Artificial vision: the effectiveness of the OrCam in patients with advanced inherited retinal dystrophies

Xuan-Thanh-An Nguyen,1 Jan Koopman,2 Maria M. van Genderen,3,4 Henk L.M. Stam4 and Camiel J.F. Boon1,5

1Department of Ophthalmology, Leiden University Medical Center, Leiden, The Netherlands
2Royal Dutch Visio, Centre of Expertise for Blind and Partially Sighted People, Amsterdam, The Netherlands
3Department of Ophthalmology, University Medical Center Utrecht, Utrecht, The Netherlands
4Bartiméus, Diagnostic Center for Complex Visual Disorders, Zeist, The Netherlands
5Department of Ophthalmology, Amsterdam UMC, Academic Medical Center, Amsterdam, The Netherlands

ABSTRACT.
Purpose: To investigate the impact of the OrCam MyEye 2.0 (OrCam) on the quality of life and rehabilitation needs in patients with advanced retinitis pigmentosa (RP) or cone-rod dystrophies (CRD). The OrCam is a wearable low-vision aid that converts visual information to auditive feedback (e.g. text-to-speech, barcode and facial recognition).

Methods: Patients with a clinical diagnosis of RP (n = 9, 45%) or CRD (n = 11; 55%), and a best-corrected visual acuity of ≤20/400 Snellen were invited to participate in this study. Questionnaires were administered at baseline and after 5.2 (standard deviation ± 1.5) weeks, which included the Dutch version of the National Eye Institute Visual Functioning Questionnaire (NEI-VFQ), the Participation and Activity Inventory (PAI) and the OrCam Function Questionnaire (OFQ).

Results: Following OrCam testing, significant improvements were observed in the ‘near activities’ subscale of the NEI-VFQ (p < 0.001); the ‘visual functioning’ subscale of the re-engineered NEI-VFQ (p = 0.001); the ‘reading’ rehabilitation goal of the PAI (p = 0.005) and the overall score of the OFQ (p < 0.001). The observed changes in questionnaire scores did not differ between phenotypes. Advantages and limitations of the OrCam were reported by patients. Three patients (15%) continued rehabilitation with the OrCam after completion of this study.

Conclusions: The OrCam mainly improves reading domains in patients with advanced stages of RP or CRD. Further improvements in the OrCam are needed to address current limitations, which may enhance its utility for patients with RP or CRD.

Key words: cone-rod dystrophies – low vision – OrCam – quality of life – retinitis pigmentosa – visual aids

Acta Ophthalmol. 2022: 100: e986–e993
© 2021 The Authors. Acta Ophthalmologica published by John Wiley & Sons Ltd on behalf of Acta Ophthalmologica Scandinavica Foundation.
This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.
doi: 10.1111/aos.15001

Introduction
Inherited retinal dystrophies (IRDs) comprise a diverse group of rare eye diseases characterized by progressive loss of photoreceptor function, ultimately leading to severe visual impairment (Cremers et al. 2018). Inherited retinal dystrophies (IRDs) can be differentiated, in part, through the order of which cells are lost (Cremers et al. 2018). In retinitis pigmentosa (RP), degeneration of rods precedes that of cones, resulting in initial symptoms of nyctalopia and peripheral visual field loss (Hamel 2006; Hartong et al. 2006; Ferrari et al. 2011). Ultimately, central vision is also lost. Conversely, in cone-rod dystrophies (CRD), the process of photoreceptor degeneration follows the opposite sequence of events than in RP, causing predominant symptoms of central vision loss, photophobia and colour vision impairment followed by peripheral vision loss and night blindness in later stages of the disease (Hamel 2007; Thiadens et al. 2011). Loss of visual function due to RP or CRD has detrimental effects on a patient’s well-being and on their ability to perform daily activities, although the extent and areas of difficulties may vary between these phenotypes (Latham et al. 2015).

For most patients with IRDs, the visual prognosis remains poor, as curative treatments are unavailable or are still under investigation. Therefore, emphasis should be on assisting patients with managing their disease, for example, through low-vision rehabilitation services (Lamoureux et al. 2007). The goal of low-vision rehabilitation is not to restore vision, but to utilize residual vision to its maximum potential (Langelaan et al. 2009). This may be achieved by low-vision centres...
through the prescription of low-vision aids (LVAs), ranging from (non-)optical aids to electronic assistive technologies. The selection of appropriate LVAs for an individual patient is complex, and several factors need to be considered prior to prescription, such as a patient’s visual and cognitive ability, disease stage, occupation and own rehabilitation goals (Das et al. 2019; Lorenzini & Wittich 2020).

The OrCam MyEye (https://www.orcam.com), or OrCam in short, is a relatively recent addition to the list of commercially available LVAs. The OrCam is a portable LVA that can be attached to the frame of a patient’s eyeglasses. It contains a small camera that converts digital or printed text to real-time auditive feedback using optical character recognition technology. As such, the intended audience for the OrCam consists of severe visually impaired or blind patients that have lost the ability to read independently. Aside from text-to-speech capabilities, the OrCam also contains colour, object, barcode, money and facial recognition. Thus, the OrCam has the potential to improve the performance of multiple daily activities in visually impaired patients. However, the impact of a single LVA remains unclear, as low-vision rehabilitation programmes typically offer multiple LVAs and multidisciplinary services over the course of rehabilitation. This makes it difficult to distinguish the contribution of a single device or service on a patient’s rehabilitation progress (Moisseiev & Mannis 2016; Waibourd et al. 2019). Insights into the effectiveness of the OrCam will provide knowledge on which patients are most likely to benefit from the device and will also inform us on which daily activities may improve when using devices such as the OrCam. In addition, as the target of interest has to be within the OrCam’s field of view, we also investigated whether the feasibility of the OrCam differed in those with different visual abilities, for example, patients with peripheral blindness or central blindness. For this purpose, this study investigated the effectiveness of the OrCam on the quality of life and the perceived difficulties in daily activities in severe visually impaired or blind patients caused by either RP or CRD.

Methods

Participants

Patients that were scheduled for one of the two Dutch low-vision rehabilitation centres, Bartiméus (Amsterdam, the Netherlands) or Royal Dutch Visio (Amsterdam, the Netherlands), were invited to participate in this study. Inclusion criteria for this study were a clinical diagnosis of RP or CRD based on full-field electroretinography data, and a best-corrected visual acuity (BCVA) of 20/200 Snellen acuity or worse. An additional inclusion criterion for patients with RP was a constricted peripheral visual field on Goldmann kinetic perimetry (<20° around point of fixation using a V4e stimulus) at the most recent examination, whereas for patients with CRD, an absolute central scotoma with residual peripheral fields was present in all. Identification of a causative gene was not a requirement for this study. Exclusion criteria for this study included the presence of other ocular diseases, significant cognitive impairment, insufficient understanding of the Dutch language and tremor-inducing conditions that could impede gesture recognition by the OrCam (e.g. Parkinson’s disease). Ethical approval for this study was obtained from the Medical Ethics Committee at the Leiden University Medical Center. The study adhered to the tenets of the Declaration of Helsinki, and informed consent was signed by all participants.

OrCam study protocol

Questionnaires were administered in patients using a personal interview-format at initial visit and at follow-up (mean follow-up: 5.2 weeks ± standard deviation [SD] 1.5). Additionally, patients underwent visual acuity testing using a Snellen letter chart and received instructions on the OrCam at first visit. Both centres followed a similar OrCam instruction protocol, performed by experienced instructors, to ensure identical training between centres. Different models of the OrCam exist, which differ in price and their available features (https://www.orcam.com). For this study, the OrCam MyEye 2.0 was tested by all patients (Fig. 1), and instructions were given on the following functions: text recognition, facial recognition, barcode recognition, object recognition, money recognition, colour recognition and telling time (Moisseiev & Mannis 2016).

The OrCam’s features are activated by pressing the touch bar located on the device itself; or hands-free via automatic target recognition, or by performing gesturing motions (e.g. pointing at a target). Following are the OrCam study protocol.

Fig. 1. OrCam MyEye 2.0 is a portable low-vision aid that can be mounted to the arms of a pair of glasses. The processor unit has an internal speaker, charge port, power button and a touch bar for activation and menu navigation (white arrow). Furthermore, the OrCam contains an optical sensor (yellow arrow), that returns scanned text or objects to auditive feedback via the internal speaker or through a Bluetooth connected earpiece. A mini flashlight is also present to aid in lower light situations. In addition to text-to-speech functions, the OrCam also contains colour (selective) barcodes, money, person and object recognition features. In order for person and object recognition features to function, it is required to scan the desired target in advance, subsequently storing this information in the internal memory of the OrCam. The OrCam is activated via the touch bar, or hands-free via automatic target recognition, or by performing gesturing motions (e.g. pointing at a target).
Questionnaires
Three questionnaires were used in this study, which included the National Eye Institute Visual Function Questionnaire (NEI-VFQ), the Participation and Activity Inventory (PAI), and the OrCam Function Questionnaire (OFQ). Patients were instructed to answer all questionnaires as if they were using their own LVAs, with the addition of the OrCam as a LVA at follow-up assessment.

The NEI-VFQ is a 25-item questionnaire with 14 supplemental items and is one of the most common vision-related quality of life questionnaires used in ophthalmic research. The NEI-VFQ is designed to evaluate aspects of daily living, which can be categorized into 12 different subscales (Mangione et al. 2001). For our study, the driving subscale was omitted, as none of the patients were permitted to drive. Answers given by patients were subsequently recoded into a 100-point scale, where a higher score represents better (visual) functioning, as suggested by the original authors (Mangione et al. 2001). An overall composite score was calculated by averaging the scores of all subscales, whilst excluding the ‘general health’ subscale.

The PAI, formerly known as the Dutch Activity Inventory, is a validated questionnaire that is used in Dutch low-vision rehabilitation centres to systematically assess the rehabilitation goals of patients (Bruijning et al. 2010a; Bruijning et al. 2010b; Elsman et al. 2018; Macnaughton et al. 2019). The PAI is based on the Activity Inventory designed by Massof and colleagues (Massof et al. 2007), which was modified in order to extend to the European population (Bruijning et al. 2010a; Bruijning et al. 2010b; Latham et al. 2015). For this study, a shortened version of the PAI was used, which included 11 rehabilitation goals related to central or peripheral vision (Table S1) (Latham et al. 2015). Patients were instructed to rate each goal on two aspects: importance and difficulty. Importance is rated on a Likert scale ranging from 0 (not important) to 3 (very important), whereas the difficulty scale goes from 0 (not difficult) to 4 (impossible). Subsequently, a priority score is calculated as the product of importance and difficulty for each included goal. The maximum achievable priority score is 12, with a higher priority score signifying a greater rehabilitation need for this specific rehabilitation goal.

The OFQ is a non-validated questionnaire that was developed solely for this study. The questionnaire contained 14 items regarding vision-related daily activities. The OFQ uses a 5-level Likert scale, with possible difficulty scores being 1 (no difficulty), 2 (some difficulty), 3 (moderate difficulty), 4 (very difficult) or 5 (impossible due to disease). The activities included on the OFQ are as follows:
1. Reading a newspaper or book.
2. Reading for longer than 30 minutes without getting tired.
3. Reading an e-mail.
4. Reading text from a distant sign such as a street sign.
5. Reading handwritten text.
6. Identifying different money bills.
7. Recognizing colours on clothing pieces.
8. Recognizing familiar objects, such as your keys or phone, at home.
9. Recognizing a familiar product in the grocery store.
10. Finding your way in the grocery store.
11. Reading a product label.
12. Recognizing familiar faces at home.
13. Recognizing familiar faces within an unfamiliar environment.
14. Telling time.

Rasch analysis
Rasch analysis was performed expeditiously on the NEI-VFQ and OFQ using the Andrich rating scale model (Winsteps 4.6.0) (Massof & Fletcher 2001; Stelmack et al. 2002; Pesudovs et al. 2010). Rasch analysis converts ordinal scores into an interval scale and provides patient’s ability and item difficulty using logit values for the underlying construct. In our study, patients with higher (visual) ability and items of greater difficulty are placed more negatively of the logit scale, whereas more positive logit values reflect patients with lower (visual) ability and items with less difficulty. For NEI-VFQ, re-engineering of the questionnaire was guided by previous authors, who proposed a two subscale structure: visual functioning and socio-emotional subscales (Table S1) (Stelmack et al. 2002; Pesudovs et al. 2010). For the OFQ, three items were removed to fit Rasch analysis, demonstrating reliable person and item separation values (reliability >0.8), scale targeting (difference between mean item and person measures <1.0 logit) and unidimensionality (variance accounted by the principal component >60%) (Table S1). Changes in person measures after OrCam rehabilitation were assessed using a stacked analysis (Anselmi et al. 2015).

Statistical analysis
Data were analysed using the SPSS version 25.0 (IBM Corp, Armonk, NY, USA). Visual acuity data were converted to Logarithm of the Minimum Angle of Resolution (logMAR) values. For hand movement vision, light perception vision and no light perception, logMAR values of 2.7, 2.8 and 2.9 were used, respectively (Talib et al. 2017). Best-corrected visual acuity (BCVA) in the better-seeing eye of included patients was categorized into two groups: severe visual impairment (SVI; 20/400 ≤ BCVA < 20/200) or blindness (BCVA < 20/400), based on criteria set by the World Health Organization (World Health Organization 2019). As data were normally distributed, a paired 2-tailed t test was used to determine significant changes in raw scores for each instrument. The effect of age, vision categories (SVI or blindness) and phenotypes (RP or CRD) on the likelihood of change were also investigated using a linear mixed model. A p-value of 0.05 or less was considered clinically significant, and correction for multiple testing using the Bonferroni method was applied where appropriate.

Results
Clinical characteristics of the patients are presented in Table 1. Twenty patients with IRD were enrolled in the study, of which nine patients were clinically diagnosed with RP (45%), and 11 patients with CRD (55%). Patients had an average BCVA of 1.5 logMAR (SD ± 0.4), which is equivalent to 20/640 Snellen visual acuity. Aside from visual field patterns, there were no differences in clinical characteristics between the two phenotypes. All patients had previously undergone low-vision rehabilitation, and the majority of patients (n = 19; 95%)...
Male (n/C6 onset.

Disease duration was defined as the difference between age at baseline and age at first symptom.

Follow-up in weeks (mean SD)

Non-optical aids (n)

Text-to-speech products (n)

Visual field pattern

Central scotoma with peripheral remnants

Optical aids (n, %)

Glasses 13 (65%) 7 (78%) 6 (55%)

Telescopes 3 (15%) 1 (11%) 2 (18%)

Hand or stand magnifiers 9 (45%) 3 (33%) 6 (55%)

Filter glasses 11 (55%) 6 (67%) 5 (46%)

Illumination control 8 (40%) 4 (44%) 4 (36%)

Braille 6 (30%) 2 (22%) 4 (36%)

White cane 13 (65%) 6 (89%) 5 (46%)

Text-to-speech products (n, %)

Screen reading software 14 (70%) 7 (56%) 9 (82%)

Daisy reader (physical or digital) 14 (70%) 7 (78%) 8 (64%)

Text-to-speech mobile applications 16 (80%) 8 (89%) 8 (73%)

p-Values were derived from the independent t-test, χ2 test or Fisher’s exact test.

BCVA = best-corrected visual acuity, logMAR = Logarithm of the Minimum Angle of Resolution, SD = standard deviation.

* Disease duration was defined as the difference between age at baseline and age at first symptom onset.

† Text-to-speech products included software (e.g. JAWS, SuperNova, Window Eyes, VoiceOver), equipment, and mobile applications that convert digital or printed text to auditive feedback (e.g. Seeing AI or KNFB reader).

included in this study were in possession of at least one LVA with text-to-speech capabilities (Table 1).

National Eye Institute Visual Function Questionnaire

At initial visit, the NEI-VFQ showed a significantly lower score on the peripheral vision subscale in patients with RP compared to patients with CRD (p = 0.014). Other subscales on the NEI-VFQ were found to be comparable between subgroups, including the overall composite score (Table S2). Rasch analysis revealed mean person measures of 0.53 (SD ± 0.64) and −0.18 (SD ± 0.59) logits for the visual functioning and socio-emotional subscales, respectively. At follow-up, significant improvements were observed in the raw scores of the near activities' subscale (+2.5, 95% CI: 13.2–33.9; p < 0.001), which was not found for other subscales after correction for multiple testing (adjusted p-value = 0.004; Fig. 2). The observed change was not affected by phenotype (p = 0.798), initial age (p = 0.089) or vision classification (p = 0.317). A significant change was also observed on the Rasch-calibrated visual functioning subscale, showing an improvement of −0.65 logits (95% CI: −0.97 to −0.32; p = 0.001) after OrCam use. No significant change was found in the socio-emotional subscale (−0.14, 95% CI: −0.40 to 0.11; p = 0.257) after rehabilitation.

Participation and Activity Inventory Questionnaire

A summary of the priority scores for each goal on the PAI is provided in Table 2. Goals with the highest priority scores, indicating goals with the highest rehabilitation needs, were ‘mobility indoors within an unfamiliar environment’ and ‘personal administration’ for patients with RP; whereas the highest priority scores were found in the ‘reading’ and ‘personal administration’ goals for patients with CRD (Table 2). Whilst the order of priority for rehabilitation goals differed between phenotypes, there was no significant difference in average scores for each goal (Table S2). Bivariate analysis revealed a correlation between the priority score of the ‘mobility indoors within an unfamiliar environment’ goal and age at initial visit (r = 0.570; p = 0.009), suggesting that the rehabilitation need for the ‘mobility indoors within an unfamiliar environment’ goal becomes greater with increasing age.

Out of the 11 rehabilitation goals included, ‘reading’ was the only goal that improved after rehabilitation with the OrCam, as shown as a lower priority score at follow-up (−2.6, 95% CI: −4.2 to −0.9; p = 0.005). When analysing the underlying tasks of the ‘reading’ goal, a significant lower priority score was found for the task ‘reading ordinary-sized print’ (−3.9, 95% CI: −6.4 to 1.3; p = 0.005), which was not found for other tasks related to the ‘reading’ goal.

OrCam Function Questionnaire

An item-person map based on Rasch analysis of the OFQ questionnaire is shown in Figure 3. Items on the OFQ that were considered most difficult for this cohort were: ‘recognizing familiar faces within an unfamiliar environment’ (−1.67 logits), reading text from a distant sign (−1.40 logits), and ‘reading a product label’ (−0.90 logit); whereas ‘reading an e-mail’ (1.18 logits) and ‘recognizing familiar objects at home’ (1.15 logits) were considered the least difficult tasks. The average person measure was 0.43 logits (SD ± 0.92), which improved significantly following OrCam rehabilitation (−1.11, 95% CI: −1.61 to 0.61; p < 0.001). The observed change did not differ between phenotypes (p = 0.696).

Overall experience with the OrCam

At final visit, patients shared their overall experience with the OrCam. Fifteen patients (75%) reported that...
the OrCam’s text recognition features functioned well in optimal light conditions. However, these features were less reliable in poorly lighted or dark rooms. Object and facial recognition features were not tested by most patients (n = 16; 80%), as patients reported that the current study period was too short to adequately test these features, or they did not consider these features necessary for their daily activities. Main advantages and limitations of the OrCam MyEye 2.0 provided by this cohort are summarized in Table 3. After completion of this study, two patients with RP (10%; aged 24 and 60) and one patient with CRD (5%; aged 51) continued with rehabilitation with the OrCam. The remaining patients (n = 17; 85%) did not resume rehabilitation with the OrCam. Reasons for not continuing with the OrCam, that were mentioned by at least five patients, were: (1) having text-to-speech products with similar functions as the OrCam (e.g. Seeing AI or KNFB reader); (2) pricing of the OrCam; (3) and lack of features that were considered important to a patient (e.g. assistance with navigation). We found no significant differences in baseline age (p = 0.845), disease duration (p = 0.258), mean logMAR BCVA (p = 0.765), visual functioning subscale score on the NEI-VFQ (p = 0.616), ‘reading’ goal priority score on the PAI (p = 0.616), or person measure score on the OFQ (p = 0.546) between those who did and those who did not resume rehabilitation with the OrCam.

**Discussion**

The objective of this study was to investigate whether the OrCam could assist in performing daily activities and subsequently improve the quality of life in patients with RP or CRD. As visual function gradually declines in patients with IRDs, so does their ability to perform daily activities, which, in turn, results in reduced vision-related quality of life (Chaumet-Riffaud et al. 2017). As such, our cohort with severely visually impaired and blind patients with IRDs presented with markedly impaired of quality of life, as measured on the NEI-VFQ.

When assessing the priority scores on the PAI, we found that the highest scores were found in the ‘mobility indoors within an unfamiliar location’ rehabilitation goal for patients with RP, whereas ‘reading’ and ‘personal administration’ were the most important rehabilitation goals in patients with CRD. These findings coincide with the different visual abilities present in patients with RP and CRD, with patients with RP most often facing challenges with mobility due to loss of peripheral vision, and patients with CRD experiencing difficulties with reading due to loss of central vision.

The Rasch-calibrated OFQ revealed that the most difficult tasks were ‘reading a distant sign’, ‘reading a product label’ and ‘recognizing familiar faces within an unfamiliar environment’, as...
they required the highest visual ability of patients. These tasks share a common theme in that they all involve visual search behaviour, which is defined as the perceptual ability to actively scan the environment to locate the target of interest amongst other visual distractors (Timmis et al. 2017). Visual search requires input from central and peripheral vision, both of which are lost, to various degrees, in our patient cohort (Sullivan et al. 2008; Timmis et al. 2017).

After rehabilitation with the OrCam, significant improvements were seen in the ‘near activity’ subscale of the NEI-VFQ. Similar results were found in a previous study with the OrCam, showing improvements in the ‘near vision’ subscale of the NEI-VFQ in patients with end-stage glaucoma (Waisbourd et al. 2019). As previous studies have demonstrated that the NEI-VFQ suffers from multidimensionality, we also obtained Rasch estimates from visual functioning and socio-emotional subscales (Stelmack et al. 2002; Pesudovs et al. 2010). Using this method, we found significant improvements in the visual functioning subscale, but no improvements in the socio-emotional subscale at follow-up. Significant improvements were also observed in the ‘reading’ goal on the PAI and the person measure score on the OFQ. These findings altogether suggest that the OrCam primarily improves reading abilities in patients with RP or CRD. The improvements

Fig. 3. Stacked person-item map of the OFQ questionnaire. Patients are shown as crosses and are mapped across the vertical line based on their (visual) ability measured in logits. For context, a patient with high abilities (i.e. no difficulty in performing activities) would be placed at the bottom of the logit scale. Similarly, item are also mapped according to their measure in logits, with the hardest items placed at the bottom of the scale. M, mean; S, 1 standard deviation from the mean; T, 2 standard deviations from the mean.
after OrCam usage did not differ between phenotypes, which may be due to our limited sample size, impeding more in-depth subgroup analysis. As suggested previously, it is possible that the level of visual acuity loss rather than visual field loss is important when selecting eligible patients for the OrCam (Waisbourd et al. 2019). Other features, such as facial and object recognition, were not tested by all patients during this relatively short follow-up, and the impact of these features on the quality of life in patients with IRDs remains uncertain. For these features, patients are required to store the person or object into the memory of the OrCam, a process that could take more than several minutes for the current version of the OrCam for each person or object, which is potentially exhaustive and time-consuming for severely visually impaired or blind patients over a study period of 5.2 weeks.

Most patients (85%) did not continue with rehabilitation, as they were in possession of other text-to-speech products, such as mobile applications with text recognition features (e.g. Seeing AI or the KNFB reader). These products share similar features with the OrCam, although, unlike the OrCam, most of these products often cannot be controlled hands-free or through gesturing motions. However, these products are typically less expensive compared to the OrCam for each person or object, which is potentially exhaustive and time-consuming for severely visually impaired or blind patients over a study period of 5.2 weeks.

Several limitations and confounding factors were mentioned by at least five patients are listed.

**Table 3. Advantages and limitations of the OrCam reported by patients with retinitis pigmentosa or cone-rod dystrophies.**

| Advantages | Limitations |
|------------|-------------|
| (+) Text recognition in optimal light conditions | (-) Difficulties with text recognition in low light conditions |
| (+) Portability | (-) Heavy and unbalanced on lightweight frames |
| (+) Hands-free | (-) Short battery life |
| (+) Colour recognition | (-) No connectivity capabilities with your smartphone |
| (+) Barcode recognition | (-) Lack of desired features* |
| (+) Bluetooth connectivity with earpieces | |

Remarks that were mentioned by at least five patients are listed.

* Example of features that were requested in this patient cohort included: assistance with navigation, voice activation, and internet connectivity.

and disadvantages of this device when prescribed to patients with RP or CRD. This knowledge may inform patients about the possibilities with the OrCam, whilst also setting realistic expectations, which, in turn, will facilitate the decision-making process regarding the OrCam. The OrCam is a useful LVA to improve reading abilities in patients with RP or CRD. Further improvements in the OrCam may enhance its utility in the rehabilitation process of patients with RP or CRD.

**References**

Anselmi P, Vidotto G, Bettinardi O & Ber-tolotti G (2015): Measurement of change in health status with Rasch models. Health Qual Life Outcomes 13: 16.

Brauijning JE, van Nispen RMA & van Rens GHMB (2010): Feasibility of the Dutch ICF Activity Inventory: a pilot study. BMC Health Serv Res 10: 310.

Brauijning J, van Nispen R, Verstraten P & van Rens G (2010): A Dutch ICF Version of the Activity Inventory: results from focus groups with visually impaired persons and experts. Ophthalmic Epidemiol 17: 366–377.

Chaumet-Riffaud AE, Chaumet-Riffaud P, Cariou A, Devisme C, Audo I, Sahel J-A & Moland-Said S (2017): Impact of retinitis pigmentosa on quality of life, mental health, and employment among young adults. Am J Ophthalmol 177: 169–174.

Cremers FPM, Boon CJF, Bujakowska K & Zeitz C. Special issue introduction: inherited retinal disease: novel candidate genes, genotype-phenotype correlations, and inheritance models. Genes 9: 215.

Das K, Gopalakrishnan S, Dalan D, Vela S, Ratra V & Ratra D (2019): Factors influencing the choice of low-vision devices for visual rehabilitation in Stargardt disease. Clin Exp Optom 102: 426–433.

Elsman EBM, van Rens GHMB & van Nispen RMA (2018): Psychometric properties of a new intake questionnaire for visually impaired young adults: The Participation and Activity Inventory for Young Adults (PAI-YA). PLoS One 13: e0201701.

Ferrari S, Di Iorio E, Barbaro V, Ponzi D, Sorrentino FS & Parmegiani F (2011): Retinitis pigmentosa: Phenotype correlations, and disease mechanisms. Curr Genomics 12: 238–249.

Hamel CP (2006): Retinitis pigmentosa. Orphanet J Rare Dis 1: 40.

Hamel CP (2007): Cone rod dystrophies. Orphanet J Rare Dis 2: 7.

Hartong DT, Berson EL & Dryja TP (2006): Retinitis pigmentosa. Lancet 368: 1795–1809.

Lamoureux EL, Pallant JF, Pesudovs K, Rees G, Hassell JB & Kefee JE (2007): The effectivenes of low-vision rehabilitation on participation in daily living and quality of life. Invest Ophthalmol Vis Sci 48: 1476–1482.
Lengelaan M, de Boer MR, van Nispen RMA, Wouters B, Moll AC & van Rens GHMB (2009): Change in quality of life after rehabilitation: prognostic factors for visually impaired adults. Int J Rehabil Res 32: 12–19.

Latham K, Baranian M, Timmis MA & Pardhan S (2015): Difficulties with goals of the Dutch ICF Activity Inventory: perceptions of those with retinitis pigmentosa and of those who support them. Invest Ophthalmol Vis Sci 56: 2381–2391.

Lorenzini M-C & Wittich W (2020): Factors

Macnaughton J, Latham K & Vianya-Estopa

Mangione CM, Lee PP, Gutierrez PR, Spritzer

McCambridge J, Witton J & Elbourne DR

Moisseiev E & Mannis MJ (2016): Evaluation

Moisseiev E & Mannis MJ (2016): Evaluation

M (2019): Rehabilitation needs and activity

M (2019): Rehabilitation needs and activity

Massof RW & Fletcher DC (2007): The Activity Inventory: an adaptive visual function questionnaire. Ophthalmic Physiol Opt 27: 1050–1058.

Massof RW, Almadian L, Grover LL, Dere- meik JT, Golstein JE, Rainey C, Epstein C & Barnett GD (2007): The Activity Inventory: an adaptive visual function questionnaire. Ophthalmic Physiol Opt 27: 1050–1058.

Massof RW & Fletcher DC (2001): Evaluation of the NEI visual function questionnaire as an interval measure of visual ability in low vision. Vision Res 41: 397–413.

McCarron J, Witton J & Elbourne DR (2014): Systematic review of the Hawthorne effect: new concepts are needed to study research participation effects. J Clin Epidemiol 67: 267–277.

Moisseiev E & Mannis MJ (2016): Evaluation of a portable artificial vision device among patients with low vision. JAMA Ophthalmol 134: 748–752.

Pestovs K, Goothwal VK, Wright T & Lamoureux EL (2010): Remediating serious flaws in the National Eye Institute Visual Function Questionnaire. J Cataract Refract Surg 36: 718–732.

Stelmack JA, Stelmack TR & Massof RW (2002): Measuring low-vision rehabilitation outcomes with the NEI VFQ-25. Invest Ophthalmol Vis Sci 43: 2859–2868.

Sullivan B, Jovancevic-Misic J, Hayhoe M & Sterns G (2008): Use of multiple preferred retinal loci in Stargardt’s disease during natural tasks: a case study. Ophthalmic Physiol Opt 28: 168–177.

Talib M, van Schooneveld MJ, van Genderen MM et al. (2017): Genotypic and phenotypic characteristics of CRIBI-associated retinal dystrophies. Ophthalmology 124: 884–895.

Thiandres AAHJ, Soerjoesing GG, Florijn RJ et al. (2011): Clinical course of cone dystrophy caused by mutations in the RPGR gene. Graefes Arch Clin Exp Ophthalmol 249: 1527–1535.

Timmis MA, Allsop J, Baranian M, Baker J, Basevitch I, Latham K, Pardhan S & van Paridon KN (2017): Visual search behavior in individuals with retinitis pigmentosa during level walking and obstacle crossing. Invest Ophthalmol Vis Sci 58: 4737–4746.

Waisbourd M, Ahmed OM, Newman J et al. (2019): The Effect of an Innovative Vision Simulator (OrCam) on Quality of Life in Patients with Glaucoma. J Visual Impair Blind 113: 332–340.

Wang BZ, Pestovs K, Keane MC, Daly A & Chen CS (2012): Evaluating the effectiveness of multidisciplinary low-vision rehabilitation. Opthom Vision Sci 89: 1399–1408.

World Health Organization (2019): International Classification of Diseases 11th Revision. Available at: https://icd.who.int/en (Accessed on 25 Dec 2020).

Received on March 25th, 2021. Accepted on August 4th, 2021.

Correspondence:

Camiel J.F. Boon, MD, PhD

Department of Ophthalmology

Leiden University Medical Center

Postal zone J3-S. Albinusdreef 2

2333 ZA Leiden

The Netherlands

Tel: +31(0)71 526 2388

Fax: +31(0)71 524 8222

Email: c.j.f.boon@lumc.nl

The authors thank Hilmar de Vries (Bartiméus, Amsterdam, The Netherlands), Rudo Poell (Royal Dutch Visio, Amsterdam, The Netherlands) and Dana Bruinenberg (Royal Dutch Visio, Amsterdam, The Netherlands) for their valuable assistance in this study. The authors also thank Michel Vloet (OrCam Technologies, Jerusalem, Israel) for his technical support on the OrCam devices. The OrCam devices were provided by OrCam Technologies (Jerusalem, Israel), and were returned upon completion of this study. The aforementioned organisations had no role in study design, data collection and analysis, decision to publish, or preparation of this manuscript.

Supporting Information

Additional Supporting Information may be found in the online version of this article:

Table S1. Questionnaires used in this study and their included items.

Table S2. Baseline scores of the NEI-VFQ, PAI and OFQ questionnaires.