Reading ability and quality of life in Stargardt disease

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ABSTRACT

Purpose: To study the reading performance of patients with Stargardt disease (STGD) and the relationship between clinical vision measurements and vision-related quality of life (VRQOL).

Methods: We studied both eyes of 16 patients with STGD. Each patient was examined for best-corrected visual acuity (Early Treatment Diabetic Retinopathy Study [ETDRS]), reading ability (MNREAD and REX charts), contrast sensitivity (Pelli-Robson charts), fixation study (MP1 micropimeter), and VRQOL (25-item National Eye Institute Visual Function Questionnaire [NEI VFQ-25]). The correlation pattern among these variables was examined and an exploratory factor analysis was used to investigate dimensionality of both visual function and VRQOL.

Results: Mean ETDRS visual acuity was about 20/160 (0.9 logMAR). All studied psychophysical measures were highly or moderately correlated with MNREAD reading speed (p<0.05 level). A similar correlation was found between psychophysical measures and VRQOL, which was higher for MNREAD measures of acuity (r = -0.75) and speed (r = 0.74). Accordingly, exploratory factor analysis suggested that a single latent dimension explained most of the variance of vision psychophysical measures as well as of VRQOL.

Conclusions: We propose that reading ability should be assessed in patients with STGD, since we found that both MNREAD reading speed and visual acuity are strong determinants of quality of life. The observed relation between reading ability and VRQOL in STGD suggests that in these patients appropriate low vision rehabilitation can improve both reading performance and consequently VRQOL. Finally, our data support the use of reading speed and visual acuity as important outcome measures for monitoring STGD progression.

Keywords: Low vision, Reading ability, Stargardt disease

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**Psychophysical testing**

Vision function testing was performed for each eye separately after optimal refraction and appropriate testing distance correction for each task.

Visual acuity was measured with Early Treatment Diabetic Retinopathy Study (ETDRS) charts at 2 meters and recorded as Snellen fraction and logMAR (7). Contrast sensitivity (CS) was measured with Pelli-Robson charts at 1 meter and recorded as Log10 CS (8). Reading ability was measured with the Italian version of the MNREAD charts at 20 cm to obtain maximum reading speed (MRS) (log10 words/minute), reading acuity, and critical print size, both recorded as logMAR, according to methods previously described (9). The effect of contrast on reading performance was defined by means of REX charts as reading contrast sensitivity (RCS), recorded as Log10 RCS (10).

**MP1 microperimetric testing**

The MP1 microperimeter (Nidek Technologies Inc., Padova, Italy) was used to study the location and stability of fixation and macular sensitivity. To study the sites of the preferred retinal locus (PRL), the patient’s task was to fixate a red circle (2° in diameter) and to maintain fixation on the center of this target for 30 seconds. The nontested eye was occluded throughout the procedure. This fixation test was followed by static perimetry using MP1. The sensitivity of the central visual field was tested with a 10° macula program during which white test lights (stimulus size Goldmann III, duration 200 ms) were presented on a dim white background (1.27 cd/m²) using a 4-2-1 threshold strategy.

The location of each subject’s PRL was referred to the fovea and the distance and direction in millimeters was converted into degrees according to strategies already described (11, 12). Fixation stability was previously defined in terms of the percentage of fixation points that fell within a 2° and 4° diameter circle during the visual field test. Fixation was classified as stable if more than 75% of the recorded fixation points fell within a 2° diameter circle and unstable if less than 75% fell within a 2° diameter circle but more than 75% fell within a 4° diameter circle, unstable if less than 75% fell within a 4° diameter circle and unstable if less than 75% fell within a 4° diameter circle. We did not record fixation stability as the bivariate contour ellipse area, since our study was assessed using Pearson correlation. Cronbach alpha was used to assess internal consistency of VRQOL subscales’ score. Analyses were conducted using Stata 13 software (StataCorp, College Station, TX, USA), accounting for correlated data within each individual since both eyes were considered in analyses, given the symmetry of visual damage in these patients.

**Statistical methods**

The correlation pattern among the variables included in our study was assessed using Pearson correlation. Cronbach alpha was used to assess internal consistency of VRQOL subscales’ score. Analyses were conducted using Stata 13 software (StataCorp, College Station, TX, USA), accounting for correlated data within each individual since both eyes were considered in analyses, given the symmetry of visual damage in these patients.

**Results**

We included 16 patients with STGD (mean age 34 years, SD 13 years, range 12-63 years).

Ten patients were female (63%) and 6 were male (38%). All patients were native Italian speakers with 10 or more years of education in 14 subjects and 5 or more in the 2 youngest patients. All patients carried 2 mutations of the ABCA4 gene. Sequence variants detected in our series are outlined in Table I. For every proband, our laboratory checked the parents and other relatives to evaluate the segregation of the ABCA4 alleles in the different members of the family, to be sure that the sequence variants lie on 2 separate alleles. We studied 31 eyes overall because one eye with visual acuity of 20/800 (1.6 logMAR) was unable to read and was excluded from analyses. When expressed as Snellen fraction, mean better eye ETDRS overall visual acuity was about 20/160 (0.84 logMAR). Electroretinogram responses could be classified as group 1 in 24 eyes, as group 2 in 6 eyes, and as group 3 in 2 eyes (6).

Electroretinogram responses were poorly related to age and genotype. The mean age of patients was 31 years in group 1, 43 years in group 2, and in group 3 there was 140-year-old patient. ABCA4 mutations of group 1 consisted of 19 missense mutations, 2 splicing mutations, and 4 nonsense mutations. Group 2 consisted of 4 missense mutations, 1 splicing mutation, and 1 deletion. The group 3 patient showed 2 missense mutations. Statistical analysis of possible differences among the 3 groups was not feasible because of the different sample sizes.

Table II reports the mean values of all variables referred to the best and the worst eye: as expected, mean values of relevant vision psychophysical variables were similar between the 2 eyes. Moreover, monocular reading ability was similar
gave instructions verbally, and provided assistance when required. The completed questionnaires were reviewed for missing data by the research staff.

The VFQ-25 comprises 25 items that require the patient to assess the levels of difficulty of particular visual symptoms or day-to-day activities. Each item is assigned to one of the following 12 subscales: general health, general vision, ocular pain, near activities, distance activities, social functioning, mental health, role difficulties, dependency, driving, color vision, and peripheral vision. The subscales are scored from 0 to 100 points, where 100 indicates the highest possible function or minimal subjective impairment. The VFQ-25 composite score is calculated as the unweighted average response to all items, excluding the questions on general health and ocular pain, which is not a consequence of STGD. We did not obtain scores for driving, since no subject had a driving license.
in the better and worse eye for all the subjects included in the study, suggesting bilateral and symmetrical functional impairment despite some differences between eyes regarding fundus appearance.

Using microperimetry, we noted that all but 3 eyes examined had an absolute scotoma; the 3 remaining eyes had a relative scotoma. For 31 eyes of 16 patients, the PRL was eccentric in 26 eyes and foveal in 5 eyes. The PRL was superior to the fovea in 18 eyes (straight up in 14 eyes, upper left scotoma in 3, and upper right in 1), nasal in 6, and inferior in 2. Fixation was unstable in 13 eyes, relatively unstable in 15 eyes, and stable in 3 eyes. Central 2 degrees and central 4 degrees mean values of fixation stability were 40% (SD 23%) and 73% (SD 24%), respectively.

Figure 1 shows NEI VFQ-25 quality of life scores. Median specific subscales scores were between 45 and 56, except for the social functioning and color vision subscales, which reached values of 71 and 100, respectively.

### TABLE I - Sequence variants detected in our series

<table>
<thead>
<tr>
<th>Patient ID</th>
<th>Mutation 1</th>
<th>Mutation 2</th>
<th>Mutation 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>c.2791G&gt;A (p.Val931Met)</td>
<td>c.1622T&gt;C (p.Leu541Pro)</td>
<td>c.3113C&gt;T (p.Ala1038Val)</td>
</tr>
<tr>
<td>4</td>
<td>c.2461T&gt;A (p.Trp821Arg)</td>
<td>c.3113C&gt;T (p.Ala1038Val)</td>
<td>c.343_381delinsGGACAA (p.Asn115_Thr127delinsGlyGln)</td>
</tr>
<tr>
<td>5</td>
<td>c.2461T&gt;A (p.Trp821Arg)</td>
<td>c.3323G&gt;A (p.Arg1108His)</td>
<td>c.4297G&gt;A (p.Val1433Ile)</td>
</tr>
<tr>
<td>6</td>
<td>c.288C&gt;A (p.Asn96Lys)</td>
<td>c.2933G&gt;A (p.Gly978Asp)</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>c.2791G&gt;A (p.Val931Met)</td>
<td>c.3322C&gt;T (p.Arg1108Cys)</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>c.5882G&gt;A (p.Gly1961Glu)</td>
<td>c.5018 + 2T&gt;C (p.?</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>c.5018 + 2T&gt;C (p.?</td>
<td>c.5898 + 5del (p.?</td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>c.4667 + 1G&gt;A (p.?</td>
<td>c.5512C&gt;A (p.His1838Asn)</td>
<td></td>
</tr>
<tr>
<td>15</td>
<td>c.4437G&gt;A (p.Trp1479X)</td>
<td>c.6419T&gt;A (p.Leu2140Gln)</td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>c.4437G&gt;A (p.Trp1479X)</td>
<td>c.6419T&gt;A (p.Leu2140Gln)</td>
<td></td>
</tr>
</tbody>
</table>

* Nonsense mutation.

### TABLE II - Mean (SD) of all psychophysical measures considered in the study, showing similar mean values between the better and worse eye

<table>
<thead>
<tr>
<th>Test (measure)</th>
<th>Better eye</th>
<th>Worse eye</th>
</tr>
</thead>
<tbody>
<tr>
<td>ETDRS visual acuity, logMAR (Snellen equivalent)</td>
<td>0.84 (0.08) (20/160†)</td>
<td>0.90 (0.1) (20/160‡)</td>
</tr>
<tr>
<td>Pelli-Robson log (contrast sensitivity)</td>
<td>19.2 (10.4)</td>
<td>16.9 (10.0)</td>
</tr>
<tr>
<td>MNREAD reading acuity logMAR</td>
<td>1.02 (0.29)</td>
<td>1.11 (0.30)</td>
</tr>
<tr>
<td>MNREAD critical print size logMAR</td>
<td>1.15 (0.30)</td>
<td>1.26 (0.26)</td>
</tr>
<tr>
<td>MNREAD reading speed (logWPM)</td>
<td>1.71 (0.37)</td>
<td>1.56 (0.34)</td>
</tr>
<tr>
<td>REX mean reading speed (logWPM)</td>
<td>1.65 (0.28)</td>
<td>1.50 (0.41)</td>
</tr>
<tr>
<td>REX reading contrast threshold (Log10 sensitivity)</td>
<td>0.88 (0.47)</td>
<td>0.81 (0.43)</td>
</tr>
<tr>
<td>MP1 mean sensitivity, dB</td>
<td>10.8 (5.3)</td>
<td>8.8 (5.7)</td>
</tr>
<tr>
<td>MP1 mean eccentricity</td>
<td>8.8 (6.2)</td>
<td>9.7 (6.8)</td>
</tr>
<tr>
<td>MP1 fixation stability within 2° (%)</td>
<td>0.80 (0.21)</td>
<td>0.66 (0.24)</td>
</tr>
<tr>
<td>MP1 Fixation stability within 4° (%)</td>
<td>0.49 (0.23)</td>
<td>0.32 (0.21)</td>
</tr>
</tbody>
</table>

ETDRS = Early Treatment Diabetic Retinopathy Study.
Correlation with maximum reading speed

Table III shows Pearson correlation between MNREAD MRS and the variables collected by means of other tests (ETDRS charts, Pelli-Robson chart, REX reading test, and MP1 microperimetry), all of which were statistically significant at a p<0.05 level. The MNREAD reading acuity yielded the highest correlation with MRS (r = -0.84), which is not surprising since they are related reading variables obtained by means of the same test. After reading acuity, ETDRS visual acuity (r = 0.78) and REX reading contrast threshold (r = 0.77) yielded the strongest correlation with MRS when compared to the remaining visual function variables (p>0.05 for all comparisons).

Correlation between quality of life subscales

Pearson correlation of all NEI VFQ-25 subscales with the reduced overall score, i.e., the total score excluding general health, was very high except for general vision (r = 0.72), peripheral vision (r = 0.76), and color vision (r = 0.55). The correlation of other subscales with the overall score was very high and ranged from 0.80 for near vision to 0.96 for mental health. Cronbach alpha was 0.93 when the subscales' scores were assessed. For this reason, the relationship of psychophysical measures with VRQOL was only assessed using the overall reduced score as a summary measure.

Association of psychophysical measures with quality of life

Table IV shows Pearson correlation between VRQOL overall score and visual psychophysical measures, with individual data presented as scatterplots for relevant variables in Figure 2. Most variables yielded high and similar Pearson correlation coefficients, particularly for MNREAD measures of acuity and speed. ETDRS = Early Treatment Diabetic Retinopathy Study; VRQOL = vision-related quality of life.

Conclusion

To our knowledge, this is the first study relating visual function to reading performance and VRQOL in a series of patients with STGD, which is relatively large given that STGD is a rare disease.

We found that macular damage in STGD causes a similar impairment of several psychophysical visual functions, including visual acuity, contrast sensitivity, and reading performance. Moreover, MRS was very strongly correlated to ETDRS acuity, differently from what is found in age-related macular degeneration (AMD) (15), where the correlation is
only moderate. Contrarily to AMD, STGD usually affects young patients, and we can speculate that they can exploit for reading function a normal midperipheral retina beyond the macular atrophy. Moreover, young patients are more able to cope with visual dysfunction and adopt reading strategies, due to higher learning skills for better brain plasticity.

Reading ability is also different from that in patients with retinitis pigmentosa, in whom reading difficulties are also determined by visual field constriction and reduced contrast sensitivity, despite the fact that visual acuity can be elevated until more advanced stages of the disease (9). Another study found good correlation between visual acuity and reading speed in patients with retinitis pigmentosa (16), but reading ability was evaluated by recalling aloud some timed sentences previously read silently and not assessed by means a specific reading test.
Using correlation analysis, the NEI VFQ-25 questionnaire seemed to explore a unique vision-related VRQOL dimension in patients with STGD. In fact, these patients are almost exclusively affected by central vision loss, and they lack peripheral visual damage, as found in our patients with STGD in all but one case. The particularly high value of the color vision subscale score in our patients may be related to partially retained color discrimination, sufficient for daily activities. Furthermore, patients with STGD are usually examined at a young age and are generally healthy, in the absence of systemic comorbidities, which could have made the subjective perception of visual health difficult to disentangle from that of compromised general health, such as in older patients. On the contrary, in patients with AMD, VRQOL score could be less accurate in describing the impact of low vision on VRQOL, as individuals have other issues in addition to their poor eyesight (17).

Consistently with the predominant role of central vision loss as the main health problem, we found that any measure of central vision function is a good proxy of VRQOL in patients with STGD. The highest correlation with NEI VFQ-25 overall score was obtained by both reading acuity and speed. This is expected since reading ability is not only a psychophysical measure, but a key activity of daily living.

The VRQOL of patients with STGD can be inferred from routine clinical ophthalmic examination. In patients in whom distance visual acuity is affected in both eyes, we propose that reading ability be assessed, since we found that both MNRAD reading speed and acuity are the strongest determinants of VRQOL.

The observed relation between reading ability and VRQOL in STGD suggests that, in these patients, appropriate low vision rehabilitation can improve both reading performance and, consequently, VRQOL (18, 19).

Finally, our data support the use of reading speed and acuity as valid and important outcome measures for monitoring the progression of the disease and its potential response to innovative treatments (20).

Disclosures

Financial support: This study was partially funded by the Fondazione Cassa di Risparmio di Firenze.
Conflict of interest: Gianni Virgili shares the patent of the Italian version of the MNREAD charts with the University of Minnesota.

References