

INITIAL EVALUATION

Publication

History and Physical Exam

Developmental motor delay, loss of function, neurological complaints Family history

Facial features; spinal deformity; limbs-joint laxity, deformity; neurological exam; proportion- short vs. average trunk length; limb shortening- rhizo, meso or acromelia

Typical Radiographs

Flexion and extension lateral C-spine, AP and lateral thoracolumbar spline, AP and lateral lower extremities hip to ankles, AP both hands and wrists





Morquio's syndrome

4 yo fraternal twins, one with achondroplasia. Note rhizometic limb shortening and normal trunk length



Spondyleopiphyseal

Dysplasia

Areas of Limb Involvement



Platyspondyly Morquio's Syndrome





Metatropic Dysplasia



ACHONDROPLASIA

Etiology

FGFR3 (fibroblast growth factor receptor 3) Autosomal Dominant

Diagnosis

Clinical diagnosis at birth can be confirmed using lab tests and radiological features

Clinical Features

Average length and weight at birth Macrocephaly, Mid-face hypoplasia Rhizomelia Genu varum

Trident hand



Clinical Problems

Foramen magnum stenosis Sleep apnea Hydrocephalus



Thoracolumbar kyphosis usually resolves Frequent otitis media Genu varum Lumbar and cervical spinal stenosis Obesity Rarely needs joint replacement





Narrowing interpediculate distance Rhizomelia, Notched Champagne glass pelvis (V shaped) physes

SPONDYLOEPIPHYSEAL DYSPLASIA

Etiology

Multiple subtypes: most common - congenital Type II collagenopathy

Diagnosis

At birth; Short trunk, short limb Radiographic features

Clinical Features

Extreme short stature Long, slim fingers Waddling gait Lumbar lordosis Genu valgum

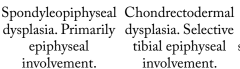




Platyspondyly, Scoliosis

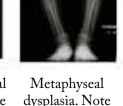


Severe epiphyseal dysplasia, coxa vara



dysplasia. Primarily dysplasia. Selective dysplasia. Note

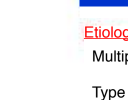
involvement.



Metaphyseal Metatropic Dysplasia. Epiphysis and tibial epiphyseal similarity to rickets. metaphysis involved.

Clinical Problems

Cervical instability **Kyphoscoliosis** Coxa vara Myopia, retinal detachment Eventual joint replacement



SKELETAL DYSPLASIA

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MULTIPLE EPIPHYSEAL DYSPLASIA

Etiology

COMP (collagen oligomeric matrix protein) abnormality Type IX collagen (less severe) Autosomal dominant

Diagnosis

Radiographic features Not usually recognized until age 5-10 yrs

Clinical Features

Mild short-stature Delayed walking Genu valgum



Appearance of bilateral Legg-Perthes is common in MED

Etiology

Sulphate transporter defect Autosomal recessive

develop early

Diagnosis

At birth; cleft palate, clubfeet, hitch hiker thumb, Symphalangism, cauliflower ears

Clinical Features

Short stature

Flexion contractures – hips knees Lateral patellar dislocation Foot deformity



Clinical Problems Painful, stiff joints Limb malalignment Eventual joint replacement



Delayed epiphyseal ossification and small epiphyses

DIASTROPHIC DYSPLASIA

Cervical kyphosis **Kyphoscoliosis** Hip dysplasia Limb Malalignment

Laryngotracheomalacia

Clinical Problems



Cervical kyphosis often resolves spontaneously



Short, thick tubular bones, delayed and fragmented epiphyses, hip dysplasia

PSEUDOACHONDROPLASIA

Etiology

COMP (collagen oligomeric matrix protein) abnormality Autosomal dominant

Diagnosis

Radiographic features Short-limbed dwarfism not usually recognized until 2-3 yrs

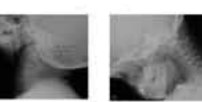
Clinical Features

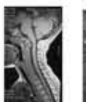
Normal face Short, stubby, hyper mobile fingers Genu varus/valus or windswept Marked ligamentous laxity



Clinical Problems

Cervical spine instability Limb malalignment Eventual joint replacement







Radiographs and MRI - C1-2 instability with cord compression





Short broad long bones with flared Platyspondyly metaphyses and small epiphyses

(odontoid hypoplasia)

MORQUIO SYNDROME

Etiology

Mucopolysaccharidosis Type IVA N-acetylgalactosamine-6-sulphatase deficiency

Autosomal recessive

Not made at birth, short stature apparent by 2 yrs Radiographic features Urine screen — keratan sulfate

Clinical Features

Awkward gait, genu valgum Barrel-shaped chest, pectus

Extreme ligamentous laxity









Diagnosis Joint replacement common

Molecular genetic testing

Normal intelligence carinatum











Platyspondyly with central beaking

