A Case Study of Severe Manifestations of Mucopolysaccharidosis type I; Hurler Syndrome

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Abstract:
Hurler syndrome (mucopolysaccharidosis type I) is a rare autosomal recessive disorder of inborn error of metabolism. It is a lysosomal storage disease in which deficiency of enzyme α-L-iduronidase leads to the accumulation of glycosaminoglycans GAGs (heparin sulfate, dermatan sulfate) in various organs of the body that leads to impairment of different organ systems. Such children appear normal at birth but later on suffer from mental and physical deterioration leading to death from cardiorespiratory failure before the second decade of life. We report a seven year old girl with severe skeletal, neurological, ophthalmologic, oro-dental and radiological findings of this disease.

Keywords: Hurler syndrome, mucopolysaccharidoses, dysostosis multiplex, endocrinology, lysosomes, GAGs.

1. INTRODUCTION
Mucopolysaccharidosis are an inborn heterogeneous group of rare metabolic disorders inherited as autosomal recessive traits, due to deficiency or absence of lysosomal hydrolase iduronidase activity. The defect has been mapped to the chromosome band 4p16.3. The most common and severe form of mucopolysaccharidosis is Hurler syndrome (MPS I - H). A wide range of phenotypes result from deficiency of this enzyme including Hurler's (severe), Scheie's (mild) and Hurler-Scheie (intermediate) syndromes. Hurler Syndrome incidence has been reported to be 1:100,000 per child birth.

2. CASE REPORT
A seven year old child presented to our healthcare department with abdominal distention and fever. Examination of the child was consistent with coarse facial features like depressed and broad nasal bridge, flaring of both nostrils, prominent supra orbital rim bilaterally, ptosis of eye balls with ocular hypertelorism, thick eye lids and full and thick lips. Other findings included developmental delay, corneal clouding, protruding tongue, abdominal distention consistent with hepatosplenomegaly and umbilical hernia.

Figure 1(a). Showing broad nasal bridge, prominent orbital rims, nostril flaring and hypertelorism
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Base line investigation showed normal TLC, platelet and retic count with low hemoglobin of 8.7mg/dl. Serum electrolytes, blood glucose levels, renal function tests and urinalysis were in normal range. Liver function tests were unremarkable except for ALP of 365 IU/L. Thyroid function tests showed low total T3 with normal TSH and free T4 levels.

Skeletal survey was consistent with dysostosis multiplex. Intraoral examination revealed a large tongue, interdental spacing with thick gingivae. Skeletal survey showed variable degree of osteopenia with macrocephaly, thick vault with ground glass capacity and J shaped sella turcica (Figure:2a). Anterior beaking aspect, abnormal shaped of vertebral bodies due to hypoplasia and kyphosis were observed in the radiographs (Figure:2b). Other radiographic findings were hip dysplasia, underdeveloped acetabula, rounded iliac wings, dysplastic femoral epiphysis and coxa valga (figure2c).

Hand wrist radiograph showed wide metacarpals with proximal pointing, irregular carpal bones, ulnar hypoplasia and increased joint space at radioulnar joints (figure:3).

Chest radiograph (PA view) showed paddle shaped ribs, shortened and thickened clavicles (figure:4). Echocardiography showed Mitral leaflet prolapse with mitral regurgitation grade II.

Ultrasound of abdomen was consistent with hepatosplenomegaly and umbilical hernia. Blood was investigated for alpha-L iduronidase enzyme activity to confirm the diagnosis of MPS I H. Fluorometric test confirmed the diagnosis of MPS I H (Hurler Syndrome).

3. DISCUSSION

Glycosaminoglycans degradation is carried out by enzyme α-L-iduronidase and its absence results in the accumulation of substrates like heparan sulphate and dermatan sulfate in lysosomes of various tissues of the body resulting in developmental delay, skeletal
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malformations, corneal clouding, organomegaly, oro-facial anomalies and problems in the cardiorespiratory system. Mucopolysaccharidosis is inherited in an autosomal recessive fashion except hunter syndrome which is X linked. The type of GAGs stored and the classification of disease depend on the specific enzyme deficiency. Table 1 shows the different types.

Table 1. Mucopolysaccharidosis table showing its different types and respective enzyme deficiency, inheritance pattern and gene locations

<table>
<thead>
<tr>
<th>MPS</th>
<th>Name</th>
<th>Increased GAGs</th>
<th>Inheritance</th>
<th>Enzyme deficiency</th>
<th>Gene location</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Hurler, Hurler-Scheie or Scheie</td>
<td>HS + DS</td>
<td>autosomal recessive</td>
<td>α-L-iduronidase</td>
<td>4p16.3</td>
</tr>
<tr>
<td>II</td>
<td>Hunter</td>
<td>HS + DS</td>
<td>X-linked recessive</td>
<td>Iduronate 2-sulfatase</td>
<td>Xq28</td>
</tr>
<tr>
<td>III</td>
<td>Sanfilippo A</td>
<td>HS</td>
<td>autosomal recessive</td>
<td>Heparan-N-sulfatase</td>
<td>17q25.3</td>
</tr>
<tr>
<td>IV</td>
<td>Sanfilippo B</td>
<td>HS</td>
<td>autosomal recessive</td>
<td>β-N-acetylglucosaminidase</td>
<td>17q21.1</td>
</tr>
<tr>
<td>V</td>
<td>Sanfilippo C</td>
<td>HS</td>
<td>autosomal recessive</td>
<td>N-acetylglucosamine 6-sulfatase</td>
<td>17q21.1</td>
</tr>
<tr>
<td>VI</td>
<td>Sanfilippo D</td>
<td>KS</td>
<td>autosomal recessive</td>
<td>N-acetylglucosamine 4-sulfatase</td>
<td>17q21.1</td>
</tr>
<tr>
<td>VII</td>
<td>Maroteaux-Lamy</td>
<td>DS</td>
<td>autosomal recessive</td>
<td>β-glucuronidase</td>
<td>4p11-q13</td>
</tr>
<tr>
<td>VIII</td>
<td>Sly</td>
<td>HS + DS</td>
<td>autosomal recessive</td>
<td>β-glucuronidase</td>
<td>7q21.11</td>
</tr>
<tr>
<td>IX</td>
<td>Naucoro</td>
<td>HS</td>
<td>autosomal recessive</td>
<td>Hyaluronidase 1</td>
<td>3p21.3</td>
</tr>
</tbody>
</table>

(V) Scheie syndrome, initially proposed as type V, was recognized to be the attenuated end of the MPS I spectrum.

(VIII) An enzyme defect was found and proposed as MPS VIII, but shortly thereafter recognized as a laboratory pitfall; the proposal was withdrawn.

IX Hyaluronic acid deficient, type X.

In severe forms of mucopolysaccharidosis, average lifespan is one to two decades. A combination of clinical picture and analysis of urinary GAGs is usually performed to achieve the diagnosis. Definitive diagnosis is usually possible through measuring enzyme activity in cultured fibroblasts or leukocytes.

Treatment consists mainly of symptomatic and supportive care. In recent years the development of new therapies like enzyme replacement therapies, substrate inhibition therapy and hematopoietic cell transplantation have changed the treatment of these patients. This has shifted treatment from supportive to curative and has improved the duration and quality of life of these patients.

Radiography features include J shaped sella turcica which is characteristic but not diagnostic for mucopolysaccharidosis. In radiographs of thorax, the main abnormality concerns the ribs, which can be “paddle-shaped” or “oar-shaped” due to anterior arch widening and tapering of the posterior arches. Other common modifications are small scapulae, usually with flattening of the glenoid cavities, a short sternum and short and thickened aspect of the clavicles. In the spine, vertebral body deformities are common with thoracolumbar gibbus. These deformities can lead to compression of spinal cord and emerging roots. The most common radiological features in the pelvis are rounded iliac wings and inferior tapering of the ileum. The long bones are often characterized by several alterations. Diaphysis are shortened and curved in the distal part; the epiphyses are slightly hypoplastic and thinned cortically with osteoporosis. In appendicular skeleton features that can be found are the notching of the proximal part of the humerus, the long and narrow aspect of the femoral neck with underdeveloped acetabula, and the hypoplasia of the lateral tibial hemiplate, resulting in genu valgum. Almost all forms of MPS show distortion of the hand and foot structure. Carpal and tarsal bones are hypoplastic and irregularly shaped; the metacarpal bones are proximally pointed, shortened and thickened. This leads to compromised functionality of hand and claw hand deformity due to failure of complete extension of fingers. At the craniovertebral junction atlantoaxial instability can also occur.

Developmental delay is a major manifestation of the disease which can be explained due to brain atrophy and neuronal death secondary to accumulation of GAG in neuronal tissues. The neuronal death could be explained by ischemic damages due to the progressive accumulation of GAG in blood vessels. Hydrocephalus may also accompany brain atrophy.

4. CONCLUSION

Knowledge of clinical and radiological features is important for diagnosis. Definitive diagnosis is usually obtained by lab analysis. Early recognition is important for monitoring the chronic and progressive course of the disease.
is also important for medical and surgical planning and for assessing the impact of therapy.

REFERENCES


