosis of the bone, and thus loosening of the
screws. For a non-tapping screw, threads are cut in the bone with a special instrument, called a bone tap (Fig-14).

PROSTHESSES

Austin-Moore prosthesis: This is used for replacement of femoral head in a case of fracture of the neck of the femur in elderly persons. The prosthesis has a head with a small neck and a stem (Fig-15a). It is available in head sizes ranging from 35 to 59 mm (odd numbers). There is a small hole at the top of the stem for the hook of the extractor, used while removing the prosthesis. The stem has two fenestrations in its middle, through which the bone supposedly grows and helps in fixation of the prosthesis. This prosthesis can thus be used only without cement because the use of cement would make its removal, if required, difficult.

Thompson prosthesis: This is a prosthesis for the head of the femur, similar to AM prosthesis (Fig-15b). It is especially indicated in cases where the neck of the femur is absorbed e.g. in old fractures of the femoral neck. It can be used with or without cement.

Charnley’s total hip prosthesis: This is a prosthesis for the replacement of both, the acetabulum and the head of the femur. The acetabulum is replaced by a plastic (polyethylene) acetabulum cup, and the head by a steel component. The diameter of the head of the prosthesis is 22 mm. Both the components are fixed to respective bones by bone cement (Polymethylmethacrylate).

Muller’s total hip prosthesis: It is essentially similar to Charnley’s prosthesis except that the size of the head of this prosthesis is 32 mm, and the stem is available in different thicknesses.

Total knee prosthesis: There are several designs available (Fig-16). Total condylar designs are most popular. In this type, the articular surfaces of femur, tibia and patella are replaced by metallic (for femur) and polyethylene (for tibia and patella) prosthesis. Common prosthesis used are Insall-Burstin knee, Freeman-Samuelson knee, etc.
A number of implants are used in orthopaedics. These may be used as a temporary device, e.g., a steel rod used for fixation of a fracture; or as a permanent device, e.g., a total hip prosthesis used for replacing a damaged hip joint. The material used for these implants is foreign to the body, and is subjected to harsh chemical environment of the body. A usual foreign body, subjected to this environment shall evoke a reaction from the body which may range from a benign to a chronic inflammatory response. To avoid this, the implant materials used in our body are so designed that they have suitable mechanical strength and are biocompatible. Implant materials can be divided in the following categories:

**Metals:** These have been used for fixation of fractures, for a long time. The most common one is stainless steel. The surgical grade stainless is SS 316L. Other metal used for fracture fixation is titanium based alloy. Titanium is stronger and lighter than steel. For manufacturing components for joint replacement, cobalt based alloys are preferred as these have high resistance to corrosion.

**Non-metals:** The most common non metal material is some form of plastic. Following non-metals are commonly used:

- **Ultra high density polyethylene (UHDPE):** This is used for making acetabular cup for total hip, and the plastic insert for knee replacement.
- **Bone cement:** This is used as an anchoring agent to fix metallic components to the bone. Chemically it is polymethylmethacrylate. On mixing the monomer, powder form of polymethacrylate with liquid methylmethacrylate, a dough like material is formed which sets in 5 to 7 minutes into a hard material. It is something like an ordinary cement which sets on adding water. This process is exothermic and irreversible.
- **Ceramics:** Ceramics have been used to design articulating surfaces of artificial joints. These are more resistant to wear, but disadvantage is that these are brittle.
- **Silicon:** Silicon implants are used for artificial interphalangeal joints in the form of silicon elastomer (silastic).
- **Polyester fiber:** This is used for manufacturing artificial ligaments.
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