Otolaryngological findings in mucopolysaccharidoses

Mucopolysaccharidoses (MPSs) are a group of lysosomal storage diseases, each of which is produced by an inherited deficiency of an enzyme involved in the degradation of acid mucopolysaccharides, now called glycosaminoglycans (GAGs). The mucopolysaccharidoses consist of a group of 7 metabolic disorders, known as mucopolysaccharidoses types I–VII. In all groups, there are clinical and otolaryngological manifestations. In MPS patients, upper airway obstruction, obstructive sleep apnea, restriction of mouth opening, middle ear effusion, hearing and breathing problems, etc. are reported as common otolaryngological findings. Increasing awareness of MPS’s among ENT doctors will be a life saving attempt for MPS suspected patients who admit an ENT doctor rather than a pediatrician. In MPS patients, tracheotomy may be difficult due to short neck. Due to mouth opening restriction, patients should be evaluated carefully before tonsillectomy and adenoidectomy operations. Airway problems must be evaluated before anesthesia. All ENT doctors should be noticed to be aware of these problems.

Keywords: Mucopolysaccharidoses (MPSs), lysosomal storage diseases, short neck, mouth opening restriction, airway problems.
trophy. Most of these children benefit from adenotonsillectomy, after which OSA significantly resolves. They experience high recurrence rate after adenoidectomy; though this is not clinically problematic. They also suffer from conductive type hearing loss due to OME, which has to be treated with ventilation tube insertion. Sometimes tracheotomy may be necessary. Tracheotomy is also highly complicated in MPS patients. GAG significantly accumulates in middle ear fluid and adenoid tissue; however, it appears not to accumulate in tonsillar tissue.

Typical facial features include coarse facial appearance, protruding or depressed frontal bone, wide eyebrows, depressed base of nose, wide nasal alae, thick lips, angled and hypoplastic mandible (micrognathia), macroglossy, distorted teeth, gingival hypertrophy and corneal opacities. Airway problems include obstructive sleep apnea (OSA), otitis media with effusion, sinusitis, frequent and recurrent respiratory infections, adenotonsillar hypertrophy, irregular nasal septum, turbinate hypertrophy, speech disorders, dyspnea, restricted temporomandibular joint motion, thickened pharyngeal wall, laryngeal abnormalities, tracheomalacia, tracheal stenosis and short neck. Otolaryngological involvement in MPS is due to accumulation of GAG in upper airway leading to obstruction and OSA, thus decreasing the quality of life of MPS patients. [3-6]

Mouth opening status had restricted due to temporomandibular joint involvement in MPSs. MPS type III had significantly better mouth opening than other MPS types. MPS type VI had significantly worse mouth opening than other types. [3] The middle ear fluid GAG level increased significantly in MPS patients. While 98.7% of MPS patients had hearing loss, 92.1% of the cases had conductive hearing loss. [3]

In this review paper, we reported otolaryngological problems (upper airway obstruction, obstructive sleep apnea, restriction of mouth opening, middle ear effusion, hearing and breathing problems, etc.) in patients with MPSs. The main aim of this paper is to ensure ENT doctors to aware of these types of diseases in their patients and sample for diagnosis (Fig. 1).

Mucopolysaccharidoses (MPSs)

Mucopolysaccharidoses (MPSs) are autosomal recessive, except for mucopolysaccharidosis type II, which is X-linked. The mucopolysaccharidoses comprise a group of 7 metabolic disorders, known as mucopolysaccharidoses types I–VII. Mucopolysaccharidosis V is defined as a form of type I and is known as mucopolysaccharidosis IS. [1]

- Mucopolysaccharidosis type I and type V (IS)
- Mucopolysaccharidosis type II
- Mucopolysaccharidosis type III
- Mucopolysaccharidosis type IV
- Mucopolysaccharidosis type VI
- Mucopolysaccharidosis type VII

1. Mucopolysaccharidosis Type I

Hurler syndrome, the most severe form of mucopolysaccharidosis type I (MPS I), is a rare lysosomal storage disease. The overall incidence of MPS I is 0.99–1.99/100,000 live births. Accumulation of glycosaminoglycans causes the progressive dysfunction of multiple organs. [7] Historically, the most-to-least severe forms are as follows: Hurler syndrome, Hurler-Scheie syndrome, and Scheie syndrome.

L-iduronidase deficiency results in an inability of the lysosome to break down GAG, namely dermatan sulfate (DS) and heparan sulfate (HS). This process is essential for normal growth and homeostasis of tissues. In this disease, GAG progressively accumulates in the lysosomes, ultimately causing cell, tissue, and organ dysfunction by largely unknown pathophysiological mechanisms. The enzyme alpha-L-iduronidase deficiency causes an increase in the urinary excretion of dermatan sulfate (DS) and heparan sulfate (HS) in patients with MPS I.

Hurler syndrome is caused by mutation in the gene (IDUA) that encodes alpha-L-iduronidase on chromosome 4. Many different mutations have been found at this locus, including mutations that cause MPS IH (Hurler syndrome), MPS IS (Scheie syndrome), and MPS IH/S (Hurler-Scheie syndrome), among others. [1]

Clinical manifestations

Clinical manifestations of mucopolysaccharidosis type I (MPS I) show a chronic multisystemic and progressive course. [6] The disease is highly heterogeneous, spanning a spectrum of severity. Children with Hurler syndrome appear normal at birth and develop the characteristic appearance over the first years of life. Symptoms across the types include facial dysmorphism, corneal clouding, hepatomegaly, valvular heart disease, obstructive airway disease, developmental delay, hearing loss, skeletal defor-
mities,\(^9\) and joint stiffness. The neck is short, and odontoid hypoplasia can occur. Vertebral subluxation with cord compression can occur. Body hair may be coarser than usual, and the skin may be thicker.

For patients with the more severe form of the disease, the most typical symptoms occur early in life. These patients typically have numerous progressively debilitating symptoms, including mental retardation. Their lifespan expectancy is less than 10 years.\(^1\)

### Otolaryngological manifestations in MPS I (Table 1)

- **Facial dysmorphism or coarse facial features:** Coarsening of the facial features is usually the first sign detected. These features often first become apparent at age between 3–6 months and may become progressively more evident. The head is large with bulging frontal bones. The skull is often scaphocephalic secondary to premature closure of the metopic and sagittal sutures. Eyes may be widely spaced, and eye sockets may be shallow, causing the eyes to slightly protrude.
- **The nasal bridge is depressed with broad nasal tip and anteverted nostrils.**
- **The cheeks are full.** The lips are enlarged, and the mouth is usually held open, particularly after age 3 years.
- **Chronic nasal discharge** is present.
- **The clinical presentation in less severe disease (MPS IS)** may be limited to mild coarsening of facial features and prognathism. A large mouth with thick lips may develop.\(^1\)
- **Hearing loss**\(^10\)

|**Table 1. Otolaryngological manifestations in MPS I.** |
|---|---|
|**Mucopolysaccharidosis type I** | |
|Head | • Large head  
• Bulging frontal bones |
|Nose | • Depressed nasal bridge  
• Anteverted nostrils  
• Chronic nasal discharge |
|Face and mouth | • Cheeks are full  
• Enlarged lips  
• Large mouth with thick lips  
• Usually open mouth  
• Coarsening of facial features and prognathism |
|Ear | • Hearing loss |

### 2. Mucopolysaccharidosis Type II

Hunter syndrome or mucopolysaccharidosis type II (MPS II) is a MPS that is another member of inherited disorders of glycosaminoglycan (GAG) catabolism.\(^11\) Each MPS is caused by a deficiency in the activity of one of the distinct lysosomal enzymes required for the stepwise degradation of the GAGs dermatan sulfate, heparan sulfate, keratan sulfate, and chondroitin sulfate.\(^7,13\) In affected patients, undegraded or partially degraded GAG accumulates within lysosomes and is excreted in excess in the urine.\(^7,12\) It is the accumulation, or storage, of GAG within lysosomes that contributes to the signs and symptoms of these disorders. All MPSs are inherited in an autosomal recessive fashion, except for Hunter syndrome, which is X-linked.

In the early 1900s, Gertrud Hurler and Charles Hunter first described patients with MPS, whose diseases now bear their names; subsequent MPSs have been assigned numbers and eponyms loosely associated with the chronology and origin of their report. MPS II was first described by Hunter in 1917. This X-linked disorder results from the deficiency of iduronate 2 sulfatase and subsequent accumulation of heparan and dermatan sulfate. In MPS II, due to the lack of iduronate 2 sulfatase (IDS), dermatan and heparan sulfate accumulate.\(^13\)

The Hunter syndrome is distinct from the other mucopolysaccharidoses in that it is an X-linked disorder. The genetic locus has been mapped to Xq28. The gene defective in this disorder encodes IDS.\(^14,15\)

The estimated incidence of MPS type II widely varies. The estimated incidence is one case per 34,000 in Israel, one case per 111,000 in British Columbia, and one case per 132,000 in the United Kingdom.\(^16-18\) Recent studies from Germany and the Netherlands report an incidence of one case in 140,000–330,000 live births, and 1.3 cases per 100,000 male births.\(^19,20\) The severe form of Hunter syndrome is typically diagnosed in children aged between 2–4 years. The attenuated form of Hunter syndrome may not be diagnosed until the teenage years or well into adulthood.

#### Clinical manifestations

Both types A and B MPS II have deficient IDS activity and are retained as terms useful in clinically describing the extremes of a disease spectrum. Children with classic type A MPS II have progressive coarsening of facial features, short stature, joint stiffness, hepatosplenomegaly, and hernias as common presenting signs and symptoms. Children with type A MPS II tend to have severe mental retardation and deafness. Other presentations include cerebral ven-
tricular dilation and dysostosis multiplex. Skin findings include hypertrichosis, thickened skin, and multiple Mongolian spots. Children with type A and B MPS II may have papular skin lesions that are ivory in color and are located on the upper back and on the lateral upper arms and thighs.[1]

Otolaryngological manifestations in MPS II (Table 2)

- Frequent upper respiratory infections occur in most patients with Hunter syndrome.
- The enlarged tongue, hypertrophic adenoids and tonsils, and skeletal changes in the jaw and neck that limit the opening of the mouth,[1,20,21] all contribute to respiratory problems, as noted in “Respiratory System”.
- Most patients have recurrent ear infections, and nearly all of them experience progressive hearing loss.[7,22–24] The hearing loss is caused by both conductive and sensorineural deficits.[22,24] Middle-ear effusion is recognized as an important contributor to hearing loss in patients with Hunter syndrome, and the evidence suggests that tympanomastoid abnormalities may also contribute to conductive hearing loss.[24] Sensorineural hearing deficit has been commonly reported.
- The teeth have been described as peg shaped, and are irregularly shaped, and gingival tissue is hyperplastic and hypertrophic.[7,20,21]
- Respiratory obstruction is secondary to the accumulation of glycosaminoglycans in the cells of the trachea.
- Patients frequently have macrocephaly. The facial features of Hunter syndrome are coarse, but the children still have faces that resemble other family members.
- Patients with Hunter syndrome tend to have short necks.

Table 2. Otolaryngological manifestations in MPS II.

<table>
<thead>
<tr>
<th>Mucopolysaccharidosis type II</th>
<th>Otolaryngological manifestations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respiratory</td>
<td>• Respiratory obstruction (accumulation of glycosaminoglycans in the cells of the trachea)</td>
</tr>
<tr>
<td>Head and neck</td>
<td>• Macrocephaly</td>
</tr>
<tr>
<td></td>
<td>• Short neck</td>
</tr>
<tr>
<td>Face and mouth</td>
<td>• Spaced teeth</td>
</tr>
<tr>
<td></td>
<td>• Enlarged tongue (more common in children older than 5 years)</td>
</tr>
<tr>
<td></td>
<td>• Hypertrophic adenoids and tonsils</td>
</tr>
<tr>
<td>Ear</td>
<td>• Hearing loss (often of mixed type but may be either conductive or sensorineural)</td>
</tr>
</tbody>
</table>

- These patients may exhibit some oral manifestations of the disease with widely spaced teeth and an enlarged tongue. The enlarged tongue is more common in children older than 5 years.[11]

3. Mucopolysaccharidosis Type III

Mucopolysaccharidosis type III (MPS III/Sanfilippo syndrome) is an inherited metabolic disorder characterized by an absence/defect of lysosomal enzymes needed to break down glycosaminoglycans (GAGs). Accumulation of GAGs leads to progressive dysfunction of cells, tissues and organs.[21–25] Incidence varies across countries with 1.21 per 100,000 babies affected in the UK.[25-27] MPS III has three phases,[28] with the first phase (1–4 years) characterized by developmental delay, the second phase (4–10 years) by behavioral disturbance, including sleep difficulties, aggressive or destructive behaviors, hyperactivity and attention difficulties, and the third/end phase (10+ years) by behavioral disturbance, including sleep difficulties, aggressive or destructive behaviors, hyperactivity and attention difficulties, and the third/end phase (10+ years) by behavioral disturbance, including sleep difficulties, aggressive or destructive behaviors, hyperactivity and attention difficulties, and the third/end phase (10+ years) by behavioral disturbance, including sleep difficulties, aggressive or destructive behaviors, hyperactivity and attention difficulties, and the third/end phase (10+ years) by behavioral disturbance, including sleep difficulties, aggressive or destructive behaviors, hyperactivity and attention difficulties, and the third/end phase (10+ years) by behavioral disturbance, including sleep difficulties, aggressive or destructive behaviors, hyperactivity and attention difficulties, and the third/end phase (10+ years) by behavioral disturbance, including sleep difficulties, aggressive or destructive behaviors, hyperactivity and attention difficulties, and the third/end phase (10+ years) by behavioral disturbance, including sleep difficulties, aggressive or destructive behaviors, hyperactivity and attention difficulties, and the third/end phase (10+ years) by behavioral disturbance, including sleep difficulties, aggressive or destructive behaviors, hyperactivity and attention difficulties, and the third/end phase (10+ years) by behavioral disturbance, including sleep difficulties, aggressive or destructive behaviors, hyperactivity and attention difficulties.

Sanfilippo syndrome results from the deficiency or absence of 4 different enzymes that are necessary to degrade the GAG heparan sulfate. Each enzyme deficiency defines a different subtype of Sanfilippo syndrome, as follows: type III A (Sanfilippo A), type IIIB (Sanfilippo B), type IIIC (Sanfilippo C), and type IIID (Sanfilippo D).[27,30] The clinical features of Sanfilippo syndrome, including the significant CNS component (brain and spinal cord), result from the progressive lysosomal accumulation of the GAG heparan sulfate.[31]

A prevalence of mucopolysaccharidosis IIIA was cited of one in 114,000 live births and the prevalence of mucopolysaccharidosis IIIB as one in 211,000 live births. Mucopolysaccharidosis IIIC and mucopolysaccharidosis IIID were much rarer, at one in 1,407,000 and one in 1,056,000 live births, respectively.[36]

Clinical manifestations

The striking clinical findings of coarse facial features and skeletal abnormalities seen in the mucopolysaccharidoses are not as apparent in patients with mucopolysaccharidosis type III. Mild facial coarsening may be present, and the
skeletal differences are typically subtle. Corneal clouding, as seen in the other mucopolysaccharidoses, is not usually present. However, classic clinical features are an abundance of coarse facial and body hair (hirsutism).\(^{[17]}\)

Otolaryngological manifestations in MPS III (Table 3)

- Dense calvaria
- Joint stiffness
- Mild coarse facial features
- Synophrys: presence of abundant hair between the eyebrows
- Hearing loss with speech delay
- Sleep disturbances\(^{[27]}\)

Table 3. Otolaryngological manifestations in MPS III.

<table>
<thead>
<tr>
<th>Mucopolysaccharidosis III</th>
<th>Head</th>
<th>Joint</th>
<th>Face and mouth</th>
<th>Ear</th>
<th>Sleep</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Dense calvaria</td>
<td>• Joint stiffness</td>
<td>• Mild coarse facial features</td>
<td>• Hearing loss</td>
<td>• Sleep disturbances</td>
</tr>
<tr>
<td></td>
<td>• Synophrys: presence of abundant hair between the eyebrows</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

4. Mucopolysaccharidosis Type IV

Morquio syndrome (mucopolysaccharidosis type IV) is a member of a group of inherited metabolic disorders collectively termed mucopolysaccharidoses. In 1929, Morquio, a pediatrician from Uruguay, and Brailsford, a radiologist from England, simultaneously described cases of what is now believed to be Morquio syndrome. Patients with Morquio syndrome (mucopolysaccharidosis type IV) can usually be clinically distinguished from patients with other MPSs because they do not have coarse facial features or mental retardation and they have additional skeletal manifestations derived from a unique spondyloepiphyseal dysplasia and ligamentous laxity. These skeletal manifestations include odontoid hypoplasia, a striking short trunk dwarfism, and genu valgus.

The enzyme deficiency in Morquio syndrome type IVA is galactosamine-6-sulfate deficiency [i.e., N-acetyl-galactosamine-6-sulfate sulfatase deficiency]. The enzyme deficiency in Morquio syndrome type IVB was β-galactosidase deficiency.\(^{[1]}\)

Compared with other patients who have MPS, those with Morquio syndrome (mucopolysaccharidosis type IV) tend to have greater spine involvement with scoliosis, kyphosis, and severe gibbus, as well as platyspondyly, rib flaring, pectus carinatum, and ligamentous laxity. Odontoid hypoplasia is the most critical skeletal feature to recognize in any patient with Morquio syndrome (mucopolysaccharidosis type IV).\(^{[28]}\)

Morquio syndrome (mucopolysaccharidosis type IV), the degradation of KS is defective because of the deficiency of either N-acetyl-galactosamine-6-sulfate sulfatase (GALNS gene) in Morquio syndrome type IVA or β-galactosidase (GLB1 gene) in Morquio syndrome type IVB. Defective GALNS also affects the catabolism of chondroitin 6-sulfate.\(^{[1]}\)

The estimated incidence of Morquio syndrome (mucopolysaccharidosis type IV) covers a wide range, including one case per 75,000 births in Northern Ireland, one case per 200,000 births in British Columbia, and one case per 263,157 births in Germany.\(^{[1]}\)

Clinical manifestations

The most dramatic finding in patients with Morquio syndrome is their skeletal habitus. This includes short stature with short trunk, pectus carinatum, kyphosis, gibbus, scoliosis, genu valgus, flaring of the lower ribs, and joint abnormalities (joints range from hypermobile to contracted). Patients can alternatively have genu varus.\(^{[1]}\)

Otolaryngological manifestations in MPS IV (Table 4)

- Mild coarsening of facial features
- Hearing difficulties,
- Carious teeth
- Thin tooth enamel\(^{[1]}\)

Table 4. Otolaryngological manifestations in MPS IV.

<table>
<thead>
<tr>
<th>Mucopolysaccharidosis IV</th>
<th>Face and mouth</th>
<th>Ear</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>• Mild coarsening of facial features</td>
<td>• Hearing difficulties</td>
</tr>
<tr>
<td></td>
<td>• Carious teeth</td>
<td></td>
</tr>
<tr>
<td></td>
<td>• Thin tooth enamel</td>
<td></td>
</tr>
</tbody>
</table>

The enzyme deficiency in Morquio syndrome type IVA is galactosamine-6-sulfate deficiency [i.e., N-acetyl-galactosamine-6-sulfate sulfatase deficiency]. The enzyme
5. Mucopolysaccharidosis Type VI

Mucopolysaccharidosis type VI (MPS VI), which is inherited as an autosomal recessive trait, results from the deficiency of N-acetylgalactosamine-4-sulfatase (arylsulfatase B) activity and the lysosomal accumulation of dermatan sulfate. MPS VI is characterized by somatic features but not by mental retardation. MPS VI is characterized by progressive connective-tissue organ involvement that results from continuous accumulation of dermatan sulfate in the skeleton, heart valves, spleen, liver, lung, dura, and cornea.

Patients appear totally normal at birth and have accelerated growth in the first year, followed by deceleration and short stature later in childhood. The diagnosis is usually established during early childhood when organomegaly, corneal clouding, coarse features, enlarged tongue, frequent respiratory illness or otitis media, and joint stiffness are all apparent. Other complications include hearing loss, chronic respiratory tract infections, sleep apnea, pulmonary hypertension, hydrocephalus, rapid-onset blindness, and cardiac valve insufficiency or stenosis.[5]

An Australian survey reported an incidence rate of one per 248,000 births from 1980–1996.[36] Reports from Germany, and northern Portugal noted birth incidences of one case per 432,610 births and one case per 238,095 births, respectively.[39,40]

Clinical manifestations

Progressive connective-tissue organ involvement that results from continuous storage of dermatan sulfate in the skeleton, heart valves, spleen, liver, lung, dura, and cornea are seen. Pulmonary hypertension, hydrocephalus, rapid-onset blindness, and cardiac valve insufficiency or stenosis are present. Hepatomegaly and splenomegaly are often present in patients with MPS VI. Umbilical and inguinal hernias are common. Growth may be normal for several years and may then stop, resulting in a final stature of 90–140 cm. A short trunk with lumbar lordosis is typically present. Corneal opacities can be detected with slitlamp examination. Joint stiffness, claw-hand deformities appear in the first few years of life. Examination of the skin frequently reveals hirsutism.[1]

Otolaryngological manifestations in MPS VI (Table 5)[1]

<table>
<thead>
<tr>
<th>Otolaryngological manifestations</th>
<th>Mucopolysaccharidosis type VI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Head</td>
<td>• Macrocephaly</td>
</tr>
<tr>
<td>Face and mouth</td>
<td>• Prominent forehead</td>
</tr>
<tr>
<td>Ear</td>
<td>• Possible coarse texture of hair</td>
</tr>
<tr>
<td>Respiratory</td>
<td>• Enlarged tongue</td>
</tr>
<tr>
<td></td>
<td>• Sleep apnea</td>
</tr>
<tr>
<td></td>
<td>• Otitis media</td>
</tr>
<tr>
<td></td>
<td>• Frequent respiratory infections</td>
</tr>
</tbody>
</table>

6. Mucopolysaccharidosis Type VII

Mucopolysaccharidosis type VII (MPS VII) is a very rare lysosomal storage disease. MPS VII, traditionally known as Sly syndrome, was characterized for the first time in a patient with skeletal features similar to those observed in other patients with MPS. Sly syndrome is caused by deficiency of the enzyme β-glucuronidase.[41] Historically, MPS VII is of interest because it was the first MPS (excluding the sex-linked gene for Hunter syndrome) for which the mutant gene was localized to chromosome 7. MPS VII is extremely rare, and few cases have been described.[1]

MPS VII the molecular defect on the gene that encodes β-glucuronidase protein (GUSB) leads to deficiency of the enzyme β-glucuronidase. This enzyme is required for the breakdown of several GAGs, including dermatan sulfate (DS), heparan sulfate (HS), and chondroitin sulfate (CS).[24] Accumulation of DS, HS, and CS takes place in the lysosome of many systems and tissues, including the CNS. The pattern of urinary excretion of HS, DS, and/or CS may vary based on the subtype of MPS VII involved.[1]

Clinical manifestations

Mucopolysaccharidosis type VII (MPS VII) is a progressive, debilitating, and often life-threatening disease that affects multiple organ system. In most severe cases, the condition presents as hydrops fetalis. Neonatal jaundice may be present at birth. Dysostosis multiplex is also associated with the severe form of Sly syndrome. Coarse facial features with macrocephaly, hepatomegaly, hepatosplenicogaly, inguinal and umbilical hernias, and repeated upper respiratory infections may be observed. Severe growth retardation may be prominent during the first 2 years of life in patients with severe disease.[1]
Otolaryngological manifestations in MPS VII

(Table 6)

- Coarse face, macrocephaly,
- Frontal prominence,
- Hirsutism
- Short neck
- Speech and language development delay due to moderate, nonprogressive mental retardation \[42\]
- Recurrent respiratory infections
- Hearing loss \[1\]

### Table 6. Otolaryngological manifestations in MPS VII.

<table>
<thead>
<tr>
<th>Mucopolysaccharidosis type VII</th>
<th>Head and neck</th>
<th>Face</th>
<th>Respiratory</th>
<th>Ear</th>
</tr>
</thead>
<tbody>
<tr>
<td>Otolaryngological manifestations</td>
<td>• Coarse face</td>
<td>• Frontal prominence</td>
<td>• Recurrent respiratory infections</td>
<td>• Hearing loss</td>
</tr>
<tr>
<td></td>
<td>• Macrocephaly</td>
<td>• Hirsutism</td>
<td></td>
<td>• Speech and language development delay</td>
</tr>
<tr>
<td></td>
<td>• Short neck</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Acknowledgements

“The preparation of this paper, including design and planning, was supported by the Continuous Education and Scientific Research Association.”

### Conflict of Interest: No conflicts declared.

### References


