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Quantitative analysis of vitreous inflammation using optical coherence tomography in patients receiving sub-Tenon's triamcinolone acetonide for uveitic cystoid macular oedema

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1	Quantitative Analysis of Vitreous Inflammation using Optical	
2	Coherence Tomography in Patients Receiving Sub-Tenon's	
3	Triamcinolone Acetonide for Uveitic Cystoid Macular Oedema	
4		
5	SUBTITLE – OCT derived measurements vitreous inflammation decrease with	
6	clinical resolution of inflammation in CMO, providing a quantitative and objective	
7	marker of disease activity in uveitis.	
8 9	KEY WORDS – uveitis, cystoid macular oedema, optical coherence tomography, imaging, outcome measures	
10		
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37 **Abbreviations**

- 38 CMO Cystoid macular oedema
- 39 EMA European Medicines Agency
- 40 FDA United States Food and Drug Administration
- 41 OCT ocular coherence tomography
- 42 RPE retinal pigment epithelium
- 43 STTA <u>Sub-Tenon's Triamcinolone Acetonide</u>

44

45 **Disclosure**

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- 53
- 54

55 **Abstract:**

56

BACKGROUND/AIMS : To evaluate the vitreous signals obtained on spectral
domain optical coherence tomography (SD-OCT) in patients with uveitic cystoid
macular oedema (CMO) and compare these signals before and after sub-Tenon's
triamcinolone acetonide injection.

61 **METHODS**

Design: Retrospective study with standardised longitudinal imaging pre- and post-62 intervention. The study cohort comprises 22 patients (22 eyes) with uveitic CMO 63 64 receiving a Sub-Tenon's Triamcinolone Acetonide (STTA) injection. Post-hoc analysis of SD-OCT images using custom software provided an "absolute" 65 measurement of vitreous signal intensity, which was expressed as a ratio to the 66 retinal pigment epithelium intensity ("VIT/RPE-relative intensity") in arbitrary units. 67 Main Outcome Measure: Difference in VIT/RPE-relative intensity before and after 68 treatment. 69

<u>RESULTS</u>: Treatment with STTA resulted in a significant reduction in VIT/RPE Relative Intensity, which was associated with both a reduction in central retinal
 thickness (CRT) and improvement in visual acuity. Mean (SD) VIT/RPE-Relative
 Intensity pre-treatment was 0.139 (0.074) vs. 0.053 (0.028) post-treatment (p=3x10⁻⁵). Mean (SD) CRT was 581µm (119µm) pre-treatment vs 333µm (95µm) post treatment (p=2x10⁻⁸); the mean reduction in CRT was 248 (95%CI: 189-306). The
 correlation coefficient between VIT/RPE-Relative Intensity and CRT was 0.534

(p=0.011) and between VIT/RPE-Relative Intensity and visual acuity was 0.702
(p=0.0001).

CONCLUSION: This study provides evidence that the OCT-derived VIT/RPE Relative intensity may be useful as a quantitative and objective marker of disease
 activity and treatment response in uveitis complicated by CMO. This first longitudinal
 study of this novel OCT parameter is an encouraging step in the development of
 sensitive objective OCT-based endpoints for trials of efficacy in uveitis.

84

86 Introduction

87

Uveitis, a group of conditions characterised by intraocular inflammation, is a major cause of blindness worldwide (1). The commonest cause of sight loss in uveitis is cystoid macular oedema (CMO), which accounts for around a third of blindness caused by the disease (2). CMO may be reversible with prompt corticosteroid treatment (3) such as with Sub-Tenon's triamcinolone acetonide (STTA), leading to visual recovery (4-6).

Vitreous inflammation can be seen on examination as a characteristic "haze", 94 caused by the presence of proteinaceous exudate in the vitreous. The level of 95 vitreous haze is considered to be a good marker of inflammation in the underlying 96 uveal tract. For this reason the National Eye Institute (NEI) "Vitreous Haze Score" is 97 the trial endpoint most commonly accepted by regulatory bodies. The NEI scale is 98 determined by examining the vitreous using an indirect ophthalmoscope with 99 comparison to standardised photographs. Weaknesses are that it is subjective, non-100 continuous, poorly discriminatory at lower levels of inflammation and has low 101 sensitivity in a clinical trial context (7-10). 102

Optical coherence tomography (OCT), an imaging modality that provides high 103 resolution, cross sectional images of ocular tissues non-invasively is well-established 104 in the measurement of macular pathology, including macular oedema (11-13). Most 105 analysis of OCT images is gualitative, but guantitative analysis - such as the 106 measurement of central macular thickness in macular oedema - provides an 107 objective and sensitive measure of deviation from normal, change over time and 108 response to therapy. We and others are using these principles to develop measures 109 all the key components of intraocular inflammation, such that the clinical assessment 110

of uveitis may become more objective and reliable (8, 14-18). We have shown in two
independent cohorts using two different OCT-platforms that measurements of
vitreous inflammation derived from OCT scans are repeatable, reliable and
correlated with clinical measures of disease activity, notably the NEI Vitreous Haze
Score (17-18). To help ensure internal standardisation, we utilised the reflective
intensity of the retinal pigment epithelium (RPE) as a reference, thereby producing a
ratio ("VIT/RPE-Relative Intensity) (17-18)..

The aim of this paper is to further validate the use of the VIT/RPE-Relative Intensity as a marker of disease activity in uveitis by assessing whether it is capable of detecting changes in the vitreous pre-/post-STTA, and whether these correlate with other signs of reduction in disease activity.

122

124 Materials and Methods

125

126 **Study Population:**

This is a retrospective, longitudinal study comprising 22 patients with uveitic CMO 127 attending a tertiary uveitis clinic at the Birmingham & Midlands Eye Centre, Sandwell 128 129 & West Birmingham Hospitals NHS Trust, United Kingdom. All patients had a complete ophthalmic assessment including visual acuity, slit lamp examination, 130 grading of anterior chamber inflammation, intraocular pressure measurement, fundus 131 examination and grading of vitreous inflammation using the NEI Vitreous Haze 132 Score. Approval for data collection and analysis was obtained from a U.K. National 133 Health Service research ethics committee and adhered to the tenets set forth in the 134 Declaration of Helsinki. All patients were consented for posterior STTA 135 administration. 136

137

138 **Procedure**:

Povidone iodine and oxybuprocaine drops were used to sterilize and anesthetize the eye before the procedure. The conjunctiva and sub-Tenons layer were lifted 10mm from the limbus superotemporally using blunt serrated forceps. The sub-Tenon's cannula was attached to a 2ml syringe containing 1ml of 40mg/ml triamcinolone acetonide, and inserted, advancing the needle 12-14mm into the posterior sub-Tenons space into which the full dose (i.e. 40mg triamcinolone acetonide) was administered (19).

146

147 **Image Acquisition:**

Optical coherence tomographic image sets were obtained using Heidelberg
Spectralis OCT (Heidelberg Engineering, Germany). The images were obtained
immediately prior to the procedure and at the first subsequent clinical review. The
volume scan images were centred on the fovea and the TruTrack Active and
AutoRescan features were used to ensure that follow-up scans were matched to the
baseline scan.. The enhanced depth protocol was not used.

154

155 **Quantitative Assessment of Vitreous Signal Intensity:**

156 As per our previously published protocol, OCT scan images were imported into 'OCTOR', a program for easy navigation and manual grading of the scans validated 157 in previous studies (20-21). Masked to all clinical data, primary graders marked out 1) 158 the uppermost extent of the vitreous space included in the scan - the "vitreous top", 159 2) the internal limiting (ILM) membrane, 3) the inner layer of the RPE, and 4) the 160 outer layer of the RPE on all the scans. This was done on five sections going 161 through the central fovea subfield of the Early Treatment Diabetic Retinopathy Study 162 (ETDRS) grid. The area between lines 1 and 2 was defined as the "vitreous 163 space"(VIT), whilst the area between 3 and 4 was defined as the "RPE space" 164 (RPE). The software then calculated the mean intensity values of all image pixels 165 contained within each space as absolute values. A relative value, the VIT/RPE-166 Relative Intensity, could then be derived to minimise the potential effects of 167 confounders such as lens opacities or anterior chamber inflammation (Figure 1). 168 169

170 Statistical Analyses:

171 Clinical and imaging data were analyzed with frequency and descriptive statistics.

172 Snellen visual acuities were converted to LogMAR (logarithm of the minimum angle

of resolution) visual acuity for the purposes of statistical analysis. Spearman's
correlation was used to assess the relationship between the VIT/RPE-Relative
Intensity and clinical/retinal imaging parameter. The Mann-Whitney U test was used
in independent samples and Wilcoxon Signed Ranks test in dependent samples.
Statistical analysis was performed using IBM SPSS software version 20.0 for
Windows (SPSS, Inc, Chicago, Illinois, USA). P values < 0.05 were considered
significant.

Results

183	
184	Baseline Characteristics:
185	The study included 22 eyes of 22 patients, before and after treatment with STTA.
186	Their baseline characteristics are listed in Table 1.
187	
188	VIT/RPE-Relative Intensity
189	Treatment with STTA was associated with a significant reduction in OCT-measured
190	VIT/RPE-Relative Intensity (Figure 2). The mean (SD) VIT/RPE-Relative Intensity
191	pre-treatment was 0.139 (0.074) vs. 0.053 (0.028) post-treatment ($p=3x10^{-5}$).
192	
193	Mean Central Retinal Thickness
194	Treatment with STTA was associated with significant reduction in the OCT-
195	measured mean central 1 mm of retinal thickness (CRT). Mean (SD) CRT was
196	580.5μm (119.4μm) pre-treatment vs 332.7μm (95.4μm) post-treatment (p=2x10 ⁻⁸);
197	the mean reduction in CRT was 247.7 (95%CI: 189.1-306.3). The correlation
198	coefficient between VIT/RPE-Relative Intensity and CRT was 0.534 (p=0.011; Figure
199	3).
200	
201	
202	Visual acuities
203	Treatment with STTA was associated with significant improvement in visual acuity
204	(p=0.0001). The number of patients with a visual acuity greater than 6/12 increased
205	from 1 (4.54%) to 17 (77.3%) with a corresponding reduction in those with 6/12 or
206	worse from 19 (86.4%) to 5 (22.7%) (Fisher exact test, p=0.0001). The correlation

207 coefficient between VIT/RPE-Relative Intensity and visual acuity was 0.702
208 (p=0.0001; Figure 4).

209

210 **Discussion**

This study provides the first 'treatment-response' data to support our proposal 211 that OCT can be used to provide an objective measure of treatment response in 212 uveitis based on changes in the vitreous. It builds on our previous feasibility study in 213 214 which we demonstrated proof of concept that the VIT/RPE-Relative Intensity could provide an objective and quantitative measure of vitreous inflammation. Both that 215 cross-sectional study and a validation study in an independent cohort showed that 216 the VIT/RPE-Relative Intensity was higher in uveitic eyes with active inflammation 217 than uveitic eyes without active inflammation or healthy controls, and that it 218 219 correlated with the clinical NEI vitreous haze score (17-18).We also showed association with other markers of disease activity such as visual acuity, AC cells and 220 221 AC flare. Importantly the VIT/RPE Relative Intensity was also shown to be a repeatable measure with high inter-grader reproducibility (17-18). 222

223

224 In this study we have demonstrated that the VIT/RPE-Relative Intensity decreases significantly in response to STTA and that this reduction was associated 225 with improvement in another measurable sign of disease activity, CMO. Critically 226 this study demonstrates VIT/RPE-Relative intensity is sensitive enough to measure 227 changes in the vitreous undetectable using the clinical NEI Vitreous Haze Score. As 228 highlighted by a number of authors, the poor discrimination of the NEI Vitreous Haze 229 Score at lower levels has led most clinical trials in this field to require subjects to 230 have a minimum NEI Vitreous Haze Score of 2+ for inclusion. This has significantly 231

limited enrolment (7-10,22). In an observational study comparing a photographic-232 based score to the NEI score, Hornbeak noted that had they used the traditional 'cut-233 off' 86% of participants would have been excluded. Although a significant proportion 234 of that cohort were scored as 0 on the NEI Vitreous Haze Score it cannot be argued 235 that all these cases were inactive as both the Hornbeak study and our current study 236 indicate that an appropriately sensitive tool is able to discriminate within this group 237 238 (9). Whereas the Hornbeak study was cross-sectional, our longitudinal study is the first to show a tool capable of detecting change in uveitis activity within these lower 239 240 levels of inflammation, even when both the pre-treatment and post-treatment clinical Vitreous Haze Score was 0. 241

242

There is an urgent need to develop sensitive objective measures of inflammation in uveitis, for use as endpoints in clinical trials and to inform treatment decisions in routine clinical practice. The FDA advises that a trial endpoint must be 'well-defined and reliable' and recommends that treatment benefit should be a measure of how a patient "survives, feels or functions". Other measures that do not capture these are regarded as "surrogate measures of benefit".

249

All measures of disease activity in uveitis are "surrogate measures". The FDA requires a surrogate to be "reasonably likely, based on epidemiologic, therapeutic, pathophysiologic, or other evidence to predict clinical benefit"., In the context of developing and assessing surrogate measures for use as trial endpoints in uveitis, we propose that they must meet two *essential criteria*: (1) The surrogate should be 'biologically relevant' given our understanding of the pathophysiology of the disease; and (2) The surrogate should be 'functionally relevant' with evidence of a downstream effect on visual function, but recognizing that this effect may be delayed
and indirect. Provided a surrogate satisfies these criteria, it should then be assessed
for *desirable criteria* such as objectivity, repeatability, and sensitivity,

260

This study provides further evidence of the biological and functional relevance of VIT/RPE relative intensity..Its biological relevance is demonstrated by its association with other markers of inflammation such as the central retinal thickness. Its functional relevance is supported by its correlation with visual recovery, however it is recognized that this is largely indirect, the primary mechanism of improvement being the restoration of central macular architecture as the oedema resolves.

267

The OCT-derived VIT/RPE-Relative intensity is the first instrument-measured marker of vitreous inflammation, and is an example of how extended applications of OCT and other imaging modalities have the potential to revolutionise our approach to the diagnosis, assessment and management of uveitis. Research into VIT/RPE-Relative intensity levels during the development of CME and its relation to vascular changes visualized on fluorescein angiography may inform our understanding of the natural history of this sight-threatening complication, and help guide treatment..

275

Furthermore the sensitivity of this small study to detect a change at a highly statistically significant level (p = 0.00003), shows how the acceptance of OCTderived objective indices could transform our approach to effectiveness trials in uveitis. The limitations of our current endpoints in uveitis provide major constraints to effectiveness trials (7-10,23), which may lead to a trial 'failing' (i.e. not meeting its primary endpoint) even in the presence of an effective therapy. This in turn 282 discourages further investment, and leads to an absence of high-quality trial data to inform clinicians, funders and policy makers with regard to main of therapies being 283 considered for use in uveitis. The high sensitivity and reproducibility of instrument-284 based measures such as the OCT-derived VIT/RPE relative intensity can provide 285 endpoints with much higher 'signal:noise' ratio than current clinical measures 286 enabling smaller, faster, cheaper trials. Such endpoints can already be adopted as 287 'signals' to inform investment decisions in early phase studies, but their adoption in 288 later-phase licensing studies will depend on achieving the further validation required 289 290 by regulatory bodies such as the FDA.

291

292 Study limitations

293 This study involves retrospective analysis of longitudinal OCT image sets obtained from a small number of patients with uveitis and OCT-confirmed CMO. It 294 therefore has the limitations of a retrospective design, and we acknowledge that 295 given the nature of this cohort the focus of the clinical assessments at the time will 296 have been directed towards the CMO, and not on accurate grading of the clinical 297 vitreous haze. We also note that visual acuity data were recorded as Snellen 298 measurements rather than the preferred LogMAR notation. It should be noted 299 however that the primary focus of this paper is on the post-hoc analysis of the OCT 300 301 image sets and their change over time, rather than on the associated clinical changes. 302

303

The design of the study was pragmatic in using scans conducted under normal macular scanning conditions. We and others have proposed a number of techniques for optimising the visualisation of vitreous using current Spectral Domain and emerging technology (8,10, 17-18). Increasing the proportion of the vitreous
which is visualised is likely to improve this technique further, enhancing sensitivity
and repeatability; it also enables anatomic localisation of foci of inflammation within
the vitreous cavity related to the distribution and type of uveitis.

311

This study is based on a small cohort, with a range of uveitic diagnoses. This heterogeneity is common in uveitis studies (8,10), and indeed the consistent performance of the VIT/RPE relative intensity tool across this range of patients is very encouraging for its future usefulness is an outcome measure. Critically, despite its size and heterogeneity, the study achieved its primary endpoint at a high level of statistical significance .

318

A barrier to the potential adoption of our technique as described in this study is that it is time-consuming, taking around 3-5 minutes per scan. Recently however we have developed a software package for automation of this process. This custom software entitled VITreous ANalysis (VITAN) can segment and annotate the scans automatically, reducing the time taken to derive measures of vitreous reflectivity to a few seconds per scan, with benefits in speed, cost, and further reduction of subjectivity or human error in marking the boundaries of anatomical structures (24).

To conclude, in this study we have further demonstrated the relevance of the OCT-derived VIT/RPE-Relative intensity as a quantitative and objective marker of disease activity and treatment response in uveitis complicated by CME. This first longitudinal study of this novel OCT parameter is an encouraging step in the

- development of sensitive objective OCT-based endpoints for trials of efficacy in
- 332 uveitis.

References: 333

- (1) Nussenblatt RB. The natural history of uveitis. Int Ophthalmol 1990 Oct;14(5-335 6):303-308. 336
- (2) Rothova A, Suttorp-van Schulten MS, Frits Treffers W, Kijlstra A. Causes and 337 frequency of blindness in patients with intraocular inflammatory disease. Br J 338 339 Ophthalmol 1996 Apr;80(4):332-336.
- (3) Levin MH, Pistilli M, Daniel E, Gangaputra SS, Nussenblatt RB, Rosenbaum JT, 340 et al. Incidence of Visual Improvement in Uveitis Cases with Visual Impairment 341 Caused by Macular Edema. Ophthalmology 2014 Feb;121(2):588-595.e1. 342
- 343 (4) Salek SS, Leder HA, Butler NJ, Gan TJ, Dunn JP, Thorne JE. Periocular triamcinolone acetonide injections for control of intraocular inflammation associated 344 with uveitis. Ocul Immunol Inflamm 2013 Aug;21(4):257-263. 345
- (5) Sen HN, Vitale S, Gangaputra SS, Nussenblatt RB, Liesegang TL, Levy-Clarke 346 347 GA, et al. Periocular corticosteroid injections in uveitis: effects and complications. Ophthalmology 2014 Nov;121(11):2275-2286. 348
- (6) Leder HA, Jabs DA, Galor A, Dunn JP, Thorne JE. Periocular triamcinolone 349 acetonide injections for cystoid macular edema complicating noninfectious uveitis. 350 351 Am J Ophthalmol 2011 Sep;152(3):441-448.e2.
- (7) Madow B, Galor A, Feuer WJ, Altaweel MM, Davis JL. Validation of a 352 353 Photographic Vitreous Haze Grading Technique for Clinical Trials in Uveitis. Am J Ophthalmol 2011 8;152(2):170-176.e1. 354
- (8) Denniston AK, Dick AD. Systemic therapies for inflammatory eye disease: past, 355 present and future. BMC Ophthalmol 2013 Apr 24;13:18-2415-13-18. 356
- (9) Hornbeak DM, Paval A, Pistilli M, Biswas J, Ganesh SK, Gupta V, Rathinam SR, 357 Davis JL, Kempen JH. Interobserver agreement in clinical grading of vitreous haze 358 using alternative grading scales. Ophthalmology. 2014 Aug;121(8):1643-8. doi: 359 10.1016/j.ophtha.2014.02.018. Epub 2014 Mar 31. PubMed PMID: 24697913; 360 PubMed Central PMCID: PMC4122589 361
- (10) Barry RJ, Denniston AK. Controversies in the Pharmacological Treatment of 362 Uveitis. Curr Pharm Des 2015 Sep 8. [Epub ahead of print] PubMed PMID: 363 26350535 364
- (11) Hee MR, Puliafito CA, Wong C, Duker JS, Reichel E, Rutledge B, et al. 365
- Quantitative assessment of macular edema with optical coherence tomography. Arch 366 Ophthalmol 1995 Aug;113(8):1019-1029. 367
- (12) Ouyang Y, Keane PA, Sadda SR, Walsh AC. Detection of cystoid macular 368
- edema with three-dimensional optical coherence tomography versus fluorescein 369
- angiography. Invest Ophthalmol Vis Sci 2010 Oct;51(10):5213-5218. 370

- (13) Alasil T, Keane PA, Updike JF, Dustin L, Ouyang Y, Walsh AC, et al.
- Relationship between optical coherence tomography retinal parameters and visual acuity in diabetic macular edema. Ophthalmology 2010 Dec;117(12):2379-2386.
- (14) Agarwal A, Ashokkumar D, Jacob S, Agarwal A, Saravanan Y. High-speed
 optical coherence tomography for imaging anterior chamber inflammatory reaction in
 uveitis: clinical correlation and grading. Am J Ophthalmol 2009 Mar;147(3):413416.e3.
- (15) Li Y, Lowder C, Zhang X, Huang D. Anterior chamber cell grading by optical
 coherence tomography. Invest Ophthalmol Vis Sci 2013 Jan 9;54(1):258-265.
- (16) Sharma S, Lowder CY, Vasanji A, Baynes K, Kaiser PK, Srivastava SK.
- Automated Analysis of Anterior Chamber Inflammation by Spectral-Domain Optical
- Coherence Tomography. Ophthalmology 2015 Jul;122(7):1464-1470.
- (17) Keane PA, Karampelas M, Sim DA, Sadda SR, Tufail A, Sen HN, et al.
- Objective measurement of vitreous inflammation using optical coherence
 tomography. Ophthalmology 2014 Sep;121(9):1706-1714.
- (18) Zarranz-Ventura J, Keane PA, Sim DA, Llorens V, Tufail A, Sadda SR, Dick AD,
 Lee RW, Pavesio C, Denniston AK, Adán A; EQUATOR Study Group. Evaluation of
 objective vitritis grading method using optical coherence tomography: influence of
 phakic status and previous vitrectomy. Am J Ophthalmol. 2015 Oct 14. pii: S00029394(15)00637-6. doi: 10.1016/j.ajo.2015.10.009. [Epub ahead of print] PubMed
 PMID: 26476212.
- (19) Venkatesh P, Kumar CS, Abbas Z, Garg S. Comparison of the efficacy and
 safety of different methods of posterior subtenon injection. Ocular Immunology &
 Inflammation 2008 Sep-Oct;16(5):217-223.
- (20) Sadda SR, Joeres S, Wu Z, Updike P, Romano P, Collins AT, et al. Error
 correction and quantitative subanalysis of optical coherence tomography data using
 computer-assisted grading. Invest Ophthalmol Vis Sci 2007 Feb;48(2):839-848.
- (21) Sadda SR, Keane PA, Ouyang Y, Updike JF, Walsh AC. Impact of scanning
 density on measurements from spectral domain optical coherence tomography.
 Invest Ophthalmol Vis Sci 2010 Feb;51(2):1071-1078.
- 401 (22) Lowder C, Belfort R, Jr, Lightman S, Foster CS, Robinson MR, Schiffman RM, et
 402 al. Dexamethasone intravitreal implant for noninfectious intermediate or posterior
 403 uveitis. Arch Ophthalmol 2011 May;129(5):545-553.
- 404 (23) Denniston AK, Holland GN, Kidess A, Nussenblatt RB, Okada AA, Rosenbaum
 405 JT, et al. Heterogeneity of primary outcome measures used in clinical trials of
 406 treatments for intermediate, posterior, and panuveitis. Orphanet J Rare Dis 2015
 407 Aug 19;10:97-015-0318-6.
- 408 (24) Keane PA, Balaskas K, Sim DA, Aman K, Denniston AK, Aslam T, et al.
 409 Automated Analysis of Vitreous Inflammation Using Spectral-Domain Optical
 410 Coherence Tomography. Transl Vis Sci Technol 2015 Sep 16;4(5):4.
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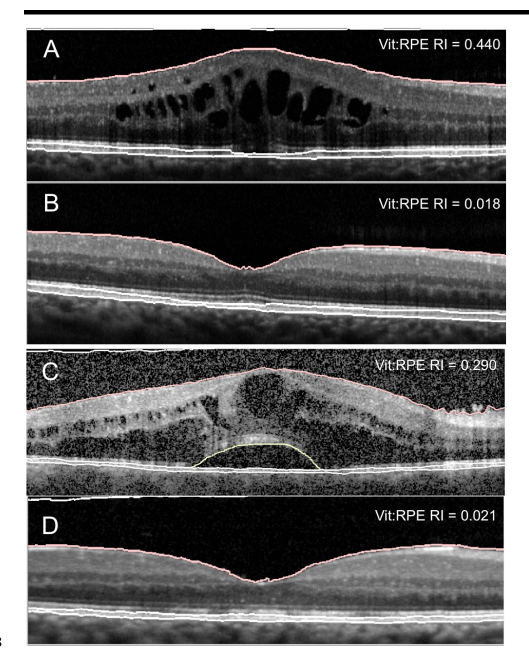
412 **Tables and figures:**

413 TABLE 1: Baseline Characteristics

Age	47.4 years (23y – 74y)
Gender	
Female	17 (77%)
Male	5 (23%)
Anatomical Site of Uveitis	
Panuveitis	10 (45%)
Intermediate Uveitis	8 (36%)
Anterior Uveitis	4 (18%)
Aetiology	
Idiopathic	14 (64%)
Sarcoidosis	4 (18%)
TINU	1 (5%)
Behcet's	1 (5%)
Reiter	1 (5%)
VKH	1 (5%)
AC Cells	

0	6
0.5+	6
1+	4
2+	4
3+	0
4+	0
Not available	2
AC Flare	
0	12 (54.54%)
0.5+	2 (9.09%)
1+	6 (27.27%)
2+	0
3+	0
4+	0
Not available	2 (9.09%)
Vitreous haze	
0	15 (68.18%)
0.5+	0
1+	2 (9.09%)
2+	1 (4.54%)
3+	0
4+	0
Not available	4 (18.18%)

7 weeks (4 – 19 weeks)
<i>,</i>



- Figure 1. Quantitative assessment of the vitreous using OCT
- demonstrated in a 47 year old male with intermediate uveitis (A,B) and a
- 422 48 year old female with panuveitis (C,D). Both patients were assessed

- ⁴²³ by standard macular-focussed OCT both before (A,C) and after (B,D)
- treatment with Sub-Tenon's Triamcinolone Acetonide (STTA), with
- 425 calculation of the Vitreous/RPE-Relative Intensity (Vit/RPE RI)..

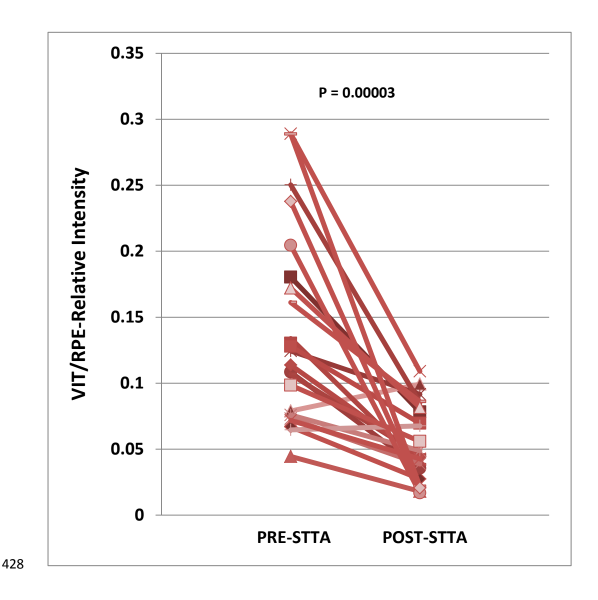


Figure 2. VIT/RPE–Relative Intensity before and after treatment with

- 430 Sub-Tenon Triamcinolone Acetonide (STTA) for Uveitic Cystoid Macular
- 431 Oedema (CMO).

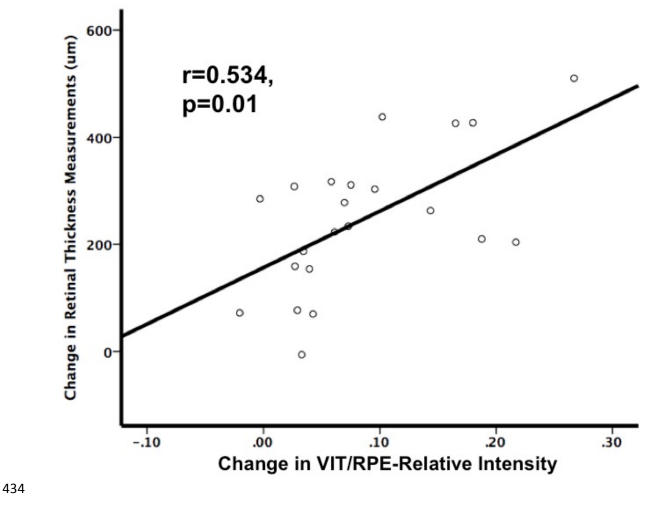


Figure 3. Correlation between change in VIT/RPE–Relative intensity and
change in mean central retinal thickness.

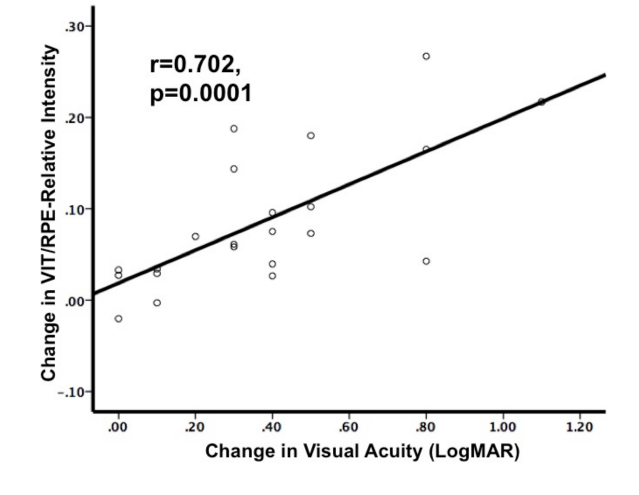


Figure 4. Correlation between change in VIT/RPE–Relative intensity and

