Questionnaire to Assess Life Impact of Treatment by Intravitreal Injections (QUALITII): Development of a patient-reported measure to assess treatment burden of repeat intravitreal injections

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ABSTRACT

Objective To understand patient burden of treatment of repeated intravitreal injections (IVI) in the management of exudative retinal diseases.

Methods and analysis Participants were sampled from a large urban retina specialty practice in Houston, Texas, USA, based on history of ongoing receipt of IVI. The 50-item Questionnaire to Assess Life Impact of Treatment by Intravitreal Injections questionnaire was developed to evaluate the patient experience including discomfort, anxiety, inconvenience and satisfaction. Categorical principal components analysis (CATPCA) was performed to assess construct validity and internal consistency. A subset of these items was used to establish a measure of total treatment burden, referred to as the IVI Treatment Burden Score (TBS).

Results 142 patients participated in this study. CATPCA analysis revealed five dimensions of patient burden: disruption of normal routine or capacity, anxiety, frequency of visits, chronicity of disease and perceived treatment value or satisfaction. Together, these dimensions accounted for 67% of variance explained. Cronbach’s alpha was 0.97. The most frequently cited cause of discomfort was the feeling after anaesthetic wore off. The most common source of anxiety was fear of injection and associated discomfort or pain. Regarding inconvenience, patients reported temporary postinjection debilitation, requiring an average of 8 hours for recovery per treatment. The most frequently identified sources of satisfaction were confidence in the provider or treatment and interactions with staff.

Conclusions Understanding and quantifying the patient burden of repeat IVI could reveal opportunities to improve delivery of care and optimise patient experiences.

Key messages

What is already known about this subject?
- Repeat intravitreal injections (IVIs) pose a substantial burden on patients, which may influence treatment adherence and thereby visual outcomes.

What are the new findings?
- We report the first phase of Questionnaire to Assess Life Impact of Treatment by Intravitreal Injections, a survey to evaluate the patient experience of receiving IVI. We present the Treatment Burden Score, a single score based on patient-reported data that serves as a validated measure of patient burden.

How might these results change the focus of research or clinical practice?
- Understanding the nature and magnitude of patient burden for repeat IVI could reveal opportunities to change practice patterns to optimise patient experiences and outcomes.

INTRODUCTION

Over the last two decades, intravitreal injections (IVIs) have become one of the most common procedures performed in medicine.12 This is in part due to the advent of antivascular endothelial growth factor (VEGF) agents and the large number of exudative retinal pathologies now recognised to be driven by upregulation of VEGF. Injection of these anti-VEGF drugs has become the cornerstone of therapy for a number of exudative retinal disorders, including neovascular age-related macular degeneration (nAMD),3 diabetic macular oedema (DME),4 diabetic retinopathy (DR)5 and retinal venous occlusive disease (RVO).6 Worldwide, the prevalence of each of these conditions ranges in the tens of millions.56 Accordingly, it is estimated that over 20 million IVIs are administered on an annual basis.1
Several randomised studies have demonstrated the efficacy of anti-VEGF agents for nAMD, DME, DR, and RVO. To manage pathological neovascularisation and exudation in these disorders, treatment typically involves repeated anti-VEGF injections over an indefinite time course. Outcomes can thereby depend greatly on patient adherence, which can in turn be influenced by patient experiences and perceptions of satisfaction versus treatment burden. There is, therefore, great interest in optimising patient experiences, towards the goal of maximising adherence.

Studies of the experiences of patients undergoing repeated IVI have used long-standing tools such as the macular disease Treatment Satisfaction Questionnaire (MacTSQ), designed to examine satisfaction with IVI in the treatment of nAMD. Interestingly, others have reported that patients with high levels of satisfaction measured by MacTSQ continued to identify several burdensome aspects of treatment. In one single-centre retrospective study, the most frequent reason patients discontinued treatment was fear of injection. Thus, careful examination of the factors underlying patient burden could yield further insight into areas of improvement in the management of exudative retinal disease.

While multiple authors have highlighted the challenging aspects of patient experiences with IVI, no single quantitative model of patient treatment burden has emerged. In an effort to balance vision benefits against associated burdens of repeated IVI for exudative retinal diseases, the exact frequency of injections needed to maintain optimal clinical outcomes has been the subject of several investigations. To facilitate this effort, it would be useful for practitioners who manage exudative retinal diseases to understand the magnitude of patient treatment burden associated with repeated IVI.

The current study uses a patient survey to characterise burdensome aspects of repeated IVI and to develop a quantitative measure of patient burden. Herein, we describe the development and psychometric analysis of the Questionnaire to Assess Life Impact of Treatment by Intravitreal Injections (QUALITII). We establish a single score of treatment burden, the IVI Treatment Burden Score (TBS) from this survey. Measurements such as the TBS could inform approaches to optimise patient experiences and adherence for a broad range of retinopathies and/or for comparative analysis of retinal therapeutics.

**MATERIALS AND METHODS**

**Participants**

Participants from a large urban retinal specialty clinic (Retina Consultants of Houston, Houston, Texas, USA) were invited by posted signage to self-administer the electronic survey on the day of presentation to the clinic. No prior medical record review was performed for this study, and no patient-specific identifying information was monitored or recorded.

**Psychometric analysis**

From the original 50-item questionnaire, psychometric analysis was performed on a subset of 20 items, estimated to address the most salient aspects of burden (online supplemental appendix 2A). These items (hereafter referred to as 20 ‘major’ items) assessed aspects of the patient experience that were hypothesised to influence perceived patient burden, including discomfort, anxiety, inconvenience and satisfaction associated with treatment. Response scales were a combination of multiple-choice items ranked on a 7-point Likert scale with graded continuum, multiple-choice items without ranked scales, and free response. The variety of response styles was selected to account for the heterogeneous nature of question items. The wide scope of subject matter further lent itself to a relatively long list of question items. Because of this length, we attempted to control for the tendency of respondents to select answers reflexively, by intermittently reversing the direction of graded response scales, where relevant.

**IVI TBS**

Following psychometric analysis as described above, a refined subset of 9 items (Q17, Q21, Q23, Q24, Q31, Q32, Q33, Q41, Q42) was used to construct the TBS, a numeric score of the perceived burden of treatment. The response scale of five of these items (Q21, Q23, Q24, Q41, Q42) directly correlated with burden, and the response...
scale of the remaining four items (Q17, Q31, Q32, Q33) inversely correlated with burden. Calculation of the TBS required arithmetic addition of the five subscores from the former group and four subscores from the latter, which were obtained by subtracting the patient-reported score from the maximum possible score on the scale provided (figure 1).

Statistical analysis and reporting

Averages reported are arithmetic means, unless otherwise described. Free response answers, along with answers involving a binned response range, were individually reviewed and converted to numerical values for statistical analysis, where relevant. Questions with specific answer choices that could not be immediately converted to a single numeric value were evaluated case by case, to determine how and whether to use these answer choices in estimating the average value of response. We tested normality of distribution of the TBS by K-S test. Other statistical tests are described in the Results section.

RESULTS

Participant demographics

A total of 142 patients completed this initial, single-centre phase of the QUALITII survey. The mean age of participants was 65.6 years, with 45% identifying as male and 55% female. Retinopathies represented among patients included AMD (31%), DME or DR (38%), RVO 9.2% or another condition (11.3%), with 10.6% reporting uncertain diagnosis. The demographic profile of participants is summarised in table 1.

Patient-reported details of IVI treatment protocols

The median number of years since diagnosis was 2.5. Most received anti-VEGF (71%), while 3.4% received dexamethasone or another medication, and 25.7% were unaware of which medication they receive. The median frequency of injection was once every 4.5 weeks, with a range of every 4–5 weeks (64.8%) up to every 13–16 weeks (1.4%), and 4.2% reporting injections on an as needed or PRN (pro re nata) basis. Treatment protocols experienced by respondents included a variety of numbing approaches, most frequently numbing gel or drops, and lasted a median of 2 hours. Table 2 summarises the characteristics of treatment protocols reported.
Patient-reported treatment burdens of repeated IVI

The broad categories of burden examined by the QUAL-ITII survey included discomfort, anxiety and burdens on time and functional capacity. A majority of participants (75.8%) experienced at least some discomfort with treatment. Most (52.3%) rated discomfort between 0 and 2 at the lower range of the response scale, but a large proportion (28.1%) rated discomfort between 4 and 6, where 6 represented very significant discomfort.

The three most uncomfortable aspects of treatment identified were use of betadine prior to injection, injection itself, and the feeling after anaesthetic wore off. Corresponding levels of anxiety experienced before, during, and after treatment were each rated at a median value of 4, on a 7-point scale, where a maximum score of 6 represented no anxiety. Therefore, patients reported moderately low but persistent anxiety throughout treatment. Among patients reporting anxiety, the most common reasons were fear of the injection itself (30.8%), fear of discomfort/pain from injection (23.1%) and fear of losing sight (12.8%).

In our study, a substantial component of patient burden arose from a combination of discomfort and the time and functional burdens imposed by treatment. This was conceptualised as disruption to normal activity or capacity. Though the average rating of discomfort was low to moderate (2.41 on a 7-point scale, where a maximum value of 6 represented very significant discomfort), a sizeable proportion (38.3%) of patients responded that injections resulted in restriction of usual activity or capacity, with a median 8-hour recovery period after each treatment.

Linear regression analysis showed that perceived burden decreased with increased satisfaction (p=0.0139, R²=0.077). The most frequently cited reasons for satisfaction in our study were confidence in the provider or treatment and pleasant interactions with staff. Interestingly, our data suggest that even patients who reported maximum satisfaction experienced non-negligible levels of distress and considerable time sacrifices. Patient responses to the 20 major question items assessing aspects of burden and satisfaction levels, are summarised in Table 2.
Psychometric analysis of the QUALITII survey
CATPCA was performed on a selection of 20 major question items from the QUALITII survey, thought to broadly represent components of perceived treatment burdens. Five dimensions of burden emerged: disruption to normal activity or capacity, anxiety, frequency of visits, chronicity of disease, and satisfaction/perceived value of treatment. Cronbach’s alpha was 0.97, suggesting a high level of internal consistency and reliability. Together, these dimensions accounted for 67% of variance explained. Proportions of variance explained by each dimension are provided in online supplemental appendix 4. In our analysis, three items did not load (Q30, Q39, Q43). A table of loading factors for each item is provided in online supplemental appendix 2B. Exclusion of all three of these items increased Cronbach’s alpha to 0.98, with variance explained rising to 76%. On individual review, these items were retained in our model due to clinically significant content.

IVI TBS
Of the 20 items underlying the five dimensions of patient burden identified by CATPCA, we identified a refined subset of nine items (Q17, Q21, Q23, Q24, Q31, Q32, Q33, Q41, Q42) whose response scales could be used as direct measures of burden. In general, response points derived from each of these items are summed towards the total TBS. The method of calculating the TBS from select QUALITII questionnaire items is provided in figure 1. Cronbach’s alpha for items included in the TBS was 0.75. In this study, the average TBS was 21.6 (actual range of 1–46), with a theoretical range of 1–54 (where 54 represents maximum burden). The SD of TBS scores across patients was 9.3, with a normal distribution. Average TBS did not differ between patients who received fewer than 20 injections vs at least 20 or greater injections (22.5 vs 20.5, student’s t-test, p=0.37). Further, logistic regression analysis showed that a higher TBS was a risk factor for the likelihood that patients would feel bothered enough by treatment to consider discontinuing (n=76, OR 1.26, 95% CI 1.0722 to 1.4885).

DISCUSSION
This paper reports the development and validation of the QUALITII survey, with the aim of understanding patient treatment burdens of repeated IVI. Principal components analysis of select survey items demonstrated five underlying dimensions of burden: disruption of normal routine or capacity, anxiety, frequency of visits, chronicity of disease and perceived treatment value or satisfaction. These items together demonstrated a high level of internal consistency and accounted for 67% of variance. The first dimension of burden identified (disruption of normal routine or capacity) consisted of items assessing severity of discomfort, length of time discomfort lasted, and perceptions of time consumed by treatment. Our results suggest that rather than the incidence of discomfort or incapacitation alone, this component of burden arises from the amount of time spent experiencing discomfort or debility. For patients who experienced any incapacitation, an average of 8 hours of time was required to recover normal activity levels. The implied functional time lost over the course of ongoing IVI could amount to substantial cumulative burden for patients.
burden is to create a postprocedure pain management plan spanning this time period.

The second dimension of burden that emerged was anxiety. In this study, patients reported moderate but persistent levels of anxiety throughout the procedure, most commonly induced by fear of the injection itself and fear of discomfort/pain, in accordance with other studies.17 25 This poses a challenge for strategising immediate interventions, because current technologies require repeat injections for optimal management. That said, patients reported that factors such as clinic delays can worsen anxiety, and factors such as adequate pain medication during the procedure and having the procedure explained can lessen anxiety. Thus, in addition to maximising pain control intraprocedurally and postprocedurally, two simple interventions to reduce patient anxiety could be to minimise treatment delays and to communicate consistently with patients through the procedure.

The third and fourth dimensions of burden in our study were frequency of visits and chronicity of disease, respectively. Question items that loaded onto these measures into estimates of burden. Ongoing treatment. As the MacTSQ15 and RetTSQ, 26 address burdensome aspects of treatment, including side effects, apprehension and pain. The TBS is a quantitative measure of patient burdens of repeated IVI that complements the above satisfaction scores, with additional emphasis on understanding the impact of treatment on quality of life. Moreover, the TBS describes the burdens of a shared treatment modality across nAMD, DME, DR and RVO. Thus, its applicability as a single score extends to a wide range of retinal disease. Such measures could be used to understand how commonly provided procedures impact patients and could potentially be used for comparisons of different retinal therapeutics.

The current study is limited by participation of patients from a single clinical practice and assessment of a predominantly English-speaking population. Further, the majority of represented patients in this study reported receiving monthly injections, perhaps introducing selection bias for patients with lower perceived burden, who could tolerate more frequent treatment. The next phases of the QUALITII study expand to multiple treatment centres and aim to diversify patient profile. Future studies may further address how burdens of IVI compare with other types of retinal procedures requiring frequent clinical visits.

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