DSS Case #6

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Disclosures

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Clinical History

• CC: 20 y/o male presented for EGD due to dysphagia and tooth pain
• PMH: Hypertension, obstructive sleep apnea, and a congenital musculoskeletal disorder
• Airway instability with subsequent desaturation during EGD
• Resuscitative efforts were unsuccessful
Autopsy Findings:

• Thin male, short stature, 75 lbs weight
• Scoliosis, pes planus, and asymmetrical muscular atrophy

Gross Brain Findings:

• Brain: 1,560 grams
• Enlarged brainstem with markedly stenotic aqueduct
• Ovoid medulla
• Internal architecture of the brainstem appeared distorted
• The dentate nuclei of the cerebellum were difficult to delineate grossly
• Obliteration of the fourth ventricle
Cross section of medulla
Discussion
Scaled cross sections of medulla
Scaled cross sections of pons
Clarification of Past Medical History......

• Fibrodysplasia Ossificans Progressiva (FOP)
  • Progressive ossification of soft tissues
  • Leads to significant disabilities and wheelchair bound by young adulthood
    • Death often due to restrictive lung disease
  • Pts with FOP have been noted to have varied neurologic symptoms such as headaches, sensory abnormalities, and movement disorders

• ACVR-1 mutation (activin receptor type 1), also known as ALK-2
  • 95% of FOP patients have a R206H mutation
ACVR1 mutations and the genomic landscape of pediatric diffuse glioma

Gelareh Zadeh & Kenneth Aldape

- ACVR-1 mutation also found in a subset of diffuse intrinsic pontine gliomas
  - Up to 27%
  - Often different point mutations than FOP

- DIPG are a member of diffuse midline glioma, harboring H3 K27M mutations
  - ACVR1 and K27M appear to be mutual
Neoplastic or Hamartomatous?

• Given the common genetic mutation as DIPG, is the brainstem mass identified in FOP patients neoplastic?
Novel asymptomatic CNS findings in patients with ACVR1/ALK2 mutations causing fibrodysplasia ossificans progressiva

Maria Savina Severino¹, Marta Bertamino², Domenico Tortora¹, Giovanni Morana¹, Sara Uccella³, Renata Bocciardi⁴,⁵, Roberto Ravazzolo⁴,⁵, Andrea Rossi¹, Maja Di Rocco²

Clinical-pathological correlations in three patients with fibrodysplasia ossificans progressiva

Kelly L. Wentworth a,⁎, Katherine Bigay a,⁎, Tea V. Chan a,⁎, Jennifer P. Ho a,⁎, Blanca M. Morales a, Joseph Connor c, Erin Brooks c, M. Shahriar Salamat c, Henry Charles Sanchez b, Geoffrey Wool f, Robert J. Pignolo d, Frederick S. Kaplan e, Edward C. Hsiao a,⁎
Imaging Insights

• High T2 & ADC values indicate low cellularity, absence of contrast, & longterm stability are consistent with hamartomas

• CNS involvement in asymptomatic children included T2 dentate nuclei abnormalities, & signal abnormalities of the dorsal pons

Summary

• Diagnosis: Glioneuronal hamartomatous proliferation of the brainstem in a patient with fibrodysplasia ossificans progressiva

• Our pathology supports radiographic interpretation of hamartoma

• ACVR1 mutations found in DIPG and FOP
  • Differing point mutations
  • Suggests that ACVR1 is probably not the driving mutation in DIPG

• Potential for misdiagnoses as DIPG
References


