International Guidelines for the Management and Treatment of Morquio A Syndrome

Christian J. Hendriksz,1* Kenneth I. Berger,2 Roberto Giugliani,3 Paul Harmatz,4 Christoph Kampmann,5 William G. Mackenzie,6 Julian Raiman,7 Martha Solano Villarreal,8 and Ravi Savarirayan9

1Salford Royal NHS Foundation Trust, Salford, UK
2New York University School of Medicine, New York
3Department of Genetics/UFRGS and INAGEMP, Medical Genetics Service/HCPA, Porto Alegre, RS, Brazil
4University of California San Francisco Benioff Children’s Hospital Oakland, Oakland, California
5University Children’s Hospital, Mainz, Germany
6Nemours/Alfred I. duPont Hospital for Children, Wilmington, Delaware
7Hospital for Sick Children, Toronto, Ontario, Canada
8Fundacion Cardioinfantil, Bogota, C/merca, Colombia
9Murdoch Childrens Research Institute and University of Melbourne, Parkville, Victoria, Australia

Manuscript Received: 25 March 2014; Manuscript Accepted: 22 September 2014

Morquio A syndrome (mucopolysaccharidosis IVA) is a lysosomal storage disorder associated with skeletal and joint abnormalities and significant non-skeletal manifestations including respiratory disease, spinal cord compression, cardiac disease, impaired vision, hearing loss, and dental problems. The clinical presentation, onset, severity and progression rate of clinical manifestations of Morquio A syndrome vary widely between patients. Because of the heterogeneous and progressive nature of the disease, the management of patients with Morquio A syndrome is challenging and requires a multidisciplinary approach, involving an array of specialists. The current paper presents international guidelines for the evaluation, treatment and symptom-based management of Morquio A syndrome. These guidelines were developed during two expert meetings by an international panel of specialists in pediatrics, genetics, orthopedics, pulmonology, cardiology, and anesthesia with extensive experience in managing Morquio A syndrome.

How to Cite this Article:

Key words: mucopolysaccharidosis IV; guidelines; symptom assessment; diagnosis; disease management

This is an open access article under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs License, which permits use and distribution in any medium, provided the original work is properly cited, the use is non-commercial and no modifications or adaptations are made.

Conflicts of interest: The authors are grateful to Ismar Healthcare NV for their assistance in writing of the manuscript, which was funded by BioMarin Pharmaceutical Inc. The current management guidelines were based on the outcome of two advisory board meetings sponsored by BioMarin Pharmaceutical Inc. Dr. Hendriksz received financial support in person or by the institution from BioMarin in the following capacities: honoraria for lectures, chairman of advisory boards, consultant on projects, research trials and travel grants. Dr. Berger has worked as a consultant for BioMarin Pharmaceutical Inc and for Genzyme. Dr. Harmatz has worked as consultant and study investigator for BioMarin Pharmaceutical Inc. and has received an honorarium. Dr. Giugliani is member of the BioMarin sponsored Morquio Program Advisory Board, and has received travel grants and speaker fees from Actelion, Amicus, BioMarin, Genzyme, Shire and Synageva. The other authors do not have any conflicts of interest.

*Correspondence to:
Dr. Christian J. Hendriksz, Consultant Transitional Metabolic Medicine, Manchester Academic Health Science Centre, The Mark Holland Metabolic Unit, Salford Royal Foundation NHS Trust, Ladywell NW2-2nd Floor Room 107, Salford, Manchester M6 8HD. E-mail: Chris.Hendriksz@srf.nhs.uk

Article first published online in Wiley Online Library (wileyonlinelibrary.com): 24 October 2014

DOI 10.1002/ajmg.a.36833
Morquio A syndrome (mucopolysaccharidosis [MPS] IVA, OMIM #253000) is a lysosomal storage disorder (LSD) inherited in an autosomal recessive fashion. It is caused by a deficiency in the enzyme N-acetylgalactosamine-6-sulfatase (GALNS) due to a mutation in the GALNS gene located on chromosome 16q24.3. This deficiency results in accumulation of the glycosaminoglycans (GAGs) chondroitin-6-sulfate and keratan sulfate (KS) in a variety of tissues [Neufeld and Muenzer, 2001]. The disease is extremely rare, with incidence rates ranging from 1 in 640,000 live births in Western Australia to 1 in 76,000 live births in Northern Ireland [Nelson, 1997; Nelson et al., 2003].

Infants with Morquio A syndrome usually appear normal at birth. However, due to the accumulation of storage material in tissues and organs, leading to cellular dysfunction, they progressively develop profound skeletal and joint abnormalities and an array of non-skeletal manifestations including respiratory disease, spinal cord compression, cardiac disease, impaired vision, hearing loss, dental problems, and to a lesser extent hepatomegaly [Montaño et al., 2007; Harmatz et al., 2013; Hendriksz et al., 2013a]. Morquio A can be distinguished from other types of MPS disorders by the typical short-trunk dwarfism with short neck (Fig. 1). The skeletal manifestations are generally more extensive and severe than in other types of MPS disorders. Hypermobility of joints is very characteristic for Morquio A syndrome and distinguishes this disease from other types of MPS. In contrast to most other types of MPS disorders, Morquio A syndrome has not been associated with cognitive impairment [Davison et al., 2013].

To date, over 220 mutations in the GALNS gene have been identified [Morrone et al., 2014]. The most common gene mutation is present in <9% of the Morquio patients, giving rise to a wide heterogeneity with regard to clinical presentation, severity of disease, and rate of progression [Tomatsu et al., 2011]. Whereas most patients present with the classical phenotype, associated with short stature and severe skeletal and joint abnormalities, some patients do not have this characteristic appearance but may show severe impairment in other domains such as cardiorespiratory disease. The specific disease manifestations depend grossly on the residual GALNS activity and the accumulation rate and location of storage material in tissues. Studies have shown similar clinical manifestations in siblings with Morquio A syndrome, providing some evidence for genotype-phenotype associations [Tylki-Szymańska et al., 1998; Rekka et al., 2012]. The accumulation of GAGs throughout the body ultimately leads to premature death in most instances, with a life expectancy ranging from 10 to 20 years of age to almost normal in some patients [Tomatsu et al., 2011].

Because of the heterogeneous and progressive nature of the disease, the management of patients with Morquio A syndrome is challenging and requires a multidisciplinary approach. However, due to the rarity of the disease, most clinicians are not familiar with the special needs of these patients. Therefore, we developed international guidelines for the management and treatment of Morquio A syndrome based on the outcome of two expert meetings. On August 2–3, 2013, an international panel of 26 specialists in pediatrics, genetics, orthopedics, pulmonology, cardiology, and anesthetics with experience in Morquio A gathered in Amsterdam for an expert meeting sponsored by BioMarin Pharmaceutical Inc. During this meeting, existing literature and clinical data on different aspects of Morquio A syndrome were reviewed and discussed and preliminary management and treatment guidelines were prepared. These were discussed in more detail by a consensus panel of nine experts during a second meeting in Barcelona on September 1, 2013.
that patients cannot be classified into different subgroups based on clinical presentation, severity of disease, and/or rate of progression. These included the observed genotypic and phenotypic heterogeneity, the limited data on natural history and pathophysiology of the disease and the observation that patients may show severe disease in one domain, but not in another (e.g., severe respiratory disease, but normal height). Because patients cannot be classified into different subgroups, management differs depending on the clinical manifestations of the patient.

**DIAGNOSIS**

Because of the progressive nature of Morquio A syndrome, early diagnosis may be critical to optimize patient outcomes. Patients with the disease generally gain clinical attention due to a cluster of progressive, multisystemic clinical and/or radiographic findings (Table I). Definitive diagnosis is facilitated by referral to a metabolic/genetics center and is dependent upon clinical diagnostic (radiographic imaging) and biochemical testing in a specialized laboratory. A diagnostic testing algorithm for Morquio A syndrome has been published in 2013 (Fig. 2) [Wood et al., 2013]. Although urinary GAG analysis and enzyme activity testing in a dried blood spot (DBS) may raise suspicion of the presence of Morquio A, a definite diagnosis usually entails demonstration of reduced GALNS activity in leukocytes or fibroblasts. Molecular analysis can be performed as a further confirmation of the diagnosis.

Obtaining a diagnosis of Morquio A remains a challenge due to the rarity of the disease and similarities with other LSDs and skeletal dysplasias. Specifically, patients that have been diagnosed with, or are undergoing evaluation for spondyloepiphyseal dysplasia, pseudoachondroplasia, multiple epiphyseal dysplasia or bilateral Legg-Calvé-Perthes disease should have Morquio A excluded because of the similarities between these diseases and Morquio A, unless there is a family history of any of these diseases [Mendelsohn et al., 2013; Lachman et al., 2014]. Also, diagnosis of Morquio A syndrome may be delayed in patients that do not exhibit the classical initial signs of the disease such as kyphosis. For example, patients with a more attenuated disease course may first present with atypical signs such as hip stiffness and pain. Furthermore, screening tests for Morquio A disease are subject to false negative results. Total urinary GAG may not be elevated in Morquio A patients, especially as patients grow older, because KS levels decrease naturally with age and can reach levels consistent with the upper range reported in unaffected individuals. Moreover, as not all Morquio A patients have elevated KS levels and as KS may also be elevated in other disorders, KS measurements should not be used as a diagnostic tool for Morquio A syndrome. Therefore, it is optional to perform an enzyme activity assay in DBS in addition to urinary GAG measurement in patients suspected of having the disease. Noteworthy, the outcome of enzyme activity testing in DBS depends on the stability of the enzyme and the storage conditions of the sample [Camelier et al., 2011]. Based on these considerations, it is recommended to proceed to enzyme activity testing in leukocytes or fibroblasts not only when screening is positive or doubtful, but even when it is negative and the clinical-radiological data are suggestive of Morquio A. In fact, if there is strong clinical suspicion, an enzyme activity test could be performed immediately, without prior urinary GAG or DBS screening. When enzyme activity analysis in fibroblasts or leukocytes is inconclusive, the test should be repeated and/or genetic analysis should be performed. In these cases with inconclusive enzyme results, identification of two known pathogenic mutations on separate alleles is required for definitive diagnosis [Wood et al., 2013]. Activity testing of other lysosomal enzymes (including β-galactosidase and another sulfatase) in conjunction with GALNS is recommended to rule out the presence of MPS IVB and other enzyme deficiencies which could have GALNS secondary deficiency, such as mucolipidosis II, III, MPS VI, and multiple sulfatase deficiency [Wood et al., 2013]. Obtaining a correct diagnosis is essential given the current availability of enzyme replacement therapy (ERT) for Morquio A syndrome (see below).

### TABLE I. Overview of Common Clinical Manifestations of Morquio A Syndrome and Prevalence of These Manifestations in the MorCAP Study (N = 325) at Baseline [Harmatz et al., 2013]

<table>
<thead>
<tr>
<th>Musculoskeletal manifestations</th>
<th>Prevalence in MorCAP (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Short stature</td>
<td>93</td>
</tr>
<tr>
<td>Short neck</td>
<td>91</td>
</tr>
<tr>
<td>Spinal abnormalities</td>
<td></td>
</tr>
<tr>
<td>Kyphoscoliosis</td>
<td>85</td>
</tr>
<tr>
<td>Odontoid dysplasia</td>
<td>65</td>
</tr>
<tr>
<td>Lumbar lordosis</td>
<td>56</td>
</tr>
<tr>
<td>Cervical spinal instability</td>
<td>49</td>
</tr>
<tr>
<td>Spinal disc disease</td>
<td>23</td>
</tr>
<tr>
<td>Hip dysplasia</td>
<td>71</td>
</tr>
<tr>
<td>Genu valgum (knock knees)</td>
<td>93</td>
</tr>
<tr>
<td>Pectus carinatum</td>
<td>97</td>
</tr>
<tr>
<td>Joint abnormalities</td>
<td></td>
</tr>
<tr>
<td>Laxity</td>
<td>87</td>
</tr>
<tr>
<td>Stiffness/pain</td>
<td>83</td>
</tr>
<tr>
<td>Contractures</td>
<td>52</td>
</tr>
<tr>
<td>Subluxation</td>
<td>47</td>
</tr>
<tr>
<td>Abnormal gait</td>
<td>94</td>
</tr>
<tr>
<td>Non-skeletal manifestations</td>
<td></td>
</tr>
<tr>
<td>Respiratory compromise</td>
<td>58</td>
</tr>
<tr>
<td>Cardiac abnormalities</td>
<td></td>
</tr>
<tr>
<td>Valve disease, small heart</td>
<td>43</td>
</tr>
<tr>
<td>Heart with low stroke volume</td>
<td>51</td>
</tr>
<tr>
<td>Increased heart rate</td>
<td></td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td></td>
</tr>
<tr>
<td>Cervical myelopathy</td>
<td>79</td>
</tr>
<tr>
<td>Cervical/thoracolumbar cord compression</td>
<td>79</td>
</tr>
<tr>
<td>Eye disorders</td>
<td></td>
</tr>
<tr>
<td>Corneal clouding</td>
<td>63</td>
</tr>
<tr>
<td>Visual impairment</td>
<td>22</td>
</tr>
<tr>
<td>Ear/labyrinth disorders</td>
<td>77</td>
</tr>
<tr>
<td>Most commonly hearing impairment, otitis media</td>
<td>77</td>
</tr>
<tr>
<td>Dental abnormalities</td>
<td></td>
</tr>
<tr>
<td>Hepatomegaly</td>
<td>69</td>
</tr>
<tr>
<td>Splenomegaly</td>
<td>17</td>
</tr>
<tr>
<td>Abdominal abnormalities</td>
<td></td>
</tr>
<tr>
<td>Other (e.g., hernias)</td>
<td>15</td>
</tr>
</tbody>
</table>
MANAGEMENT GUIDELINES
Treatment Modalities to Provide the Deficient Enzyme

Until recently, treatment of Morquio A syndrome was limited to supportive, symptomatic care, including symptom-based medications, physical therapy, surgery, and rehabilitation. However, in light of the pathophysiology of the disease, with a deficiency in a single enzyme giving rise to an array of clinical manifestations, a therapy that restores or replaces the deficient enzyme is desirable. Current experience with hematopoietic stem cell transplantation (HSCT), which has shown to be valuable in MPS IH (Hurler syndrome), is very limited and not encouraging for patients with Morquio A syndrome [Tomatsu et al., 2011; Algahim and Almassi, 2013]. More data are required to provide evidence of its efficacy in these patients.

ERT with recombinant human GALNS (elosulfase alfa) has recently been approved for Morquio A syndrome, providing a systemic treatment approach. Elosulfase alfa has shown to be effective with a favorable safety profile. A multicenter, phase 1/2, open-label, dose-escalation study including 20 Morquio A patients aged 5–18 years showed sustained improvements in endurance, as measured using a 6-min walk test (6MWT) and a 3-min stair climb test (3MSCT), and respiratory function using elosulfase alfa 1.0 mg/kg/week and 2.0 mg/kg/week, which persisted after approximately 2 years of treatment [Hendriksz et al., 2012]. Urinary KS also decreased with treatment, with the largest decreases observed when dosing at 2.0 mg/kg/week. Based on the outcome of the phase 1/2 study, a multicenter, double-blind, placebo-controlled phase 3 study was performed to assess the efficacy and safety of infusions with elosulfase alfa 2.0 mg/kg every week and every other week (N = 176; aged ≥ 5 years) [Hendriksz et al., 2014a]. The study showed significant improvement in endurance of 22.5 m in 6MWT distance during 24 weeks of treatment with elosulfase alfa at 2.0 mg/kg/week as compared with placebo (P = 0.0174). No significant impact was observed with the every other week dosing regimen. There was no significant effect of any of the dosing regimens on the number of stairs climbed in a 3MSCT. Both dosing regimens led to rapid and sustained reductions in urinary KS, with the greatest effect seen with the weekly dosing regimen. The study also showed numerical improvements over placebo after 24 weeks of treatment with elosulfase alfa 2.0 mg/kg/week for most exploratory endpoints, including maximum voluntary ventilation (MVV; P = 0.094) and forced vital capacity (FVC; P = 0.304). A composite measure developed by Delphi strategy based on an equal weighting of the change from baseline to week 24 of the z-scores of three component measures, i.e., 6MWT, 3MSCT, and MVV, also showed improvement. Both doses of elosulfase alfa had favorable safety profiles, generally similar to that of other ERTs. Most adverse events were mild or moderate infusion-associated reactions (IARs, occurring after infusion onset and within 1 day after infusion end) such as vomiting and pyrexia (fever), which were generally manageable with symptomatic treatment and/or infusion rate modification. The frequency of IARs was higher during the first 12 weeks of treatment and tended to occur less frequently with time. As anaphylactic reactions may occur during infusion, patients should be closely observed during and after administration of elosulfase alfa.

Elosulfase alfa treatment should be implemented as soon as the diagnosis of Morquio A syndrome has been confirmed to replace the deficient GALNs enzyme. The recommended dose of elosulfase

Early Assessments

Early recognition of clinical manifestations of Morquio A syndrome allows timely intervention and may help prevent irreversible damage. Therefore, patients with Morquio A syndrome should undergo a comprehensive multisystem evaluation of physical manifestations of disease, functional ability and disease burden at diagnosis. Table II illustrates the recommended assessments organized by organ system along with a schedule for longitudinal assessment. Each patient should also be referred to appropriate allied health care professionals such as a physiotherapist, occupational therapist, and audiologist depending on the disease burden at diagnosis. More details on these assessments will be discussed below.

Multidisciplinary Approach

The diverse spectrum of disease manifestations of Morquio A warrants a multi-disciplinary management approach, involving an array of specialists coordinated by a physician experienced in working with patients with LSDs. The role of this coordinating physician, usually a pediatrician, metabolic physician, or clinical geneticist, is to continuously monitor the evolution of disease, to refer to a specialist as needed, to assist in coordinating the care/ recommendations from the multidisciplinary team, and to serve as a medical home for the patient. In addition, this coordinating physician needs to regularly educate other health professionals (other specialists, dentists, and physiotherapists) about the disease and discuss the risks and benefits of interventions and necessary precautions with treatments/evaluations. The coordinating physician should provide guidance to the patient and family and encourage correct follow-up of therapies [Martins et al., 2009]. In order to stimulate continuity of care, it is also important that patients and families are educated about the disease and its possible complications and risks.

It is important that the coordinating physician of Morquio A patients organizes an experienced multidisciplinary team of specialists, concentrating all experience in a limited number of physicians per subspecialty. This requires referral of a critical mass of MPS patients to the same specialists, allowing them to accumulate experience in treating these patients and encourage their education on the disease.

SYMPTOM-BASED MANAGEMENT

Musculoskeletal Manifestations

Overview. Skeletal and joint abnormalities are the most apparent and prevalent disease manifestations of Morquio A syndrome. The majority of patients show short stature with corresponding short trunk and neck, abnormalities in spine, upper extremities, thorax, hips and/or lower extremities, and joint and gait abnormalities [Montaño et al., 2007; Aslam et al., 2013; Dha-wale et al., 2013; Harmatz et al., 2013]. Radiographic findings (dysostosis multiplex) suggesting Morquio A syndrome include abnormally shaped vertebral bodies with anterior beaking, posterior scalloping, platyspondyly and dens hypoplasia, thoracolumbar kyphosis, short, broad metacarpals with proximal rounding, irregular carpal bones, rounded iliac wings, acetabular dysplasia, coxa valga, genu valgum, ankle valgus, pectus carinatum, paddle-shaped ribs and short, and thick clavicles (Fig. 3), although only some of these may be present at diagnosis and missed by the non-expert [Hendriksz et al., 2013b].

Typical features of Morquio A not occurring in other types of MPS disorders are joint hypermobility and deformity in the wrists, leading to floppy wrists with weak grip and loss of fine motor skills [Aslam et al., 2013]. Patients may also show subluxation of the hip joints and joint instability in the knees, which can exacerbate genu valgum, patella dislocation, and gait abnormalities. Dens hypoplasia in combination with ligamentous laxity can lead to atlantoaxial instability and subsequently to spinal canal stenosis and spinal cord compression [Solanki et al., 2013].

Evaluation. Each Morquio A patient should be referred to an orthopedic surgeon with experience treating MPS disease at diagnosis. Regular assessments of the hips, lower extremities, and spine are recommended for optimal outcomes (Table II) [Solanki et al., 2013; White et al., 2014]. Radiographic assessments of the hips and lower extremities should focus on the presence of progressive hip dysplasia (shallow acetabulum with subluxation), genu valgum (tibiofemoral angle), and ankle valgus. Information on gait and mobility can be obtained by a simple physical exam and interrogation of patients. The clinical utility of gait analysis has not been established at present.

Evaluation of the spine should focus on the presence of spinal stenosis, instability and cord compression. Detailed instructions for detecting spinal cord compression in patients with Morquio A and indications for surgery have been recently published [Solanki et al., 2013]. Briefly, plain radiography of the cervical spine (anteroposterior, lateral neutral, and flexion extension) and of the thoracolumbar spine (anteroposterior and lateral) should be performed every 1–3 years, depending on clinical history. Magnetic resonance imaging (MRI) of the whole spine (neutral position) should be done annually, focusing on all three potential sites of cord compression, i.e., occipitocervical, cervicothoracic, and thoraco-lumbar. If available, flexion-extension MRI of the cervical spine is recommended every 1–3 years depending on signs, symptoms and radiographic instability. Computed tomography (CT) of the spine can be valuable for pre-operative planning.

Each patient with Morquio A syndrome also requires regular assessment of upper limb function (fine motor skills). Valuable standardized upper extremity function tests include evaluation of...
TABLE II. Recommended Schedule of Assessments in Patients With Morquio A Syndrome

<table>
<thead>
<tr>
<th>Assessment</th>
<th>At diagnosis</th>
<th>Follow-up frequency</th>
<th>As clinically indicated</th>
<th>Pre-ERT&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medical history</td>
<td>X</td>
<td>Every visit</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Physical examination</td>
<td>X</td>
<td>Every visit</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Upper limb function</td>
<td></td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Standardized upper extremity function test</td>
<td>X</td>
<td>Annually</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Hips and lower extremities</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Hips/pelvis: AP pelvis radiograph</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Lower extremities: standing AP radiographs</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Spine/spinal cord compression</td>
<td>X</td>
<td>Every 1–3 years</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Plain radiograph spine</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>MRI spine</td>
<td>X</td>
<td>Annually</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>CT neutral region of interest</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Cardiac function</td>
<td>X</td>
<td>Every 1–3 years&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>ECG&lt;sup&gt;d&lt;/sup&gt;</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Echocardiogram</td>
<td>X</td>
<td>Every 2–3 years&lt;sup&gt;c&lt;/sup&gt;</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Heart rate</td>
<td>X</td>
<td>Annually</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Respiratory function</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>FVC</td>
<td>X</td>
<td>Annually&lt;sup&gt;f&lt;/sup&gt;</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>MVV</td>
<td>X</td>
<td>Annually&lt;sup&gt;f&lt;/sup&gt;</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Respiratory rate</td>
<td>X</td>
<td>Annually&lt;sup&gt;f&lt;/sup&gt;</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Oxygen saturation&lt;sup&gt;g&lt;/sup&gt;</td>
<td>X</td>
<td>Annually&lt;sup&gt;f&lt;/sup&gt;</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Overnight sleep study</td>
<td>X</td>
<td>Annually&lt;sup&gt;f&lt;/sup&gt;</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Neurological function</td>
<td></td>
<td>Every visit (minimally every 6 months)</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Neurological exam</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Ophthalmological function</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Slit-lamp biomicroscopy of cornea</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Intraocular pressure</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Refractive error</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Examination of posterior segment</td>
<td>X</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Scotopic and photopic electroretinogram</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Hearing</td>
<td>X</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Audiology assessment (multimodal)</td>
<td>X</td>
<td>Annually</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Dental evaluation</td>
<td>X</td>
<td>Annually</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Evaluation of oral health by dentist</td>
<td>X</td>
<td>Annually</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Endurance</td>
<td>X</td>
<td>Annually</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>GMWT, T25FW&lt;sup&gt;e&lt;/sup&gt;</td>
<td>X</td>
<td>Annually</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Height and length</td>
<td>X</td>
<td>Every visit</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Weight</td>
<td>X</td>
<td>Every visit</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Head circumference (≤3 years)</td>
<td>X</td>
<td>Every visit</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Pubertal stage (age 9 until mature)&lt;sup&gt;j&lt;/sup&gt;</td>
<td>X</td>
<td>Every visit</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Disease burden</td>
<td>X</td>
<td>Every 6 months</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Pain assessment</td>
<td>X</td>
<td>Every 6 months</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Ool. questionnaire</td>
<td>X</td>
<td>Annually</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Functional test/ADL questionnaire</td>
<td>X</td>
<td>Annually</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>Evaluation by physiotherapist</td>
<td>X</td>
<td>Annually</td>
<td>X</td>
<td>X</td>
</tr>
</tbody>
</table>

<sup>6</sup>GMWT, 6-min walk test; ADL, activities of daily living; AP, anteroposterior; ECG, electrocardiogram; ERT, enzyme-replacement therapy; FVC, forced vital capacity; GAG, glycosaminoglycans; MRI, magnetic resonance imaging; MVV, maximum voluntary ventilation; Ool., quality of life; T25FW, timed 25-foot walk.

<sup>a</sup>If not done within 3–6 months, these assessments should be done before treatment with ERT is started.

<sup>b</sup>For example pre-operative planning.

<sup>c</sup>ECG and echocardiogram at diagnosis and after 1 year. If no signs of cardiac involvement, assessments can be repeated every 3 years, otherwise follow-up in expert centers according to standard of care.

<sup>d</sup>In symptomatic patients (e.g., suspicious ECG) or post-pubertal patients, prolonged ECG [Holter monitoring for 5–7 days including normal exercising] should be done in expert centers at diagnosis and every 1–3 years.

<sup>e</sup>Heart rate, respiratory rate, and oxygen saturation should be measured before and after each endurance test; choice of endurance measure depends on patient’s physical and developmental abilities (for the GMWT consistently use the same hallway).

<sup>f</sup>Annual follow-up only required until children stop growing or when patient is on treatment. Once growth has stopped, testing frequency can be decreased to every 2–3 years provided that respiratory symptoms remain unchanged.

<sup>g</sup>Oxygen saturation can be determined either by pulse oximetry or by arterial blood gas analysis.

<sup>h</sup>Screening studies should be done in home on an annual basis. Full polysomnography should be performed at diagnosis in an expert center, then every 3 years, unless clinically indicated (or before major surgery). Patients with a positive test and those who need ventilatory support should be evaluated by a sleep expert.

<sup>i</sup>For example pre- and post-operatively.

<sup>j</sup>Pubertal stage can be assessed using two scores: genitalia (male), breast (female), pubic hair (male and female) as described by Marshall and Tanner [Marshall and Tanner, 1969, 1970].

<sup>k</sup>For example GMWT/T25FW, pinch/grip test and functional dexterity test.

<sup>l</sup>For example 6MWT, 6-min walk test; ADL, activities of daily living; AP, anteroposterior; ECG, electrocardiogram; ERT, enzyme-replacement therapy; FVC, forced vital capacity; GAG, glycosaminoglycans; MRI, magnetic resonance imaging; MVV, maximum voluntary ventilation; Ool., quality of life; T25FW, timed 25-foot walk.

<sup>m</sup>If not done within 3–6 months, these assessments should be done before treatment with ERT is started.

<sup>n</sup>For example pre-operative planning.

<sup>o</sup>ECG and echocardiogram at diagnosis and after 1 year. If no signs of cardiac involvement, assessments can be repeated every 3 years, otherwise follow-up in expert centers according to standard of care.

<sup>p</sup>In symptomatic patients (e.g., suspicious ECG) or post-pubertal patients, prolonged ECG [Holter monitoring for 5–7 days including normal exercising] should be done in expert centers at diagnosis and every 1–3 years.

<sup>q</sup>Heart rate, respiratory rate, and oxygen saturation should be measured before and after each endurance test; choice of endurance measure depends on patient’s physical and developmental abilities (for the GMWT consistently use the same hallway).

<sup>r</sup>Annual follow-up only required until children stop growing or when patient is on treatment. Once growth has stopped, testing frequency can be decreased to every 2–3 years provided that respiratory symptoms remain unchanged.

<sup>s</sup>Oxygen saturation can be determined either by pulse oximetry or by arterial blood gas analysis.

<sup>t</sup>Screening studies should be done in home on an annual basis. Full polysomnography should be performed at diagnosis in an expert center, then every 3 years, unless clinically indicated (or before major surgery). Patients with a positive test and those who need ventilatory support should be evaluated by a sleep expert.

<sup>u</sup>For example pre- and post-operatively.

<sup>v</sup>Pubertal stage can be assessed using two scores: genitalia (male), breast (female), pubic hair (male and female) as described by Marshall and Tanner [Marshall and Tanner, 1969, 1970].

<sup>w</sup>For example GMWT/T25FW, pinch/grip test and functional dexterity test.
grip strength and pinch strength, the 9-hole peg test and the functional dexterity test [Aaron and Jansen, 2003; Poole et al., 2005]. In order to obtain consistent measurements, it is important to record the position of the wrist and whether the wrist was supported or not during testing. Passive and active assessments of the range of motion of elbows and shoulders can be useful, but generally do not have therapeutic consequences.

**Interventions.** Interventions for managing hip subluxation, knee valgus, and ankle valgus in patients with Morquio A syndrome have been recently discussed in detail by White et al. [2014]. Briefly, depending on age and severity of the deformity, hip subluxation can be managed using pelvic and femoral osteotomy, shelf acetabuloplasty, or total hip arthroplasty [Tassinari et al., 2008; Pryce Lewis and Gibson, 2010; Dhawale et al., 2012; White, 2012]. Correction of knee valgus using guided growth (8-plate hemiepiphysiodesis) is effective in the growing child and may prevent hip problems and ankle valgus at a later age [Dhawale et al., 2012]. Knee osteotomy or knee arthroplasty can be an option for older patients [Atinga and Hamer, 2008]. Ankle valgus can generally be managed by orthotics, but may sometimes require surgical correction (e.g., guided growth or osteotomy) [Dhawale et al., 2012]. Recurrence of the knee and/or ankle valgus is not uncommon.

Interventions for spinal cord compression in patients with Morquio A syndrome include spinal decompression, fusion or a combination of both [Ransford et al., 1996; White, 2012; Solanki et al., 2013]. Different approaches for managing spinal cord compression at the axial, subaxial, and thoracolumbar spine and the associated risks have been discussed by Solanki et al. [Solanki et al., 2013]. It is important to be aware that instability and stenosis may develop again in the region of previous fusion after surgery.

**FIG. 3.** Typical radiographic features (dysostosis multiplex) in patients with Morquio A syndrome: (A) Rib cage from a 12 year-old female Morquio A patient showing paddle shaped ribs. (B) Typical dysostosis multiplex changes in the pelvis and hips of an 8 year-old Morquio A patient showing dysplastic femoral epiphyses and narrowed inferior ilia sloping into the acetabular roofs. (C) Knee valgus in a 7 year-old Morquio A patient. Reproduced from Dhawale et al. [Dhawale et al., 2012] with permission from Lippincott Williams & Wilkins. (D) Cervical spine showing dens hypoplasia, which may be associated with atlantoaxial instability. Reproduced from Solanki et al. [Solanki et al., 2013] with permission from Springer. (E) Thoracolumbar spine changes including platyspondyly, anterior beaking, thoracolumbar kyphosis, and posterior vertebral scalloping. Reproduced from Solanki et al. [Solanki et al., 2013] with permission from Springer.
Surgical interventions requiring anesthesia warrant pre-operative evaluation of anesthetic risk factors and should be performed by a team experienced in managing these cases. Peri-operative care and anesthesia is discussed in more detail below.

Physical therapy and pain medication can be beneficial to manage musculoskeletal manifestations in some patients. A walking aid or wheelchair can help improve mobility and pain. However, efforts should be made to keep patients independently mobile as long as possible as the quality of life (QoL) drops dramatically when patients become wheelchair dependent [Hendriksz et al., 2014b].

Respiratory Manifestations

Overview. Respiratory impairment is the leading cause of morbidity and mortality in patients with Morquio A syndrome and can be due to obstructive or restrictive disease [Montano et al., 2007; Berger et al., 2012]. The upper and lower airways of Morquio A patients can be narrowed and tortuous due to a combination of GAG deposition in airway walls (Fig. 4), abnormalities in the skull or spine, tracheal distortion, tracheobronchomalacia, and thickened secretions. Patients may develop airway occlusion upon neck flexion and adopt a "sniff position" to preserve airway patency. Restrictive disease can develop due to a small and abnormally shaped thoracic cage or impaired diaphragmatic motility [Berger et al., 2012].

Evaluation. Forced vital capacity (FVC) and maximum voluntary ventilation (MVV) should be assessed annually until children stop growing. Once growth has stopped, the frequency of testing can be decreased to every 2–3 years provided that respiratory symptoms remain unchanged. Additional testing should be performed if respiratory symptoms change or if intercurrent illnesses occur. In addition, evaluation before and after initiation of ERT is recommended. Because of the growth impairment associated with Morquio A, expressing FVC and MVV as % of normal is of limited value. However, absolute values can be used to monitor respiratory function in patients over time.

In order to detect reduced exercise capacity due to respiratory restriction, it is also recommended to measure respiratory rate and arterial oxygen saturation before and after annual endurance testing (e.g., utilizing the 6MWT). Oxygen saturation can be obtained with a pulse oximeter along with heart rate. Reduction in oxygen saturation during exercise should prompt additional evaluation (e.g., pulmonary function and chest radiograph). Treatment targeted to the etiology for the desaturation may be administered and use of supplemental oxygen can be considered particularly if the oxygen saturation falls below 88–90%. In addition, the presence of inordinate tachycardia during exercise should prompt cardiac evaluation (e.g., assessment of valve and myocardial function and assessment for cardiac arrhythmia).

Routine physical exam can also identify signs of potential respiratory problems such as an enlarged tongue or sniff position. Fiber-optic laryngoscopy may be considered in patients showing obstructive symptoms and prior to surgery. Evaluation of respiratory function is also recommended before any planned air travel to ensure safety during the flight [Ahmedzai et al., 2011; Shrikrishna and Coker, 2011].

In order to timely detect SDB, in-home screening sleep studies that monitor oxygen saturation are recommended on an annual basis. A sleep study with full polysomnography in an expert center is recommended at diagnosis and then every 3 years, unless clinically indicated or before major surgery. Patients with a positive test and those who need ventilatory support should be evaluated by a sleep expert.

Interventions. The management of respiratory manifestations in patients with MPS has been discussed by Berger et al. [Berger et al., 2012]. Briefly, patients may benefit from supportive therapies such as regular influenza and pneumococcus vaccinations, bronchodilators and aggressive and prompt treatment...
of upper respiratory infections [Berger et al., 2012]. Tonsillectomy and/or adenoidectomy are frequently required in patients with obstructed upper airways. SDB can generally be managed successfully by continuous positive airway pressure (CPAP, for patients with OSA) or non-invasive ventilator support (e.g., bilevel positive airway pressure to treat sustained hypoventilation). Tracheostomy may be required if ventilator support is ineffective or in patients with airway obstruction during wakefulness. However, as many complications may occur during and after tracheostomy placement, this procedure should be performed only in centers with experience in Morquio A disease [Berger et al., 2012].

Cardiovascular Manifestations

Overview. Unlike other MPS disorders [Braunlin et al., 2011], cardiac valve abnormalities in Morquio A patients are generally very mild. Occasionally, older patients may develop clinically important cardiac disease post-pubertally. Detailed evaluation of cardiac data performed in Morquio A patients enrolled in the multicenter, cross-sectional Morquio A Clinical Assessment Program (MorCAP) [Harmatz et al., 2013] demonstrated variable incidence of valvular regurgitation and stenosis. In addition, Morquio A patients typically have a small left ventricular diameter and an abnormally low stroke volume (personal communication Prof. Dr. C. Kampmann). As compensation, these patients may demonstrate an abnormally high heart rate and high myocardial work index. In addition, blood pressure in these patients can increase dramatically if they switch from a supine to a sitting position.

Evaluation. Because cardiac pathology in Morquio A patients is generally mild, electrocardiography (ECG) and echocardiography every 3 years are sufficient, if there are no signs of cardiac involvement (Table II). Heart rate and blood pressure should be measured before and after annual endurance testing. Resting blood pressure and heart rate (before endurance testing) can indicate presence of arterial hypertension or resting tachycardia. Repeat measurement directly after endurance testing provides insight into a patient’s cardiovascular reserve and might detect cardiovascular incompetence. The left ventricular diameter generally does not increase after cessation of growth unless significant valve disease is present. In symptomatic patients (e.g., suspicious ECG) or post-pubertal patients, prolonged ECG (Holter monitoring for 5–7 days including normal exercising) should be performed at diagnosis and every 1–3 years to detect cardiac arrhythmias, which may be suspected based on experience in other MPS types.

Interventions. It is important to be aware that an elevated heart rate in Morquio A patients may be needed to compensate for a small cardiac stroke volume. Therefore, treatment of tachycardia with beta blockers should be avoided. Also ACE inhibitors should be used with caution as they may result in a disproportional increase in heart rate, especially in subjects with marked changes in blood pressure between supine and sitting positions.

Valve replacement may be considered for patients that present with or progress to severe aortic or mitral valve disease [Nicolini et al., 2008; Pagel and Almassi, 2009].

Neurological Manifestations

Overview. Morquio A patients can develop neurological symptoms due to myelopathy secondary to spinal cord compression [Harmatz et al., 2013; Solanki et al., 2013]. The MorCAP baseline data (N = 325) showed nervous system disorders or cord compression in 51% of patients, including 30% with cervical myelopathy, 14% with cervical cord compression, and 13% with thoracolumbar cord compression [Harmatz et al., 2013]. The prevalence of these disorders increased with age, i.e., 36% in 0–4 year old patients, 48% in the 5–11 year olds, 58% in 12–18 year olds, and 55% in >18 year olds. Cervical cord compression can lead to unsteady gait, upper and lower extremity weakness, dysesthesias, urinary dysfunction, paralysis and sudden death [Lipson, 1977; White, 2012]. At the thoracolumbar level, cord compression may result in lower back pain, radiating leg pain and paraplegia with insidious onset and associated consequences such as lower limb weakness, sensory anomalies and disturbed bladder function [Dalvie et al., 2001].

Evaluation. In order to identify patients with spinal cord compression in an early stage, neurological examinations at maximum intervals of six months are recommended (Table II). A neurological examination in patients with Morquio A syndrome may reveal hyperreflexia, increased muscle tone, pyramidal tract signs (ankle clonus and Babinski sign) and/or proprioceptive deficits [Solanki et al., 2013]. However, neurological assessment can sometimes be difficult due to lower limb function involvement. Also, neurological signs and symptoms may underestimate the severity of spinal cord compression seen on MRI. Clinical and neurological findings should therefore be correlated with imaging studies of the spine [Solanki et al., 2013]. In patients with multisegmental myelopathy, it can be difficult to determine the level of compression responsible for the observed neurological deficit. If available, it can be worthwhile to measure somatosensory evoked potentials (SEPs) and motor evoked potentials (MEPs) on an outpatient basis. However, the value of these studies in the evaluation of spinal cord compression still needs to be established [Solanki et al., 2013].

Interventions. MRI may over or underestimate the risk of cord compression in patients with Morquio A syndrome, therefore a neurosurgeon should be part of the multidisciplinary team to assist with making this assessment. Neurological monitoring during surgical interventions requiring anesthesia is recommended in cases where spinal cord compression is a concern. The treatment of spinal cord compression has been discussed previously (see musculoskeletal manifestations).

Ophthalmological Manifestations

Overview. Diffuse corneal clouding and refractive error problems (astigmatism, myopia, and hyperopia) are very common findings in Morquio A patients and may lead to reduced visual acuity and photosensitivity [Couprie et al., 2010; Summers and Ashworth, 2011; Harmatz et al., 2013; Hendriksz et al., 2013a]. Although corneal clouding worsens with age, it tends to be less severe in Morquio A syndrome than in MPS I and MPS VI [Ashworth et al., 2006]. Cataract, open-angle glaucoma, retinopathy, optic disc swelling and optic nerve atrophy, and pseudoxophthalmos due to shallow orbits have also been reported sporadically.
in patients with Morquio A syndrome [Dangel and Tsou, 1985; Cahane et al., 1990; Iwamoto et al., 1990; Olsen et al., 1993; Käsmann-Kellner et al., 1999; Ashworth et al., 2010; Couprie et al., 2010; Hendriksz et al., 2013a].

**Evaluation.** Ophthalmological function should be assessed in each Morquio A patient at diagnosis (Table II) [Fahnehjelm et al., 2012; Hendriksz et al., 2013a]. Afterwards, a basic evaluation of vision/ocular abnormalities should be part of the general physical examination that is recommended at every visit. Referral to an ophthalmologist is only required in case of clinical abnormalities.

**Interventions.** Patients with impaired vision may benefit from refractive correction or low vision aids; filtering glasses and hats can be used to manage photosensitivity [Tomatsu et al., 2011]. Corneal clouding can be managed surgically by corneal transplantation [Leslie et al., 2005; Tomatsu et al., 2011], but reopacification of corneal grafts may occur similar to experience in other MPS disorders [Käsmann-Kellner et al., 1999]. Concomitant retinopathy, glaucoma, or optic nerve atrophy can also limit restoration of vision with corneal transplantation. Patients with cataracts may benefit from cataract surgery.

**Audiological Manifestations**

**Overview.** Sensorineural or mixed conductive and sensorineural hearing loss commonly develop in Morquio A patients in the first decade of life [Bredenkamp et al., 1992; Riedner and Levin, 1977]. Hearing loss may result from recurrent respiratory tract infections or otitis media, deformity of the ossicles, and/or abnormalities of the inner ear [Bredenkamp et al., 1992; Riedner and Levin, 1977; Tomatsu et al., 2011; Hendriksz et al., 2013a]. A study including 18 patients with Morquio A syndrome showed conductive hearing loss in all three patients <8 years and mixed or sensorineural hearing loss in 14 out of 15 patients ≥8 years [Riedner and Levin., 1977]. Ten patients required a hearing aid. In the MorCAP study, ear and labyrinth disorders (mostly hearing impairment and otitis media) were reported by 77% of patients [Harmatz et al., 2013].

**Evaluation.** Hearing impairment is an underestimated issue in Morquio A patients. Therefore, age-adjusted audiology assessments should be done at diagnosis and on an annual basis thereafter (Table II) (e.g., acoustic emissions in neonates, visual reinforcement audiometry in children from 8 months up to 3 years of age, conventional ear checks for patients of 3 years or older) [Hendriksz et al., 2013a].

**Interventions.** Conductive hearing loss due to retained middle ear fluid can be treated using ventilation tubes [Hendriksz et al., 2013a]. Long-lasting types of tympanostomy tubes are preferable to use on the first occasion considering the anesthetic risks and the risk of the reoccurrence of the middle ear fluid [Hendriksz et al., 2013a]. Post-aural hearing aids may be most appropriate if a progressive neurosensory element to hearing loss is present.

**Abdominal Manifestations**

**Overview.** Abdominal manifestations of Morquio A include umbilical, inguinal or bilateral diaphragmatic hernias, hepatomegaly, splenomegaly (less common), and other gastro-intestinal disorders (e.g., chronic constipation, diarrhea) [Nursal et al., 2000; Harmatz et al., 2013; Hendriksz et al., 2013a]. However, gastro-intestinal manifestations tend to be less prominent than in other types of MPS disorders.

**Evaluation.** Assessment of gastro-intestinal problems in Morquio A patients should be part of routine clinical evaluation.

**Interventions.** Hernias can be surgically repaired by herniorrhaphy and frequently recur.

**Dental Abnormalities**

**Overview.** Patients with Morquio A syndrome tend to have small, widely spaced teeth, often with thin, structurally weak enamel and small pointed cusps, spade-shaped incisors, pitted buccal surfaces, and other developmental abnormalities of primary and permanent dentition (Fig. 5) [Kinirons and Nelson, 1990; Rolling et al., 1999; James et al., 2012]. Therefore, they are vulnerable to caries formation [James et al., 2012].

**Evaluation.** Patients should be referred to a dentist at diagnosis. Close monitoring of dental development (at least annually) and regular dental care is important to prevent caries and attrition of the teeth [Tomatsu et al., 2011; Hendriksz et al., 2013a].

**Interventions.** In order to prevent caries, Morquio A patients should receive fluoride supplementation; fissure sealing of dentition may be considered in some cases.
Overall Assessments

**Endurance.** Patients with Morquio A syndrome may show reduced endurance due to impaired cardiac, respiratory, musculoskeletal, and/or neurological function, which may impact significantly on functional status/mobility and QoL [Harmatz et al., 2013]. Endurance testing at diagnosis and annually thereafter is recommended to follow up overall disease progression (Table II). Information on endurance can be obtained by routinely asking endurance questions during clinical exam (e.g., How far can you walk? How many stairs can you climb?) or by using an endurance test.

The 6-min walk test (6MWT) is considered the most appropriate endurance test for patients with Morquio A syndrome, as it is readily available and is a combined assessment of musculoskeletal and cardiopulmonary capacity. This standardized test measures how far a person can walk on a hard, flat surface in 6 min and has been used to assess endurance in several MPS (including Morquio A) clinical trials [American Thoracic Society, 2002; Wraith et al., 2004; Harmatz et al., 2005; Harmatz et al., 2006; Muenzer et al., 2006; Hendriksz et al., 2012]. Detailed instructions for the 6MWT have been published by the American Thoracic Society [American Thoracic Society, 2002]. In order to allow comparison of test outcomes, the 6MWT should always be performed at the same location. Walking aids used during the 6MWT should be documented and should be consistent over time (i.e., do not change type of aid or perform with/without aid over time). In patients with limited ambulation who are unable to do the 6MWT, endurance can be assessed using a timed 25-foot walk test (T25FW). The T25FW, which has originally been developed as a component of the Multiple Sclerosis Functional Composite (MSFC), measures the time needed to cover 25 feet [Polman and Rudick, 2010]. For Morquio A patients, the test has been adapted in order to allow patients to cover the 25 feet distance by crawling or rolling if walking is impossible. Detailed instructions for the T25FW for Morquio A patients, based on the instructions developed for multiple sclerosis patients, are given in Supplementary Appendix 2. Blood saturation, heart rate and respiratory rate should be measured immediately before each endurance test, upon test completion, and 2 min after test completion. Compromised vision and hearing should be taken into consideration when implementing these tests in Morquio A patients. Endurance testing is also recommended before and regularly after initiation of ERT as a measure of treatment efficacy.

**Growth.** The majority of adult patients with Morquio A syndrome are below the 3rd centile in height on the Center for Disease Control growth charts and shorter than 120 cm [Harmatz et al., 2013]. They also tend to have a higher body mass index than normal (>85th centile) [Montaño et al., 2008]. Growth velocity generally starts to fall below normal levels after the age of 1 year [Montaño et al., 2008]. Growth charts for patients with Morquio A syndrome have been published in 2008 [Montaño et al., 2008].

Standing height (sitting height if the patient is unable to stand), length (supine) and weight, head circumference (up to 3 years of age), and stage of puberty (from age 9 years until maturity) [Marshall and Tanner, 1969, 1970] should be documented at diagnosis and at each visit to follow the relative change in individuals over time (Table II). Height and weight should also be measured before and regularly after initiation of ERT to evaluate the impact of treatment. One should be aware that height may vary (and even decrease) over time due to skeletal changes such as kyphosis or knee valgus. Because obesity may exacerbate lower limb and other problems, it is important to monitor weight (prevent excessive increase in BMI) in patients with Morquio A syndrome. Assessment of bone density is difficult to interpret in MPS patients given the lack of appropriate normative data for comparisons, and therefore not recommended on a routine basis [Fung et al., 2010].

**Pain.** Pain is a major, but underreported, symptom of Morquio A syndrome [Harmatz et al., 2013]. It is mainly related to musculoskeletal problems (e.g., joint pain and muscular pain) and may occur in many parts of the body, including the spinal area, upper and lower extremities, and the head and neck area. A patient-reported outcomes study in 27 adults and 36 children/adolescents with Morquio A showed that pain can interfere profoundly with activities of daily living (ADL) and can be a driver for wheelchair use [Hendriksz et al., 2014b]. Although increased mobility (less frequent wheelchair use) was found to be associated with more severe and widespread pain, pain interference with ADL was most severe in patients using a wheelchair all the time.

Pain severity should be assessed at each clinical visit by specifically asking patients about their analgesic use, or about recent improvement or worsening of pain (Table II). A simple, self-completed question for rating pain/discomfort is included in the EQ5D-5L (www.euroqol.org). This standardized questionnaire is developed to measure health-related QoL in a wide range of health conditions and currently available in 106 languages. External factors and conditions should be considered when interpreting pain assessment, for example pain might be increased after a long flight.

**Quality of Life.** Many factors may affect QoL in Morquio A patients, including reduced endurance or mobility, difficulties in ADL, dependence on caregivers, frequent surgical interventions, pain and fatigue [Harmatz et al., 2013; Hendriksz et al., 2013a]. It is important to be aware that although patients using a wheelchair all of the time experience less pain severity and fatigue due to limited functional activity, they tend to have a considerably poorer HRQoL [Hendriksz et al., 2014b]. Regular assessments of QoL and ADL are recommended (Table II). QoL can be assessed using reproducible and age-appropriate questionnaires such as the EQ-5D-5L. There is need for QoL questionnaires for adult patients, covering adult-specific needs such as employment, financial problems, and sexuality and reproduction. Information on the disease impact on ADL can be obtained using previously discussed functional tests (e.g., 6MWT/T25FW, pinch/grip test and functional dexterity test), open-ended questions, and reproducible, age-appropriate ADL questionnaires such as the MPS Health Assessment Questionnaire (MPS HAQ) [Harmatz et al., 2013].

Simple interventions may considerably improve the functional capacity and QoL of patients with Morquio A syndrome. For school children, communication between school staff, physiotherapists and health professionals about the inclusion of the patient in activities at school, specialized equipment, and walking aids is of major importance. It may be valuable to educate schools, including teachers and peers, on the disease and the fact that children may be...
absent for infusions and may have some limitations. This can enhance understanding, stop the potential for bullying, and stimulate peer tolerance and respect. When mobility is decreasing, adequate advice can be valuable for the patient in order to achieve a good balance between independence and maintaining mobility. Regular assessment of vision and hearing is important to ensure that deficits in these areas are being addressed.

**Urinary GAG.** In the absence of a well validated, readily available KS assay, measurement of urinary KS has a very limited role in the diagnosis of Morquio A patients and, while it is a robust marker of pharmacodynamic activity of elosulfase alfa, its clinical utility as a surrogate marker for elosulfase alfa efficacy has not been established. In the phase III study, urinary KS demonstrated a pharmacodynamic effect of elosulfase alfa in the weekly 2 mg/kg study population resulting in a 41% decrease in urinary KS at week 24 of the study [Hendriksz et al., 2014a]. However, changes in urinary KS did not correlate with clinical efficacy measures. Urinary KS can be measured prior to and just after starting elosulfase alfa to determine the pharmacodynamic effects of ERT treatment. Annual follow-up assessments are of limited use. Furthermore, the standard dye-based quantitative total GAG tests are predicted to be even less sensitive than the urinary KS test, and there is no current data to suggest that this testing is of any use in clinical management of Morquio A [Wood et al., 2013].

**Peri-Operative Care and Anesthesia**

Patients with Morquio A syndrome may be at high anesthetic risk due to cervical instability, compromised respiratory function or cardiac problems [Theroux et al., 2012; Walker et al., 2013]. Therefore, any elective surgery requires pre-operative evaluation of anesthetic risk factors and should be performed by an experienced team at centers familiar with MPS disorders. A spectrum of airway management equipment should be available during and after the procedure. The anesthetic problems and detailed instructions for pre-, peri- and post-operative care of patients with MPS in general and for patients with Morquio A have been published in 2013 in two papers by Walker et al. and Theroux et al. [Theroux et al., 2012; Walker et al., 2013].

**Physiotherapy**

As mentioned previously, Morquio A patients require regular evaluation of upper limb function by for example a physiotherapist. Referral to a physiotherapist on a routine basis (Table II) can also be valuable to encourage patients to perform routine exercises to better preserve joint function and fine motor skills [Tomatsu et al., 2011] and to participate in activities such as swimming or hippotherapy (occupational and physical therapy using the movements of a horse). Physiotherapy can be personalized to each patient and directed to address those joints and/or functions that are most impacted for that individual at that time.

**Transition From Pediatric to Adult Care**

During late adolescence, the relationship of patients with Morquio A syndrome with the medical team changes from a passive (with the parents mostly communicating with the medical team) to an active one (i.e., direct communication between patient and healthcare provider) [Hendriksz, 2013c]. This process, during which patients become increasingly independent from their parents and start managing their own health care, can be difficult, particularly in countries where the organization and facilities of the healthcare system require transition from pediatric to adult services. In order to guide this process and to ensure that the adult providers are knowledgeable in managing these patients and that patients are not lost to follow-up, there is need for a formal, site-specific, transition strategy. For example, joint visits with the pediatrician and the physician coordinating the adult patient for a few years or input of patient information by the pediatrician into a registry accessible for the adult team may guide adult care after transition. Alternatively, patients may carry their own health records. In any case, it is important to involve the patient and his/her family in this transition process.

**FUTURE DIRECTIONS**

Insights into the natural history and pathophysiology of Morquio A syndrome have grown rapidly over the past decade thanks to large observational studies, mutation analyses, and basic research in the framework of the development of ERT [Tomatsu et al., 2005; Montano et al., 2007; Tomatsu et al., 2008; Harmatz et al., 2013]. Nevertheless, current knowledge on the disease remains relatively limited. Future studies providing additional insight into genotypic-phenotypic correlations and biomarkers for disease activity are warranted and may allow the development of individualized management strategies according to the clinical manifestations of a patient.

As for other types of MPS disorders, implementations in symptom management and the recent introduction of ERT for Morquio A syndrome may create new challenges for the future management of these patients. As ERT can have a positive impact on mobility and endurance, it may encourage patients to adopt a more active life style. This may also create new expectations for the future and lower the threshold for orthopedic surgery. If the life expectancy of patients increases due to better treatment options, the previously-discussed needs of adult patients will gain even greater importance. A longer life expectancy is likely to be associated with an increase in the number of surgical interventions in patients with Morquio A and with increasing anesthetic challenges when patients grow older.

Another challenge for the future will be improving the convenience of ERT. Weekly visits to the hospital for infusions can be perceived as very bothersome by patients and families, especially if they have to travel far. Creating the opportunity for remote/home infusions may provide an answer to this problem.

**ACKNOWLEDGMENTS**

The authors are grateful to all specialists involved in the creation of these management guidelines. The expert panel involved in the consensus meeting in Barcelona consisted of Christian J Hendriksz, Salford Royal NHS Foundation Trust, Salford, UK; Kenneth I Berger, New York University School of Medicine, NY, USA; Roberto Giugliani, Medical Genetics Service/HCPA, Department of
REFERENCES


Alegre, Porto Alegre, Brazil; Edward W Fong, Children's Hospital & Research Center Oakland, Oakland, CA, USA; Tai-Tong Wong, University of Hong Kong, Hong Kong, China. Both expert meetings were coordinated and funded by BioMarin Pharmaceutical Inc.


Hendriksz CJ. 2013c. Transition services for adolescents with lysosomal storage disorders. CML-Lysosomal Storage Diseases 11:69–76.


Hendriksz CJ. 2013c. Transition services for adolescents with lysosomal storage disorders. CML-Lysosomal Storage Diseases 11:69–76.


SUPPORTING INFORMATION

Additional supporting information may be found in the online version of this article at the publisher’s web-site.