Decrease in Choroidal Blood Flow After Half and One-Third Dose Verteporfin Photodynamic Therapy For Chronic Central Serous Chorioretinopathy

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Research Article

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Abstract

Background: The effect of various reduced doses of verteporfin photodynamic therapy (PDT) on choroidal blood flow in chronic central serous chorioretinopathy (CSC) remains unclear. Therefore, this study aimed to evaluate choroidal blood flow after half-dose PDT (1/2PDT) and one-third dose PDT (1/3PDT) with verteporfin for chronic CSC using laser speckle flowgraphy (LSFG) and spectral-domain optical coherence tomography.

Methods: Twenty-seven eyes of 27 patients with serous retinal detachment (SRD) caused by chronic CSC for more than 6 months were included in this study. Patients were divided into the 1/2PDT (n=12; January 2018 to July 2019) and 1/3PDT (n=15; August 2016 to December 2017) groups based on the treatment period. The best-corrected visual acuity (BCVA), central foveal thickness (CFT), central choroidal thickness (CCT), and mean blur rate in the macular area (m-MBR) and optic nerve head (ONH-MBR) were obtained using LSFG and evaluated at baseline (pre-treatment), and 2 weeks, 1 month, 3 months, and 6 months after treatment.

Results: SRD disappeared after one month in 92% and 93% eyes in the 1/2PDT and 1/3PDT groups, respectively. Recurrence of SRD was observed in one eye at the 6-month follow-up after 1/2PDT and 2 eyes at the 3-month follow-up after 1/3PDT. No significant improvement was observed in baseline BCVA in the 1/3PDT and 1/2PDT groups. The average m-MBR against baseline significantly decreased at 2 weeks, 1, 3 and 6 months in the 1/2PDT group. The average m-MBR against baseline decreased significantly only at the 2 week follow-up in the 1/3PDT group.

The average rate of change in the CCT against baseline decreased significantly throughout 6 months in the 1/2PDT group and for up to 3 months in the 1/3PDT group. No significant fluctuation was observed in the ONH-MBR.

Conclusions: PDT significantly affected choroidal blood flow depending on the verteporfin dose in chronic CSC.

Trial Registration: This trial was retrospectively registered (UMIN000026850; Approval date 03/04/2017).

Background

Recent advancements in retinal imaging technology have enabled novel methods of investigating choroidal pathology in retinal diseases. The choroid performs highly important physiological functions such as providing vascular supply, nutritional support, and oxygen to the outer retina (1).

Central serous chorioretinopathy (CSC) is one of the most common retinal diseases, which presents with serous retinal detachment (SRD) at the posterior pole, and causes central scotoma, blue vision and metamorphopsia, and (especially) micropsia. Although the clinical course of CSC is usually benign, 30% to 50% of patients experience repeated recurrence and some of these patients develop chronic CSC,
which results in poor visual acuity (2, 3). The pathophysiology of acute and chronic CSC is not fully understood, although several studies have revealed that retinal pigment epithelium (RPE) and choroidal dysfunction could play an important role (4-6). Gass suggested that a focal increase in the permeability of the choriocapillaris layer was the primary cause of damage to the overlying RPE in patients with CSC. (7) Focal leakage on fluorescein angiography (FA) and dilated choroidal vessels and focal choroidal vascular hyperpermeability on indocyanine green angiography (ICGA) are suggestive of the increase in choroidal hydrostatic pressure and defects in tight junctions of the RPE cells in CSC (8).

Photodynamic therapy (PDT) with verteporfin, which was introduced as a promising therapeutic approach for CSC, is based on remodelling of the capillaries in the underlying damaged RPE, which decreases the hyperpermeability of the dilated choroidal vessels (9) (10). Thereafter, studies have reported the efficacy of reduced doses of verteporfin PDT in minimising the angiogenetic complications responsible for severe visual dysfunction in CSC. Studies on half-dose PDT have shown favourable results with comparable effects and fewer side-effects than those with full-dose PDT (11) (12). Moreover, 30%-PDT has been shown to reduce retinal thickness, with a good fluorescein angiography–based improvement rate for acute CSC (13).

Laser speckle flowgraphy (LSFG), a commercially available device, provides a non-invasive quantitative method of determining ocular blood flow, including choroidal blood flow (14). Saito et al. reported a significant decrease in blood flow against the baseline in acute CSC (15). However, the effect of various reduced doses of verteporfin PDT on choroidal blood flow in chronic CSC remains unclear.

The purpose of this study was to investigate the changes in the choroidal blood flow and thickness after half and one-third dose PDT in patients with chronic CSC.

Patients And Methods

The institutional review board of Toho University Omori Medical Center approved the protocol of this retrospective study review (approval number: 27-277). All patients provided informed consent for participation after they received an explanation of the nature and possible consequences of the treatment in accordance with the tents of the Declaration of Helsinki. This study was registered with the UMIN clinical trial registry (UMIN ID: 000026850).

Patients

Consecutive cases of patients with chronic CSC who were admitted to Toho University Medical Center Omori Hospital were recruited between August 2016 and July 2019. All participants were Asian. All patients underwent basic ophthalmic examinations, including best-corrected visual acuity (BCVA), intraocular pressure (IOP), fundus examination, spectral-domain optical coherence tomography (SD-OCT), and LSFG. FA and ICGA were performed at the initial visit. BCVA was obtained using Landolt C charts. These values were subsequently converted to the logarithm of the minimum angle of resolution (logMAR) equivalent for statistical comparisons. The exclusion criteria included patients with a history of retinal
photocoagulation, presence of choroidal neovascularization (CNV) or other ischaemic retinopathies, a history of intravitreal injections of anti-vascular endothelial growth factors, and those taking medications such as corticosteroids, adrenergic agonists, or adrenergic antagonists.

**Diagnosis of chronic CSC**

Patients were diagnosed with CSC based on clinical findings such as the presence of subretinal fluid (SRF) and macular choroidal vasodilatation on optical coherence tomography (OCT) imaging, fluorescein leakage below or near the central fovea of the macula on FA and diffuse vascular permeability on ICGA. Figure 1 depicts a representative case of CSC.

Chronic CSC was defined as persistence of SRF for more than 6 months with no improvement after conservative treatment or expectation of spontaneous resolution.

Each case of CSC was divided into active and resolved eyes based on the presence of SRD in the macular area. The presence of any fluid was designated as active CSC.

**One third and half dose reduced photo dynamic therapy**

Patients were divided into the one-third dose (verteporfin 2 mg/m²) PDT group (1/3PDT) and half-dose (verteporfin 3 mg/m²) PDT group (1/2PDT) according to the treatment period: those who underwent PDT between August 2016 and December 2017 received 1/3PDT and those who underwent PDT between January 2018 and July 2019 received 1/2PDT.

The treatment parameters for PDT included infusion of verteporfin (2 mg/m² or 3 mg/m²) verteporfin for 10 minutes, followed by activation with a diode laser of 689-nm wavelength (Visulas 690 S; Carl Zeiss Meditec, Dublin, California, USA) with a radiation dose of 50 J/cm² for 83 s at the site of hyperpermeability depicted by ICGA. The PDT-irradiated area was determined using ICGA to identify the area of choroidal vasodilatation and congestion, and the area with extravasation in the macula.

**OCT measurements**

SD-OCT (Spectralis; Heidelberg Engineering, Heidelberg, Germany) images were acquired before treatment (i.e. at baseline) and 2 weeks and 1, 3, and 6 months after treatment to analyse the retinal and choroidal thickness. Central retinal thickness was defined as the distance between the inner limiting membrane and the RPE in the fovea on the SD-OCT images. Choroidal thickness was obtained with enhanced-depth imaging (EDI)-OCT. The requisite parameters on the vertical and horizontal scans were measured manually using a built-in calibre tool, and the values were averaged. Choroidal thickness was defined as the distance between the line corresponding to Bruch’s membrane beneath the RPE and the chorioscleral interface under the fovea.

**Laser Speckle Flowgraphy**
The LSFG-NAVI™ (Softcare Co., Fukuoka, Japan) was used to acquire the LSFG images before treatment and 2 weeks and 1, 3, ad 6 months after treatment. The determination of optic nerve head (ONH) circulation with LSFG has been described in detail previously (16). Previous studies have reported that the mean blur rate (MBR) is an indicator of retinal blood flow. The original MBR values were continuously recorded at 118 frames within 4 s, followed by averaging the entire data set to synthesise a still image corresponding to the duration of one heartbeat. The average MBR was determined as the mean value of the synthesized MBR histogram during one heartbeat. An offline analysis software (LSFG Analyzer, version 3.0.47.0) combined all the images and transformed each pixel into a colour-coded map to which the calculated mean blur ratio was assigned, which is a quantitative indicator of the relative blood flow rate.

We calculated the macular MBR (m-MBR) and ONH-MBR using the LSFG Analyzer software.

The MBR of the optic disc (ONH-MBR) was measured to evaluate the retinal blood flow. The MBR at the macular area (m-MBR), which is the retinal avascular region, was used to evaluate the choroidal blood flow. The m-MBR was measured to evaluate the changes in the MBR: the relative MBR was calculated against the baseline and expressed as the rate of change in relative blood flow velocity (23). LSFG was performed after the participants had rested for 10 min in a quiet room maintained at 24°C. The MBR was evaluated by calculating the rate of change against the first baseline measurement.

Measurement of other systemic and ocular parameters

Pulse rate, systolic blood pressure (SBP), and diastolic blood pressure (DBP) were measured before PDT. The mean arterial pressure (MAP) was calculated by the formula of MAP = DBP + 1/3 (SBP - DBP). Ocular perfusion pressure (OPP) was calculated as OPP = 2/3 MAP-IOP (23).

Statistical analysis

All values were presented as the mean and standard deviation, unless specified otherwise. The unpaired t-test and Chi-squared test was used to compare the characteristics of the two groups. The rate of change of m-MBR and ONH-MBR in each group was evaluated using the Steel test, and the Wilcoxon test was used to compare the rate of change of m-MBR and ONH-MBR in the 1/2PDT and 1/3PDT groups.

p-values < 0.05 were considered statistically significant. The JMP (version 11) statistical analysis software (SAS Institute, Inc., Cary, NC, USA) was used to analyse the data.

Results

No systemic and ocular adverse events associated with verteporfin injection were observed throughout the study period. The baseline demographic and clinical data of the two treatment groups are summarised in Table 1. A total of 27 eyes from 27 patients with chronic CSC were considered, of which 15 eyes were treated with 1/3PDT and 12 eyes were treated with 1/2PDT. No significant differences were found in the sex (p=0.78), baseline BCVA (1/2PDT, 0.13±0.25, 1/3PDT, 0.21±0.25; p=0.35), duration of...
symptoms (1/3PDT, 18.4±16.4, 1/2PDT, 18.8±12.2 ; p=0.59), CRT at baseline (1/2PDT, 334.2±94.5; 1/3PDT, 275.1±94.5; p=0.06), choroidal thickness at baseline (1/2PDT, 451.2±132.5; 1/3PDT, 452.2±85.1; p=0.88), and PDT spot size (1/2PDT, 2888.3±550.5; 1/3PDT, 2558.9±710.7; p=0.22), except for age (1/2PDT, 51.5±5.7; 1/3PDT, 46.3±7.2; p=0.04) between the two treatment groups.

Table 1 Baseline demographic and clinical data of the 1/2PDT and 1/3PDT groups

|                          | 1/2 PDT          | 1/3 PDT          | P value |
|--------------------------|------------------|------------------|---------|
| Age                      | 51.5±5.7         | 46.3±7.2         | 0.04    |
| sex (male:female)        | 9:03             | 12:03            | 0.78    |
| BCVA (logMAR BCVA)       | 0.13±0.25        | 0.21±0.25        | 0.35    |
| duration of symptoms (months) | 16.8±12.2      | 18.5±16.4        | 0.59    |
| IOP(mmHg)                | 15.0±1.8         | 13.5±3.7         | 0.06    |
| OPP(mmHg)                | 51.3±7.5         | 52.3±12.6        | 0.90    |
| CCT (µm)                 | 334.2±94.5       | 275.1±94.5       | 0.67    |
| CRT (µm)                 | 451.2±132.5      | 452.2±85.1       | 0.88    |
| PDT spot size (µm)       | 2888.3±550.5     | 2558.9±710.7     | 0.22    |

BCVA: best-corrected visual acuity, IOP: intraocular pressure, OPP: ocular perfusion pressure, CFT: central foveal thickness, CCT: central choroidal thickness, 1/2PDT: half-dose photodynamic therapy, 1/3PDT: one-third dose photodynamic therapy, logMAR: logarithm of the minimum angle of resolution

The disappearance rate of SRF 1 month after treatment was 92% (11/12 eyes) in the 1/2PDT and 93% (14/15 eyes) in the 1/3PDT group.

Recurrence was observed in one eye at the 6-month follow-up in the 1/2PDT group and two eyes at the 3-month follow-up in the 1/3PDT group over the 6-month follow-up period.

No significant improvement was observed in the BCVA compared to baseline (0.13±0.25 and 0.21±0.25, respectively) at two weeks (0.13±0.21 and 0.19±0.25, respectively), 1 month (0.13±0.22 and 0.13±0.23, respectively), 3 months (0.07±0.19 and 0.12±0.27, respectively) and 6 months (0.02±0.19 and 0.13±0.30, respectively) after PDT in the 1/2PDT and 1/3PDT groups.

Figure 2 depicts the time course of the rate of change in the m-MBR against baseline in the 1/2PDT and 1/3PDT groups. In the 1/2 PDT group, the average m-MBR against baseline (100%) significantly decreased to 85.3% (p=0.004), 85.0% (p=0.037), 83.1% (p=0.015) and -82.7% (p=0.015) 2 weeks and 1, 3 and 6 months after treatment, respectively. In the 1/3PDT group, the average m-MBR against baseline
(100%) decreased to 87.2% (p<0.001), 98.5% (p=0.79), 100.2% (p=0.99), and 92.5% (p=0.14) 2 weeks and 1, 3 and 6 months after treatment, respectively. The rate of change in the m-MBR differed (statistically) significantly between the 1/2PDT and 1/3PDT groups at 2 weeks and 1 month and 3 months after treatment.

### Foveal choroidal thickness

Figure 3 shows the time course of the rate of change in the CCT against baseline in the 1/2PDT and 1/3PDT groups. In the 1/2PDT group, the average of rate of change of CCT against baseline (100%) significantly decreased to 83.1% (p<0.001), 80.6% (p<0.001), 78.1% (p<0.001) and 79.6% (p<0.001) 2 weeks and 1, 3 and 6 months after treatment, respectively. In the 1/3PDT group, the average of the rate of change of CCT against baseline (100%) decreased to 90.3% (p=0.001), 92.2% (p=0.001), 90.7% (p=0.001), and 92.8% (p=0.5) 2 weeks and 1, 3 and 6 months after treatment, respectively. The rate of change in CCT differed (statistically) significantly between the 1/2PDT and 1/3PDT groups 2 weeks and 1, 3 and 6 months after treatment.

The ONH-MBR showed no significant fluctuation at 106% after 2 weeks, 104% after 1 month, 101% after 3 months, and 102% after 6 months in the 1/2PDT group. The ONH-MBR showed no significant fluctuation at 99% after 2 weeks, 96% after 1 month, 96% after 3 months, and 99% after 6 months in the 1/3PDT group.

Moreover, no significant change was observed at any time-point after treatment compared to baseline.

The OPP also showed no significant fluctuation at 98% for 2 weeks, 102% for 1 month, 103% for 3 months, and 110% at 6 months in the 1/2PDT group and at 97% for 2 weeks, 107% for 1 month, 102% for 3 months, and 104% for 6 months in the 1/3PDT group. There was no significant change in the fellow eyes at any time after treatment compared to baseline.

### Discussion

Hyperpermeability of the choroidal blood vessels and the consequent increase in choroidal hydrostatic pressure comprise the current principal hypothesis to explain the pathophysiological mechanism of CSC (17). ICGA depicts arterial filling delay and subsequent hyperpermeability in patients with CSC (18). We found that sub-foveal choroidal circulation decreased significantly after the administration of reduced doses of verteporfin PDT in patients with chronic CSC as depicted by the significant disappearance of SRF in the 1/2PDT and 1/3PDT groups on LSFG. There was no significant correlation between MBR and OPP, which suggests that systemic circulation did not affect the m-MBR.

To the best of our knowledge, this is the first report that investigated the alteration in choroidal blood flow using LSFG after PDT in chronic CSC using two clinically representative concentrations of verteporfin.

The salient feature of this study is that it evaluated the blood flow directly using LSFG. The majority of the MBR in LSFG is thought to be derived from choroidal blood flow. Moreover, the medium and large
blood vessels of the choroid in the macular area are clearly visualised, which is the retinal avascular area. Therefore, the m-MBR is considered to reflect the pathological condition in CSC (19). Saito et al. reported that the mean rate of the change in m-MBR decreased significantly at 6 months with spontaneous remission of SRF and visual recovery in patients with acute CSC (15).

Damage to the vascular endothelium, hypoperfusion, and choroidal hyperpermeability due to the formation of free radicals associated with radiation is thought to be the therapeutic mechanism of action of PDT in CSC (10). Our finding, i.e. the notable decrease in m-MBR after PDT, supports this hypothesis. Interestingly, the significant decrease in the m-MBR in the 1/3PDT group was observed for only 1 month, in contrast to that in the 1/2PDT group, which was sustained for up to 6 months. Moreover, we found that the average time for the recurrence of SRD was shorter in the 1/3PDT group than that in the 1/2PDT group. These results could indicate that the choroidal blood flow would be affected in verteporfin-dose dependent manner. PDT with verteporfin is a promising therapeutic modality for CSC. However, it is accompanied by the risk of serious complications such as angiographic closure following treatment (20). Therefore, various regimens, including reduced dose, reduced power (21), reduced time (22), and reduced verteporfin (3) (23) (24) have been devised. Verteporfin concentrations affect normal choroidal vessels in a dose-dependent manner. Zhao et al. studied the treatment of acute CSC with PDT at verteporfin concentrations of 70%, 60%, 50%, 40%, 30%, 20%, and 10%, and found that the 30% dose of verteporfin (1.8 mg/m²) is the minimum required therapeutic dose (25). Moreover, they also reported that the 50% dose of verteporfin may be more effective in resolving SRF and fluorescein leakage with better visual outcomes in acute CSC compared to the 30% dose (26) (13). Our findings also support the notion that 50% of verteporfin is more beneficial with respect to the efficacy and safety of the modified PDT protocol.

Choroidal thickening is considered to be one of the most characteristic clinical signs of CSC and is associated with choroidal vascular hyperpermeability (27). (28) The subfoveal choroid layer was thicker in the affected and fellow eyes in patients with CSC than that of the normal control eyes (29). Numerous OCT studies have reported structural changes after PDT. Maruko et al. reported that the subfoveal choroid thickness decreased significantly after a transient increase following half-dose verteporfin PDT. (30) Moreover, Kinoshita et al. reported that the decreased CCT may be attributed to the reduction in the dilation of the outer choroidal vessels in half-dose verteporfin PDT (31). Izumi et al. reported that subfoveal intrachoroidal structure showed greater alteration than that of the choriocapillaris and medium choroidal vessels after half-dose PDT for CSC (32). We found that the CCT decreased after half and one-third PDT and was accompanied by a decreased in the m-MBR, which may indicate that the decreased blood flow in the choroidal vessels resulted in choroidal thinning. Simultaneous evaluation of blood flow measurements using LSFG and choroidal structural changes using OCT and ICGA could explain and depict the pathophysiology of CSC. However, the relationship between m-MBR and choroidal thickness in retinal-choroidal disease is not uniform in other diseases. Although the choroid thickness increases in inflammatory diseases such as Harada disease (19) and acute zonal occult outer retinopathy (33) the m-MBR reportedly decreases owing to the statis of circulation in the acute phase of inflammation.
We found no significant change in visual acuity after the resorption of the SRF, which could be attributed to the relative longer disease duration (18.4 ± 16.4 months) in our chronic CSC population. The persistence of SRF in chronic CSC is thought to be associated with irreversible progressive photoreceptor damage, leading to loss of visual function (34) (35). Clinical features such as later age of onset, cystoid macular degeneration, CNV, and disruption of the ellipsoid zone are reportedly associated with poor visual acuity in chronic CSC (35). Further longitudinal studies are needed to investigate choroidal blood flow and these clinical findings.

The limitations of our study included the small sample size and short follow-up period. Moreover, we did not investigate the relationship between the choroidal blood flow measured by LSFG and increased permeability of the choroidal vessels obtained using ICGA. Hyperpermeability of choroidal blood vessels and the consequent increase in choroidal hydrostatic pressure are considered to be the key pathogenic factors for CSC. The association between choroidal blood flow and localised hyperpermeability of the choroid in the subretinal space may be an indicator of disease activity, and requires further investigation.

**Abbreviations**

CSC: central serous chorioretinopathy

LSFG: laser speckle flowgraphy

FA: fluorescein angiography

ICGA: indocyanine green angiography

1/2PDT: half-dose photodynamic therapy

1/3PDT: one-third dose photodynamic therapy

SRD: serous retinal detachment

SD-OCT: spectral-domain optical coherence tomography

BCVA: best-corrected visual acuity

CFT: central foveal thickness

CCT: central choroidal thickness

MBR: mean blur rate

m-MBR: macular MBR

ONH: optic nerve head
IOP: intraocular pressure
RPE: retinal pigment epithelium
LogMAR: logarithm of the minimum angle of resolution
CNV: choroidal neovascularization
SRF: subretinal fluid
OCT: optical coherence tomography
EDI-OCT: enhanced-depth imaging OCT
SBP: systolic blood pressure
DBP: diastolic blood pressure
MAP: mean arterial pressure
OPP: ocular perfusion pressure

Declarations

Ethics approval and consent to participate

The experimental protocol was approved by the Ethics Committee of Toho University (#27-277 Approval date 10/3/2016).

Trial Registration: This trial was retrospectively registered (UMIN000026850; Approval date 03/04/2017)

Consent for publication: Not applicable.

Availability of data and materials. The datasets generated and/or analysed during the current study are not publicly available due to data privacy concerns but are available from the corresponding author on reasonable request.

Competing interests: The authors declare that they have no competing interests.

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Authors’ contributions

SK conceived of the presented idea. SK and ST wrote the main manuscript text and TI prepared figures 1-3. AT and TK verified the analytical methods. YH supervised the findings of this work. All authors
reviewed the manuscript.

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