Spinal Dysraphism

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Disclosure

• Nothing to disclose
Open Neural Tube Defect
Myelomeningocele

• Myelomeningocele is an example of open neural tube defect.
• It is a defect in primary neurulation.
• It can be seen on prenatal labs with a high alpha-fetaprotein and acetylcholinesterase.
• Can also be seen on prenatal ultrasound.
  – Lemon sign
  – Banana sign
Myelomeningocele
Myelomeningcele

- The incidence is 0.7-0.8 per 1,000 live births.
- The overall lifetime mortality is 24-60%
  - Usually because of the Chiari II malformation and unrecognized shunt malfunction.
- A normal IQ is seen in 70-75% of patients.
- Roughly 60% of patients will require a VP shunt.
  - That number is significantly reduced from previous reports of close to 80%
Myelomeningocele

• There are three management schemes for myelomeningocele:
  – Initial stabilization and closure of the myelomeningocele
  – Assessment and treatment of hydrocephalus.
  – Long term management of shunt malfunction, Chiari II, tethered cord, syringomyelia
Myelomeningocele

• Initial Management:
  – Closure within 48-72 hours
  – Prophylactic antibiotic coverage (ampicillin and gentamicin)
  – Head Ultrasound
  – Renal Ultrasound
  – X-ray of spine (babygram)
  – Keep the defect covered with moist non-adhesive dressing
  – Dedicated muscle group testing to determine the affected level.
Myelomeningocele

- Chiari II malformation:
  - Caused by underdevelopment of the posterior fossa bone and crowding by the structures of the rhomboencephalon (Unified Theory)
  - Herniation of the brainstem, cerebellar vermis, 4th ventricle through the foramen magnum
    - Lower cranial neuropathies
    - Swallowing dysfunction
    - Disordered breathing
Myelomeningocele

- Prenatal closure before 26 weeks gestation compared to postnatal closure.
  - Delivered at 37 weeks
- The study showed decreased shunt rate and a better functional outcome in the prenatal surgery group.
- There was an increase in pre-term labor in the prenatal group though.
During prenatal counseling, care should be exercised in recommending prenatal surgery when the ventricles are 15 mm or larger because prenatal surgery does not appear to improve outcome in this group.
Myelomeningocele
Closed Neural Tube Defects
Cutaneous Markers

• Cutaneous markers are seen in the general population 3% of the time; however, 80% of patients with spinal dysraphism have cutaneous markers.
  – It is very common for two or more of these markers to coexist.
• Cutaneous markers occur:
  – In the midline of the back
  – Above the level of the coccyx
  – Commonly overlie the spinal lesion
• Neurologic
  Presentation:
  – Lower extremity weakness
  – Muscle Atrophy
  – Radicular pain
  – Abnormal gait

• Orthopaedic:
  – Leg length discrepancy
  – Foot deformities

• Urologic:
  – Recurring UTI
  – Change or regression
• Imaging:
  – MRI is the study of choice
  – Ultrasound can be useful to find the level of the conus in children less than 6 months
  – CT can be an adjunct in those diseases with bony abnormalities
Meningocele

• Congenital herniation of the dura and arachnoid through a bony defect in the spine.
  – Nerve roots can be involved, but typically not the spinal cord.

• Incidence is 0.3-0.4 per 1,000 births.

• Can be anterior or posterior
  – Posterior meningoceles- secondary neurulation
  – Anterior meningoceles- mesenchymal abnormalities.
Spinal Lipoma

Premature Disjunction

- CE
- NC
- Mesenchyme invading into the neural groove

Formation of “Fusion Line”

- Healed cutaneous ectoderm
- Fibrofatty stalk
- “Fusion line” of fat, spinal cord, and pia-arachnoid

Distorted spinal cord

Image: MRI scan showing a measurement of 13.63 mm.
Spinal lipoma

• Classification:
  – Dorsal:
    • Lipoma–cord interface is entirely on the dorsal surface of the lumbar spinal cord, always sparing the distal conus
  – Terminal:
    • Unlike the dorsal and transitional types, terminal lipomas insert into the caudal extremity of the conus without blending with the spinal cord or its root entry zones
Spinal Lipoma

• Classification:
  – Transitional:
    • The lipoma–cord interface thus created may be undulating and tilted so that the neural placode is rotated to one side or even spun into a parasagittal edge-on orientation, but the neural tissue is always ventral to this interface.
  – Chaotic:
    • Its caudal portion is *ventral* to the neural placode and does engulf neural tissue and nerve roots.
Spinal Lipoma

• Treatment:
  – Two schools of thought:
    • Operate prophylactically to prevent symptoms.
      – Usually around 3 months of age
      – Preferred method in America
    • Wait until the patient is symptomatic.
      – Preferred method in Europe
Dermal Sinus Tract

• Usually seen in one in every 2500 births.
• Children are usually neurologic normal at birth but there is increased risk of deficit with age if untreated
Although there is no consensus about the timing of surgery in uncomplicated, asymptomatic patients with dermal sinus tracts, it seems logical to proceed with surgery whenever the infant is physiologically ready for the procedure.
Split Cord Malformation

- SCMs can be divided into two main types, classically termed *diastematomyelia*.
- Usually present with a Faun’s tail.
Split Cord Malformation

• Type 1 SCM consists of two hemicords separated by an osteocartilaginous median septum, each housed in a separate dural sheath.

• Type 2 SCM consists of two hemicords contained within the same dural envelope and separated by a fibrous septum.
Thank you