**Psychology Health & Medicine – Methodological issue**

**Title page**

*Title:* **Establishing the usefulness of the GO-QOL in a UK hospital-treated population with thyroid eye disease in the CIRTED trial**

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On behalf of the CIRTED investigators

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**Abstract**

Thyroid eye disease (TED) is a potentially sight-threatening and cosmetically disfiguring condition arising in 25-50% of patients with Graves' hyperthyroidism. CIRTED is the first study to evaluate the long-term role of radiotherapy and prolonged immunosuppression with azathioprine in treating TED, one aim of which was to validate the use of the English version of GO-QOL in an UK population with TED. In a three stage design over a 48 week period, the GO-QOL was tested and compared to a general measure of quality of life (WHOQOL-Bref). In stage 1 utilising a standard 14 day test-retest design both GO-QOL subscales achieved Cronbach’s alphas demonstrating excellent validity and internal reliability (Visual Function 0.929 and 0.931; Appearance 0.888 and 0.906). In stage 2, Repeated Measures ANOVA demonstrated longitudinal validity, with both subscales of the GO-QOL showing significant change over time (Visual Function, η2=0.114, p<.001; Appearance, η2=0.069, p<.002). In stage 3 the GO-QOL showed discriminant validity at the week 48 time point, with the visual function subscale being able to detect changes in groups identified by clinicians (using BCCOM ratings of improvement or deterioration), while both subscales could detect group differences when based on participants’ subjective ratings of TED noticeability and severity. The results of this project provide support for the English translation of the GO-QOL as an outcome measure for patients with moderately severe active Graves' orbitopathy/TED.

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**Keywords**: thyroid eye disease (TED), Graves’ ophthalmopathy (GO), GO-QOL, adults, psychometrics, reliability, validity

**Introduction**

Graves’ ophthalmopathy or orbitopathy (GO) also known as thyroid eye disease (TED) is a potentially sight-threatening and cosmetically disfiguring condition arising in 25-50% of patients with Graves' hyperthyroidism (Bahn & Heufelder, 1993). The condition is rare and causes redness and grittiness of the eye and can lead to disfiguring swelling of the eyelids, proptosis (abnormal protrusion or displacement of the eye) and even blindness (Weetman, 1991). GO is an autoimmune disorder, linked to thyroid autoimmunity by autoantigens shared between the thyroid and the orbit of the eye (Perros, Crombie & Kendall-Taylor, 1995). Inflammatory processes are activated and fibroblasts in the eyes’ orbital tissue become stimulated leading to orbital tissue swelling, hyaluronan production, and expansion of the extraocular muscles, retro-orbital fat and connective tissues (Khoo & Bahn, 2007).

Currently the management of GO/TED is considered suboptimal, and available treatments do not specifically target the underlying pathogenic process (Bartalena et al, 2016). The Combined Immunosuppression and Radiotherapy in Thyroid Eye Disease (CIRTED) trial was designed to assess the effect of using radiotherapy and the immunosuppressive drug azathioprine in combination with standard prednisolone treatment (Rajendram et al, 2008). CIRTED is the first study to evaluate the long-term role of radiotherapy and prolonged immunosuppression with azathioprine in treating GO/TED.

It is well established that TED can have a major impact on quality of life, in particular disfiguring changes to the eyes and face which can have a direct impact on psychological health (Coulter, Frewin, Krassas & Perros, 2007). As the aim of treating GO/TED is to improve patients’ visual function as well as making them look and feel better, it is important to assess the patients’ perception of these markers as part of a clinical trial. The GO-QOL questionnaire was developed by Terwee, Gerding, Dekker, Prummel & Wiersinga (1998) as a TED specific quality of life questionnaire that can be used as an outcome measure for studies and may also be of use in clinical practice. Marcocciet al (2011) tested the use of selenium in mild GO and used the GO-QOL to evaluate quality of life outcomes. They showed a correlation that indicated that as participants’ improved with selenium treatment their quality of life also improved, as measured by the GO-QOL.

The aim of this three stage project was to validate the use of the English version of GO-QOL in an UK population with TED. The GO-QOL is available in both English and Dutch; the Dutch version having previously been validated in the Netherlands (Terwee et al, 1999). In the first stage of the work, the internal validity and test-retest study, the objective was to assess the consistency of the GO-QOL in measuring functional and appearance-related issues resulting from TED over a 14 day period where the expectation is that scores on both administrations should be correlated.

In the second stage, we measured longitudinal validity i.e. the responsiveness of the GO-QOL to changes in TED post-treatment against a more general measure of quality of life (in this case, the WHOQOL-Bref; The WHOQOL Group, 1998). This work had two aspects:

1. If the GO-QOL is valid (i.e. sensitive to changes in visual functioning and appearance as a result of TED) we would expect larger effect sizes for the GO-QOL than for the general quality of life measure (WHOQOL-Bref).
2. Furthermore, changes in clinical characteristics relating to visual functioning and appearance, as indicated by transitional variables, should be more closely associated with changes in scores on the relevant subscales of the GO-QOL than with the WHOQOL-Bref.

The third and final stage was to explore the extent to which the GO-QOL can demonstrate discriminant validity. That is, whether the GO-QOL can distinguish between patient populations based on either clinician ratings of improvement or participants’ subjective measures. Although it should be noted that subjective severity does not always correlate well with objective measures of disease and physicians’ assessments (Bessell, Dures, Semple & Jackson, 2012).

**Methods**

***Design***

The study protocol has been described in detail previously (Rajendram et al, 2008). In brief, CIRTED is a 2x2 factorial design, double-masked, multi-centre, randomized controlled trial (see Figure 1 for a diagram of the trial design).

***Ethical approval***

Ethical approval was from the UK’s National Health Service South West Central Bristol Research Ethics Committee (REC reference: 05/Q2006/62). Clinical Trial Authorisation was given by the Medicines and Healthcare products Regulatory Agency (MHRA, reference: 03299/0003/001-0001; ISRCTN22471573) with the University of Bristol acting as the legal sponsor. Research governance and local Research and Development approvals were obtained across all sites prior to the start of recruitment. All participants gave written informed consent.

[Figure 1 near here]

***Materials*** (see also Table 1)

The Graves' ophthalmopathy quality of life assessment (GO-QOL) questionnaire consists of two subscales, each comprising eight questions, on visual function and the psychological impact of changed appearance (Terwee et al, 1998).

The WHOQOL-Bref is a widely used and previously validated measure of quality of life consisting of 28 items which cover subjective overall quality of life and subjective overall health, plus items relating to domains of physical, psychological, social relationships and environment (The WHOQOL Group, 1998). The WHOQOL-Bref has been widely used with a range of populations and is reported to have good psychometric properties (Skevington, Lotfy & O’Connell, 2004).

Two transitional variables relating closely to the subscales of the GO-QOL were included at follow up as an external standard to identify changes post-treatment (labelled “T1” and “T2”). These variables were agreed with the authors of the GO-QOL as being suitable for this purpose. In order to explore the impact of TED and its treatment on psychological adjustment and daily functioning more broadly, Visual Analogue Scales (VAS) were also included in the study. VAS scales are easy for respondents to complete and are often used in clinical assessments (Carr, 1997).

Clinician ratings of disease severity and activity were also included. The Binary Composite Clinical Outcome Measure (BCCOM) a system of major and minor criteria used in previous TED trials (Prummel et al, 2004; Marcocci et al, 2001; Mourits et al, 2000). It is a clinician rating of improvement in the CIRTED trial used at 1 year post treatment to classify study participants’ treatment as being successful or not (see Appendix 1 for information on its derivation).

[Table 1 near here]

Protocol & Participants

All participants referred to a trial centre during the duration of the study were considered for inclusion. Participants were prescribed a high dose of tapering prednisolone at their initial enrolment visit. If they were eligible, responded to steroids and not excluded, participants were randomized into one of four trial arms (see Figure 1, Table 2, and CIRTED study protocol – Rajendram et al, 2008).

For validating test-retest reliability, participants completed the GO-QOL twice at a two-week interval, time point 1 (-2 weeks, i.e. enrolment into the study) and time point 2 (0 weeks, i.e. randomization into the study). Two weeks have previously been used by Terwee et al (1999) for assessing test-retest reliability, as it is long enough to avoid recall bias and short enough for patients not to experience clinically important changes in their condition. Data from participants for the longitudinal validity testing was collected at all four time points.

[Table 2 near here]

**Results**

The results are reported in the three stages that the work was undertaken. All analyses were undertaken using SPSS version 23.

***Stage 1 GO-QOL validation: internal validity and test re-test***

167 participants attended for study enrolment, while 133 attended the randomization appointment. Of these, 142 participants completed the GO-QOL at study enrolment, and 126 completed the GO-QOL at randomization (see Table 3).

[Table 3 near here]

Item response tables for time points 1 (-2 weeks, enrolment) and 2 (0 weeks, randomisation) were generated to identify patterns of responses and are shown in Tables 4 & 5.

[Tables 4 & 5 near here]

Not all participants cycled or drove, hence the lower number of responses to questions 1 and 2 on the Visual Function subscale at both time points. As per the questionnaire authors’ instructions (Terwee et al, 1998), adjustments for missing data were made when totalling the raw scores for each subscale prior to transforming them into a total score out of 100 for subsequent analyses.

Internal validity was assessed using Cronbach’s alpha calculations; at time 1 (-2 weeks, enrolment) the visual function subscale achieved a Cronbach’s alpha score of 0.929 (CI 0.909 – 0.944) while the appearance subscale recorded 0.888 (CI 0.857 – 0.912). At time 2 (week 0, randomization), the alpha results for visual function remained virtually unchanged at 0.931 (CI 0.912 – 0.946), while the appearance subscale improved slightly to 0.906 (CI 0.880 – 0.926). This indicates good internal validity of the subscales at both time points, with values over 0.7 generally acceptable for psychometric questionnaires (BPS, 1992).

Pearson correlation coefficients for both subscales were found to be highly significant (visual function, r=0.774, p<.001; appearance, r=0.862, p<.001), indicating the robust test-retest reliability of the GO-QOL subscales.

***Stage 2: Longitudinal validation of GO-QOL***

Longitudinal validation was performed using data from the 126 participants that were randomised into the trial (Table 6).

[Table 6 near here]

In this second stage, we measured the responsiveness of the GO-QOL to changes in TED post-treatment against the WHOQOL-Bref (a more general measure of quality of life). We hypothesized that, if the GO-QOL is sensitive to changes in visual functioning and appearance as a result of TED we would expect larger effect sizes for the GO-QOL than for the general measure (WHOQOL-Bref). Effect size was quantified using partial eta squared where the following thresholds can be used to determine effect size interpretation: η2 of less than 0.01 indicates an inconsequential effect, η2  between 0.01 and 0.09 a small effect, η2 between 0.09 to 0.25 a medium effect, η2 from 0.25 to 0.50 a large effect, and η2 over 0.50 a very large effect (Cohen, 1988).

Of the 126 randomized participants, 108 participants provided enough completed questionnaire data at the 12 week trial appointment, while 100 of those participants who attended the 48 week appointment completed the study measures (see Table 7).

[Table 7 near here]

For the sake of completeness, internal validity was assessed again using Cronbach’s alpha calculations; at both time 3 (short term 12 week trial appointment) and time 4 (long term 48 week appointment). At time 3 the visual function subscale achieved a Cronbach’s alpha score of 0.904 (CI 0.854 – 0.937) while the appearance subscale recorded 0.918 (CI 0.891 – 0.938). At time 4 the alpha result for visual function had reduced slightly to 0.887 (CI 0.823 – 0.928), while the appearance subscale remained largely unchanged at 0.915 (CI 0.886 – 0.937). As before, with all values over 0.7 this indicates good internal validity of the subscales at both time points.

Repeated measures ANOVA (with time as 3 level factor, i.e. -2, 12, and 48 weeks) revealed significant changes over time for both GO-QOL subscales (Visual Function, *p*=0.001, Appearance, *p*=0.002), and for the psychological domain of the WHOQOL (p=0.002). Effect sizes varied between the subscales for the GO-QOL with only Visual Function recording a medium effect size (η2=0.114); the Appearance subscale showed a small effect size (η2=0.069). Effect sizes for the WHOQOL subscales were all small (psychological, η2=0.064; physical, η2=0.037; social, η2=0.041; environment, η2=0.043) (Table 8).

[Table 8 near here]

We also hypothesized that changes in clinical characteristics relating to visual functioning and appearance, as indicated by the transitional variables, should be more closely associated with changes in scores on the relevant subscales of the GO-QOL than with the WHOQOL-Bref. As expected since all study participants were taking steroid treatment, correlation calculations showed positive correlations between both the transitional variables (T1 and T2) and the GO-QOL appearance subscale at the 12-week time point, while the Visual Function subscale only correlated with T1 (Table 9). Similarly, three of the WHOQOL-Bref subscales significantly correlated with T1, while all four subscales significantly correlated with T2. At 48-weeks significant correlations were observed only for the appearance subscale of the GO-QOL with the transitional variables.

We included two Visual Analogue Scales to explore the impact of treatment for TED on psychological adjustment and daily functioning more broadly. At 12 weeks significant correlations were observed between both VAS scales and the appearance subscale of the GO-QOL. Three of the four domains of the WHOQOL-Bref significantly correlated with the VAS for participant perceived noticeability of TED (psychological, physical and environment), while a different trio of domains significantly correlated with the VAS for participant ratings of severity of TED (physical, social and environment). At 48 weeks, the only significant correlations observed were between both VAS scales and the appearance subscale of the GO-QOL.

[Table 9 near here]

***Stage 3 Exploring discriminant validity and the relationships between subjective and objective measures of disease severity***

The third and final stage was to explore the discriminant validity of the GO-QOL as well as the relationships between the subjective and objective clinical measures for TED used in this study.

Clinician ratings on BCCOM at the 48 week time point were used to split participants into two groups: condition has deteriorated (-1, N=58), or improved (+1, N=36; Table 10). Analyses were then undertaken to determine any differences in mean GO-QOL scores between these groups. It might be expected that there would be differences between the mean changes (pre/post-treatment) in GO-QOL scores depending on whether clinicians reported visual functioning/appearance having either deteriorated or improved; i.e. we would expect larger increases in scores on the GO-QOL for those participants where clinician’s reported an improvement compared to those for whom they did not.

[Table 10 near here]

Independent samples t-tests were undertaken to determine differences between the groups on both the GO-QOL and WHOQOL measures while effect sizes were calculated using Cohen’s *d* (Cohen, 1988). For broad interpretation purposes threshold values for statistically significant effects for the statistic, *d* are: 0 < *d* < 0.1 a trivial effect; 0.1 < *d* < 0.2 a small effect; 0.2 < *d* < 0.5 a moderate effect; 0.5 < *d* < 0.8 a medium size effect; 0.8 < *d* < 1.3 a large effect; 1.3 < d < 2.0 a very large effect; while *d* > 2.0 is a huge effect. However, these are, at best, guidelines and the value of *d* is very much context dependent.

The independent samples t-tests showed a significant difference for the Visual Function subscale at 48-weeks (*p*=0.006) with a medium effect size (d=0.6; Table 11). Although no similar statistically significant difference was seen for the Appearance subscale at the same time point a moderate effect size was recorded (d=0.3). Similarly, no statistically significant differences between WHOQOL domain scores were observed for the BCCOM groups at the same time point, and effect sizes ranged from small (d=0.2 for the physical and environment domains) to moderate and medium (d=0.4 and 0.5 for the psychological and social domains, respectively).

[Table 11 near here]

The cohort was also split in relation to the scores achieved on the two Visual Analogue Scales, where 5.1 was used as the cut off to identify those of the 100 (where we had these data at week 48) who rated themselves as still having TED that was either noticeable or severe. When analysed according to these groupings, the subscale domains of the WHOQOL still showed no significant differences, but in relation to perceived noticeability of TED both subscales of the GO-QOL showed statistical significantly differences (p<.001) with a medium effect size for the Visual Function subscale (d=0.8) and a very large effect size for the Appearance subscale (d=1.6; Table 12). In relation to groupings based on perceived severity of TED both subscales of the GO-QOL showed statistical significantly differences (p<.001), and again a medium effect size for the Visual Function subscale (d=0.8) and a very large effect size for the Appearance subscale (d=1.2; Table 12).

[Table 12 near here]

So it would seem that the GO-QOL does indeed demonstrate discriminant validity, although there is some suggestion in these data, that it may be dependent on the grouping variable utilised. The clinician ratings (BCCOM) and participant subjective ratings (VAS scales) were broadly similar in their results for the Visual Function component of the GO-QOL, but to distinguish in relation to appearance issues, it would seem that using participant ratings results in stronger, statistically significant differences.

**Discussion**

The results of this project support both the internal validity and reliability of the English translation of the GO-QOL as an outcome measure for patients with moderately active Graves' orbitopathy. Longitudinal validity has also been confirmed, with the GO-QOL being more sensitive to changes in TED over time than the more general WHOQOL-Bref., whilst also being associated with generally larger effect sizes, with both measures show an improving trend for the study participants over time. This longitudinal pattern of change for the subscales of the GO-QOL has also been shown over a 24 week period in a recent study testing Teprotumumab for thyroid-associated ophthalmology (Smith et al, 2017). While both subscales of the GO-QOL showed significant change over time in Smith et al’s study, the Visual Function subscale was the one with the greatest change over time, as also suggested in our data.

The analysis with the transitional variables is more equivocal. The hypothesized changes in clinical characteristics were generally associated with significant correlations for both quality of life measures with the transitional variables at the 12 week time point. By the 48 week time point only the GO-QOL appearance subscale recorded significant correlations with the T1 (my eye condition causes me pain and discomfort) and T2 (my eye condition limits my ability to do the things I want to do). Given the focus of the T1 and T2 variables on what might be considered to be issues more associated with visual function, it is curious that the observed correlations are with the appearance subscale. The Visual Analogue Scales in the study are more focused on what might be considered appearance issues – the perceived noticeability and severity of the TED. It is probably no surprise that these were significantly correlated with the appearance subscale of the GO-QOL at both the 12 week and 48 week study time points.

The GO-QOL has also demonstrated discriminant validity, with the visual function subscale being able to detect changes in groups identified by clinicians (using BCCOM ratings of improvement or deterioration), while both subscales could detect group differences when based on participants’ ratings of TED noticeability and severity. It is worth noting that BCCOM is constructed to include factors such as diplopia which would correlate much better with function than with appearance (Appendix 1). It has been suggested that patients with TED overrated the extent to which their appearance was affected, while endocrinologists underrated it (Terwee et al, 2003). Of course, the measures employed do not necessarily take into account the reference point, i.e. patients are possibly comparing themselves prior to TED, or with other people who do not have the disease at all, whereas clinicians could be comparing across individuals who have TED. Without interviewing the participants and clinicians concerned it is impossible to know.

Historically health care professionals have found it difficult to engage with patients’ concerns regarding their looks and body image (Bessell et al, 2012), quantifying these concerns with a measure like the GO-QOL may help doctors’ engagement with their patients concerns and target therapy accordingly.

**Competing interest statement:** The authors of this study have no competing interests to declare.

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**APPENDICES**

**Appendix 1**

**Binary composite clinical outcome measure (BCCOM)**

The BCCOM is a binary outcome with a positive result with no deterioration vs no change or deterioration in any component of the score.

***Major Criteria***

▪ An improvement of ≥ 1 grade in diplopia score\*

▪ An improvement of >8 degrees of eye movement in any direction#

▪ A reduction of ≥ 2 mm in proptosis

***Minor Criteria***

▪ A reduction of ≥ 2 mm in lid aperture

▪ An improvement of ≥ 1 grade in soft tissue involvement†

▪ An improvement in best-corrected visual acuity of ≥ 1 line on the Snellen chart·

▪ Subjective improvement^

***Response to Treatment***

*Very good*: ≥ 2 major criteria

*Good*: 1 major criterion

*Fair*: ≥ 2 minor criteria

*No Change:* 1 minor criterion

*Worse:* Deterioration in at least 1 major or 2 minor criteria

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^ Marcocci, C., Bartalena, L., Tanda, M. L., Manetti, L., Dell-Unto, E., Rocchi, R., … Pinchera, A. (2001). Comparison of the effectiveness and tolerability of intravenous or oral glucocorticoids associated with orbital radiotherapy in the management of severe Graves' ophthalmopathy: results of a prospective, single-blind, randomized study. *Journal of Clininical Endocrinology & Metabolism,* 86(8), 3562-7. doi: <http://dx.doi.org/10.1210/jcem.86.8.7737>

Table 1. Overview of the standardised questionnaires.

|  |  |
| --- | --- |
| **Measure** | **Description** |
| **Standardised questionnaires** | |
| GO-QOL | - TED specific quality of life measure, validated in the Netherlands  - 2 subscales: ‘visual function’ & ‘appearance’ comprising 8 questions each  - each item is scored as follows: 1= not impaired; 2=a little impaired; 3=severely impaired  - raw scores are transformed to give a total out of 100 for each subscale  - higher scores indicate greater quality of life |
| WHOQOL-Bref | - general quality of life measure, validated for use in the UK  - 4 subscales: psychological, physical, social and environmental; 28 questions in total  - scored on a 5-point Likert scale  - raw scores are transformed to give a total out of 100 for each subscale  - higher scores indicate greater satisfaction with life |
| **Transitional variables** | |
| T1 | - is a single item: “My eye condition causes me physical pain/discomfort”  - scored as follows: 1= never/almost never; 2=sometimes; 3=often; 4=almost always |
| T2 | - is a single item: “My eye condition limits my physical ability to do the things I want to do”  - each item is scored as follows: 1= never/almost never; 2=sometimes; 3=often; 4=almost always |
| **Visual Analogue Scales** | |
| Noticeability | - is a single item: “How noticeable do you feel your thyroid eye disease is to other people?”  - scored on a 10 cm line with the following anchors:  Not at all noticeable |..........| Very noticeable  - scored 0-10  - higher scores indicate greater distress |
| Severity | is a single item: “How severe do you feel your thyroid eye disease is?”  - scored on a 10 cm line with the following anchors:  Not very severe |..........| Extremely severe  - scored 0-10  - higher scores indicate greater distress |
| **Clinical rating** | |
| Binary Composite Clinical Outcome Measure (BCCOM) | - clinician-rating  - binary composite outcome score with a positive result (improvement with no concomitant deterioration) versus no change or any deterioration (see Appendix 1)  - deteriorated (-1), improved (1) |

Table 2. Data collection time points and their relation to participant trial appointments, plus study measures used at each time point, and number of potential participants attending each appointment (N)

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  |  | Trial Appointment | Study measures | N |
| Study time points | **1** | -2 weeks Enrolment | GO-QOL, WHOQOL, VAS, demographic data | 167 |
| **2** | 0 weeks Randomization | GO-QOL | 133 |
| **3** | 12 weeks  Short term | GO-QOL, WHOQOL, transition variables, VAS | 108 |
| **4** | 48 weeks  Long term | GO-QOL, WHOQOL, transition variables, VAS, BCCOM | 102 |

***Table 3. Participant characteristics for Stage 1 GO-QOL Validation: internal validity and test re-test***

|  |  |  |
| --- | --- | --- |
|  | **Study time point** | |
| **Variable** | **Enrolment (-2 weeks)** | **Randomization (week 0)** |
| **Sample size** | 142 | 126 |
| **Age** | 48.02±11.44 | 48.27±11.40 |
| **Sex** (male/female) | 41/101 | 34/92 |
| **Ethnicity** |  |  |
| Caucasian/Black | 99/16 | 87/14 |
| Asian/Oriental | 11/4 | 10/4 |
| Other (or not stated) | 12 | 11 |

***Table 4.******Item response table for GO-QOL at time point 1 (-2 weeks, enrolment).***

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | **N** | **Min**  **score** | **Max**  **score** | **Mean**  **score** | **SD** |
| **Visual Function subscale** | | | | | |
| VF1 Cycling | 87 | 1 | 3 | 2.43 | 0.80 |
| VF2 Driving | 103 | 1 | 3 | 2.19 | 0.81 |
| VF3 Walking indoors | 139 | 1 | 3 | 2.71 | 0.54 |
| VF4 Walking outdoors | 141 | 1 | 3 | 2.49 | 0.64 |
| VF5 Reading | 140 | 1 | 3 | 2.01 | 0.70 |
| VF6 Watching TV | 141 | 1 | 3 | 2.12 | 0.68 |
| VF7 Hobbies | 122 | 1 | 3 | 2.29 | 0.77 |
| VF8 Interference with daily life | 141 | 1 | 3 | 2.10 | 0.77 |
| **Appearance subscale** | | | | | |
| App9 Change in appearance | 142 | 1 | 3 | 1.42 | 0.54 |
| App10 Feeling watched | 142 | 1 | 3 | 2.14 | 0.77 |
| App11 Unpleasant reactions | 140 | 1 | 3 | 2.49 | 0.68 |
| App12 Impact on self-confidence | 142 | 1 | 3 | 1.73 | 0.72 |
| App13 Feeling of social isolation | 142 | 1 | 3 | 2.50 | 0.69 |
| App14 Influence on friendships | 142 | 1 | 3 | 2.46 | 0.73 |
| App15 Less often in photos | 141 | 1 | 3 | 1.90 | 0.85 |
| App16 Camouflaging appearance | 142 | 1 | 3 | 2.04 | 0.82 |

Key: N= number of responses; SD = standard deviation

***Table 5*. *Item response table for GO-QOL at time point 2 (week 0, randomisation)***

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
|  | **N** | **Min**  **Score** | **Max**  **Score** | **Mean**  **score** | **SD** |
| **Visual Function subscale** | | | | | |
| VF1 Cycling | 75 | 1 | 3 | 2.59 | 0.68 |
| VF2 Driving | 93 | 1 | 3 | 2.38 | 0.75 |
| VF3 Walking indoors | 125 | 1 | 3 | 2.72 | 0.47 |
| VF4 Walking outdoors | 126 | 1 | 3 | 2.61 | 0.61 |
| VF5 Reading | 126 | 1 | 3 | 2.23 | 0.69 |
| VF6 Watching TV | 126 | 1 | 3 | 2.36 | 0.66 |
| VF7 Hobbies | 110 | 1 | 3 | 2.47 | 0.69 |
| VF8 Interference with daily life | 126 | 1 | 3 | 2.28 | 0.73 |
| **Appearance subscale** | | | | | |
| App9 Change in appearance | 126 | 1 | 2 | 1.50 | 0.50 |
| App10 Feeling watched | 126 | 1 | 3 | 2.17 | 0.76 |
| App11 Unpleasant reactions | 125 | 1 | 3 | 2.58 | 0.60 |
| App12 Impact on self-confidence | 126 | 1 | 3 | 1.83 | 0.75 |
| App13 Feeling of social isolation | 126 | 1 | 3 | 2.59 | 0.65 |
| App14 Influence on friendships | 126 | 1 | 3 | 2.45 | 0.77 |
| App15 Less often in photos | 126 | 1 | 3 | 2.01 | 0.83 |
| App16 Camouflaging appearance | 126 | 1 | 3 | 2.05 | 0.81 |

Key: N= number of responses; SD = standard deviation

Table 6. CIRTED trial allocation groups.

|  |  |  |
| --- | --- | --- |
| **Group** | **Allocation** | **N=126 randomised** |
| 1 | Radiotherapy and Azathioprine | 31 (24.6%) |
| 2 | SHAM Radiotherapy and Azathioprine | 31 (24.6%) |
| 3 | Radiotherapy and PLACEBO | 32 (25.3%) |
| 4 | SHAM Radiotherapy and PLACEBO | 32 (25.3%) |

***Table 7. Participant characteristics for Stage 2: Longitudinal validation of GO-QOL***

|  |  |  |
| --- | --- | --- |
| **Variable** | **Study time point** |  |
|  | **Short term (12 weeks)** | **Long term (48 weeks)** |
| **N** | 108 | 101 |
| **Age** | 49.48±10.82 | 49.88±10.46 |
| **Sex** (male/female) | 29/79 | 27/73 |
| **Ethnicity** |  |  |
| Caucasian/Black | 79/13 | 72/10 |
| Asian/Oriental | 8/4 | 9/4 |
| Other (or not stated) | 4 | 5 |

***Table 8: Mean scores* ± *standard deviations for G0-QOL and WHOQOL-Bref across study time points, results for RM ANOVA with effect sizes***

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| *Study measure* | *Enrolment (-2 weeks)* | *Short term*  *(12 weeks)* | *Long term (48 weeks)* | *RM ANOVA F & p value* | *Effect size*  η2 |
| **GO-QOL subscales:** | | | | | |
| Visual Function | 65.94 ±28.49 | 72.49 ±26.64 | 76.11 ±24.73 | 11.034  p<0.001 | 0.114 Medium |
| Appearance | 54.1  ±26.54 | 58.81 ±27.25 | 60.84 ±28.75 | 13.061  p=0.002 | 0.069  Small |
| **WHOQOL-Bref subscales:** | | | | | |
| Physical | 58.69 ±22.64 | 60.93 ±21.01 | 63.46 ±20.67 | 3.18  p=0.044 | 0.037  Small |
| Psychological | 52.66 ±21.35 | 52.14 ±23.04 | 57.79 ±20.20 | 6.53 p=0.002 | 0.064  Small |
| Environment | 66.60 ±20.03 | 67.81 ±18.92 | 69.15 ±17.13 | 1.97 p=0.143 | 0.043  Small |
| Social Relationships | 66.08 ±20.81 | 60.83 ±21.53 | 65.54 ±21.77 | 2.99 p=0.054 | 0.041  Small |

Table 9. Transition scores and visual analogue scales correlated with GO-QOL and WHOQOL scores.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  | **T1** | **T2** | **VAS**  **Noticeability** | **VAS**  **Severity** |
| **Week 12** GO-QOL subscales: | | | | |
| Visual Function | -0.413\*\* | 0.015 | -0.108 | -0.117 |
| Appearance | -0.409\*\* | -0.580\*\* | -0.587\*\* | -0.552\*\* |
| **Week 12** WHOQOL-Bref subscales: | | | | |
| Psychological | -0.284\*\* | -0.475\*\* | -0.399\*\* | -0.127 |
| Physical | -0.451\*\* | -0.612\*\* | -0.383\*\* | -0.466\*\* |
| Social | -0.193 | -0.307\*\* | -0.167 | -0.262\* |
| Environment | -0.295\*\* | -0.508\*\* | -0.346\*\* | -0.253\*\* |
|  |  |  |  |  |
| **Week 48** GO-QOL subscales: | | | | |
| Visual Function | -0.017 | -0.162 | 0.014 | -0.046 |
| Appearance | -0.389\*\* | -0.574\*\* | -0.723\*\* | -0.678\*\* |
| **Week 48** WHOQOL-Bref subscales: | | | | |
| Psychological | 0.19 | 0.017 | -0.037 | 0.148 |
| Physical | 0.149 | -0.005 | -0.106 | -0.024 |
| Social | 0.158 | 0.083 | -0.151 | 0.089 |
| Environment | 0.199 | 0.025 | -0.07 | 0.006 |

Key: \*\* = significant at p<.001, \*= significant at p<.05

***Table 10. Participant characteristics for Stage 3: Exploring discriminant validity based on BCCOM***

|  |  |  |
| --- | --- | --- |
| **Variable** | **Study time point 48 weeks** |  |
|  | **-1 deteriorated** | **+1 improved** |
| N | 58 | 36 |
| Age | 50.31±10.72 | 48.72±10.53 |
| Sex (male/female) | 12/46 | 12/24 |
| **Ethnicity** |  |  |
| Caucasian/Black | 46/4 | 22/6 |
| Asian/Oriental | 5/2 | 4/1 |
| Other/missing | 1/0 | 3/0 |

***Table 11. Comparing means of GO-QOL & WHOQOL subscales according to BCCOM group (improved n=36, deteriorated n=58) at 48 week time point***

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **GO-QOL** | **BCCOM group** | **Mean±sd** | **p value** | **Cohen’s d (CI)** |
| Visual Function | +1 (improved) | 83.06±21.06 | 0.006 | 0.6 medium (CI 0.19 – 1.04) |
|  | -1(deteriorated) | 67.86±26.46 |
| Appearance | +1 (improved) | 64.89±28.33 | 0.152 | 0.3 moderate (CI -0.13 – 0.73) |
|  | -1 (deteriorated) | 55.89±28.26 |
| **WHOQOL** |  |  |  |  |
| Psychological | +1 (improved) | 51.73±22.32 | 0.138 | 0.4 moderate (CI -0.03 – 0.83) |
|  | -1(deteriorated) | 59.61±19.29 |
| Physical | +1 (improved) | 58.08±22.27 | 0.537 | 0.2 small (CI -0.23 – 0.63) |
|  | -1(deteriorated) | 61.63±22.21 |
| Social | +1 (improved) | 56.30±23.75 | 0.083 | 0.5 medium (CI -0.89 – -0.05) |
|  | -1(deteriorated) | 67.34±22.96 |
| Environment | +1 (improved) | 65.08±19.66 | 0.356 | 0.2 small  (CI -0.23 – 0.63) |
|  | -1(deteriorated) | 69.74±19.68 |

***Table 12. Comparing means of GO-QOL subscales according to VAS group at 48 week time point***

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **GO-QOL** | **VAS noticeability** | **Mean±sd** | **p value** | **Cohen’s d (CI)** |
| Visual Function | <5.0 (improved, n=45) | 84.63±20.72 | 0.001 | 0.8 medium (CI 0.41 – 1.19) |
|  | >5.1(noticeable, n=54) | 66.37±26.36 |
| Appearance | <5.0 (improved, n=45) | 79.86±18.17 | 0.001 | 1.6 very large (CI 1.17 – 2.03) |
|  | >5.1(noticeable, n=55) | 45.23±25.11 |
|  | **VAS severity** |  |  |  |
| Visual Function | <5.0 (improved, n=53) | 84.00±20.46 | 0.001 | 0.8 medium  (CI 0.41 – 1.19) |
|  | >5.1(severe, n=45) | 64.54±26.71 |
| Appearance | <5.0 (improved, n=53) | 74.29±23.14 | 0.001 | 1.2 large (CI 0.77 – 1.63) |
|  | >5.1(severe, n=46) | 46.06±25.60 |