Fabry disease: Guidelines for the evaluation and management of multi-organ system involvement

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Fabry disease is an X-linked metabolic storage disorder due to the deficiency of lysosomal α -galactosidase A, and the subsequent accumulation of glycosphingolipids, primarily globotriaosylceramide, throughout the body. Males with classical Fabry disease develop early symptoms including pain and hypohidrosis by the second decade of life reflecting disease progression in the peripheral and autonomic nervous systems. An insidious cascade of disease processes ultimately results in severe renal, cardiac, and central nervous system complications in adulthood. The late complications are the main cause of late morbidity, as well as premature mortality. Disease presentation in female heterozygotes may be as severe as in males although women may also remain asymptomatic. The recent introduction of enzyme replacement therapy to address the underlying pathophysiology of Fabry disease has focused attention on the need for comprehensive, multidisciplinary evaluation and management of the multi-organ system involvement. In anticipation of evidence-based recommendations, an international panel of physicians with expertise in Fabry disease has proposed guidelines for the recognition, evaluation, and surveillance of diseaseassociated morbidities, as well as therapeutic strategies, including enzyme replacement and other adjunctive therapies, to optimize patient outcomes. **Genet Med 2006:8(9):539–548.**

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When one or more of the enzymes required for breakdown of complex macromolecules is deficient, pathologic accumulation occurs within lysosomes, progressively impairing normal cellular function and leading to clinical symptoms. Fabry disease, an Xlinked disorder, is a lysosomal storage disorder caused by deficient activity of α -galactosidase A. The primary disease process, i.e., pathologic accumulation of predominantly globotriaosylceramide (GL-3), involves vascular endothelial cells and other cell types such as neural cells, cardiomyocytes, and all types of renal cells,¹ leading to cellular dysfunction, tissue remodeling, fibrosis, ischemia and, ultimately to severe end-organ damage (Table 1). The classical phenotype is seen in most males and rarely in females. Early disease manifestations such as acropa-

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resthesias, angiokeratoma, and hypohidrosis become apparent in the pediatric years, whereas later in life, kidney, cardiac and central nervous system (CNS) complications develop. Although the pathologic processes are initiated in childhood, the rate of progression from sub-clinical processes to clinical symptoms can differ for each organ. In addition, there are some phenotypes in which not all classically affected organs are involved. The cardiac variant with residual enzymatic activity is seen in males and a variety of attenuated phenotypes are seen in women.¹

review

The gene encoding α -galactosidase A has been isolated and sequenced,^{2,3} and currently more than 245 different mutations (missense, nonsense, splice, deletion and insertion errors) have been identified.⁴ Attempts have been made to correlate genotype with clinical outcome, however these efforts have been confounded by the fact that there a very few recurrent mutations and, even within families, the phenotype may vary. The variability in females may be partly accounted for by nonrandom X-chromosome inactivation, but other factors such as genetic modifiers may play a role in both males and females.

Recently, the clinical manifestations, diagnostic strategies, and early efficacy of enzyme replacement therapy (ERT) for Fabry disease have been reviewed.^{1,5–8} Detailed knowledge of the natural history of Fabry disease provides the basis to measure the effects of treatment options. Several registries of clinical data are currently active for Fabry disease, of which the

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Organ system	Cell types potentially affected by GL-3 deposits	Spectrum of pathophysiologic findings
Kidney	Podocytes, mesangium, glomerular endothelium, epithelium of Bowman's capsule, loops of Henle and distal tubule, arterial and arteriolar smooth muscle and endothelium, interstitial cells ^{11,92}	Glomerular sclerosis, tubular atrophy, interstitial fibrosis ^{13,92}
Cardiac	Cardiomyocytes, conduction system cells, vascular endothelial and smooth muscle cells, valvular fibrocytes ⁹³	LV-hypertrophy; heart failure; stenosis or diffuse narrowing of epicardial vessels; atherosclerotic plaques; coronary vasospasm; thrombotic and thromboembolic complications ^{9,10}
Neurologic	Neurovascular endothelial cells, vasa vasorum, neurons within central and peripheral nervous system, including dorsal root and autonomic ganglia ^{94,95}	Ischemic injury (prothrombotic and occlusive abnormalities) and metabolic failure resulting in: Functional disruption of neuronal cells, loss of small myelinated and unmyelinated fibers; ischemic small vessel multifocal leukoencephalopathy; large vessel ectasia ^{51,78,94}
Dermatologic	Vascular endothelial cells, smooth muscle cells, fibroblasts, perineurium, eccrine sweat glands, including epithelium ^{64,96,97}	Weakening of capillary wall and vascular ectasia within (epi-)dermis; narrowing of small blood vessels around the eccrine sweat glands ^{64,65,98}
Ophthalmologic	Epithelial cells in the cornea, lens, ⁷² vascular endothelial cells ⁷⁵	Streaks in corneal epithelium; vasculopathy of the conjunctival and retinal vessels; central retinal artery occlusion; reduced lacrimal secretion ^{75,78}
Pulmonary	Airway epithelial cells, vascular endothelial cells, smooth muscle cells ^{62,63}	Airway narrowing, capillary blockage ⁶²
Gastrointestinal	Vascular endothelial and perithelial cells in the small intestine, colon and rectum; smooth muscle cells; autonomic nerve ganglia in the intestinal wall; small unmyelinated neurons and perineurial cells ^{55,99}	Narrowing of mesenteric small blood vessels ⁵⁵
ENT	Vascular endothelial cells, smooth muscle cells, ganglion cells ⁴⁸	Narrowing or total occlusion of cochlear vessels ⁴⁸ ; ischemic auditory neuropathy ⁴⁸

 Table 1

 Pathophysiologic findings in Fabry disease

Fabry Registry (www.fabryregistry.com) is the largest. For the registries to provide aggregate data on natural history and treatment response for all participants, a relatively uniform practice standard would provide a useful framework. Moreover, as the overall care of Fabry patients moves beyond the realm of genetic subspecialists, there is a need for management guidelines for a rare condition in which there may be only a single patient per medical practice.

An international panel of physicians with extensive expertise in the clinical management of patients with Fabry disease collaborated in developing guidelines for the therapeutic management of this condition. The authors, who represent eight subspecialties, performed a comprehensive review of the relevant medical literature on Fabry disease. The outcome of this review was supplemented by the experts' unpublished clinical experiences within their specialties. As prospective, longitudinal registry data are obtained and analyzed, opinion-based guidelines presented here may be modified.

Disease manifestations and management recommendations

The progressive nature of Fabry disease reflects different cascades of disease processes which may precede clinical symptoms by a significant time period. Lysosomal accumulation can lead to lysosomal and cellular dysfunction, which trigger tissue responses, such as inflammation, that ultimately lead to irreversible damage and renal, cardiac, and CNS complications. Vascular endothelial cell storage may lead to luminal encroachment and occlusion, as well as to altered vascular reactivity and a prothrombotic state,^{9,10} leading to ischemic events affecting various sites. Pathophysiologic observations in tissue specimens of Fabry disease patients are summarized in Table 1. As there is no fixed combination and order of organ involvement and disease progression, all relevant organ systems need to be assessed, treated, and monitored individually, as discussed below.

Renal involvement

Chronic kidney disease (CKD) is a prominent feature of classical Fabry disease and the main cause of premature death in the classical phenotype. Initial manifestations include impaired filtration rate (GFR), proteinuria, and tubular derangements.¹¹ Microalbuminuria and blunted concentrating ability have been reported in adolescents. Typically, reduced GFR and proteinuria associated with GL-3 accumulation develop during the third decade in affected males.^{1,12,13} Glomerular sclerosis, tubular atrophy, and interstitial fibrosis result in progressive kidney involvement and failure, at a rate comparable to diabetic nephropathy. Before the availability of dialysis and transplantation, death from kidney failure occurred early in the fifth decade in classically affected males.^{1,12,13} In contrast, a much lower proportion of females suffer progressive renal dysfunction to the point of end-stage renal disease or transplantation.14

Kidney outcome may be more directly related to interstitial and glomerular fibrosis and scarring rather than GL-3 deposits in parietal and visceral glomerular epithelial cells; epithelial

deposits have been demonstrated in heterozygous females with little derangement in kidney function.^{15–18} GL-3 deposits in podocytes have not been directly related to the magnitude of proteinuria or the rates of progression or severity of the kidney involvement in Fabry disease.

The evaluations of kidney function that should be carried out in every Fabry patient are outlined in Table 2 and should include estimates of GFR, total protein, (micro)albumin excretion, and urinary sodium excretion. In early stages of kidney involvement, quantitative estimates of GFR are necessary. While several 24 hour

Organ system	Assessment	Recommendation
General	General status, school or work performance, sports, depression, anxiety, drug use, pedigree update, somatic growth	Baseline (at first visit), every 6 months
	Complete physical examination SF-36® Health Survey, or PedsQL TM Measurement Mode	Baseline, every 6 months
	Genetic counseling	Baseline, every 6 months for new issues
	Genotype	If not previously determined
Kidney ^{<i>a</i>}	Serum electrolytes, creatinine, BUN; 24 hour urine or spot urine for total protein/creatinine, albumin/creatinine, sodium, creatinine, and (optional) GL-3	Baseline. Every 3 months if CKD stage 1 or 2 and >1 g/day of proteinuria or CKD stage 4 Every 6 months if CKD stage 3 Every 12 months if CKD stage 1 or 2 and <1 g/day of proteinuria
Cardiac	Palpitations, angina	Baseline, every 6 months
	Blood pressure, rhythm	Every evaluation visit
	ECG, echocardiography 2-D with Doppler	Baseline. Every other year for patients ≤35 years of age, every year thereafter
	Holter monitoring, 30-day event monitoring MRI, strain rate imaging	If an arrhythmia is suspected or palpitations are present Optional
	Coronary angiography	If clinical signs of angina
Neurologic	Acroparesthesias, fatigue, fever, sweating, heat and cold tolerance, joint pains, stroke-related symptoms, TIA	Baseline, every 6 months
	Neurologic exam, Brief Pain or McGill Pain Inventory	Baseline, every 6 months
	Brain MRI without contrast	Baseline. At time of a TIA or stroke event or in females
	Magnetic resonance angiography Cold and heat intolerance, pain, vibratory thresholds, sweat output, post-ganglionic sudomotor function, superficial skin blood flow	If cerebral vasculopathy should be excluded When available
	Comorbid stroke risk factors: Cholesterol (Total, LDL, HDL), triglycerides Lipoprotein A, total plasma homocysteine, factor V Leiden (G1691A), Protein C, Protein S, prothrombin G20210A, antithrombin III, anticardiolipin antibody, lupus anticoagulant	Anually At baseline, single assessment
ENT	Tinnitus, hearing loss, vertigo, dizziness	Baseline, every 6 months
	Audiometry, tympanometry, otoacoustic emissions	Baseline, and yearly thereafter
Ophthalmologic	Visual disturbances, light sensitivity	Baseline, every 6 months
	General ophthalmologic exam (slit-lamp, direct ophthalmoscopy, best corrected visual acuity, visual fields)	Baseline, every 12 months
	Retinal dysfunction testing (ERG, color vision testing, visual evoked potentials, retinal angiography), tear secretion testing	If clinically indicated
Pulmonology	Cough, exertional dyspnea, wheezing, exercise intolerance	Baseline, every 6 months
	Spirometry, including response to bronchodilators, treadmill exercise testing, oximetry, chest X-ray	Baseline, every 2 years or more frequently for clinical indications
Gastrointestinal	Postprandial abdominal pain, bloating, diarrhea, nausea, vomiting, early satiety, difficulty gaining weight	Baseline, every 6 months
	Endoscopic or radiographic evaluations	If symptoms persist or worsen despite treatment
Skeletal	Bone mineral density	Baseline

Table 2

^ackd stages: 1 gfr (mL/min/1.73m²), >90; 2 gfr, 60-89; 3 gfr 30-59; 4 gfr, 15-29; 5 gfr, <15 = kidney failure (dialysis or transplantation).

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urine collections would be ideal, the utility of "spot" urine protein/creatinine ratios and estimated GFR with the modification of diet with renal disease (MDRD) equation has been established. Assessment of proteinuria and GFR can be used for the staging of CKD, as described in the Kidney Disease Outcomes Quality Initiative (K/DOQI CKD) Guidelines of the United States National Kidney Foundation (Table 2).¹⁹ Kidney biopsies may be useful as a baseline assessment and in patients with atypical presentations, including a repeat kidney biopsy when disease is progressing despite therapy.

Therapeutic management primarily focuses on the control of blood pressure, lipids, and proteinuria.^{20,21} Angiotensinconverting enzyme (ACE) inhibitors and angiotensin receptor blockers (ARB) are important therapeutic options in all forms of proteinuric renal disease. The most important goals for adjunctive therapy in patients receiving ERT are stabilization of kidney function and reduction of urinary protein and albumin excretion to the absolute minimum, and not necessarily reduction of the systemic blood pressure to a preset goal. Patients may be unable to tolerate significant doses of ACE and ARB blood pressure medications even when they are uptitrated slowly, in part because of the relatively low baseline blood pressures and substandard exercise tolerance. Renal transplantation is undertaken routinely for Fabry disease patients who have reached end-stage renal disease.^{22,23} Of note, GL-3 accumulation does not occur in the transplanted kidney. For patients on dialysis, a pharmacokinetic study showed that agalsidase beta is not lost during standard hemodialysis, and these patients can be infused with agalsidase beta during the course of their routine hemodialysis.24

Cardiac involvement

The heart is frequently involved in the various phenotypic variants of Fabry disease, including women with clinical disease and males with the cardiac variant. In the latter, cardiac manifestations of Fabry disease are prominent and may or may not be accompanied by other manifestations.²⁵ In various surveys, 3–12% of patients with unexplained left ventricular hypertrophy (LVH) have Fabry disease,^{26–28} indicating that Fabry disease should be strongly considered if onset is over 40 years of age, with concentric hypertrophy, and if evidence of major outflow obstruction or long-standing hypertension are absent.

Cardiac events are a key cause of premature death, either due to primary Fabry-related cardiac disease or driven by cardiovascular disease seen in most patients on dialysis, regardless of underlying disease. Among dialysis patients, Fabry patients have increased morbidity and mortality.²⁹ The earliest heart abnormalities are shortened PR interval, resting bradycardia, mild valvular insufficiency, and diastolic dysfunction^{30,31} Involvement of the conduction system leads to potentially lethal arrhythmias. Permanent cardiac pacing is needed for about 10-20% of patients.^{32,33}

Myocardial GL-3 deposits represent <1% of the actual myocardial mass.²⁵ Other mechanisms, including inflammation and neuro-hormonal dysregulation have been proposed as stimuli to cardiac hypertrophy.²⁵ Hypertrophic cardiomyopathy can occur with or without left ventricular outflow tract obstruction, sometimes leading to significant intraventricular pressure gradients. Left ventricular enlargement and reduced ejection fraction can be seen in advanced stages of the disease, accompanied by clinical signs of congestive heart failure. The use of cardiac transplantation in a Fabry patient has been reported.³⁴

Damage to the coronary vascular bed may lead to angina pectoris, variant angina, and myocardial infarction and does not necessarily follow the typical patterns observed in atherosclerosis. Involvement of small, penetrating vessels is predominant.⁹ Both LDL- and HDL-cholesterol are commonly elevated.³⁵ Disturbance of the vasodilatative/vasoconstrictive mechanisms favoring the latter may account for the lymphedema in the absence of hypoproteinemia or varices.³⁶ Raynaud's phenomenon and vasospastic reactions leading to variant angina are common observations in Fabry patients.

Patients with Fabry disease should be monitored by routine ECG and echocardiography as specified in Table 2. Echocardiography is helpful to document concentric or eccentric LVH with or without left ventricle outflow tract (LVOT) obstruction, presence and grade of diastolic dysfunction, and rarely in advanced stages of the disease, dilated cardiomyopathy and significant valvular regurgitation. Holter (24-hour) monitoring may detect rhythm abnormalities, but 30-day event or loop monitoring can be required. When LVH is present, magnetic resonance imaging (MRI) can help determine the amount of scarring present. Coronary angiography is necessary when clinical signs of angina are present.

Treatment remains mainly symptomatic. Endothelial dysfunction with vasospasm and thrombotic events are treated with ACE-inhibitors, calcium channel blockers and anti-platelet drugs (aspirin, clopidogrel). Dyslipidemia (most commonly, hypercholesterolemia) should be treated with statins.³⁷ Permanent cardiac pacing in patients with higher degrees of A-V blocks is used. Left ventricular (LV) outflow tract obstruction leading to significant pressure gradients can be managed conservatively with verapamil. Interventional techniques such as ethanol induced percutaneous transluminal septal myocardial ablation (PTSMA) can be applied where conservative measures fail to bring relief. The use of beta blockers in cases of coronary artery disease should be monitored carefully because of the risk of worsening of bradycardia.38 In a typical patient, an ACE or ARB for afterload reduction is the most appropriate, unless there is frank outflow obstruction.

Neurological involvement

Neurological morbidity is composed of early symptoms related to dysfunction of the peripheral and autonomic nervous systems, as well as late symptoms due to vascular complications in the central nervous system. However, autonomic dysfunction might contribute to vascular events, and primary storage in neurons might also be involved in loss of neurons in the brain.³⁹

Prothrombotic and occlusive abnormalities, as well as large vessel ectasias, may compromise vascular reactivity and lead to ischemic-hypoxic damage. Given the intraneuronal GL-3 storage, primary cellular metabolic failure may also play a role in the pathophysiology of clinical stroke. In a cohort of 33 Fabry patients, 24% suffered at least one stroke at first occurrence mean age of 29 years.⁴⁰ Onset of CNS symptoms has been shown to occur earlier in males (age 34) than in females (age 40).⁴¹ The burden of CNS complications increases with age and one study showed that all patients older than 54 years had cerebrovascular involvement.⁴² Mild dementia can appear late secondary to diffuse leukomalacia, multiple strokes or possibly to lipid storage in hippocampal and frontal lobe neurons.⁴³

The earliest symptoms of Fabry disease are often episodic acroparesthesias which have been reported as early as age 2 and relate to small-fiber involvement in the peripheral nervous system. These can occur in both genders as constant burning pain and tingling especially in the feet and hands. Episodic pain crises of extremely severe attacks of sharp neuropathic pain⁴⁴ can last from a few minutes to several weeks and can be accompanied by fatigue, low-grade fever and joint pains. Pain may be triggered by physical activity, exercise, cold or heat exposure, fever, and stress. Additional early manifestations related to autonomic dysfunction include hypo- or anhidrosis, and poor temperature and exercise tolerance, that have been reported as early as age 645 and may also be present in females.46 Other symptoms include abnormalities of blood pressure regulation, poor cerebrovascular reactivity, impaired pupillary constriction, abnormal gastrointestinal motility with pain, and decreased tear and saliva formation.

Comprehensive neurological evaluation is essential, particularly before the institution of ERT, to assess disease extent and severity. History of pain, including frequency and severity, should be assessed using tools such as the Brief Pain Inventory or the McGill Pain Inventory. Clinical investigations include brain imaging by CT or, preferably by MRI with T1, T2 and FLAIR-weighted images. Magnetic resonance angiography (MRA) may be indicated to exclude cerebral vasculopathy. Additional MRI sequences (DWI, ADC) can assist in evaluation at time of transient ischemic attack (TIA) or stroke event. Test methods for monitoring nerve fiber dysfunction⁴⁷ (Table 2) are useful, but not widely available. Laboratory evaluation of comorbid stroke risk factors may identify patients with significantly elevated homocysteine, with a vitamin deficiency state, or with other genetic prothrombotic risk factors (Table 2).

Symptomatic therapy continues to play an important role in the clinical management of Fabry disease.⁴⁴ Phenytoin, carbamazepine oxcarbazepine, gabapentin and/or topiramate are used for treatment of painful crises and acroparesthesias. Patients are advised to self-monitor their activity in order to avoid factors that precipitate pain, limit their exposure to extreme temperatures and maintain hydration. Spraying the skin with water can temporarily help compensate for hypohidrosis.

Aspirin and other platelet-inhibiting agents (such as clopidogrel and ticlopidine) are often recommended as prophylaxis to minimize the risk of stroke. Clopidogrel should be used if aspirin is not tolerated, and both drugs may be indicated in patients who have had a thrombotic stroke or TIA. Proper hydration and avoidance of hypotension and hypertension are important for the maintenance of adequate cerebral perfusion.

ENT involvement

Until recently,⁴⁸ the otologic manifestations in Fabry disease were underappreciated, although progressive hearing loss predominantly in the high tone range, sudden deafness (often asymmetric), tinnitus, and vertigo with dizziness and nausea were reported decades ago.^{49,50} Hearing loss should be looked for actively since it may not be reported spontaneously.⁵¹ One study documented self-reported hearing loss in 41% of 98 male Fabry patients. An abnormal audiogram was obtained in 78% of these patients.⁵²

Study of cochlear function including conventional audiometry, tympanometry, auditory brainstem response (ABR) audiometry, and otoacoustic emissions in 22 male patients disclosed a high prevalence of progressive hearing loss, both of sensorineural origin (23%) and sudden deafness (32%), as well as tinnitus aurium (27%).⁴⁸

A detailed medical history of tinnitus, hearing loss, and vertigo should be carefully documented. Audiometry should be performed at baseline and if clinical symptoms appear. Sudden hearing loss can be managed with infusion of vasodilators and steroids. Hearing aids or cochlear implants should be considered in case of advanced deafness. Episodes of vertigo can be severe and associated nausea may require trimethobenzamide, prochlorperazine, or other antinausea drugs.

Gastrointestinal manifestations

Gastrointestinal symptoms may include postprandial abdominal pain and bloating followed by multiple bowel movements and chronic or intermittent diarrhea, nausea, vomiting and early satiety.⁵³ Many affected males have difficulty gaining weight⁵⁴ and are in general smaller in height and weight than their unaffected sibs. The symptoms typically begin in adolescence, sometimes before the age of 10 and often as primary symptom.⁵⁵ In a cohort of 25 pediatric patients, 80% experienced symptoms of nonspecific enteropathy (e.g., abdominal pain, diarrhea).⁵⁶ Generally, symptoms are more frequent and appear earlier among males.^{57,58}

The medical history should focus on bowel habits, weight gain and dietary aspects. Radiographic or endoscopic evaluation may be indicated. Delayed gastric emptying and dyspepsia have been successfully treated symptomatically, to some extent, with metoclopramide⁵⁹ and H-2 blockers, respectively.

Pulmonary manifestations and exercise intolerance

Pulmonary involvement may manifest as chronic cough, dyspnea with exercise, and wheezing. Chronic cough is a common complaint in men as well as women.⁶⁰ While the cough may be worsened by reactive airway disease or tobacco use, it can occur in individuals without these risk factors.^{61,62} Spirometry may show signs of obstructive ventilatory impairment,^{61,62} particularly with decreased FEV₁, FEV₁/FVC, or FEF_{25–75}.⁶⁰

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Exercise intolerance in Fabry disease is typically attributed to hypohidrosis, pulmonary disease, or cardiomyopathy, but can begin early in childhood and is not uncommon in female patients. A recent study revealed a decrease in diastolic blood pressure equal to or >10 mmHg during exercise testing in approximately half of the patients studied, with the largest declines occurring in females.⁶⁰

Spirometry and exercise testing with monitoring of blood pressure should be performed at baseline and every two years thereafter. This testing, along with the ECG and echocardiogram, is especially helpful for distinguishing causes of exercise intolerance and fatigue. If pulmonary involvement is significant, induced sputum may show typical Fabry lamellar inclusions,⁶³ but bronchoscopy and/or a lung biopsy may be required to rule out other etiologies for the pulmonary disease.

Dermal manifestations

Dermatological disturbances include the cutaneous vascular lesions (angiokeratomas), and, as previously discussed, anhidrosis, or more commonly hypohidrosis. Angiokeratomas, caused by weakening of the capillary wall and vascular ectasia within the epidermis and dermis,^{64,65} are one of the earliest clinical signs and may be seen in both genders. They usually manifest between 5 and 13 years of age.⁶⁶ They initially appear as small, slightly raised, purplish-red, non-blanching angiectases. With age, there is progressive increase in the number and size of these lesions. The hips, back, thighs, buttocks, penis, and conjunctiva.⁶⁷ The large genital lesions are particularly susceptible to thrombosis⁶⁸ and bleeding. Among female carriers, frequencies of occurrence of angiokeratomas up to 35% have been reported.⁶⁹

The results of various laser methods (e.g., variable pulse width 532-nm Neodymium, and flash lamp-pumped dye laser) in treating angiokeratomas of Fabry disease have been mixed or discouraging.⁷⁰ In addition, this treatment did not prevent formation of new lesions. More pedunculated lesions may be treated with a series of liquid nitrogen treatments prior to laser therapy. Topical moisturizers can be helpful in preventing skin fissures due to hypohidrosis and secondary infections.

Ophthalmologic manifestations

The ocular findings play an important role in the early recognition of Fabry disease. Corneal manifestations are reported to be present in over 90% of the patients.⁷¹ The best-known ocular sign of Fabry disease is the pattern of pale, spiral streaks in the corneal epithelium known as "cornea verticillata."⁷²; Mild forms have been observed in 88% of a cohort of pediatric patients.⁵⁶ The streaks may vary in color from pale gray to brownish or yellowish. In some patients, the corneal involvement is quite homogeneous, appearing as a diffuse, even corneal haze, with little or no evident streaking. The pattern of streaks is often most prominent in heterozygous females.^{73,74} The corneal haze or streaks associated with Fabry disease rarely, if ever, cause visual symptoms. In some cases, there may be a history of difficulties seeing under conditions of severe veiling glare, as when driving toward oncoming headlights.

Fabry disease patients are also susceptible to the development of opacities of the crystalline lens, or cataracts. The most characteristic lesion is a unique pattern of branching opacities lying just inside the posterior lens capsule.⁷² These dendritic opacities do not impair visual acuity, and do not represent an indication for cataract extraction. They are best visualized after dilation of the pupils, viewing the posterior lens capsule in the reflected red light of the fundus with a slit-lamp biomicroscope.

Vasculopathy often manifests in the conjunctival blood vessels, most commonly those in the inferior sectors of the bulbar conjunctiva.⁷² The most typical conjunctival lesion is the presence of small numbers of saccular aneurysms, best visualized with the slit-lamp biomicroscope. The prominent conjunctival vessels do not cause discomfort, and are not unusually prone to hemorrhage. Vascular involvement of the retina is common, leading to narrowing of the retinal arterioles, dilation and irregularity of the retinal veins, and exaggerated tortuosity of the retinal vessels.⁷⁵ Vision in Fabry disease patients is rarely impaired due to the retinal vasculopathy but sudden visual loss due to central retinal artery occlusion^{72,76} and ischemic optic neuropathy have been encountered.⁷⁷ Finally, lacrimal secretion may be reduced as a consequence of vascular involvement or GL-3 deposition in the lacrimal gland.⁷⁸

The ocular stigmata of Fabry disease rarely if ever cause significant impairment of vision and, as a rule, do not require treatment.⁷⁹ Best corrected visual acuity and visual fields should be tested periodically. If clinically indicated, retinal dysfunction testing (electroretinogram (ERG), color vision testing, visual evoked potentials, retinal angiography) and tear secretion testing may confirm ophthalmologic pathology.

Skeletal manifestations

A recent prospective study investigated bone involvement in 23 male patients with a mean age of 31 years. Bone mineral density (BMD) measurement of the lumbar spine and femoral neck revealed either osteopenia or osteoporosis in 88% of the patients.⁸⁰ It is recommended to perform initial assessment and long-term follow-up of BMD, as well as to prevent osteoporotic fractures.

Enzyme replacement therapy

Clinical research programs of enzyme replacement therapy (ERT) for Fabry disease led to commercial availability of agalsidase alpha (Replagal[®], Shire Human Genetic Therapies, Inc.)^{81,82} and agalsidase beta (Fabrazyme[®], Genzyme Corp.)^{83,84} in most European countries in 2001, and for agalsidase beta in the US in 2003. Infused α -galactosidase A is taken up by vascular endothelial and parenchymal cells into lysosomes, presumably through the mannose-6 phosphate, mannose and asialoglycoprotein receptors on the surface of the target cells. Both products are very similar, with comparable specific activities per mg and glycosylation patterns,⁸⁵ as well as complete cross-reactivity of IgG anti-

bodies to these products.86 Reduction of symptoms and prevention of late complications are key therapeutic goals for ERT. Organ systems must be assessed separately and at regular intervals in order to monitor the effectiveness of treatment (Table 2).

Treatment with both agalsidase alpha⁸² and agalsidase beta⁸⁴ have been shown to reduce GL-3 inclusions. Patients receiving agalsidase beta for at least 5 years showed no decline in renal function unless there was prominent renal sclerosis at baseline. However, TIAs and strokes continued to occur, emphasizing the need for adjunctive therapies. The peripheral neuropathy and hypohidrosis improve slowly. Late complications in older patients may be forestalled. In a randomized, placebo-controlled trial, 51 Fabry patients representing both genders received biweekly agalsidase beta infusions of 1 mg/kg, while 31 patients received placebo. All patients had mild-to moderate renal disease at baseline and median treatment duration was 18.5 months. Agalsidase beta significantly reduced the risk of reaching the composite endpoint of major renal, cardiac, cerebrovascular events, or death by 61% (P = 0.034) in the per protocol population, after adjusting for the baseline imbalance in proteinuria between the treatment groups. Beneficial effects on estimated GFR and serum creatinine levels were more pronounced in patients who began agalsidase beta at less advanced stages of renal dysfunction.87,88

In the authors' experience, proteinuria may be reversible if treatment is started early, but it is also clear that lost renal function is not recovered. Predictive factors for therapeutic outcome may include advanced renal pathology (e.g., interstitial fibrosis, tubular atrophy, glomerulosclerosis), impaired GFR, and severity of baseline proteinuria. A reduction of severe cardiac events appears consistent with earlier findings. ERT has been shown to reduce myocardial storage of GL-3,84 and a significant reduction in LV mass and improvement in intracardiac conduction (QRS complex) were observed with agalsidase beta therapy.89 Another study demonstrated significant improvement of myocardial regional function and reduction of LV mass with agalsidase beta.90 The latter observation was also made in a separate study of agalsidase alpha in females.91

Consensus guidelines for both adjunctive therapies and preventive measures are presented in Tables 3 and 4. The patient's treatment and follow-up evaluations should be supervised by a physician experienced in the management of patients with Fabry disease or other inherited metabolic diseases. Based on current knowledge, every male Fabry patient should be offered ERT, irrespective of the stage of CKD (Table 3).

Fabry disease in females

The mean age of onset of disease complications in females and the age of death appear to be later than in males.14 Affected females may display all of the classical symptoms but previously unsuspected Fabry disease has also been found in women with isolated organ involvement.28 The frequency of hypertrophic cardiomyopathy and of cerebrovascular accidents (stroke, transient ischemic attacks) may be increased in females over the age of 40.57 DNA-based diagnosis rather than enzyme ac-

Current guidelines for instituting enzyme replacement therapy in Fabry disease patients				
Fabry Population	Guideline for Instituting ERT			
Adult males (>16 y)	At time of diagnosis of Fabry disease			
Pediatric males	At time of development of significant symptoms ^a or			
	If asymptomatic, consider at 10–13 y			
Females (all ages)	Monitor; institute if significant symptoms" or evidence of progression of organ involvement			

Table 3

^aSee text for explanation.

tivity testing is required in females with either a positive family history or symptoms suggesting Fabry disease. Once the diagnosis is confirmed by molecular methods, the disease burden needs to be assessed clinically. A biopsy of the affected organ (the kidney, or more rarely, the heart) may be required before starting ERT to document the extent of GL-3 storage and to exclude other etiologies. Asymptomatic women can be evaluated every year, but symptomatic women should be monitored every six months, as is recommended for males.

Females with Fabry disease should be offered ERT if they manifest significant symptoms or show evidence of progressive end organ Fabry involvement including: chronic acroparesthesias resistant to conventional therapy, persistent proteinuria (>300 mg/24 hours), GFR below 80 mL/minute/1.73 m², clinically significant cardiac involvement, a previous cerebrovascular accident or history of transient ischemic attacks, or ischemic changes on brain MRI (Table 3). Other possible considerations include symptoms unresponsive to other forms of medical management, including chronic, disabling gastrointestinal dysfunction, pulmonary involvement, or abnormal blood pressure regulation during exercise testing.

Pediatric Fabry disease

Until recently, descriptions of the early, often subtle, presentations of Fabry disease were scarce, partly because the diagnosis was usually made well into adulthood. Recent reports have helped to clarify the natural history. In one study, onset of symptoms occurred earlier in males (mean age 10.9 years) than in females (22.6 years). The mean delay between first symptoms and diagnosis was 13.7 years for males and 16.3 years for affected women.⁵⁷ Most often, a positive family history leads to diagnosis in childhood. For affected children, particular attention should be paid to pain, hypohidrosis, gastrointestinal symptoms, school performance, level of activity, and development of proteinuria. Physical findings are likely to be limited to a few angiokeratoma and the characteristic eye findings.

When to initiate ERT in children is an additional area in which clear, evidence-based recommendations cannot be given at present. The authors agree that, once the diagnosis is confirmed, it is important to institute treatment at the onset of clinically significant symptoms. Boys who have few symptoms

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Table 4

Adjunctive therapies and preventive measures in Fabry disease patients for common Fabry related r	morbidities
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Symptomatic treatment and preventive measures				
Kidney	Proteinuria	Angiotensin enzyme converting (ACE) inhibitor or angiotensin receptor blocker (ARB)		
	Renal failure	Dialysis or kidney transplantation (donor screened negative for Fabry disease)		
Cardiac	Blood pressure	ACE inhibitor, calcium channel blocker to treat endothelial dysfunction with vasospasms		
	Dyslipidemia	Statins		
	Significant left ventricular outflow tract obstruction	Verapamil, percutaneous transluminal septal myocardial ablation (PTSMA)		
	Higher degrees of A-V block, or significant brady- or tachyarrhythmias	Permanent cardiac pacing (or implantable dual chamber defibrillator)		
	Significant coronary artery stenosis	Percutaneous transluminal coronary angioplasty (PTCA) or aorto-coronary bypass grafting (CABG)		
	Cardiac failure	Cardiac transplantation		
Neurologic	Painful crises and acroparesthesias	Avoidance of physical exertion (over-heating) or exposure to circumstances that provoke attacks, air conditioning, wetting clothes, cooling vests Phenytoin, carbamazepine, oxcarbazepine, gabapentin or topiramate		
	Stroke prophylaxis	Aspirin and other platelet inhibiting agents (enteric coated aspirin, 81 mg qd at age 30 for males and age 35 for females) Clopidogrel if aspirin is not tolerated Consider both drugs in case of stroke or TIA Promote adequate intake of Vitamins B12, B6, C, and folate		
	Depression, anxiety, drug use	Psychiatry referral, pain center referral, social work, serotonin reuptake inhibitor		
ENT	Vertigo-related nausea	Trimethobenzamide, prochlorperazine		
	Hearing loss (moderate)	Hearing aids		
	Profound deafness	Cochlear implants		
Dermatologic	Angiokeratomas	Consider laser methods (e.g., variable pulse width 532nm Neodymium, and flash lamp-pumped dye laser) for problematic lesions, especially if on ERT. Liquid nitrogen treatments prior to laser therapy if lesions are more pedunculated		
Ophthalmologic		Treatment rarely required		
Pulmonology		Cessation of smoking		
	Airway obstruction	Bronchodilators provide relief in some		
Gastrointestinal	Delayed gastric emptying	Small and frequent meals, metoclopramide		
	Dyspepsia	Dietary restrictions, H-2 blocker		

should be considered for ERT around the age of 10–13 years, but some patients begin treatment earlier. There is increasing evidence that suggests that ERT is most beneficial in terms of clinical outcome when initiated early in the disease process, i.e., before irreversible changes occur in key tissues and organs, and it may be anticipated that the incidence of serious complications may be reduced.⁸⁸ The timing of ERT initiation in boys is a collective decision between medical professionals, family, and the patient (Table 3).

Perspectives

Therapeutic management of Fabry disease requires a multidisciplinary approach by medical specialists experienced in treating this rare condition. Such a team approach necessitates active participation and communication between the geneticist, nephrologist, cardiologist, neurologist, and others.

Comprehensive monitoring of Fabry patients, regardless of age, sex, or treatment with ERT, should be conducted at regular intervals. Our proposed guidelines for these assessments are

summarized in Table 2. Data collected in patients representing the entire disease spectrum, and contributed to Fabry disease patient registries, will allow for further clarification of the response to treatment and better define the optimal care for these complex patients.

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