Subretinal fluid morphology in chronic central serous chorioretinopathy and its relationship to treatment: a retrospective analysis on PLACE trial data

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ABSTRACT.

Purpose: To explore subretinal fluid (SRF) morphology in chronic central serous chorioretinopathy (cCSC) after one session of either high-density subthreshold micropulse laser (HSML) treatment or half-dose photodynamic therapy (PDT).

Methods: We retrospectively obtained optical coherence tomography (OCT) scans from a subset of patients from a randomized controlled trial on treatment-naïve eyes with cCSC allocated to either HSML treatment or half-dose PDT. OCT scans were evaluated prior to treatment and 6–8 weeks post-treatment, where we measured maximum SRF height and width, calculated the maximum height-to-maximum width-ratio (maxHWR) and calculated the total SRF volume.

Results: Forty-one eyes of 39 cCSC patients were included. SRF morphology ranged from flat to dome-shaped, quantified as maxHWR ranging between 0.02 and 0.12. SRF volume was median 0.373 µl (range: 0.010–4.425 µl) and did not correlate to maxHWR (rho = -0.004, p = 0.982). Half-dose PDT was superior to HSML treatment in complete SRF resolution (RR 0.373, p = 0.004) and in morphological changes of SRF (ΔmaxHWR p = 0.001; Δmax width p < 0.001; Δvolume p = 0.025). SRF resolved completely in 19/22 PDT-treated eyes (86%) and 5/19 HSML-treated eyes (26%). SRF volume increased in five eyes (26%) after HSML treatment, and in none of the eyes after half-dose PDT. SRF morphology at baseline did not predict treatment outcomes.

Conclusion: SRF morphology changed after both HSML treatment and half-dose PDT in cCSC, with SRF disappearing in most PDT-treated patients, whereas SRF volume increased in a sizeable proportion of HSML-treated patients. Baseline SRF characteristics measured in this study were unable to predict outcomes after either HSML treatment or half-dose PDT.

Key words: central serous chorioretinopathy – imaging – optical coherence tomography – subretinal fluid – volume analysis

Introduction

Central serous chorioretinopathy (CSC) is a common chorioretinal disease and an important cause of loss of central vision. Its key clinical manifestation is the accumulation of subretinal fluid (SRF) in the macula and it is etiologically linked to dysfunction of the blood-retina barrier, that is leakage from the choroid and defects in the retinal pigment epithelium (Daruij et al. 2015). The incidence of CSC peaks at age 30–50 years and is up to eight times more prevalent among males (Kitzmann et al. 2008). The disease has been associated with corticosteroid use, stressful life or endogenous hypercortisolism, and a certain genetic predisposition (Daruij et al. 2015; van Dijk et al. 2016; van Dijk et al. 2017; Schellevis et al. 2019; van Dijk et al. 2019; van Rijssen et al. 2019c; Mohabati et al. 2020). Acute CSC cases are often observed, in which SRF usually spontaneously regresses in typically 3–4 months, without the need of treatment (van Rijssen et al. 2019c). Chronic CSC (cCSC) is considered to require treatment, because chronic persistence of SRF leads to progressive photoreceptor damage and vision loss, which in turn significantly impairs the quality of life of the patients (Breukink et al. 2016). It is currently unknown if acute CSC and cCSC are either a
continuum or different disease entities, despite being part of the same pachychoroid disease spectrum, with overlapping genetic risk factors (Cheung et al. 2019; van Rijsen et al. 2019c; Mohabati et al. 2020).

For cCSC, several treatment options are available (van Rijsen et al. 2019c). Treatments that are most often prescribed are half-dose or half-fluence photodynamic therapy (PDT), oral mineralocorticoid antagonist treatment (eplerenone or spironolactone), and high-density subthreshold micropulse laser (HSML) treatment. In the first randomized controlled treatment trial studying the efficacy of oral eplerenone treatment versus placebo, eplerenone was not superior to placebo (Lotery et al. 2020). The half-dose PDT versus high-density subthreshold micropulse laser treatment in patients with chronic central serous chorioretinopathy (PLACE) trial was the first large investigator-initiated multicentre randomised controlled trial in cCSC, comparing half-dose PDT to HSML as a treatment for treatment-naive cCSC patients (Breukink et al. 2015; van Dijk et al. 2018). The PLACE trial showed that half-dose PDT was superior to HSML treatment in achieving a complete resolution of SRF on optical coherence tomography (OCT) as well as change in retinal sensitivity on microperimetry (van Dijk et al. 2018). To date, little is known on how SRF morphology changes with treatment, and whether SRF morphology at baseline can predict SRF morphology after treatment.

In this study, we obtained data from a subsample of the PLACE trial and analysed OCT data for SRF morphology using both two- and three-dimensional measures of SRF. We then compared the treatment effects of one session of half-dose PDT to one session of HSML treatment.

Methods

Study design

This was a retrospective clinical study based on the data that were collected prospectively in the PLACE trial (ClinicalTrials.gov: NCT01797861, EudraCT: 2012-004555-36). The PLACE trial was an investigator-initiated, open-label, multicentre, randomized controlled trial, which compared half-dose PDT to HSML treatment (Breukink et al. 2015; van Dijk et al. 2018). Informed consent was obtained from all participants. Institutional review board approval was obtained from all participating centres and all aspects of this study adhered to the tenets of the Declaration of Helsinki. Details of the study design are available in the previously published protocol (Breukink et al. 2015). For the current study, we obtained data from one study centre (Leiden University Medical Center, Leiden, the Netherlands). We only included data from eyes of which the available OCT scans contained the entire area of SRF accumulation. This was to allow accurate estimates of SRF morphology, especially in terms of SRF width and SRF volume which otherwise would be impossible if only a part of the SRF was visible on the available OCT scan. No meaningful a priori power calculation was possible for this exploratory imaging study.

Eligibility of participants and eyes

The PLACE trial included adults with cCSC. In brief, for inclusion in the PLACE trial, visual symptoms had to be present for at least six weeks, in combination with presence of SRF including the fovea on OCT, hyperfluorescent changes typical of cCSC on indocyanine green angiography (ICGA), and one or more ill-defined hyperfluorescent leakage areas on fluorescein angiography (FA) with retinal pigment epithelium window defect(s) compatible with cCSC (Breukink et al. 2015). Eyes were excluded in case of previous treatment for active CSC, recent (defined as current or within three months before the baseline visit of the current study) use of either topical or systemic corticosteroids, presence of SRF that could have been caused by other ocular diseases, best-corrected visual acuity (BCVA) of Snellen 20/200 or worse, profound central chorioretinal atrophy, myopia of six diopters or more, continuous and/or progressive visual loss for more than 18 months or serous detachment on OCT for more than 18 months, presence of intraretinal fluid, contraindications for any procedures related to the study, presence of soft drusen in treated or follow eye or any macular neovascularization (Breukink et al. 2015; van Dijk et al. 2018).

Treatment and follow-up evaluation

Eligible patients were randomly assigned to either half-dose PDT or HSML treatment. The area to be treated was determined by a central reading centre and was based on the hyperfluorescent changes on ICGA (Breukink et al. 2015; van Dijk et al. 2018). This approach was used because ICGA-guided therapy targets the primary affected tissue (the choroid) in CSC, while decreasing the risk of undertreatment because ICGA generally reveals more extensive abnormalities compared to FA (van Rijsen et al. 2019a; van Rijsen et al. 2019b). Overlying focal area(s) of leakage on FA, which generally correspond to SRF accumulation on OCT, were always included in the treatment area. Specific details regarding half-dose PDT and HSML treatment are described in detail elsewhere (Breukink et al. 2015; van Dijk et al. 2018; van Rijsen et al. 2020).

Prior to therapy, patients were examined using BCVA using Early Treatment of Diabetic Retinopathy Study (ETDRS) charts, retinal microperimetry, a quality of life questionnaire, ophthalmoscopy, fundus photography, fundus autofluorescence, retinal OCT, FA, and ICGA (Breukink et al. 2015; van Dijk et al. 2018). Importantly for this study, these examinations included a macular OCT using the Heidelberg HRA-Spectralis Special Domain (Heidelberg Engineering, Heidelberg, Germany) with the enhanced-depth imaging mode used to capture at least 25 B-scans with a 20° x 20° field (Breukink et al. 2015; van Dijk et al. 2018). This scan was performed with the use of the high-resolution module. Each B-scan in this mode consisted of a minimum of 20 averaged scans (ART) and is registered over the infrared reflectance image. OCT imaging was performed at baseline and 6–8 weeks after either HSML treatment or half-dose PDT (Breukink et al. 2015; van Dijk et al. 2018).

Morphological analysis of the subretinal fluid

All individual OCT scans of each eye were measured for SRF height and SRF width (Fig. 1). Within one stack of OCT scans, we scrolled through the OCT scans to measure the SRF width.
from left to right, and measured the visually observed highest point of SRF height (from retinal pigment epithelium layer to photoreceptor layer). The highest measured values of SRF height and SRF width within the OCT scans were defined as the maximum SRF height and the maximum SRF width, respectively. We then used these measures to calculate the SRF maximum height-to-maximum width-ratio (maxHWR). We used a previously described method to calculate the volume of the SRF (Subhi & Sørensen 2018). Briefly, we extracted OCT images to an image processing program, identified SRF as regions of interest whereby such regions were coloured manually, and imported these images to an image-stack using IMAGEJ version 1.51a (http://www.imagej.net) (National Institute of Mental Health, Bethesda, MA, USA) (Schneider et al. 2012). Here, coloured regions of interest in stack images can be extrapolated into volumes using the plug-in MEASURE STACK version 1 (OptiNav Inc., Bellevue, Washington, USA) if the distance between the images is known (Fig. 1). The exact distance between the images was extracted for each scan from the Heidelberg Eye Explorer software (Heidelberg Engineering, Heidelberg, Germany). 

**Data analysis and statistics**

We first explored the relationships between maximum SRF height, maximum SRF width, maxHWR and SRF volume in treatment-naïve cCSC eyes. BCVA was converted from ETDRS letters to logMAR and used for correlation studies to the measured morphological variables. Then, we evaluated the treatment effect on SRF morphology. Finally, we compared SRF morphology change after one session of treatment with HSML treatment versus one session of treatment with half-dose PDT. Categorical values were reported in numbers and percentages. Continuous variables were reported in mean and standard deviation (SD) and compared using parametric tests if normal distribution was present, otherwise these non-normally distributed variables were reported in median and interquartile range (IQR) and compared using non-parametric tests. Correlation analyses were made using Pearson’s correlation if both variables were normally distributed, otherwise Spearman’s correlation was employed. All statistical analyses were made using IBM SPSS 26 (IBM, Armonk, NY, USA). Figures were made using GRAPHPAD PRISM 9.0.0 (GraphPad Software, La Jolla, CA, USA). p values < 0.05 were considered statistically significant.

**Results**

**Study sample**

We screened data from 72 eyes of 61 cCSC patients, of which 41 eyes (39 patients) fulfilled the eligibility criteria for this study. Nineteen eyes (18 patients) were allocated to the HSML treatment, and 22 eyes (21 patients) to half-dose PDT. Patient characteristics did not differ significantly between the study groups (Table 1).

**Morphology of subretinal fluid at baseline**

Looking at all 41 eyes, we measured a mean maximum SRF height of 185 μm (SD: 102 μm, range: 36–582 μm) and a mean maximum SRF width of 2942 μm (SD: 1399 μm, range 798–6471 μm). The interrelationship between maximum height and maximum width was quantified by looking at the maxHWR, which when low indicates a flatter SRF and when high indicates more dome-shaped SRF (Fig. 2). This unit-free measure had a mean of 0.07 (SD: 0.03) and ranged from 0.02 to 0.12, exhibiting that SRF certain morphological variation. We observed a moderate correlation between the maxHWR and both the maximum SRF height (rho = 0.381, p = 0.014, Pearson’s correlation) and maximum SRF width (rho = –0.345, p = 0.027, Pearson’s correlation) (Fig. 2).

The SRF volume was measured to be a median of 0.373 μl (IQR: 0.133–0.881 μl, range 0.010–4.425 μl). Although SRF volumes strongly correlated with both maximum SRF height (rho = 0.838, p < 0.001, Spearman’s correlation) and maximum SRF width (rho = 0.868, p < 0.001, Spearman’s correlation), no correlation was observed to the maxHWR (rho = –0.004, p = 0.982, Spearman’s correlation) (Fig. 3).

The BCVA at baseline did not correlate significantly with the maximum SRF height (rho = 0.055, p = 0.734, Pearson’s correlation), the maximum SRF width (rho = –0.102, p = 0.527, Pearson’s correlation), the maxHWR (rho = 0.025, p = 0.879, Spearman’s correlation), and the SRF volume (rho = –0.029, p = 0.858, Pearson’s correlation).

**Figure 1.** Methodological approach to measuring subretinal fluid morphology in a chronic central serous chorioretinopathy patient. (A) We imported a stack of images of optical coherence tomography scans with subretinal fluid (SRF). (B) Each scan was measured for SRF height (top), SRF width (middle) and SRF area for volume calculation (bottom). From the measured SRF heights and widths, highest values were noted for the measures of maximum SRF height and maximum SRF width. (C) By summarizing the SRF areas in each scan and with the distance between the scans known, it is possible to estimate the three-dimensional structure of the SRF and to calculate its volume.
Figure 2. The maximum height-to-maximum width-ratio of subretinal fluids in chronic central serous chorioretinopathy patients. (A) A histogram of the calculated maximum height-to-maximum width-ratio (maxHWR) in this study highlights the morphological variation of subretinal fluid (SRF) in chronic central serous chorioretinopathy. As shown here on the left, a low maxHWR indicates relatively flatter SRF, whereas the example on the right shows a case of high maxHWR with relatively more dome-shaped SRF. (B) A strict correlation between maxHWR and the maximum height or the maximum width would indicate that the variation of maxHWR was strictly a question of SRF size. However, correlation analyses using Pearson’s correlation indicated that maxHWR only moderately correlated with the maximum SRF height and the maximum SRF width.

Table 1. Characteristics of study groups at baseline.

|                          | HSML treatment group (N = 19 eyes) | Half-dose PDT group (N = 22 eyes) | p value |
|--------------------------|----------------------------------|----------------------------------|---------|
| **Demographics**         |                                   |                                  |         |
| Age, years, mean (SD)    | 48.4 (7.8)                       | 47.5 (7.6)                       | 0.69*   |
| Male gender, N (%)       | 17 (89%)                         | 21 (95%)                         | 0.46*   |
| **SRF characteristics at baseline** |                                |                                  |         |
| Maximum height, µm, mean (SD) | 175 (76)                       | 194 (122)                        | 0.54*   |
| Maximum width, µm, mean (SD) | 2.834 (1.552)                  | 3.036 (1.282)                    | 0.66*   |
| MaxHWR, median (IQR)     | 0.07 (0.06–0.09)                 | 0.07 (0.05–0.08)                 | 0.51†   |
| Volume, µl, median (IQR) | 0.387 (0.178–0.874)             | 0.352 (0.111–0.888)             | 0.92†   |
| BCVA at baseline, logMAR, mean (SD) | 0.10 (0.20)                   | 0.15 (0.16)                      | 0.44*   |

BCVA = best-corrected visual acuity; HSML = high-density subthreshold micropulse laser; IQR = interquartile range; MaxHWR = maximum height-to-maximum width-ratio; N = number; PDT = photodynamic therapy; SD = standard deviation; SRF = subretinal fluid.

* Independent samples t-test.
† Mann–Whitney U-test.
‡ χ²-test.

Treatment effect on subretinal fluid morphology

One session of treatment resulted in complete SRF resolution in a total of 24 out of 41 eyes (59%); in 19 out of 22 eyes treated with half-dose PDT (86%), and in five out of 19 eyes treated with HSML treatment (26%). This difference in favour of half-dose PDT was statistically significant (RR = 3.28, 95% CI: 1.52–7.09, p = 0.003). Differences in SRF morphology characteristics between half-dose PDT treatment and HSML treatment were statistically significant in favour of half-dose PDT for all measured parameters (Fig. 4).

In the half-dose PDT group, the three eyes without complete SRF resolution had all experienced a reduction in SRF quantified as a decrease in SRF height (cases: −60 µm, −216 µm, −510 µm), a decrease in SRF maximum width (cases: −286 µm, −2749 µm, −3588 µm), and a decrease in SRF volume (cases: −0.025 µl, −0.890 µl, −4347 µl). The maxHWR decreased in all cases (cases: −0.02, −0.02, −0.07). The case with the largest SRF volume decrease was also the case with the largest maxHWR decrease.

Out of the 14 remaining eyes without complete SRF resolution on OCT in the HSML treatment group, eight (57%) experienced a decrease in SRF accumulation (Δmaximum height median −52 µm (IQR: −32 to −119 µm), Δmaximum width median −393 µm (IQR: −195 to −1401 µm), Δvolume median −0.195 µl (IQR: −0.037 to −0.525 µl)), one (7%) did not experience any significant change (Δmaximum height −1 µm, Δmaximum width −6 µm, Δvolume 0.000 µl); and five (36%) experienced SRF increase (Δmaximum height median 50 µm (IQR: 27 to 84 µm), Δmaximum width median 497 µm (IQR: 214 to 841 µm), Δvolume median 0.717 µl (IQR: 0.093 to 0.845 µl)). The ΔmaxHWR decreased in six out of seven (86%) eyes with SRF volume decrease (median −0.01; IQR: −0.02 to −0.002), decreased very slightly (−0.0004) in the only eye without any significant change in SRF volume, and increased in all five eyes with SRF volume increase (median 0.003; IQR: 0.0006–0.02).

Across groups, change in BCVA from baseline to first follow-up (ΔBCVA) correlated significantly with Δmaximum height (rho = 0.310, p = 0.048, Spearman’s correlation), Δmaximum width...
Since BCVA is analysed in logMAR, these correlations state that a decrease in SRF height, SRF width or SRF volume correlate with an improvement in BCVA (Fig. 5).

We did not find any convincing evidence of a predictive value of the measured SRF morphology characteristics at baseline, since none of the measures were clearly associated with complete SRF resolution after treatment (Table 2).

**Discussion**

In this study, we explored SRF morphology in cCSC using OCT scans from one centre in the PLACE trial. We document the extent of the variation of SRF morphology in cCSC when quantified as maximum height, maximum width, maxHWR, and volume. We found that documenting SRF morphology gives a more detailed insight into changes after treatment. A key finding of this study is that HSML treatment is not only associated with a higher proportion of cases without complete SRF resolution as compared to half-dose PDT, which was previously known (van Dijk et al. 2018), but also to a quantifiable SRF increase in five out of 19 eyes (26%). Hence, HSML treatment often did not provide the desired treatment effect, but even increased the SRF volume in a significant proportion of patients. This
Table 2. Comparison of baseline subretinal fluid morphology characteristics between chronic central serous chorioretinopathy patients who did and those who did not experience complete resolution of the subretinal fluid after treatment.

| Eyes allocated to | Complete resolution of SRF after treatment | No complete resolution of SRF after treatment | p value |
|-------------------|--------------------------------------------|-----------------------------------------------|---------|
| HSML treatment    |                                            |                                               |         |
| Maximum height, μm, mean (SD) | 170 (66) | 187 (107) | 0.76* |
| Maximum width, μm, mean (SD) | 2717 (1519) | 3161 (1777) | 0.64* |
| MaxHWR, median (IQR) | 0.08 (0.06–0.09) | 0.06 (0.06–0.07) | 0.34† |
| Volume, μl, median (IQR) | 0.475 (0.088–0.874) | 0.265 (0.219–0.854) | 0.89† |
| Eyes allocated to half-dose PDT |                                            |                                               |         |
| Maximum height, μm, mean (SD) | 175 (91) | 318 (235) | 0.40* |
| Maximum width, μm, mean (SD) | 2962 (1.229) | 3501 (1.812) | 0.66* |
| MaxHWR, median (IQR) | 0.08 (0.07–0.11) | 0.07 (0.05–0.08) | 0.16† |
| Volume, μl, median (IQR) | 0.893 (0.088–4.425) | 0.330 (0.111–0.861) | 0.41† |

HSML = high-density subthreshold micropulse laser; IQR = interquartile range; MaxHWR = maximum height to maximum width ratio; PDT = photodynamic therapy; SD = standard deviation; SRF = subretinal fluid.

* Independent samples t-test.
† Mann-Whitney U-test.

Our study also revealed further structural insight into the SRF dynamics after HSML treatment and half-dose PDT. A single session of half-dose PDT resulted in a complete SRF resolution in all but three eyes. In the remaining three eyes, SRF volume decreased, in a fashion where the SRF decrease lead to a flatter SRF structure, that is a decrease in both SRF maximum height and SRF maximum width but with a relatively stronger decrease in SRF maximum height leading to a lower maxHWR. For the HSML-treated patients, we saw a similar pattern in that an SRF volume decrease leads to a lower maxHWR and therefore a flatter SRF structure in all but one eye. However, this group also had eyes with SRF volume increase, wherein the SRF obtained a higher maxHWR and therefore a more dome-shaped structure.

The clinical importance of these SRF dynamics in relation to therapy in cCSC remains largely unknown and warrants further studies. Of importance, our correlation analyses revealed that a BCVA improvement after therapy was correlated with a reduction of SRF maximum height, SRF maximum width, and SRF volume. Interestingly, the correlation coefficient was −0.3 for ΔmaxHWR and Δvolume, which indicate a low/small correlation, whereas the correlation coefficient was −0.5 for ΔmaxHWR, which indicates a moderate correlation. Thus, our study suggests that an improvement of the SRF width may have a larger influence on the BCVA than the improvement of either SRF height or SRF volume.

In conclusion, we here document the extent of SRF morphology in cCSC, and explore changes after a single session of either HSML treatment or half-dose PDT. Analysis of SRF morphology revealed that a significant proportion of HSML-treated cCSC patients have a persistent or even increased SRF volume after treatment, while half-dose PDT induces a complete resolution of SRF in the vast majority of patients.

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