A 16-year-old Girl with Morquio Syndrome: A Case Report

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Abstract

Morquio syndrome is a rare autosomal recessive disorder of mucopolysaccharide metabolism, also called mucopolysaccharidosis type IV. We report a case of Morquio syndrome in a 16-year-old girl of normal intelligence, who got herself admitted in Bangabandhu Sheikh Mujib Medical University, Dhaka, Bangladesh. The patient had short stature and skeletal deformity and she belonged to a non-consanguineous marriage of her parents. She was diagnosed on the basis of clinical features, typical radiological changes and positive urinary mucopolysaccharide screening test.

Key words: Morquio syndrome, Bangladesh, mucopolysaccharidosis

Introduction

Morquio syndrome is a rare inherited disorder of mucopolysaccharide metabolism. Among seven different types, Morquio syndrome is type IV mucopolysaccharidosis (Type IV-A and Type IV-B) which caused by deficiency of N-acetylgalactosamine-6-sulfatase and ã-galactosidase respectively, each resulting in a defective degradation of keratan sulfate.\textsuperscript{1} The internationally estimated incidence of Morquio syndrome covers a wide range of 1 case among 75000 to 250000 in different parts of Europe.\textsuperscript{2} In 1929, Mr. Morquio, a pediatrician in Uruguay and an English radiologist Mr. Brailsford independently and simultaneously described what is known as Morquio-Brailsford syndrome.\textsuperscript{3,4}

In children with Morquio syndrome, body cannot breakdown sugar chains called glycosaminoglycan, that helps to build bone, cartilage, eyes, cornea, skin, connective tissue (tendons and ligaments) etc. due to lack or inactivity of lysosomal enzymes. As a result, glycosaminoglycans accumulate in cells, blood, bones and connective tissue and causes damage over time.

Case Report

A 16-year-old girl got herself admitted in Internal Medicine Department of Bangabandhu Sheikh Mujib Medical University (BSMMU) with growth failure, swelling and deformity of back, low back pain and occasional shortness of breath on exertion for 2 years. She was delivered by normal vaginal delivery at full term and was healthy with average weight. Both antenatal and postnatal periods were uneventful. Her parents were of normal height and there was no history of consanguinity. Her only younger brother had normal growth and development. History did not reveal any familial incidence of similar disorder. The girl had normal growth and development till the age of 5 years after that she did not grow as much as other children of the same age.

General physical examination revealed that the girl looked disproportionately short with short neck and trunk. Limbs were normal in length though they appeared long in relation to the trunk. There were lumbar...
kyphosis and gibbus, mild genu valgus and pectus carinatum. Her pulse was 100 b/min with water hammer character, blood pressure was 90/60 mm Hg and temperature was normal. Anthropometry revealed head circumference 53 cm, height 127 cm, arm span 145 cm, weight 24 kg, body mass index (BMI) 15 kg/m². The CDC (Centers for Disease Control and Prevention) growth chart showed height for age and weight for age both were below 5th centile.

There was lumbar gibbus and mild restriction of movement of lumbar spine and hyper mobility of joints. Mild pectus carinatum with reduced chest expansion and an early diastolic murmur in the aortic area was also noted. Ophthalmological examination revealed her vision 6/9 on both eyes and mild corneal clouding was noted on slit lamp biomicroscopy. Fundoscopy showed cherry red macula. Her I.Q. (Intelligence Quotient) and neurological examination was normal.

She had bilateral sensory neural deafness as evidenced on pure tone audiometry. Oral examination showed multiple decayed teeth with over retained tooth pieces of primary teeth. OPG (orthopantomogram) showed decayed teeth.

Radiological features
Definite flattening of vertebral bodies with central beaking were seen in cervical and lumbar regions with kyphoscoliosis (Figure 1,2). There was mild genu valgum and ribs were broad and wide (Figure 3,4). Pelvic radiograph showed shallow acetabulum and deformed irregular femoral head with dislocation (Figure 5). Those findings are strongly suggestive of Morquio syndrome.

Laboratory findings
Her haemoglobin was 12.3 mg/dl, ESR 10 mm in 1st hour, WBC 7400/cumm with polymorphonuclear leukocyte and intracytoplasmic stipling (Reilly bodies), RBS 5.6 mmol/L, S. calcium 8.2 mg/dl, S. alkaline phosphatase 423 U/L, S. creatinine 4 mg/dl, S. inorganic phosphate 4.9 mg/dl, S. parathyroid hormone 26.5 pg/ml, and vitamin D (25 hydroxy) 8.5 ng/ml which was below normal range. ECG showed T inversion in V₁ to V₆. Echocardiogram showed mild cardiomegaly and aortic regurgitation, ultrasonogram of whole abdomen showed mild hepatomegaly.

Urinary keratan sulfate was 38 mg/L, that was more than normal (control 10 mg/L). Direct enzymatic assay was not done due to unavailability of the test in our country.
Finally, the patient was diagnosed as Morquio syndrome on the basis of clinical, radiological and positive urinary keratan sulfate level. We treated the patient conservatively with NSAIDS (Nonsteroidal anti-inflammatory drugs) and vitamin D supplement. Enzyme replacement was not given as we could not do enzymatic assay and this sort of treatment is not available in our country.

Discussion
Diagnosis of Morquio syndrome is usually made by characteristic clinical feature, radiological findings, positive urinary mucopolysaccharide and confirmed by definite enzyme assay. Our patient had characteristic clinical features in addition to typical radiological features of Morquio syndrome with positive urinary mucopolysaccharide level. Enzyme test was not done as this investigation is not available in Bangladesh.

We did not find any previously published case report for this condition in our country. Few cases were reported in India and the clinical features were almost identical with our case, though some of the cases from India showed sibling involvement, our patient had a brother who was completely normal.\textsuperscript{5,6}

Our patient had several skeletal changes consistent with Morquio syndrome. The syndrome is characterized by several skeletal changes, which include short stature, microcephaly, short neck, cervical scoliosis, cervical spine subluxation with a very short torso, thoracolumbar kypho-scoliosis, hypoplasia of the odontoid process, a prominent curvature of the spine (scoliosis or kyphosis), flat bridged nose, bulging forehead, barrel chest with pectus carinatum, pot belly, genu valgum and hypermobile joints.\textsuperscript{7} Extra skeletal manifestations include aortic valve disease which was evident in our case clinically and later proved by echocardiograph. Patient can also have mild hepatosplenomegaly, corneal clouding, coarse facial features, prognathism and dental abnormalities. Our patient had dental decay which is common for the syndrome.\textsuperscript{8,9} The airway issues of these patients are frequently complicated by narrow and flat trachea, chronic respiratory disease, recurrent pulmonary infection, obstructive sleep apnoea leading to restrictive lung disease. Ophthalmologically such patients manifest slight cloudiness of cornea. The corneal involvement has little effect on vision. Other features are inguinal hernia and loss of nerve function below neck due to cervical myelopathy.\textsuperscript{10,11}
This syndrome may become complicated by atlantoaxial instability, compression of cervical spinal cord, complications during endotracheal intubation and restrictive lung disease. Characteristic radiological findings include odontoid hypoplasia, hypoplastic lumbar vertebra which may give rise to lumbar kyphosis, platyspondyly of vertebra with central beaking, flaring of ileum, shallow acetabulum, flattening and irregular femoral head with subluxation, coxa and genu valga deformity. Historically type IV-A was considered to have more severe manifestations than type IV-B. Life expectancy for patients with the type IV-A is normally less than 30 years, but isolated cases of long survival has been documented. These patients survive into adult life, possibly because they developed ossification of the odontoid peg and C1 ring to a variable extent. The diagnosis is usually made on the physical and radiological features of bones, blood enzymes and excessive urinary levels of keratin sulfate which we found to be positive in our case. Once diagnosed it requires a multidisciplinary approach to patient care.

**Conclusion**

The typical clinical presentation of Morquio syndrome makes the diagnosis easier for this rare condition. Patient can present to physicians of different specialties starting from Dentist, ENT specialist to Internist and respiratory physicians. Treatment is multi-disciplinary and challenging. Enzyme assay is necessary for confirmation and to detect its type which will help in offering enzyme replacement therapy to the sufferer. Respiratory complications are most common cause of mortality and should be sorted in every cases for early intervention.

**Conflict of interest:** Nothing to declare.

**References**