A comparison of the NEIVFQ25 and GQL-15 questionnaires in Nigerian glaucoma patients

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Aim: To compare two vision-specific quality of life (QOL) instruments – the disease-specific 15-item Glaucoma Quality of Life questionnaire (GQL-15) and the nonglaucoma-specific 25-item National Eye Institute Visual Function Questionnaire (NEIVFQ25).

Methods: The QOL of 132 glaucoma patients being managed in Lagos University Teaching Hospital and an equal number of controls matched for age and sex was assessed using two vision-specific instruments: GQL-15 and the NEIVFQ25. The categorization of the severity of glaucoma into mild, moderate, and severe disease was determined using the degree of visual field loss. The Statistical Package for the Social Sciences (version 15; SPSS Inc, Chicago, IL) software program was used for analyzing the data obtained. Spearman’s correlation coefficients were used to assess the correlation between the scores from the two questionnaires.

Results: Patients had the greatest difficulty with activities affected by glare and dark adaptation in the GQL-15. Driving and general vision were the factors most affected in the NEIVFQ25. The Spearman rho values showed strong correlations (rho: > 0.55) between the NEIVFQ25 and GQL-15 QOL scores for the total number of participants (rho: −0.83), and the mild (rho: −0.76), moderate (rho: −0.75), and severe (rho: −0.84) cases. There was a moderate correlation (rho: −0.38) for QOL scores of controls. Cronbach’s alpha was 0.94 for the GQL-15 and 0.93 for the NEIVFQ25, showing high internal consistency for both questionnaires.

Conclusion: The GQL-15 and the NEIVFQ25 questionnaires showed high internal consistency, correlated strongly with each other, and were reliable in the assessment of glaucoma patients in this study.

Keywords: primary open-angle glaucoma, quality of life questionnaires, GQL-15, NEIVFQ25

Introduction

The term “quality of life” (QOL) is the subjective perception of well-being and wholeness. It is a broad concept affected in a complex way by a person’s physical health, psychological state, level of independence, social relationships, and his or her relationship to the salient features of his or her environment. It reflects the difference between a person’s hopes and expectations and his or her present experience. Thus, QOL is influenced by the social, health, economic, and environmental conditions that affect human and social development.

The main goal of treatment in glaucoma patients is to improve clinical outcomes thereby improving the patient’s QOL. Measures of visual acuity (VA), intraocular pressures (IOP), and visual fields have been utilized over the years to assess treatment efficacy in glaucoma. The achievement of therapeutic goals based on clinical indicators...
Vision-related QOL instruments may be disease-specific or nonspecific. The nondisease-specific vision-related QOL instruments include: the National Eye Institute Visual Function Questionnaires (NEIVFQ51, NEIVFQ25), the 14-item Visual Functioning Index (VF-14), Assessment of Function Related to Vision (AFREV), the Activities of Daily Vision Scale (ADVS), and the Visual Activities Questionnaire (VAQ). Quality of life assessment using a nondisease-specific instrument facilitates the direct comparison of the QOL of patients with different eye problems.

Some specific QOL instruments have been developed for glaucoma, such as the Glaucoma Quality of Life 15 (GQL-15), the Symptom Impact Glaucoma Score (SIG), the Glaucoma Health Perceptions Index (GPHI), the Glaucoma Symptom Scale (GSS), and the Comparison of Ophthalmic Medication for Tolerability (COMTOL). These instruments are thought to be more accurate and specific for glaucoma with more visual field considerations.

The NEIVFQ25 is a questionnaire developed by the Research and Development Corporation. It is a shorter version of the longer questionnaire, which has 51 items (NEIVFQ51), with reduced administration time. Studies suggest that the NEIVFQ is a reliable tool for the evaluation of QOL in glaucoma patients. As it is a broader instrument used to assess all visual problems, newer and disease-specific tools can be compared to it.
was carried out on all participants; this examination included gonioscopy, dilated stereoscopic optic nerve head assessment at the slit-lamp with a Volks +78 diopter sphere (DS) lens, and standard achromatic perimetry using a full threshold central 30-2 strategy (Optifield Sinemed Inc, Benicia, CA).

Visual field defects were classified into mild, moderate, and severe visual field loss using mean deviation (MD) indices and a modification of the Hodapp–Parrish–Anderson classification27 in the more severely affected eye. Mild visual field loss was defined as a mean deviation of less than or equal to −6 decibels (dB), moderate visual field loss was defined as MD greater than −6 db but less than −12 dB, and severe visual field defect as MD greater than −12 dB.

**Eligibility criteria for cases**
Patients with POAG on medical therapy or those who had undergone trabeculectomy at least 3 months before the study were eligible. The purpose of the 3-month period was to ensure that the surgical experience was not recent and would not significantly influence QOL responses.

**Exclusion criteria for cases**
The presence of refractive errors greater than 5 DS or 2 diopters Cylinder (DCyl), visually significant cataracts (greater than Stage 2 LOCS III classification),28 or any other ocular pathology were criteria for exclusion from the study. Patients with secondary glaucoma or angle-closure glaucoma were also excluded. All patients who had undergone trabeculectomy, or any intraocular surgery, within the 3 months preceding the study were excluded from the study. Patients with mental or systemic illness, and hearing, cognitive or mobility defects were excluded from the study.

**Eligibility criteria for controls**
Age and sex-matched patients with refractive errors corrected to at least 6/9, normal optic discs and visual fields, and no underlying ocular pathology or systemic co-morbidities were recruited as controls. The degree of refractive error acceptable was less than 5 DS from emmetropia or less than 2 DCyl of astigmatism.

**Exclusion criteria for controls**
Patients with suspicious optic discs, abnormal visual fields, ocular hypertension (defined as IOP greater than 21 mmHg in either eye), visually significant lens opacities, high refractive errors, and corrected visual acuity less than 6/9 were excluded. Participants who had a history of glaucoma or unexplained blindness in a first-degree relative or who were being managed for systemic illness were also excluded from the study.

**Quality of life assessment**
Quality of life assessment was done using the interviewer-administered format of the NEIVFQ25 and the GQL-15 questionnaires. Two optional utility questions rating general health and vision on a linear scale from the NEIVFQ25 appendix were included in the questionnaire. The NEIVFQ25 was administered first, followed by the GQL-15, administered by the same interviewer in a single sitting. Participants were not informed that two different QOL instruments were being administered.

**Translation of QOL assessment questionnaires**
For respondents who were not literate in English, the NEIVFQ25 and the GQL-15 questionnaires were translated into Yoruba, Igbo, and Hausa, which are the three most prominent indigenous Nigerian languages. Care was taken to reduce any bias that could occur because of translation. Translation and back-translation methods were used by competent bilingual translators to ensure accuracy.

The scoring method designed by the Research and Development Corporation for the NEIVFQ25 was used. Individual items were scored. The minimum score was zero and the maximum score for each item was 100. A higher score signified a better QOL. The average score for each domain was calculated from the individual item scores. The summary NEIVFQ25 score was obtained by calculating the average of the domain scores with the exclusion of the general health domain score.

With the GQL-15, higher scores indicate poorer QOL. A scale of 0 to 5 was used to code the item level responses: 5 represented severe difficulty due to visual reasons, 1 indicated no difficulty with performing the activity, and 0 signified abstinence from activity for nonvisual reasons. Item response scores of the GQL-15 were added to obtain the summary scores.

To calculate the subscale scores for the four domains of the GQL-15, the item level responses were scored on a numerical interval scale ranging from 0, indicating no difficulty, to 100, indicating severe difficulty. The subscale score for each domain was calculated using an average of the scores generated for the component item-level responses. Poorer QOL and increasing difficulty with vision-related activities were associated with higher subscale scores.
A measure of the degree of difficulty in performing visual tasks outlined in the GQL-15 was depicted by corresponding visual performance. This score was obtained by first subtracting the subscale scores from the total possible score × 100 (1500). The product was thereafter converted to a percentage.

**Data entry and statistical analysis**

Data was entered into Microsoft Excel for Windows XP Professional and was later uploaded and analyzed using the Statistical Package for the Social Sciences version 15 (SPSS, Chicago III). Cronbach’s alpha was used to assess the internal consistency of the questionnaires. The range, mean, median, and standard deviation of QOL scores for the two questionnaires were calculated and compared for the subgroups of participants. Spearman’s rank correlation coefficients were used to assess the correlation between the GQL-15 and the NEIVFQ25. A modest or weak correlation was defined as rho values < 0.31, values ranging from 0.32–0.55 were defined as moderate correlation, and rho values > 0.55 showed strong correlation. *P*-values equal to or less than 0.05 were considered statistically significant.

**Results**

Administration of the NEIVFQ25 questionnaire took about 12 minutes, on average, while administration of the GQL-15 lasted about 5 minutes. The Cronbach’s alpha for the NEIVFQ25 was 0.93, and 0.94 for the GQL-15. Eight participants (3%) responded to the translated versions of the NEIVFQ25 and GQL-15 questionnaires. Mild glaucoma patients had a mean age of 52.81 years. For patients with moderate and severe glaucoma, the mean age was 59.11 and 62.67 years, respectively. Most of these patients had bilateral disease (90.9%), and binocular visual field defects were demonstrated in 93.94% of the patients.

**NEIVFQ25 and GQL-15 scores in cases and controls**

The mean QOL score using the NEIVFQ25 was higher in controls, 96.7 (SD ± 2.34), than in cases, 85.2 (SD ± 16.07). Using the GQL-15, the mean QOL for controls was 15.75 (SD ± 1.85) and 24.07 (SD ± 12.4) for cases. The corresponding visual performance was 84.54% (SD ± 21.65) for cases and 98.75% (SD ± 3.02) for controls (see Figure 1). All these differences had a *P*-value of <0.001 and thus were statistically significant. Females, younger participants, and participants who had received tertiary education reported better QOL with both the GQL-15 and the NEIVFQ25.

**NEIVFQ25 and GQL-15 scores in varying degrees of disease severity**

The average QOL scores in patients with moderate and severe glaucoma were significantly different (*P* < 0.001). This was also the case for participants with severe and mild glaucoma. This difference was noted for the two QOL questionnaires. Both questionnaires, however, did not detect a significant difference in the QOL scores of patients with mild and moderate glaucoma. A comparison of the GQL-15 QOL scores of patients with mild and moderate glaucoma using the Mann–Whitney *U* test did not show a statistically significant difference (*P* = 0.25). A comparison of the NEIVFQ25 QOL scores for mild and moderate glaucoma was also not statistically significant (*P* = 0.92). The decrease in QOL from controls (n = 132) across mild (n = 44), moderate (n = 44), and severe (n = 44) glaucoma cases for the NEIVFQ25 and the GQL-15 are depicted in the form of line plots in Figures 2 and 3.

**NEIVFQ25, GQL-15 scores and correlation with visual function**

The NEIVFQ25 scores correlated best with the mean logarithm of the minimum angle of resolution (logMAR) Visual Acuity (VA) better eye (rho: 0.54, *P* < 0.001), mean contrast sensitivity score (rho: 0.47, *P* < 0.0010 and mean logMAR VA worse eye (rho: 0.46, *P* < 0.001). The GQL-15 scores correlated best with mean deviation in both eyes (MD OU), (rho: 0.43, *P* < 0.001), mean contrast sensitivity score (rho: 0.43, *P* < 0.001), and mean log MAR VA worse eye (rho: −0.42, *P* < 0.001). Only a modest or weak correlation existed between the pattern standard deviation of
patients and both the NEIVFQ25 (rho: $-0.13$, $P = 0.16$) and GQL-15 (rho: $0.13$, $P = 0.15$).

**Correlation between NEIVFQ25 and GQL-15 scores**

Correlation between the NEIVFQ25 and GQL-15 QOL scores was assessed using Spearman’s correlation coefficient. The scores had strong correlations when the scores for total number of participants ($-0.753$), total number of cases ($-0.827$), and mild ($-0.762$), moderate ($-0.748$), and severe cases ($-0.836$) were compared. There was a moderate correlation for QOL scores of controls with the Spearman’s correlation coefficient ($rho = -0.378$).

**Correlation between NEIVFQ25 and GQL-15 subscales/domains**

Using the GQL-15, the glare and dark adaptation subscale was the most affected, while the NEIVFQ25 revealed that the general vision and driving subscales were the most affected. Figure 4 clearly shows that participants had the greatest difficulty with tasks affected by glare and dark adaptation.

There were statistically significant correlations between the subscales of the NEIVFQ25 and the GQL-15; these results are shown in Table 1. The outdoor mobility domain of the GQL-15 had the highest correlation with the peripheral vision subscale of the NEIVFQ25 ($rho: -0.620$, $P < 0.001$). The glare and dark adaptation domain of the GQL-15 correlated best with the distance activities subscale of the NEIVFQ25 ($rho: -0.656$, $P < 0.001$). The central and near vision domain of the GQL-15 correlated best with the near activities subscale of the NEIVFQ25 ($rho: -0.731$, $P < 0.001$). The peripheral vision domain of the GQL-15 correlated best with the distance activities subscale of the NEIVFQ25 ($rho: -0.709$, $P < 0.001$) but also had a strong correlation.
Table 1: Correlation between NEIVFQ25 items and GQL-15 domain items

<table>
<thead>
<tr>
<th>NEIVFQ25 items</th>
<th>Outdoor mobility</th>
<th>Glare dark adaptation</th>
<th>Central and near vision</th>
<th>Peripheral vision</th>
<th>Total GQL-15 score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gen health</td>
<td>−0.198**</td>
<td>−0.235**</td>
<td>−0.185**</td>
<td>−0.280**</td>
<td>−0.245**</td>
</tr>
<tr>
<td>Gen vision</td>
<td>−0.395**</td>
<td>−0.447**</td>
<td>−0.433**</td>
<td>−0.545**</td>
<td>−0.492**</td>
</tr>
<tr>
<td>Ocular pain</td>
<td>−0.199**</td>
<td>−0.325**</td>
<td>−0.305**</td>
<td>−0.264**</td>
<td>−0.348**</td>
</tr>
<tr>
<td>Near activities</td>
<td>−0.489**</td>
<td>−0.591**</td>
<td>−0.731**</td>
<td>−0.621**</td>
<td>−0.635**</td>
</tr>
<tr>
<td>Distance activities</td>
<td>−0.510**</td>
<td>−0.656**</td>
<td>−0.655**</td>
<td>−0.709**</td>
<td>−0.685**</td>
</tr>
<tr>
<td>Social functioning</td>
<td>−0.614**</td>
<td>−0.541**</td>
<td>−0.688**</td>
<td>−0.584**</td>
<td>−0.548**</td>
</tr>
<tr>
<td>Mental Health</td>
<td>−0.434**</td>
<td>−0.565**</td>
<td>−0.535**</td>
<td>−0.542**</td>
<td>−0.603**</td>
</tr>
<tr>
<td>Role difficulty</td>
<td>−0.614**</td>
<td>−0.508**</td>
<td>−0.570**</td>
<td>−0.565**</td>
<td>−0.577**</td>
</tr>
<tr>
<td>Dependency</td>
<td>−0.596**</td>
<td>−0.508**</td>
<td>−0.570**</td>
<td>−0.565**</td>
<td>−0.577**</td>
</tr>
<tr>
<td>Driving</td>
<td>−0.452**</td>
<td>−0.644**</td>
<td>−0.526**</td>
<td>−0.488**</td>
<td>−0.633**</td>
</tr>
<tr>
<td>Colour vision</td>
<td>−0.585**</td>
<td>−0.416**</td>
<td>−0.515**</td>
<td>−0.461**</td>
<td>−0.418**</td>
</tr>
<tr>
<td>Peripheral vision</td>
<td>−0.620**</td>
<td>−0.520**</td>
<td>−0.588**</td>
<td>−0.639**</td>
<td>−0.565**</td>
</tr>
<tr>
<td>Total NEIVFQ25 score</td>
<td>−0.451**</td>
<td>−0.707**</td>
<td>−0.615**</td>
<td>−0.668**</td>
<td>−0.753**</td>
</tr>
</tbody>
</table>

Notes: Spearman’s rho values. **Correlation significant at 0.01 level (2 tailed).
Abbreviations: NEIVFQ25, 25-item National Eye Institute Visual Functioning Questionnaire; GQL-15, 15-item Glaucoma Quality of Life questionnaire.

Discussion

The two questionnaires had good reliability indices. The high Cronbach’s alpha obtained for the NEIVFQ25 in this study (0.93) suggests internal consistency. This differs from the findings of Mangione et al11 in which a Cronbach’s alpha ranging from 0.71 to −0.85 was obtained. We obtained a Cronbach’s alpha of 0.94 for the GQL-15 which is only slightly higher than that obtained for the NEIVFQ25 (0.93) but is very close to that obtained by Nelson et al16 for the GQL-15 (0.95).

In this study, QOL worsened with increasing severity of glaucoma. This has been reported by other researchers. A combination of a generic health-related QOL tool and a vision-specific questionnaire was used in some of the studies, unlike in the present study, in which two vision-specific QOL questionnaires were used.

The NEIVFQ25 and the GQL-15 showed good group validity in that they were able to differentiate between cases and controls. They were also able to discriminate between some patients with different degrees of glaucoma severity. The NEIVFQ25 and GQL-15 QOL scores were significantly different for patients with mild and severe glaucoma as well as for patients with moderate and severe glaucoma. However, the QOL scores obtained for patients with mild glaucoma and moderate glaucoma did not differ significantly either for the NEIVFQ25 (P = 0.92) or the GQL-15 (P = 0.25).

In a study by Goldberg et al,29 mean GQL-15 QOL scores of mild, moderate, and severe glaucoma patients were significantly different. Nelson et al,16 on the other hand, found a statistically significant difference in the GQL-15 QOL scores of mild and severe glaucoma patients but did not find a statistically significant difference in scores of patients with mild and moderate glaucoma, as in the present study, or in those with moderate and severe glaucoma. The fact that there is no universally acceptable gold standard for categorizing disease severity in glaucoma could account for this.

The line plot in Figure 1 shows decreasing NEIVFQ25 scores with increasing severity of glaucoma, signifying worsening...
of QOL. This decrease is evident even between controls (n = 132) and cases of mild glaucoma (n = 44). The plot in Figure 2 shows increasing GQL-15 scores (worse QOL) with increasing severity of disease. There is also a clear decrease in QOL when mild cases are compared to controls.

The line plots clearly show that the slope of the plot for the QOL from mild to moderate is much less than the slope of the plot from control to mild or moderate to severe for both questionnaires.

This suggests that the belief that patients with mild glaucoma do not have a reduction in QOL until a visual threshold is reached is probably erroneous and corresponds with recent studies that suggest that QOL is reduced in glaucoma patients who are not even aware that they have the disease and in those who are in early stages of the disease. The Los Angeles Latino Eye Study findings suggest that there is no normal glaucoma patient. They found that QOL measured by the 25-item Visual Function Questionnaire (VFQ25) and the physical component of the Medical Outcomes Study Short-Form 12 (SF-12) was lower in participants with glaucoma compared to those without. This persisted in 75% (n = 160) of cases who did not know they had glaucoma when answering the questions, and therefore, the reduced QOL was clearly a disease effect.

This finding that QOL is reduced significantly in patients with mild glaucoma may also explain why the questionnaires used in this study (NEIVFQ25 and GQL-15) could not differentiate between moderate and mild glaucoma patients. This finding may not be a flaw or weakness in these questionnaires but perhaps suggests that the absence of a significant difference in the QOL of these two groups of patients is because the decrease in QOL had occurred earlier. This suggests that assessment of QOL should be done at the onset of the disease whether or not there is documentation of visual loss (preperimetric glaucoma).

**NEIVFQ25, GQL-15 scores and correlation with visual function**

Both questionnaires correlated only weakly with the PSD visual field index and had a moderate correlation with the MD OU and MD worse eye. The rho value for the correlation between the NEIVFQ25 and the MD worse eye was 0.38 (P < 0.001), and it was −0.38 (P < 0.001) for the GQL-15. For MD OU, the rho for the NEIVFQ25 was 0.41 (P < 0.001) while for the GQL-15, it was 0.43 (P < 0.001).

This shows that both questionnaires appeared to correlate nearly equally with the visual field indices in the present study. This differs from the finding of Pourjavan et al\(^\text{11}\) in which the NEIVFQ25 had a stronger correlation with visual field indices (rho: 0.61, 0.55) than the GQL-15, which had a moderate correlation only (rho: −0.47, 0.38).

**Correlation between NEIVFQ25 and GQL-15 scores**

There was a strong correlation between the overall scores of both questionnaires (rho: −0.753, P < 0.001) and both questionnaires showed statistically significant correlations between their subscales/domains. It is important to note that the outdoor mobility domain of the GQL-15 had the highest correlation with the peripheral vision subscale of the NEIVFQ25 (rho: −0.620, P < 0.001) and that the GQL-15 central and near vision domain correlated best with the NEIVFQ25 near activities subscale (rho: −0.731, P < 0.001). Similarly, the GQL-15 peripheral vision domain had a strong correlation (rho: −0.639, P < 0.001) with the NEIVFQ25 peripheral vision subscale. This suggests a high degree of correlation between similar domains or subscales of both instruments.

The NEIVFQ25 driving subscale had the highest correlation with the GQL-15 glare and dark adaptation domain. This is important, as the participants had the greatest difficulty carrying out tasks that were affected by glare and poor dark adaptation such as general vision tasks and driving. One can therefore infer that glare and poor adaptation to darkness may account for some of the difficulties glaucoma patients experience with driving. Therefore, its assessment should be included in the clinical management of glaucoma patients and steps taken to reduce optic glare in the patient’s work and home environments and by environmental modification.

Glare and dark adaptation subscale scores of the GQL-15 increased with increasing severity of disease (mild: 11.46 ± 15.12; moderate: 11.98 ± 18.62; severe: 36.18 ± 28.74) implying increasing difficulty with visual tasks affected by glare and dark adaptation with progression of glaucoma. The glare and dark adaptation score had a moderate correlation with the MD (rho: 0.374).

The fact that there were an equal number of controls and cases as well as an equal number of cases with mild, moderate, or severe glaucoma strengthened the design of this study. Though only eight participants (3%) responded to the translated version of the GQL-15 and the NEIVFQ25 questionnaires, which have not been validated, the entire data was re-analyzed excluding these eight participants. A comparison showed no significant difference in the results after excluding these participants so the study results are presented without excluding them. The authors plan to validate these versions in a subsequent multicenter study.
Conclusion

The NEIVFQ25 and the GQL-15 both appeared reliable in the assessment of QOL in this study. They showed similarities in their correlation with visual function, visual field indices, and their ability to differentiate between patients with varying severity of glaucoma. The NEIVFQ25, however, is a broader instrument better evaluating psychosocial wellbeing than the GQL-15, while the GQL-15 is a shorter, simpler instrument, which could be easily administered in clinical practice. More studies may further assess the QOL of Nigerian glaucoma patients and the reliability and validity of available QOL instruments in the Nigerian setting. However, it should be noted that, though useful and invaluable, QOL assessment is complementary to clinical evaluation of glaucoma patients and cannot replace it.

Disclosure

The authors report no conflict of interest in this work.

References

NEIVFQ25 and GQL-15 in POAG patients in Lagos, Nigeria