Skeletal Dysplasia (Non - Sclerosing dysplasias – Part II)

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Introduction

In continuation with Sclerosing Dysplasias (Part I), non sclerosing dysplasias constitute a major group of skeletal lesions including dwarfism syndromes. The following list includes most of the common dysplasias either involving epiphysis, metaphysis, diaphysis or epimetaphysis. These include multiple epiphyseal dysplasia, spondyloepiphyseal dysplasia, metaphyseal dysplasia, spondylometaphyseal dysplasia and epimetaphyseal dysplasia (Hindigodu Syndrome). These are included in dwarfism syndromes.

Dwarfism indicates a short person in stature due to genetic or acquired causes. It is defined as an adult height of less than 147 cm (4 feet 10 inches).

The average height of Indians is men- 5ft 3 ½ inches and women–5 ft 0 inches much less than average American or Chinese people.

Dwarfism syndromes consist of 200 distinct medical entities. In Proportionate Dwarfism, the body appears normally proportioned, but is unusually small. In disproportionate dwarfism, one or more body parts are relatively large or small in comparison to those of the body parts of an average-sized adult. The growth variations in specific areas apparent. This may be divided into

Rhizomelic = root, e.g., bones of the upper arm or thigh, Mesomelic = middle, e.g., bones of the forearm or lower leg, Acromelic = end, e.g., bones of hands and feet and Micromelic = entire limbs are shortened.

Out of 70 dwarfism syndromes, the most common form of dwarfism is achondroplasia. It is a proportionate dwarfism and is rhizomelic in 70% of the patients. It is autosomal dominant. There is rhizomelic type of short limbs with increased spinal curvature. Skull abnormalities are also noted. The radiographic features are mentioned in Table I.

Table I: Achondroplasia- Radiographic features
- Large skull with prominent frontal bones and a narrow base.
- The interpedicular distance decreases caudally in lumbar region but with normal vertebral height.
- Posterior scalloping
- The pelvis is square with small sciatic notches and the inlet configuration with classical champagne glass appearance
- Trident hands
- Delayed appearance of carpal bones
- Dumb bell shaped limb bones

Fig. 1a: Achondroplasia - large skull and narrow base
Fig. 1bc: Achondroplasia – b. Body gram of an infant with short limbs, c. Typical dumbbell shaped femur.

Fig. 1de: Achondroplasia – Spine- Interpedicular spaces. Shorter from lumbar to lumbosacral junction.

Fig. 1fg: Achondroplasia - Adult spine. Note the posterior scalping of vertebral bodies with a narrow canal.

Fig. 1hi: Pelvic inlet shows champagne Glass appearance

Fig. 1jk: Achondroplasia – Spinal canal stenosis better demonstrated on CT

Fig. 1lm: infant - Short and stubby metacarpals and phalanges, m- child with Trident hand
PSEUDO ACHONDROPLASIA

This type of dwarfism is rare but simulates achondroplasia. The children are of normal intelligence with short limbs. The children are invariably normal at birth, and they usually present either around 2 years of age with a delay in walking or a little later with an abnormal waddling gait. The radiographic features are listed in Table II.

Table II: Pseudo Achondroplasia

<table>
<thead>
<tr>
<th>Radiographic Features</th>
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<tbody>
<tr>
<td>Shortened bones, proximal more than distal, suggesting rhizomelic type of dwarfism, with flared and irregular metaphysis.</td>
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<tr>
<td>Epiphyses are small, irregular, and often fragmented with delayed appearance (the femoral capital and humeral epiphysis are most affected).</td>
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<td>Medial beaking of the femoral neck is one of the characteristic features.</td>
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<tr>
<td>The hand and foot bones (metacarpals, metatarsals and phalanges) are broad and shortened with small and rudimentary epiphysis.</td>
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<tr>
<td>Pelvis appears squared with broad iliac wings and narrow sacrosciatic notches. The acetabulum is poorly formed with horizontal roofs.</td>
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<tr>
<td>Skull and facial bones are normal</td>
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<tr>
<td>Odontoid dysplasia</td>
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<tr>
<td>Interpedicular distance is characteristically normal</td>
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</tbody>
</table>

Fig. 2ab: Pseudo Achondroplasia – a. skull & facial bones are normal, b. Spine Interpedicular distance is characteristically normal.

Fig. 2cd: Pseudo Achondroplasia – c. Pelvis similar to achondroplasia, d. Hypoplastic L1.
Lethal Forms of Dwarfism –

Are diagnosable in utero by ultrasonography
- Homozygous achondroplasia
- Achondrogenesis
- Thanatophoric dwarfism
- Metatrophic dysplasia

Achondrogenesis, it is autosomal recessive, almostly with short trunk and short limbs. Radiographic features (fig. 3) include defective /absent ossification of the vertebrae, pubis and Ischium, grossly shortened tubular bones with metaphyseal irregularities, thin ribs which may show fractures and crescent shaped iliac wings.

Thanatophoric Dysplasia

It is almost always lethal. Radiological features (fig. 4ab) include flattening of the vertebral bodies and widened disc spaces against a back drop of well formed neural arches-‘H’ or inverted U appearance on AP view.

Metatrophic dysplasia – Can live for a short time

Characterized by short extremities ,normal trunk at birth,& short trunk with kyphoscoliosis later in life. Radiographic features (fig. 5ab) include platyspondyly, progressive kyphoscoliosis with hypoplastic odontoid process.

Other Syndromes are mentioned in table III

Table III
- Mucopolysaccharodosis – MPS 1 to 7
- Spondyloepiphyseal dysplasia
- Spondylometaphyseal dysplasia
- Osteogenesis imperfecta
- Pycnodysostosis
- Chondrodysplasia calcificans congenita
- Chondroectodermal dysplasia
- Hypothyroidism
- Hypochondroplasia
- Diastrophic dysplasia
- Noonan / Turner syndromes
- Primordial dwarfism
- Other storage disorders
Mucopolysaccharidosis (MPS) (Several Types)

The mucopolysaccharidoses (MPS) are lysosomal storage disorders caused by the deficiency of enzymes required for the stepwise breakdown of glycosaminoglycans (GAGs), previously known as mucopolysaccharides. An estimated total incidence of all types of MPS of approximately 1 in 20,000 live births. Morquio’s syndrome is estimated to occur in 1 of every 200,000 births.

Morquio Syndrome (MPS type IV)

It is most common MPS and is autosomal recessive. Deposition of excess keratosulfate due to deficiency of N-acetyl galactosamine-6-sulfatase. There is excess keratan-sulfate in urine which differentiates from Hurler’s. Normal intelligence which also differentiate from Hurler’s. It is short trunk dwarfism with universal platyspondyly with anterior central beaking which is different from Hurler’s (fig. 6abcd). There are widened disc spaces. Frontal bossing in the skull with normal sella and depressed nasal bridge. This is also different from Hurler’s. There is delayed appearance of ossification centers.

Hurler syndrome - MPS 1 is next common disorder. The affected child is mentally retarded. Radiographic features include “J” Shaped sella with platyspondyly and inferior beaking of the vertebra (fig. 7abcde).

The other six types of Mucopolysaccharidosis show varied clinical features with different enzymes in the urine. Radiological features are not as distinctive as Morquio and Hurler.
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**Fig. 7cd**: Hurler – Platy spondyly with inferior beaking.

**Fig. 7e**: Hurler – Hand shows tilting of the ulna towards and bullet shaped proximal ends of metacarpals

Spondyloepiphyseal Dysplasia (SED) – Congenita (Figs. 8)

Children are of normal intelligence, mild lordosis with short spine. The limb bones are long.

**Fig. 8**: SED

Spondyloepiphyseal Dysplasia – Tarda

Radiologically, platy spondyly is noted with a posterosuperior hump in the vertebral bodies (Fig. 9).

**Fig. 9**: SED Tarda – The postero superior hump is characteristic.

Spondyloepiphyseal Dysplasia with Peripheral Arthropathy

In several SED patients the peripheral joints of hands and feet are also involved. Radiologically the metacarpal and metatarsal heads are larger with osteoporosis of the bones simulating rheumatoid arthritis. Hence, some authors called is Pseudo-
rheumatoid spondyloepiphyseal dysplasia (fig. 10).

![Fig. 10: SED with peripheral arthropathy simulating rheumatoid arthritis.](image1)

**Spondylo Metaphyseal Dysplasia (SMD):**

The children are of short stature with normal intelligence. Radiologically, platyspondyly is noted with irregularity of the metaphysis and anterior beaking. The articular margins are irregular (fig. 11). In the wrist the findings may simulate healing rickets (fig. 12ab).

![Fig. 11: SMD – Platspondyl with anterior beaking and irregular articular margins.](image2)

**Table IV**

1. **Infantile**
   - Thantophoric dwarfism
   - Metaphoric dwarfism
   - Osteogenesis imperfecta type 2

2. **Childhood**
   - Morquio’s & Hurler’s
   - Spondyloepiphyseal dysplasia congenita
   - Spondyloepiphyseal dysplasia tarda

Acquired causes of platyspondyly are noted in Table V.

**Table V**

**Acquired - Malignant Disorders**

**Multiple myeloma**
- Metastatic Disease
- Round Cell Tumors
- Leukemia

**Benign Disorders**
- Histiocytosis X
- Infection (TB)
- Trauma
- Steroids
- Hemangioma
- Paget’s
- Kummel’s
Handigodu disease is a new syndrome of familial spondyloepimetaphyseal dysplasia. It is inherited at an autosomal dominant trait. The disease is prevalent in Hundigodu village in Karnataka, South India. This is one of the monogenic disorders. On the basis of detailed clinical, anthropometric and radiological investigations of 234 effected individuals, a new diagnosis is reported. Hypocalcitonemia is noted. Radiologically, a spectrum of deformities is noted in the joints, particularly in the hips and vertebral column (Fig. 13abc).

There are four types
- I - Mild, Blue sclera
- II - Perinatal, Lethal
- III - Uncommon - Severe
- IV - Uncommon - Resembles I

Radiological features include a variable spectrum. Intra uterine recognition by ultrasound is known. Radiological features include fractures, lucent bones, unossified skull bones with Wormian bones (fig. 14a). Basilar impression is often noted. Popcorn mineralization is rarely observed in old fractures (fig. 14b). Fractures of long bones with abundant callus are noted (fig. 14cd). The spine shows compressed fractures (fig. 14e).
CLEIDOCRANIAL DYSPLASIA

The children are of short stature generally with normal mentation. Radiological features include wide anterior fontanelle, hypoplastic / absent clavicles and mid line defects. These features are similar to pycnodysostosis except the bones are not dense (fig. 15ab).

Chondrodysplasia Punctata – Conradi-Hunermann Syndrome

It is of two types, one is autosomal dominant with heterogenous expression & normal life expectancy. Radiological features include Epiphyseal dysplasia.

Punctate or stippled calcification of multiple epiphyseal centers and can be diagnosed in utero during the 2nd & 3rd trimesters by ultrasonography.

The second form is autosomal recessive which is potentially lethal with symmetrical limb shortening contractures of joints, mental retardation, optic atrophy, cataracts and high arched palate. Radiological features are same as form one (fig. 16ab).

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Chondroectodermal dysplasia (Ellis–Van Creveld syndrome)

Clinically, short limbed dwarfism with ectodermal dysplasia, polydactyly and congenital heart disease. Absent hypoplastic nails, dysplastic teeth and upper lip anomalies.

Radiologically, polydactyly, syndactyly and syncarpus in the hand (fig. 17ab). Long bones are short. Heart anomalies include ASD or single atrium.

Acroperipheral Dysostosis

A rare entity with short bones of the hands and feet. No other abnormalities are noted. radiologically, short metacarpals and phalanges are noted (Fig. 18).

Fibrous dysplasia is the most common dysplasia which is a mixture of sclerosing and nonsclerosing types. This entity will be dealt in the next article.

Fig. 18: Acroperipheral Dysostosis - Note the short bones of both hands.

Summary

Non – sclerosing dysplasias constitute a spectrum of disorders including dysplasias, enzymatic disorders and dysostoses. Some are lethal and can be diagnosed by ultrasonography of the fetus. Radiological characteristics of various disorders are described and illustrated. A rare variety of spondyloepimetaphyseal dysplasia, Handigodu disease from India is also included. Genetic studies and enzymatic studies are essential for proper diagnosis and prognosis, genetic counselling is very important.

Further Readings


3. Amaka C. Offiah and Christine M. Hall, Radiological diagnosis of the constitutional disorders of bone. As


