Observation of the vitreous in diabetic macular edema using swept-source OCT

Abstract

Purpose: This study describes the relationship between the vitreous and diabetic macular edema (DME) using swept source OCT (SS-OCT).

Method: We examined 81 eyes of 58 consecutive patients diagnosed as DME. The condition of the posterior wall of the posterior precortical vitreous pocket (PPVP) was classified into 4 groups based on 12-mm horizontal and vertical scans through the macula using SS-OCT. Classifications were: Group 1, no posterior vitreous detachments (PVDs); Group 2, perifoveal PVD; Group 3, macular PVD; and Group 4, complete PVD. DME types, which included cystoid macular edema (CME) and serous retinal detachment (SRD), were also investigated in each group. The control group consisted of 46 eyes of 39 diabetic patients without DME as shown by OCT.

Result: The number of eyes diagnosed as DME for each of the groups was 18, 35, 6 and 22, respectively. OCT images revealed there were 35 (43%) out of 81 eyes with perifoveal PVD, that was accompanied by CME (94%) with or without SRD, which was statistically significant as compared with the other groups (P=0.0005). In Group 2, there were 21 (60%) out of 35 eyes with proliferative diabetic retinopathy (PDR), with this ratio significantly higher when compared to the other groups (P=0.034).

Conclusion: A significant correlation exists between CME associated with DME and perifoveal PVD.

Keywords: swept-source OCT, DME, CME, vitreous

Abbreviations: DME, diabetic macular edema; SS-OCT, swept source optical coherence tomography; PPVP, posterior precortical vitreous pocket; CME, cystoid macular edema; PVD, posterior vitreous detachments; SRD, serous retinal detachment; PDR, proliferative diabetic retinopathy; DR, diabetic retinopathy; VEGF, vascular endothelial growth factor

Introduction

Diabetic macular edema (DME) is the most common cause of visual loss in patients with diabetic retinopathy (DR) and is responsible for a decreased vision-related quality of life in working-aged people. Visual loss occurs due to the leakage of blood constituents from microaneurysms and the dilated macular capillary bed into the surrounding intraretinal or subretinal space, which leads to retinal thickening and macular edema with image distortion. Several treatment interventions have been utilized, including photocoagulation, vitrectomy, steroid, and anti-vascular endothelial growth factor (VEGF) agents. In addition to the breakdown of the blood-retinal barrier, the pathological changes in the vitreomacular interface play an important role in the persistent thickening or distortion of the neuroglial tissue in the macula. This has been shown to lead to the peeling of the posterior wall of the posterior precortical vitreous pocket (PPVP) during pars plana vitrectomy. Vitrectomy removes both the VEGF protein that is derived from ischemic retinal tissues and the cytokines involved with the inflammatory responses. Although the specific details of the condition between the DME and vitreous remain unclear, vitrectomy has been thought to be effective for DME.

Recent advances in optical coherence tomography (OCT) technology have enabled more detailed and precise quantitative assessments of optical structural changes. Swept-source OCT (SS-OCT) is a new generation of OCT that provides higher penetration into the choroid and sclera. SS-OCT also enables clearer visualization of the vitreous even if a patient has progressed cataract, although SS-OCT can observe the surface of the retina and the vitreous to a degree. Itakura et al. recently reported being able to make a precise and detailed observation of the PPVP when using SS-OCT. The current study used SS-OCT with the aim of examining the relationship the morphologic features of DME and the condition of the posterior wall of the PPVP. We also investigated whether SS-OCT is useful for observing of the surface of the retina and the vitreous in DME.

Method

Patients

The protocol of the study was approved by the Institutional Review Board and Ethics Committee of Gunma University Graduate School of Medicine, and all investigations were conducted in accordance with the Declaration of Helsinki. Informed consent was obtained from all patients prior to all examinations. All subjects were examined at Gunma University from August 2012 to November 2013. In this study, our definition of DME is a condition of diabetic retinopathy with central foveal thickness more than 300 mm. The eyes of diabetic patients who were examined by OCT in our department and found to be without DME were designated as the control group. The cases with proliferative tissues observed using indirect funduscopy were divided into Proliferative diabetic retinopathy (PDR). No inflammation was observed in any of the eyes based on indirect funduscopy and slit-lamp biomicroscopy examinations.

SS-OCT imaging

The OCT images of the eyes were obtained at the initial patient...
visit by SS-OCT (DRI OCT-1 Atlantis, Topcon, Tokyo, Japan) and used to investigate the posterior wall of the PPVP. This particular OCT system has an A-scan repetition rate of 100,000 Hz, with a light source that operates in the 1-μm wavelength region. The light source consists of a wavelength-tunable laser centered at 1,050 nm with a 100 nm tuning range. This system used an 8μm axial resolution and a 20 μm lateral resolution, with a 2.3 mm imaging depth in the tissue. The ocular fundus was examined by performing 12 mm horizontal and vertical scans through the fovea. Based on these scans, the condition of the posterior wall of the PPVP was classified into the following four groups: Group 1, no PVDs (Figure 1); Group 2, perifoveal PVD (Figure 2); Group 3 macular PVD (Figure 3); and Group 4, complete PVD (Figure 4).

![Figure 1](image1.png) **Figure 1** Representative fundus photo and OCT image for a Group 1 (no PVD) case (A and B). The condition of the posterior wall of the PPVP is indicated by the arrows.

![Figure 2](image2.png) **Figure 2** Representative fundus photo and OCT image for a Group 2 (perifoveal PVD) case (A and B). The condition of the posterior wall of the PPVP is indicated by the arrows.

![Figure 3](image3.png) **Figure 3** Representative fundus photo and OCT image for a Group 3 (macular PVD) case (A and B). Vitreofoveal separation with persistent attachment to the optic disc is observed. The condition of the posterior wall of the PPVP is indicated by the arrows.

![Figure 4](image4.png) **Figure 4** Representative fundus photo and OCT image for a Group 4 (complete PVD) case (A and B). The posterior wall of the PPVP is not observed.

The proportion of serous retinal detachment (SRD) and CME was also determined in each group. When cystic changes in the retina were found within 2mm from the fovea without serous retinal detachment, cases were classified CME group. Because the retinal thickness was increased to an extent in all cases, the morphological investigations in this study focused on only SRD and CME. We excluded all cases associated with vitreous hemorrhage, neovascular glaucoma, epiretinal membrane and proliferative membrane that generated obvious traction to the fovea.

**Citation:** Akiyama H, Itakura H, Kashima T, et al. Observation of the vitreous in diabetic macular edema using swept-source OCT. *Adv Ophthalmol Vis Syst.* 2015;2(2):49–53. DOI: 10.15406/aovs.2015.02.00037
Statistical analysis

Statistical analysis was performed using the Excel 2012 software (Microsoft Corporation, Remond, WA, USA). The Mann-whitney U test and Student’s t-test were used to determine the significance of the differences between DME and No DME (control) groups. Statistical analysis of the CME, SRD, and PDR incidence was performed using Chi-square tests, with P<0.05 considered significant.

Result

Clinical characteristics

This study examined 81 eyes of 58 consecutive Japanese patients (35 males, 23 females) diagnosed with DME and 46 eyes of 39 diabetic patients (24 males, 15 females) without DME (Table 1). The mean central macular thickness of patients±SD was 434±139 mm (range, 302 to 800) in the patients with DME and 259±19 mm (range, 222 to 295) in the controls (P<0.0001). The Table 2 shows the prevalence of the different stages of PVD in patients with or without DME. There were no significant differences in the prevalence of each of the vitreous condition found between the series of patients with or without DME.

Table 3 shows characteristics of patients with DME. The age of patients in group 4 was significantly higher than those of the patients in group 2 (P=0.0002). There were 47 cases diagnosed as non-proliferated diabetic retinopathy (NPDR) and 34 diagnosed as proliferative diabetic retinopathy (PDR). In Group 2, 21 (60%) out of 35 eyes were classified as PDR, with this ratio shown to be significantly higher than that observed in the other groups (P=0.034).

At the initial visit, 50 of the 81 eyes with DME were found to have undergone pan retinal photocoagulation (PRP). There were no DM patients at their initial visit who were found to have received focal grid laser treatment, anti-VEGF injections, or steroid injections for DME. Cataract surgery had been performed prior to the initial visit in 13 of the 81 eyes with a median duration between the surgery and OCT examination of 33 months.

Table 1 Characteristics of patients, Diabetes, and diabetic retinopathy in subjects with diabetic macular edema (DME) and No DME

| Variable                                      | DME   | No DME | P Value |
|-----------------------------------------------|-------|--------|---------|
| No. of patients                               | 58    | 39     |         |
| No. of eyes                                   | 81    | 46     |         |
| Sex of patients                               |       |        | 0.906   |
| Male                                          | 35    | 24     |         |
| Female                                        | 23    | 15     |         |
| Age                                           | 60.8±12.7 | 60.3±11.9 | 0.74   |
| HbA1c                                         | 7.8±1.7 | 7.9±2.0 | 0.638   |
| Mean duration of diabetes±SD (years)          | 13.3±6.9 | 13.0±8.4 | 0.831  |
| Best-corrected visual acuity                  |       |        |         |
| Range                                         | 60/2000~20/16 | 20/32~20/16 | < 0.0001 |
| Type of diabetic retinopathy (No. of eyes)     |       |        | < 0.001 |
| No diabetic retinopathy                       | 0     | 16     |         |
| Non-proliferative diabetic retinopathy         | 43    | 26     |         |
| Proliferative diabetic retinopathy            | 38    | 4      |         |
| Laser treatment (No. of eyes)                 |       |        | < 0.0001 |
| None                                          | 31    | 72     |         |
| Panretinal photocoagulation                    | 50    | 9      |         |
| Focal laser                                   | 0     | 0      |         |
| Central macular thickness (mean±SD, mm)        | 434±139 | 259±19 | < 0.0001 |

Citation: Akiyama H, Itakura H, Kashima T, et al. Observation of the vitreous in diabetic macular edema using swept-source OCT. Adv Ophthalmol Vis Syst. 2015;2(2):49–53. DOI: 10.15406/aovs.2015.02.00037
Table 2 Prevalence of the different Stages of posterior vitreous detachment (PVD) in patients with diabetic Macular Edema (DME) and No DME

| Condition of PPVP | DME% of series (No. of eyes) | No DME % of series (No. of eyes) |
|-------------------|-----------------------------|----------------------------------|
| No PVD            | 22 (18 years)               | 35 (16 eyes)                     |
| Perifoveal PVD    | 43 (35 eyes)                | 22 (10 eyes)                     |
| Macular PVD       | 8 (6 eyes)                  | 8 (4 eyes)                       |
| Complete PVD      | 27 (22 eyes)                | 35 (16 eyes)                     |

PPVP, posterior precortical vitreous pocket; PVD, posterior vitreous detachment

Table 3 Characteristics of patients with diabetic macular edema (DME), age, HbA1C, classification of diabetic retinopathy (DR), photocoagulation, and the serious systemic diseases found in each group at the patient’s initial visit

| Age (average) | HbA1C (average) | Duration of Diabetes (Range)(Year) | DR classification | Prior treatment of PC | Time between PC and initial Visit (Month) | Previous severe illness at initial visit |
|---------------|-----------------|-----------------------------------|-------------------|----------------------|------------------------------------------|----------------------------------------|
| (Group 1) PVD(-) | 62.5±6.0       | 8.3±2.1                           | 15.4±9.1 (2~30)   | 14                   | 11                                       | 7                                      | AMI 2                                  |
| (Group 2) Perifoveal PVD | 54.9±14.3     | 7.8±1.8                           | 13.2±6.9 (3~35)   | 14                   | 21*                                      | 11                                     | 24                                    | AMI 2 Collagen disease I Chronic renal failure I |
| (Group 3) Macular PVD | 65.3±9.3       | 8.5±0.9                           | 10.8±5.8 (5~20)   | 4                    | 2                                       | 4                                      | 27.3±15.0                             | Cerebral infarction I |
| (Group 4) Complete PVD | 67.5±11.8      | 7.2±1.1                           | 12.9±5.8 (5~22)   | 15                   | 7                                       | 7                                      | 25.3±30.5                             | Cerebral infarction I |

*P=0.034

PC, photocoagulation; AMI, acute myocardial infarction

Anatomical results

For Groups 1 through 4, the number of eyes in each group was 18, 35, 6 and 22, respectively (Table 4A&B). The number of patients in each group was 12, 27, 5 and 14, respectively. OCT images indicated that 35 (43%) out of the 81 eyes had perifoveal PVD (Group 2). In contrast to the other groups, the Group 2 eyes were significantly accompanied by CME (94%) with or without SRD (P<0.05).

Table 4A Morphological characteristics in each group

| Morphology | Total |
|------------|-------|
| SRD        | CME   | SRD +CME |
| (Group 1) PVD(-) | 5     | 8        | 5       | 18     |
| (Group 2) Perifoveal PVD | 2     | 25*     | 8       | 35     |
| (Group 3) Macular PVD   | 0     | 4        | 2       | 6      |
| (Group 4) Complete PVD  | 13    | 6        | 3       | 22     |

Discussion

Previous studies have reported that breakdown of the BRB and cytokines such as VEGF might play an important part in the development of CME. The findings of our current study additionally suggest that CME and perifoveal PVD might be more closely related than previously thought. SS-OCT, which is a new generation of OCT that provides higher penetration into the choroid and sclera, has enabled more detailed and precise quantitative assessments of DME and the vitreous. In the current study, we defined DME as diabetic retinopathy with a central foveal thickness more than 300 mm. Since SS-OCT showed that CME or SRD existed in almost all of the cases treated clinically for these DMEs, we decided to focus on the CME or SRD configurations in DME. This enabled us to observe the condition of the vitreous and retina in detail regardless of the progressed cataract. In addition, the use of SS-OCT without indirect funduscopy and slit-lamp biomicroscopy made it possible to classify all of the DME cases into 4 groups. Gaucher et al. 18 and Ghazi et al. 19 have previously used OCT-1 or Stratus OCT to determine the relationship between DME and the vitreous, the resolution of these earlier devices is obviously inferior as compared to SS-OCT. While these previous studies included some cases of DME with ERM, our current study excluded ERM cases from our analysis, as we were attempting to investigate the specific relationship between the DME configurations and the vitreous.

Yamaguchi et al. 20 reported finding three cases of CME associated with DME that were caused by vitreous traction. Their results showed that a resolution of the cystic cavities and restoration of the foveal depression occurred after a spontaneous separation of the hyaloid at the fovea. The OCT images in our current study revealed the presence of macular PVD in 6 of 81 eyes, with all of these cases confirmed to have CMEs. However, it was not possible to determine when the adhesion of the posterior wall of the PPVP to fovea has been released. In addition, we also could not completely follow the natural course of these cases due to the intravitreal injection of anti-VEGF or because of interruptions of the observation periods at our hospital. In our study, the 21 (60%) of the 35 eyes that showed perifoveal PVD...
were classified as PDR, with this ratio significantly higher than that observed in the other groups. There have been several other reports that have examined the occurrence of PVD following panretinal photocoagulation for diabetic retinopathy.\(^{21,22}\) For example, Tagawa et al.\(^{21}\) reported that complete PVD occurred primarily in NPDR, while partial PVD was primarily observed in PDR as the proliferation interfered with its development. In the current study, we found that the DMEs exhibiting CME were primarily accompanied by perifoveal PVD, thereby meeting the PDR classification of diabetic retinopathy. One potential limitation of our current study was that both eyes of the same subject were used as independent observations. In addition, it has been speculated that there may be numerous other factors involved in the mechanism of DME. Thus, further investigations will need to be undertaken in order to conclusively define the mechanism of DME.

### Table 4B

| PVD(−) | Perifoveal PVD | Macular PVD | Complete PVD |
|--------|----------------|-------------|--------------|
|        |                |             |              |

| PVD=posterior vitreous detachment |
|----------------------------------|
*P=0.0005

### Conclusion

Although we are unable to definitively determine the extent of the traction between the posterior wall of the PPVP and fovea, we can speculate that, in addition to vascular hyperpermeability, the vitreous might also be able to influence the development of CME associated with DME. We would like to emphasize the basis of CME or SRD is speculated that, in addition to vascular hyperpermeability, the vitreous might be able to lead to morphologic changes of DME occasionally. Furthermore, SS-OCT appears to be useful for observing the surface of the retina and the vitreous in DME.

### Acknowledgments

None.

### Conflicts of interest

Author declares that there is no conflict of interest.

### References

1. Moss SE, Klein R, Klein BE. The 14-year incidence of visual loss in a diabetic population. *Ophthalmology*. 1998;105(6):998−1003.
2. Otani T, Kishi S, Maruyama Y. Patterns of diabetic macular edema with optical coherence tomography. *Am J Ophthalmol*. 1999;127(6):688−693.
3. Murakami T, Nishijima K, Sakamoto A, et al. Foveal cystoid spaces are associated with enlarged foveal avascular zone and microaneuerysms in diabetic macular edema. *Ophthalmology*. 2011;118(2):359−367.
4. Nasrallah FP, Jalkh AE, Van Coppenolle F, et al. The role of the vitreous in diabetic macular edema. *Ophthalmology*. 1988;95(10):1335−1339.
5. Foos RY, Kreiger AE, Forsythe AB, et al. Posterior vitreous detachment in diabetic subjects. *Ophthalmology*. 1980;87(2):122−128.
6. Lewis H, Abrams GW, Blumenkranz MS, et al. Vitrectomy for diabetic macular traction and edema associated with posterior hyaloidal traction. *Ophthalmology*. 1992;99(5):753−759.
7. Gandrafer A, Messmer EM, Ulbig MW, et al. Resolution of diabetic macular edema after surgical removal of the posterior hyaloid and the inner limiting membrane. *Retina*. 2000;20(2):126−133.
8. Harbour JW, Smiddy WE, Flynn HW, et al. Vitrectomy for diabetic macular edema associated with a thickened and taut posterior hyaloid membrane. *Am J Ophthalmol*. 1996;121(4):405−413.
9. Ikeda T, Sato K, Katano T, et al. Vitrectomy for cystoid macular oedema with attached posterior hyaloid membrane in patients with diabetes. *Br J Ophthalmol*. 1999;83(1):12−14.
10. Haller JA, Qin H, Apte RS, et al. Vitrectomy outcomes in eyes with diabetic macular edema and vitreomacular traction. *Ophthalmology*. 2010;117(6):1087−1093.
11. Flaxel CJ, Edwards AR, Aiello LP, et al. Factors associated with visual acuity outcomes after vitrectomy for diabetic macular edema: Diabetic retinopathy clinical research network. *Retina*. 2010;30(9):1488−1495.
12. Aiello LP, Avery RL, Arrigg PG, et al. Vascular endothelial growth factor in ocular fluid of patients with diabetic retinopathy and other retinal disorders. *N Engl J Med*. 1994;331(22):1480−1487.
13. Adams AP, Miller JW, Bernal MT, et al. Increased vascular endothelial growth factor levels in the vitreous of eyes with proliferative diabetic retinopathy. *Am J Ophthalmol*. 1994;118(4):445−450.
14. Watanabe D, Sazuma K, Matsui S, et al. Erythropoietin as a retinal angiogenic factor in proliferative diabetic retinopathy. *N Engl J Med*. 2005;353(8):782−792.
15. Funatsu H, Yamashita H, Noma H, et al. Increased levels of vascular endothelial growth factor and interleukin-6 in the aqueous humor of diabetics with macular edema. *Am J Ophthalmol*. 2002;133(1):70−77.
16. Itakura H, Kishi S, Li D, et al. Observation of posterior pre cortical vitreous pocket using swept-source optical coherence tomography. *Invest Ophthalmol Vis Sci*. 2013;54(5):3102−3107.
17. Haritoglou C, Kook D, Neubauer A, et al. Intravitreal bevacizumab (avastin) therapy for persistent diffuse diabetic macular edema. *Retina*. 2006;26(9):999−1005.
18. Gaecher D, Tadayoni R, Erginay A, et al. Optical coherence tomography assessment of the vitreoretinal relationship in diabetic macular edema. *Am J Ophthalmol*. 2005;139(5):807−813.
19. Ghazi NG, Ciralsky JB, Shah SM, et al. Optical coherence tomography findings in persistent diabetic macular edema: The vitreomacular interface. *Am J Ophthalmol*. 2007;144(5):747−754.
20. Yamaguchi Y, Otani T, Kishi S. Resolution of diabetic cystoid macular edema associated with spontaneous vitreofoveal separation. *Am J Ophthalmol*. 2003;135(1):116−118.
21. Tagawa H, McMeel JW, Furukawa H, et al. Role of the vitreous in diabetic retinopathy. I. Vitreous changes in diabetic retinopathy and in physiologic aging. *Ophthalmology*. 1986;93(5):596−601.
22. Sebag J, Buzney SM, Belyea DA, et al. Posterior vitreous detachment following panretinal laser photocoagulation. *Graefes Arch Clin Exp Ophthalmol*. 1990;228(1):5−8.