Threshold Levels of Visual Field and Acuity Loss Related to Significant Decreases in the Quality of Life and Emotional States of Patients with Retinitis Pigmentosa

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Retinitis pigmentosa · Quality of life · Emotional state · Visual field · Visual acuity

Abstract

Introduction: Retinitis pigmentosa (RP) is an inherited retinal disorder, characterized by photoreceptor degeneration inducing progressive vision loss. This study evaluates its impact on quality of life (QOL) and emotional states of patients affected by RP. Methods: A cross-sectional study was conducted on 60 RP patients diagnosed with rod-cone dystrophy and on 20 control subjects. The RP population has been divided into 3 groups according to visual field (VF) and visual acuity (VA) impairments. Concurrently, scores of self-reported QOL (25-item National Eye Institute Visual Functioning Questionnaire) and of the Hospital Anxiety and Depression Scale for anxiety/depression assessments were collected. Results: For the QOL composite score, we noticed consistent differences between all VF and VA affected groups and their control group. We also found significant differences between both the most affected VF group (VF1: ØVF <20°) and VA group (VA1: VA <0.3) compared to other VF and VA groups. For anxiety/depression scores, consistent differences have been found between the control group and VF1 and VA1, respectively. Conclusions: This work determines that, for RP patients, a significant QOL and emotional state deterioration correlates with a residual VF diameter below 20° and a VA lower than 0.3. It introduces, for the first time, thresholds to be used in visual restoration or visual preservation therapies to improve QOL of RP patients.

Introduction

Retinitis pigmentosa (RP) is a group of inherited retinal diseases which induce photoreceptor (rods and cones) degeneration leading to progressive visual loss [1, 2]. These diseases are characterized by a great heterogeneity from both genetic and clinical points of view [1, 3–5]. In most cases, RP provokes rod photoreceptor degeneration, responsible for defective dark adaptation (night vision).
blindness). The major handicap induced by RP, however, relates to the later loss of cone photoreceptors, which ineluctably results in peripheral visual field (VF) impairment. Finally, the disease progression can affect the central vision and induce total blindness.

In addition to sensory damage, vision loss generates many social and psychological repercussions impacting the quality of life (QOL) of individuals with visual impairment. Evaluation of the QOL has gradually taken its rightful place in ophthalmology with the development of several questionnaires since 1980. It has increasingly been implemented to evaluate the real impact of visual function troubles on patients’ daily lives and to improve patient management.

The evaluation of the QOL becomes particularly useful in the early phase clinical trial (https://clinicaltrials.gov) where patients have limited residual vision to assess the positive contribution or the side effects of new therapies. It is mandatory to know the real clinical impacts of a significant change in a surrogate end point such as a VF or visual acuity (VA) improvement. Previous studies have demonstrated correlations between the degree of VF, VA loss and QOL, showing that QOL decreases proportionally to the loss of visual function [6–9].

In our study, we have extended the correlation analyses between visual loss and QOL in order to determine a transitional level of vision that can represent a threshold for preservative therapeutic approaches or restorative strategies.

**Methods**

A cross-sectional study has been conducted on 60 patients diagnosed with typical RP ‘rod-cone dystrophy’ who were referred to the Reference Center for Hereditary Retinal Dystrophy/Clinical Investigation Center (Inserm CIC 1423/Quinze-Vingt Hospital, Paris [10]). A control group of 20 age-matched healthy subjects with no systemic or ocular disease in terms of VF impairment (VF groups) and VA loss (VA groups) have been set. Participants were all 18 years old or more, French speakers with no systemic or ocular disease apart from the ones studied. Functional ocular evaluations such as monocular and binocular VF assessments and VA measurements have been performed. The 25-item National Eye Institute Visual Functioning Questionnaire (NEI-VFQ-25), QOL survey and the Hospital Anxiety and Depression Scale (HADS) questionnaire assessment of the patients’ emotional states were self-administered and completed by each participant (or when needed by one of their relatives).

This study follows the tenets of the Declaration of Helsinki. Informed consent was obtained from all patients and normal subjects after explanation of the study. The protocol has been approved by an institutional review board: CPP (Persons Protection Committee in French) Ile de France V, Project 06693, EUDRACT 2006-A00347-44.

**Visual Field**

VF was assessed by monocular and binocular VF Goldmann kinetic manual perimetry [11]. The study population included patients with monocular and binocular VF restricted to the central area (i.e. patients lacking some peripheral retinal sensitivity) using isopter V4 for monocular and isopter III4 for binocular VF, respectively. Patients have been classified into VF groups according to their binocular VF. The choice of VF without peripheral islands allowed to have homogenous patient groups and to avoid the too great variability that can be induced by the diversity of sizes, locations and sensibilities of remaining peripheral islands. We have considered that normal binocular VF consists of a horizontal 120-degree-wide area bordered on each side by a monocular 30-degree-wide visual area. Binocular VF assessment is legal reference in France to determine functional capacity in a forensic or professional context, according to Decree No. 921216 published in the Official Journal of November 4, 1993.

The RP population has been divided into the following 3 VF groups, each including 20 subjects: VF1, with a residual binocular VF of less than 20° in diameter (ØVF <20°); VF2, with a residual binocular VF between 20 and 40° in diameter (20° ≤ ØVF ≤ 40°); VF3, with a residual binocular VF larger than 40° in diameter (ØVF >40°). The VF groups were compared to the control group of 20 healthy subjects without any VF restrictions (VFC group).

**Visual Acuity**

The Early Treatment Diabetic Retinopathy Study (ETDRS) logarithmic scale has been used to evaluate VA at a distance of 4 m. Except in a few cases where a slight difference was observed, the VA was comparable in both eyes. The RP patients have been divided into the following 3 VA groups, based on best eye VA: VA1 where VA ≤ 0.3 (15 patients); VA2 where 0.3 < VA ≤ 0.7 (21 patients); VA3 where VA >0.7 (24 patients). The VA groups have been compared to the control group of 20 healthy subjects where VA was equal to 1 (VAC group).

The 25-Item National Eye Institute Visual Function Questionnaire

The NEI-VFQ-25 was developed in the late 1990s to assess various dimensions of QOL in subjects with visual loss, including psychological and subjective aspects. It is reliably and significantly correlated with the NEI-VFQ-51 original version [12, 13]. The NEI-VFQ-25 was validated for various eye diseases, and it is among the most used and valuable psychometric tools [14–18].

It is made up of 25 questions that address 12 aspects of daily living: general health, general vision, near vision, distance vision, driving, peripheral vision, color vision, ocular pain, role limitation, dependency, social function and mental health. We used a French translated version validated for glaucoma and ocular hypertension [19]. Patients answered the questionnaires themselves without any interviewer. The answer to each of the 25 questions has been converted into a 100-point scale, in which 100 represents the best possible score. Results were distributed according to 12 subscores covering each of the 12 individual domains and a composite score (CS) representing the mean score of all subscores, except for general health which is not directly related to the visual condition [20].

The Hospital Anxiety and Depression Scale

The HADS questionnaire evaluates patients’ anxiety-depression scores to determine their emotional state [21]. The reliability
of the questionnaire has been confirmed in numerous studies, among them two reviews [22, 23] including 200 and 747 relevant studies, respectively. We used a French translated version introduced in a validation study by Lépine et al. [24]. The questionnaire is composed of 14 questions (7 for the anxiety score and 7 for the depression score). Each item is noted on a 0–3 scale according to a reading grid. The anxiety or depression diagnosis is negative below 7, low between 8 and 10, and positive above 11.

**Statistical Analysis**

Analysis of the data has been performed using the Statistical Package for the Social Sciences (SPSS, version 19.1, IBM Corporation). All statistical tests were carried out with significant level of risk set at $\alpha = 5\%$, or less in case of repeated Mann-Whitney, using Bonferroni correction tests, in order to preserve a global risk of $\alpha = 0.05$. Except for age, analyses of data showed they did not comply with normal distributions, so we used nonparametric tests.

The test of Spearman correlations was performed between all subscores of NEI-VFQ-25.

Comparisons of the frequency distribution for categorical variables of sex were performed with Pearson’s $\chi^2$ test between VF and VA groups. Age comparison was performed with a 1-factor analysis of variance.

A Kruskal-Wallis test was carried out to compare ranks of variables (CS, anxiety and depression scores) between VF and VA groups. In case of a significant difference between the 4 groups, post hoc tests were conducted.

In addition, within each VF group, QOL scores were compared according to patients’ VA groups with the Kruskal-Wallis test. Similarly, within each VA group, QOL scores were analyzed according to patients’ VF groups with the Kruskal-Wallis test.

**Results**

The demographic analysis of populations showed no significant difference within VF or VA groups in terms of age and gender. The average population age was 44 years (±14, SD) for RP and 41 years (±13) for the control population.

| Scores | VF1 (n = 20) | VF2 (n = 20) | VF3 (n = 20) | VFC (n = 20) |
|--------|-------------|-------------|-------------|-------------|
| CS     | 37.7 [19.7] | 55.9 [22.6] | 60.8 [24.4] | 93.7 [7.8]  |
| Anxiety| 10.0 [3.5]  | 9.50 [6.3]  | 9.0 [4.3]   | 6.50 [2.3]  |
| Depression| 7.0 [7.0] | 4.50 [6.3]  | 4.0 [5.5]   | 2.00 [3.5]  |

Values are presented as medians with interquartile ranges in square brackets. RP VF groups: VF1, ØVF <20°; VF2, 20° ≤ ØVF ≤ 40°; VF3, ØVF >40°; VFC = VF control group.

The CS and anxiety/depression scores of VF groups (median and interquartile range) are shown in table 1.

When comparing the CS values between the VFC group and the RP VF groups (table 1), a significant difference was found between groups (Kruskal-Wallis test for CS; $p < 0.001$). It appears, in figure 1, that CS in VFC is markedly above RP VF groups, and, in all VF groups, CS gradually decreased with severity of VF impairment. There was a significant difference in CS between groups VF1 and VF2 ($p = 0.013$; table 2) that also revealed marked differences between VF1 and groups VF3 and VFC.

A significant difference was evaluated between groups VF3 and VFC ($p < 0.001$; table 2). The data distributions shown in figure 1 suggest significant differences also between groups VF2 and VFC. The CS values for groups VF2 and VF3 did not differ significantly.

For anxiety and depression scores, an overall significant difference has been seen between VF groups and VFC ($p = 0.004$ and $p = 0.004$, respectively). The median scores for the groups are shown in table 1. Between-group comparisons of anxiety scores (table 2) showed significant differences between groups VF2 and VFC ($p = 0.004$). Consequently, the anxiety score was different between groups VF1 and VFC. No significant difference was revealed between groups VF1 and VF3 and between
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Table 2. Results of post hoc tests (Mann-Whitney) comparing CS between VF groups and control group

| Scores | VF1–VF2 | VF1–VF3 | VF2–VF3 | VF1–VFC | VF2–VFC | VF3–VFC |
|--------|---------|---------|---------|---------|---------|---------|
| CS     | 0.013*  | –       | 0.070   | –       | –       | <0.001* |
| Anxiety| –       | 0.355   | –       | –       | 0.004*  | 0.020   |
| Depression| –       | 0.087   | –       | 0.001*  | 0.045   | 0.022   |

RP VF groups: VF1, ØVF <20°; VF2, 20° ≤ ØVF ≤ 40°; VF3, ØVF >40°. * Significance threshold: p < 0.0167 for CS and anxiety score and p < 0.0125 for depression score. Post hoc test: Mann-Whitney.

Table 3. Scores of the 3 RP groups according to VA level and control group

| Scores | VA1 (n = 15) | VA2 (n = 21) | VA3 (n = 24) | VAC (n = 20) |
|--------|--------------|--------------|--------------|-------------|
| CS     | 37.4 [19.2]  | 56.1 [18.6]  | 59.8 [21.7]  | 93.73 [7.8] |
| Anxiety| 11.0 [7.0]   | 10.0 [4.0]   | 9.0 [4.0]    | 6.50 [2.3]  |
| Depression| 7.0 [6.0]   | 6.0 [7.0]    | 4.0 [6.3]    | 2.00 [3.5]  |

Values are presented as medians with interquartile ranges in square brackets. RP VA groups: VA1, VA <0.3; VA2, 0.3 < VA ≤ 0.7; VA3, VA >0.7; VAC = control group.

Data distribution in figure 2 also shows a difference between groups VA2 and VAC. Groups VA2 and VA3 did not show any significant difference.

Fig. 2. Distribution of CS in the VA and VAC groups. RP VA groups: VA1, VA <0.3; VA2, 0.3 < VA ≤ 0.7; VA3, VA >0.7. The line in the box plot is the median, which is surrounded by a box the top and bottom of which indicate the interquartile range (the limits within which the middle 50% of observations fall). The distance between the extreme values and box plot represent 25% higher values and 25% lower values.
CS medians based on VF groups and VA groups are shown in Table 5 with the results from the Kruskal-Wallis test comparing subgroups. Within VF groups, a significant difference between CS of VA subgroups was only revealed for group VF2 (p = 0.021). Within VA groups, a significant difference between CS of VF subgroups was shown for the groups VA2 (p = 0.048) and VA3 (p = 0.016).

**Discussion and Conclusion**

VF and VA are considered the most influential psychophysical parameters impacting a patient’s daily life [6–9]. Szlyk et al. [6] have shown that a patient’s difficulties in performing everyday activities were correlated with VF and VA, and that the patient’s self-assessments were more strongly related to the VA factor. Similar correlations between the NEI-VFQ-25 scores and the American Medical Association’s functional visual scores were reported by Seo et al. [7], showing that both VA and VF are important factors impacting QOL. Sugawara et al. [8] studied the effects of VF loss on the QOL of RP patients with preserved VA (≥0.7) and demonstrated a significant negative correlation between the peripheral visual field loss and vision-related QOL scores (NEI-VFQ-25). Similar were the correlations between VF impairment and QOL in patients with low VA (≤0.3) associated with various ocular diseases such as glaucoma, macular degeneration, diabetic retinopathy and RP [9].

The results of our study are consistent with those of these previous studies. By selecting a homogeneous group of patients with a tubular VF, we have avoided the variability of the scores of QOL that could be related to residual peripheral islands of retinal sensitivity with different levels, sizes or locations. We clearly showed that, in patients with severely reduced VF or VA, the QOL is much lower than in less affected groups. These results point to a threshold size of central residual VF of a diameter of 20° and a threshold level of VA of 0.3, below which the QOL is dramatically altered. A similar VF limit has been reported by Haymes et al. [25] concerning the mobility alteration of RP patients, which explains in part the worsening of QOL scores.

When analyzing the influence of the VA and VF parameters to each other, we observed that in the group VF1

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**Table 4.** Results of post hoc tests (Mann-Whitney) comparing CS between VA groups and control group

| Between groups | VA1–VA2 | VA1–VA3 | VA2–VA3 | VA2–VAC | VA3–VAC |
|----------------|---------|---------|---------|---------|---------|
| CS             | 0.0047* | –       | 0.109   | –       | <0.001* |
| Anxiety        | –       | 0.503   | –       | –       | 0.0065* |
| Depression     | –       | 0.055   | –       | 0.0033* | 0.081   |

RP VA groups: VA1, VA <0.3; VA2, 0.3 < VA ≤ 0.7; VA3, VA >0.7; VAC = control group. * Significance threshold: p < 0.0167 for CS and depression score and p < 0.025 for anxiety score.

**Table 5.** CS in VA-VF subgroups

| Groups | VA1 | VA2 | VA3 | Kruskal-Wallis test |
|--------|-----|-----|-----|---------------------|
| VF1    | (n = 8) 34.28 [9.83] | (n = 7) 43.03 [16.72] | (n = 5) 48.83 [22.44] | 0.460 |
| VF2    | (n = 5) 27.77 [17.58] | (n = 4) 52.51 [20.28] | (n = 11) 62.80 [13.47] | 0.021* |
| VF3    | (n = 2) 51.25 [2.58] | (n = 11) 60.80 [12.97] | (n = 7) 72.77 [24.70] | 0.175 |
| Kruskal-Wallis test | 0.234 | 0.048* | 0.016* |

Values are presented as medians with interquartile ranges in square brackets. * Significance threshold: p < 0.05.
(smaller residual VF, regardless of the VA level), the CS did not differ. Similarly, in the group with VA below 0.3 (VA1), regardless of the VF size, the QOL scores did not change. Thus, it appears that when a parameter is very low, the improvement of the other parameter should have little or no impact on the QOL of patients.

Independently from the degree of vision loss, being sick can affect patient QOL due to the perception of his or her abilities to perform certain daily activities. Some previous studies have correlated the QOL scores with the emotional state of patients. Owslley and McGwin [26] suggested that depressed patients may have low NEI-VFQ-25 scores independently of the impact of vision problems. For Hahm et al. [27], there was a negative correlation between the depression score (Beck Depression Inventory scale) and the NEI-VFQ-25 CS; however, the Beck Depression Inventory score was not correlated with any objective measure of visual function. It seems that it is also important to consider emotional states because they can affect the variability of visual inspections. Indeed, the variability of VF and VA measures is partly related to the severity of the disease, but these can also vary due to stress and anxiety generated by the evaluation of visual performance itself, especially on patients whose emotional state is affected by visual loss [28–30]. This is even truer when RP is in advanced and severer stages [31]. Thus, the results of ophthalmic tests and the emotional states of RP patients influence each other. Some authors, such as Rozanski et al. [30], have attempted to counter this bias; they developed a theoretical model showing that VF variability is linked to the patients’ negative emotional states, providing a basis for the development of coping strategies resolving this problem.

The results of the present work show that RP has a major impact on a patient’s emotional state, and this is even more pronounced when the degree of visual impairment becomes greater. In our study, patients with VF diameters less than 20° (group VF1) and VA less than 0.3 (group VA1) report anxiety and depression significantly higher than the control group, thus confirming previously found thresholds.

Numerous studies have evaluated the natural course of visual field progression in RP patients and discovered the various time periods over which half of the remaining field area would be lost (half-life) [32–36]. Their results showed half-lives ranging from 4.5 years [33] to 11 years [32]. It would be interesting in future studies to specify this loss rate according to genetics in order to better predict the visual prognosis of patients and include the progression profile as an important patient selection criterion for clinical trials.

In conclusion, the results of our study confirm that QOL is influenced by having RP, and it correlates in a significant way with the VF size and the VA impairment. Furthermore, they suggest a threshold approximation (VF diameter of 20° and VA equal to 0.3) below which QOL is very deteriorated and the emotional state is highly affected. Thresholds that could represent the minimal therapeutic effects have to be targeted by therapists aiming at visual restoration, or not to be reached by those aiming at preservation. Translating a functional benefit, induced by a therapeutic action in terms of improvement of QOL, is indeed determinant.

Subjective and objective data can facilitate the doctor-patient dialogue [37–39] and help multidisciplinary teams of specialized therapists to adapt appropriate strategies for patient reeducation/readaptation and social compensations [40, 41].

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Conflicts of Interest

All the authors hereby certify that there are no conflicts of interest with any financial issue.

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