Improved ways to screen for patients with Fabry disease, involving optometry in a multidisciplinary approach

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Introduction

Although Fabry disease has been known for more than a century (1898), this lysosomal storage disorder remains poorly recognized. With a prevalence of 1/40,000 to 1/117,000 live male births, Fabry is considered one of 7,000 known rare diseases that exist in the US and reported. However, this number of patients seems to be underestimated.

Many heterozygous subjects are affected without being diagnosed, no one seeing the globality of their symptoms. These are quite variable and unspecific, often leading to confusion with rheumatoid diseases or chronic inflammatory conditions. In some cases, the condition remains subclinical or is not characteristic of the full spectrum of the disease. A typical patient’s odyssey means multiple visits to more than ten different medical specialists before he or she achieves a confirmatory diagnosis.

Conclusion:

En se basant sur les résultats obtenus, le modèle de dépistage mis en place a été évalué comme positif. Il confirme le rôle crucial des optométristes dans le dépistage des maladies systémiques ayant des implications oculaires. L’éducation continue est essentielle à la remise à jour et en perspective de notions apprises il y a longtemps mais qui ne se rencontrent pas au quotidien dans les pratiques. De plus, ceci suggère que l’optométriste peut être impliqué dans des équipes multidisciplinaires visant le dépistage de patients à risque de maladies, notamment celles qui entraînent des manifestations oculaires.
Fabry disease is considered a rare disease, based on its prevalence. It is recognized, however, that there are many individuals affected who are unscreened. This article aims to demonstrate how optometrists can help to define improved ways to screen patients affected by this rare metabolic disorder, in a multidisciplinary perspective.

A screening model, based on continuous education for optometrists was developed. Under this model, suspect patients identified by optometrists are referred to Université de Montréal’s vision clinic (EOUM) for further testing and assessment. Should ocular manifestations and/or case history prove relevant to these rare diseases, a urinary test is then performed to find related biomarkers. When suspicions narrow to probable diagnosis, more affected, before any major systemic involvement occurs. Because ocular manifestations are among the first to appear in Fabry patients, early in their life, it becomes interesting to consider optometrists as key primary care players to detect and screen for Fabry disease to a greater extent. This article aims to explain how it can be done effectively, in a multidisciplinary perspective.

**Fabry disease explained**

Fabry disease is an X-chromosome linked disease, and counts 431 different mutations for the GLA gene. It is characterized by a deficiency of the lysosomal enzyme alpha-galactosidase A, (GLA or α-gal A). Consequently, normal degradation and catabolism processes of membrane glycosphingolipids, namely globotriaosylceramide (also known as GL-3, CTH, or Gb₃) can no longer be processed in nearly all cells of the human organism. GL-3 substrates cause deposits within the blood vessels. Its distribution is heterogeneous, with a preference for organs that naturally accumulate the greatest amount of it (the heart and kidneys), it also favours the vascular endothelial cells, renal dorsal root ganglion cells, the cornea and the skin.

**Purpose:**

Fabry disease, the subjects are referred to metabolic disorder specialists for complete DNA testing and medical follow-up of their condition.

**Methods:**

Continuous education lectures were given across Quebec, reaching nearly 60% of the province’s optometrists.

**Results:**

Sixteen months following the model’s implementation, ten suspected patients were referred. Of these, two new Fabry patients were confirmed, leading to the diagnosis of five other relatives with the disease. Two additional persons, diagnosed as Fabry patients, but lost to medical follow-up for many years, were once again placed under the care of Fabry experts. To this point, because of optometric involvement, seven new patients of Fabry were diagnosed and two were brought back under experts care.

**Conclusion:**

Continuous education lectures were given across Quebec, reaching near 60% of the province’s optometrists.

**Key words:** Fabry disease, screening program, corneal pigmentation, urine biomarkers, lysosomal storage disorder.
During its course, the most severe (referred to as "classic") form of Fabry disease leads to multiple organ damage but clinical presentations typically vary from one patient to another (Table 1). On the other hand, milder or later-onset variants, with manifestations circumscribed to one organ, can be seen in patients showing some residual enzyme activity.13-14

In its classic form affecting hemizygotes, the severity of the clinical picture correlates positively with the person’s age6, but not with genotype - except where vessel tortuosity is present.15-16

### Table 1: Signs and symptoms associated with the classic form of Fabry’s disease

<table>
<thead>
<tr>
<th>Symptoms (most appears in early childhood and adolescence)</th>
<th>Signs</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Acroparesthesia (numbness, tingling of the extremities)</td>
<td>• Angiokeratomas (bathing trunk area, umbilicus, oral mucosa, fingers, thorax)</td>
</tr>
<tr>
<td>• Joint and abdominal pain</td>
<td>• Ocular manifestations</td>
</tr>
<tr>
<td>• Hypohydrosis</td>
<td>• Facial minor dysmorphic features</td>
</tr>
<tr>
<td>• Fever</td>
<td>• Renal dysfunction</td>
</tr>
<tr>
<td>• Heat/exercice intolerance</td>
<td>• Cardiac complications</td>
</tr>
<tr>
<td>• Hearing loss and vertigo</td>
<td>• Cerebrovascular disorders (TIA, strokes)</td>
</tr>
</tbody>
</table>

### Ocular manifestations related to Fabry disease

Ocular manifestations can be identified very early in childhood, by age three15 or even earlier.17 These typically occur at the same time as systemic symptoms appear, especially in hemizygous patients.

Classic ocular manifestations include vortex pigmentation of the cornea (verticillata), lens opacities (anterior whitish opacities and posterior subcapsular cataract), conjunctival vessel anomalies (tortuosity and micro-aneurysms) and retinal vessel tortuositites (Figures 1 to 4),13 although such manifestations do not usually impair vision but create visual symptoms such as photophobia.

### Systemic treatment

For decades, therapy was symptomatic. Since 2001 (Europe) and 2003 (USA), enzyme replacement therapy (ERT) has become an available option in the causal treatment, and been shown to be clinically beneficial.18 This treatment helps to mitigate signs and symptoms of the disease (Table 2) and would potentially reduce the Gb, deposition that leads to irreversible organ damage.3,19-20 ERT also provides an improvement in quality of life.20 In addition to ERT, the most recent pharmacological approach uses genetic therapy or chemically-induced pharmacological chaperones 1-deoxygalactonojirimycin (DGJ) and galactose to stabilize the human α-GAL glycoprotein and consequently to increase enzyme activity within lyzozomes.21

Without treatment, male Fabry patients usually die 20 years earlier than the general male population, due to renal failure, progressive cardiomyopathy and/or cerebrovascular events.22 Women also have a shorter life expectancy of 15 years compared with the general population.22
One way to reverse this natural course is to implement efficient screening strategies to identify suspects early in the disease process, and to refer them to Fabry specialists in a timely manner. Such a screening strategy can start with the involvement of those who see these potential patients on a daily basis. Considering that ocular manifestations are among the first to appear and the easiest to assess, eye care professionals should be targeted as key players.

**Methods**

**Defining the model**

In 2009, a collaborative pattern for the ocular follow-up of diagnosed Fabry patients, under the requirements of the Canadian Fabry Disease Initiative, was established between Université de Montréal, École d’optométrie (EOUM), and the genetic center of one of its university hospital (Hôpital du Sacré-Cœur de Montréal), treating most of the Fabry patients in the province. The first step in responding to CFDI’s request was to review their requirements for oculo-visual examination and follow-up (time, equipment, standards, etc.). Secondly, a faculty member (LM) from EOUM was designated to take charge of the project, based on expertise in anterior segment. The third step was to secure a formal referral pathway that would function reciprocally between EOUM and the CFDI research team (under Dr. Bichet, of Hôpital du Sacré-Cœur de Montréal). The fourth step was then developed. It involved recruiting practising optometrists to screen for Fabry patients on a large scale. Several continuing education lectures were conducted across the province. During these events, emphasis was made on the clinical course of the disease (age of onset and natural evolution), the systemic symptoms and ocular manifestations related to Fabry and the proposed screening referral pathway. Written documentation (manuscripts, posters with photos) and DVDs about Fabry and other lysosomal storage disorders were also provided as reminders for in-office use.

Under the developed model, a patient presenting with specific ocular manifestations and/or symptoms of Fabry is considered a suspect. The situation is explained to the patient who is offered to be

### Table 2: Summary of the effects of enzyme replacement therapy (ERT) on Fabry’s patients

<table>
<thead>
<tr>
<th>Symptoms</th>
<th>ERT outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Gastrointestinal disturbance/abdominal pain</td>
<td>• Alleviated</td>
</tr>
<tr>
<td>• Renal function (overall)</td>
<td>• Stabilize or slow the decline</td>
</tr>
<tr>
<td>• Proteinuria</td>
<td>• Not improved</td>
</tr>
<tr>
<td>• Cardiac complications</td>
<td>• Beneficial: reduction in left ventricular mass</td>
</tr>
<tr>
<td>• Hearing</td>
<td>(if hypertrophy present before tx)</td>
</tr>
<tr>
<td>• Cerebrovascular events</td>
<td>• Stabilize or improve (enzyme alpha)</td>
</tr>
<tr>
<td></td>
<td>• No improvement if tx started after event</td>
</tr>
<tr>
<td></td>
<td>• Unclear to prevent stroke on long term</td>
</tr>
</tbody>
</table>
were confirmed at EOUM’s vision clinic for confirmation and further assessment. If the clinical findings prove relevant (presence of corneal pigmentation and at least manifestation of one systemic symptom), or in the case of a positive genetic background, a urine sample collected on filter paper is sent to the CHUS Expertise Centre in Clinical Mass Spectrometry (Dr. Auray-Blais Waters’ laboratory) for tandem mass spectrometry analysis for Gb3 levels (see below). In the event of patients living outside of Montreal area, urinary test kits (filter papers and request forms) are sent to local optometrists to be administered. Based on clinical and lab results, if enough suspicion points to Fabry (cornea verticillata, systemic symptoms and positive urinary test result for Gb3 accumulation), a referral is made to the nearest genetic centre for complete DNA testing and follow-up.

Ocular data collected from confirmed patients are part of the CFDI registry and constitute the basis of our current longitudinal study on ocular manifestations related to Fabry.

**Urinary biomarker testing**

A method for screening high-risk individuals was developed to detect both male and female Fabry disease patients. The methodology is reliable, efficient and specific, if done after the age of 6 years. It is based on an analysis of urinary Gb3 using tandem mass spectrometry, in a laboratory in Sherbrooke, QC. Urinary excretion of Gb3 is normalized to creatinine and patients are always age-matched to controls. Nevertheless, it must be taken into account that patients with cardiac variant mutations who have residual enzyme activity do not excrete excessive amounts of Gb3 in their urine.

**Results**

**Continuous education**

During the first 16 months of the project, 750 Quebec optometrists (out of 1,300 – 57.6%) participated in continuing education events, during evening and weekends.

**Patient referrals**

The first examination of a patient referred to the EOUM clinic was made in September 2009. Since that time, 10 patients have been seen, (Table 3), referred by optometrists who attended continuing education events.

The urinary test was performed on every suspect (6 out of 10) who showed signs (3 suspects) or symptoms (3 suspects) that suggested Fabry. This test

**Table 3**: Demographics of the referred patients to U de M vision clinic for screening

<table>
<thead>
<tr>
<th>Patient Sent by ODs</th>
<th>Sex</th>
<th>Age</th>
<th>Corneal pigmentation seen</th>
<th>Positive Case History (Systemic symptoms)</th>
<th>Urinary test ordered</th>
<th>Patient sent for DNA testing</th>
<th>Fabry confirmed</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>F</td>
<td>18</td>
<td>Yes–typical*</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>2</td>
<td>M</td>
<td>22</td>
<td>Yes (1)</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>3</td>
<td>†</td>
<td>34</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>4</td>
<td>F</td>
<td>51</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>5</td>
<td>M</td>
<td>39</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>6</td>
<td>M</td>
<td>23</td>
<td>Yes–typical</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>7</td>
<td>F</td>
<td>62</td>
<td>Yes (2)</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>8</td>
<td>F</td>
<td>51</td>
<td>Yes-atypical</td>
<td>Yes(a)</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>9</td>
<td>M</td>
<td>46</td>
<td>Yes (3)</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>10</td>
<td>†</td>
<td>35</td>
<td>Yes (4)</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>

- Typical means bilateral pigmentation, verticillata type, asymetrical
- Hudson-Stahli type of pigmentation– monocular-Not related to Fabry
- Pigmentation secondary to amiodarone – symetrical OU
- Pigmentation secondary to chloroquine
- Pigmentation secondary to scarring
- Relatives affected (cousins)
allowed for the detection of 2 out of 3 suspects who showed corneal pigmentation and symptoms. The discovery of these patients led to cascade screening and the detection of other Fabry patients (5) in the family, including two young homozygous patients five and seven years old, respectively. The last patient with corneal signs had a negative result on the urinary test. However, because of her positive family background for the disease (cousins were known Fabry patients), she was referred for DNA testing which was also negative.

In addition to these patients, four additional suspects living in rural areas were co-managed with their local optometrists. Urinary tests were negative for the disease and consequently they were not referred for DNA testing.

Aside from these 19 individuals (10 suspects seen + 5 relatives + 4 co-managed patients), two other patients who had been lost to follow-up for many years were brought back under the care of either Dr. Bichet's team or a local geneticist in Quebec City, following an optometric examination.

In total, 7 new patients (2 + 5 relatives) were diagnosed as a direct consequence of the screening model we have established, and 2 more were brought back under appropriate care for their systemic condition.

**Discussion**

In the past, several strategies have been proposed to increase screening for patients with rare diseases. These strategies have included developing and launching an interactive public website to inform and communicate with the public and with medical professionals; a mail-based survey of all primary care physicians of a known area; systematic targeted screening of the relatives of patients already diagnosed, and a newborn/population-based urine testing program should biomarkers be detected. For Fabry disease in particular, very few other initiatives have been implemented worldwide to increase the screening rate for patients. Some initiatives have attempted to look at specific groups of patients considered to be "at risk" based on their signs or symptoms. For example, studies have been conducted among hemodialysis patients, with a positive-identification success rate of 0.2 to 1.2%. A similar approach involving patients presenting with cryptogenic stroke or unexplained left ventricular hypertrophy have yielded a 3-6% success rate.

In Argentina, blood sampled on filter paper (dry blood testing) from patients with signs and symptoms has been systematically sent for analysis, with a yield of detection of 4.96%. Using this approach, 70 patients were found positive for the disease within 2.5 years of implementation of the protocol. One other experiment was conducted in Germany in 2003: 615 ophthalmologists were recruited and 125,908 patients were examined. Out of these, 44 subjects (3.5%) were suspects and 21 (1.75%) were confirmed as Fabry patients. This result was not considered as successful as other strategies, due to the time and effort needed to screen a small number of patients.

As mentioned previously, in order to increase the overall positive results from screening strategies, it is preferable to develop a targeted protocol and to constitute an interdisciplinary group to identify confirmed patients. In our case, several aspects of other screening programs were put into place with the uniqueness to include optometry in the multidisciplinary team and to be able to rely on biomarker testing at an affordable cost, compared to the higher cost of DNA testing. Referral patterns were set up considering available resources.

The innovative feature of our approach was to involve optometrists, on a large scale, as team members for the screening effort. In North America, optometrists represent the most accessible resources in primary eye care. In the province of Quebec specifically, 1,300 optometrists see 30-35% of the population every year. This high level of population penetration represents a unique opportunity for screening a large population for Fabry disease.
Limitations of our analysis
We cannot estimate our rate of success as defined by the percentage of patients screened based on the total number of patients who consulted an optometrist who attended a continuing education event. We cannot estimate the male/female ratio of patients who consulted with a trained optometrist during this 16 months period of time. It is therefore difficult to compare our data with the German experiment or other screening projects.

On the other hand we can roughly approximate the value of the model. Since we know that those who attended continuing education events represent 60% of Quebec optometrists, we can assume that they have seen as many as 2.4 million people, in theory, over the last 16 months (3 million/year/total ODs x 60% x 1.33 years). Considering that the prevalence of Fabry is 1/200,000 in the general population we can estimate that 12 Fabry patients (2.4 million x 1/200,000) should have been seen. The screening process helped to identify 7 individuals who were not diagnosed, priorly, and 2 that were lost to follow-up. In such a perspective, 9 of 12 potential patients were identified. This gives an idea of the value of the screening effort.

On the other hand, most of the patients referred to EOUM for confirmation of the disease were not found positive for Fabry, based on the lack of symptoms or due to other corneal pigmentation causes. This suggests that the educational process can be improved, or repeated. Also, sensitivity of the screening could be improved by modifying the referral criterion to only include patients with corneal deposits (unexplained by other commonly known drugs or other common causes of corneal pigmentation) and the presence of systemic symptoms or a family history.

Updated Data
The continuous education seminars were also conducted across Canada. As of December 2012, 18 other patients and relatives were found by optometrists and confirmed as Fabry patients. This brings the total of patient screened up to 25 in the last 2.5 years.

Conclusion
Overall, a year and a half after its implementation, this screening program met its goals, as it involved optometry and helped uncover new Fabry patients in Quebec.

This screening program was effective because it was based on a defined protocol, a multidisciplinary approach and a major effort to provide up-to-date information to optometrists as primary care providers.

Our ability to rely on an accessible resource, everywhere in the territory, helped immeasurably. In this sense, optometrists should be considered key players in the development of any large-scale screening program for rare diseases involving ocular manifestations.

References


