Variations of posterior vitreous detachment

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Abstract

Aims—To identify variations in posterior vitreous detachment (PVD) and establish a clinical classification system for PVD.

Methods—400 consecutive eyes were examined using biomicroscopy and vitreous photography and classified the PVD variations—complete PVD with collapse, complete PVD without collapse, partial PVD with thickened posterior vitreous cortex (TPVC), or partial PVD without TPVC.

Results—In each PVD type, the most frequently seen ocular pathologies were as follows: in complete PVD with collapse (186 eyes), age related changes without vitreoretinal diseases (77 eyes, 41.4%) and high myopia (55 eyes, 29.6%); in complete PVD without collapse (39 eyes), uveitis (23 eyes, 59.0%) and central retinal vein occlusion (8 eyes, 20.5%); in partial PVD with TPVC (64 eyes), proliferative diabetic retinopathy (30 eyes, 46.9%) and in partial PVD without TPVC (111 eyes), age related changes without vitreoretinal diseases (62 eyes, 55.9%). This PVD categorisation was significantly associated with the prevalence of each vitreoretinal disease (p<0.0001, χ² test on contingency table).

Conclusions—PVD variations can be classified into four types, which is clinically useful because each type corresponds well to specific vitreoretinal changes.

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Results of previous studies have shown that characteristic types of posterior vitreous detachment (PVD) are observed with specific vitreoretinal diseases. For example, partial PVD is a characteristic vitreous change in proliferative diabetic retinopathy; and complete PVD is observed in typical age related PVD. The morphological configurations of PVD vary, and the variations strongly affect the prognosis of the vitreoretinal diseases. For example, preretinal neovascularisation in diabetic retinopathy rarely occurs when there is complete PVD. However, the causes of these PVD variations are unknown. Therefore, we classified the PVD types found in the clinical cases and correlated the various vitreoretinal diseases associated with these PVD types.

Patients and methods

To clarify the typical vitreous changes occurring in retinal diseases, we examined 400 consecutive eyes with PVD (321 patients; mean age 58.6 years, SD 16.6) by biomicroscopy and vitreous photography using the El Bayad-Kajura preset lens, the +90 dioptre preset lens, or both. This technique provides strong dynamic observation of the vitreous more easily and is more comfortable for the patients. These patients had been referred to the department of vitreous study at Asahikawa Medical College for further evaluation of vitreoretinal changes.

The following ophthalmic conditions were seen in the 400 study eyes: age related change without significant vitreoretinal disease, 141 eyes (104 patients; mean age 67.5 (SD 8.0) years); uveitis, 74 eyes (55 patients; mean age 47.3 (20.1) years); high myopia (refractive error greater than −8.0 dioptres), 66 eyes (60 patients; mean age 51.2 (17.1) years); proliferative diabetic retinopathy, 33 eyes (29 patients, mean age 52.8 (17.6) years); non- proliferative diabetic retinopathy, 23 eyes (17 patients; mean age 66.4 (11.6) years); branch retinal vein occlusion, 27 eyes (25 patients; mean age 66.3 (11.6) years); central retinal vein occlusion, 20 eyes (20 patients, mean age 59.5 (16.1) years); retinal breaks, nine eyes (seven patients, mean age 52.7 (16.1) years); and retinitis pigmentosa, seven eyes (four patients, mean age 48.8 (19.3) years).

The morphological variations of PVD were classified into four types:

1. Complete PVD with collapse, in which the vitreous gel exhibits liquefaction and, therefore, a large retrocortical space and smooth movement of the detached vitreous with ocular movement are observed. In this type, the posterior vitreous cortex has a characteristically sigmoidal shape when the patient is sitting (Fig 1). Weiss’s ring is typically observed on the posterior vitreous cortex.
(2) Complete PVD without collapse, in which the vitreous gel is minimally liquefied and, therefore, movement of the detached vitreous is limited. When vitreous shrinkage is advanced, the retrocortical space is clearly defined, and the posterior vitreous cortex is convex (Fig 2). Weiss’s ring is also typically observed on the posterior vitreous cortex.

(3) Partial PVD with a thickened posterior vitreous cortex (TPVC), in which the detached posterior vitreous cortex is thickened, taut, and anchored at two points (Fig 3). Some variations in this category are seen, because the two points of attachment vary—that is, at the vitreous base, the optic disc, along the vascular arcade, the neovascular complex, and the macula.

(4) Partial PVD without TPVC, in which the posterior vitreous cortex is not thickened (Fig 4). In the last category, we also included vitreous gel attachment to the macular area through a round defect in the posterior vitreous cortex (Fig 5). This pathology was first described by Jaffe,8 characterised and photographed by Sebag and Balazs,16 17 and documented photographically and videographically in clinical cases by Kakehashi and associates.18 19

The $\chi^2$ test and the Student’s $t$ test were used for data analysis. The level of statistical significance was $p < 0.05$.

**Results**

The mean patient ages in years in each group were as follows: complete PVD with collapse, 61.6; complete PVD without collapse, 44.6;
partial PVD with TPVC, 51.9; and partial PVD without TPVC, 62.5. The mean patient ages were significantly higher in patients with complete PVD with collapse and partial PVD without TPVC than in the other groups (p = 0.0001 by the Student’s t test).

Table 1 shows the prevalence of ocular disease in each PVD type. All cases in this study were included in this table. The PVD categorisation was statistically significantly associated with the prevalence of each vitreoretinal disease (p<0.0001, by goodness of fit for $\chi^2$, 4 x 9 contingency table). In each PVD type, the most frequently seen ocular pathologies were as follows: in complete PVD with collapse (186 eyes), age related change without vitreoretinal diseases (77 eyes, 41.4%) and high myopia (55 eyes, 29.6%); in complete PVD without collapse (39 eyes), uveitis (23 eyes, 59.0%) and central retinal vein occlusion (eight eyes, 20.5%); in partial PVD with TPVC (64 eyes), proliferative diabetic retinopathy (30 eyes, 46.9%); and in partial PVD without TPVC (111 eyes), age related change without vitreoretinal diseases (62 eyes, 55.9%). The prevalence of ocular disease was statistically different among the PVD types (p<0.0001, between complete PVD with collapse and complete PVD without collapse, complete PVD with collapse and partial PVD with TPVC, complete PVD without collapse and partial PVD with TPVC, partial PVD with TPVC and partial PVD without TPVC; p<0.001, complete PVD with collapse and partial PVD without TPVC).

![Figure 3](image1.png)

Figure 3  (A) Schematic sketch of partial posterior vitreous detachment (PVD) with a thickened posterior vitreous cortex (TPVC), which exhibits minimal movement with ocular movement. (B) In a patient with proliferative diabetic retinopathy, the fundus examination by indirect ophthalmoscopy revealed tractional retinal detachment. Biomicroscopic vitreous examination revealed vitreous traction upon the detached retina (arrowhead) with a condensed posterior vitreous cortex (arrows). This case was classified as partial PVD with TPVC.

![Figure 4](image2.png)

Figure 4  (A) Schematic sketch of partial posterior vitreous detachment (PVD) without a thickened posterior vitreous cortex (TPVC), which exhibits some mobility with ocular movement. (B) In a patient who complained of flashes in his right eye, the fundus examination by indirect ophthalmoscopy did not reveal retinal disease. The biomicroscopic vitreous examination showed localised PVD in the upper quadrants of the right eye. The vitreous gel was moderately liquefied, and with ocular movement, the posterior vitreous cortex (arrow) was not condensed and had a smooth, wavy motion. This case was classified as partial PVD without TPVC.
The prevalence of each PVD type in common vitreoretinal diseases found in the present study is shown in Table 2. The prevalence of each PVD type in each common vitreoretinal disease was statistically significantly different except for that found between uveitis and central retinal vein occlusion (p>0.5), between branch retinal vein occlusion and non-proliferative diabetic retinopathy (p>0.1), and between non-proliferative diabetic retinopathy and central retinal vein occlusion (p>0.1). In other words, uveitis and central retinal vein occlusion had a similar pattern of the prevalence of each PVD.

### Discussion

Some PVD classifications have been proposed previously, the most practical of which was suggested by Tolentino and colleagues. These authors classified PVD based on whether or not detachment of the posterior vitreous cortex is complete or incomplete and if the detached cortex collapsed. They also classified atypical PVD into three types, funnel-shaped, hammock-shaped, and PVD with a dehiscent cortex.

Even though their classification accurately represents the PVD variations, it does not correspond well to specific vitreoretinal changes. For example, both partial PVD with a condensed posterior vitreous cortex resulting from proliferative diabetic retinopathy and partial PVD without a condensed posterior vitreous cortex resulting from age related changes are classified as incomplete PVD without collapse. Therefore, we refined the old classification and made a more precise classification and correlated it with specific vitreoretinal changes.

Three main factors are associated with PVD—that is, vitreous liquefaction, vitreous shrinkage, and weakening of vitreoretinal adhesion. Vitreous liquefaction frequently is observed in association with age related vitreous changes and in highly myopic eyes. The mechanism of age or disease related vitreous liquefaction is uncertain. However, some investigators have suggested that free radicals cause hyaluronan depolymerisation which leads to destruction of the gel structure (liquefaction). Other studies showed that in

**Table 1** Prevalence of ocular diseases by posterior vitreous detachment (PVD) (% of eyes)

<table>
<thead>
<tr>
<th>Condition</th>
<th>CPVD with collapse</th>
<th>CPVD without collapse</th>
<th>PPVD with TPVC</th>
<th>PPVD without TPVC</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>77 (41.4)</td>
<td>1 (2.6)</td>
<td>1 (1.6)</td>
<td>62 (55.9)</td>
<td>141</td>
</tr>
<tr>
<td>Uveitis</td>
<td>21 (11.3)</td>
<td>23 (59.0)</td>
<td>1 (1.6)</td>
<td>8 (7.2)</td>
<td>74</td>
</tr>
<tr>
<td>High myopia</td>
<td>55 (29.6)</td>
<td>2 (5.1)</td>
<td>1 (1.6)</td>
<td>8 (7.2)</td>
<td>66</td>
</tr>
<tr>
<td>PDR</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>30 (46.9)</td>
<td>3 (2.7)</td>
<td>33</td>
</tr>
<tr>
<td>BRVO</td>
<td>8 (4.3)</td>
<td>1 (2.6)</td>
<td>10 (15.6)</td>
<td>8 (7.2)</td>
<td>27</td>
</tr>
<tr>
<td>NPDR</td>
<td>11 (5.9)</td>
<td>2 (5.1)</td>
<td>2 (3.1)</td>
<td>8 (7.2)</td>
<td>23</td>
</tr>
<tr>
<td>CRVO</td>
<td>6 (3.2)</td>
<td>8 (20.5)</td>
<td>2 (3.1)</td>
<td>4 (3.6)</td>
<td>20</td>
</tr>
<tr>
<td>Retinal break</td>
<td>5 (2.7)</td>
<td>2 (5.1)</td>
<td>0 (0)</td>
<td>2 (1.8)</td>
<td>9</td>
</tr>
<tr>
<td>RP</td>
<td>3 (1.6)</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>4 (3.6)</td>
<td>7</td>
</tr>
<tr>
<td>Total</td>
<td>186</td>
<td>39</td>
<td>64</td>
<td>111</td>
<td>400</td>
</tr>
</tbody>
</table>

*Age indicates age related change without evidence of vitreoretinal disease.

CPVD = complete posterior vitreous detachment; PPVD = partial posterior vitreous detachment; TPVC = thickened posterior vitreous cortex; PDR = proliferative diabetic retinopathy; NPDR = non-proliferative diabetic retinopathy; BRVO = branch retinal vein occlusion; CRVO = central retinal vein occlusion; RP = retinitis pigmentosa.

p<0.0001; CPVD with collapse – CPVD without collapse; CPVD with collapse – PPVD with TPVC; CPVD without collapse – PPVD with TPVC; CPVD without collapse – PPVD without TPVC; PPVD with TPVC – PPVD without TPVC.

p<0.001; CPVD with collapse – PPVD without TPVC.
diabetes glucose binds the collagen and induces collagen fibril crosslinking and aggregation with concomitant liquefaction.\textsuperscript{23,24} Vitreous shrinkage frequently is observed in disease-related vitreous change. Blood components and cell mediation are thought to be the primary mechanisms of vitreous shrinkage in disease-related PVD.\textsuperscript{25–28} Free radical induced crosslinking of collagen fibres also is thought to cause vitreous shrinkage.\textsuperscript{29,30} The mechanism of weakening of vitreoretinal adhesion is uncertain. Recent reports have suggested that changes in the extracellular matrix combined with the thickness of the internal limiting lamina and the hyalocytes found in the posterior vitreous cortex may be associated with changes in vitreoretinal adhesion.\textsuperscript{31,32}

Complete PVD with collapse is typical of the age-related type, which is characterised by reduced vitreoretinal adhesion, vitreous gel liquefaction, and mild vitreous shrinkage. Highly myopic eyes also frequently have complete PVD with collapse. Diffuse chorioretinal atrophy in high myopia might induce vitreous liquefaction and somehow reduce vitreoretinal adhesion.

Complete PVD without collapse was most frequently associated with uveitis and central retinal vein occlusion in the present study. Increased blood components or inflammatory cells induce vitreous gel shrinkage. The vitreous gel did not exhibit as much liquefaction because these patients are younger. Vitreous shrinkage induces PVD with minimal vitreous liquefaction, resulting in complete PVD without collapse in these patients.

Partial PVD with TPVC typically was observed in association with proliferative diabetic retinopathy, in which new vessels grow out of the retina and the optic disc into the vitreous cortex.\textsuperscript{33} Strong adhesion results from cell migration and proliferation in the vitreous. Chronic leakage of the blood components induces vitreous shrinkage, especially in the posterior vitreous cortex in these cases. The concurrence of strong vitreoretinal adhesion and vitreous shrinkage produces this kind of PVD. Clinically, detecting this type of PVD is very important. In diabetic retinopathy, it is a precursor of tractional retinal detachment. Partial PVD without TPVC was found mostly in patients without retinal disease. This may be a transitory phase of complete PVD with collapse or another result of age-related vitreous change, because the difference in patient ages between complete PVD with collapse and this type was not significant. Two types of partial PVD without TPVC were found in the present study, that in which the posterior vitreous cortex is not taut (Fig 4) and in which the vitreous gel is attached to the macular area through a round defect in the posterior vitreous cortex (Fig 5). The latter type may be easily misdiagnosed as complete PVD unless the examiner carefully scans the posterior fundus. This type of PVD is more common than the former type and has to be diagnosed correctly.\textsuperscript{10–15} because vitreous attachment is an important factor affecting the prognosis of macular diseases such as macular breaks,\textsuperscript{34,35} or preretinal membranes.\textsuperscript{18,36}

In summary, our new clinical classification of PVD is precise and corresponds well with the presence of various types of retinal disease. This classification of PVD might be useful for evaluating vitreoretinal disorders.

### Table 2 Prevalence of posterior vitreous detachment (PVD) by ocular diseases (no of eyes (%))

<table>
<thead>
<tr>
<th>CPVD with collapse</th>
<th>CPVD without collapse</th>
<th>PPVD with TPVC</th>
<th>PPVD without TPVC</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age*</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>77 (54.0)</td>
<td>1 (0.7)</td>
<td>1 (0.7)</td>
<td>62 (44.0)</td>
<td>141</td>
</tr>
<tr>
<td>Uveitis</td>
<td>21 (28.4)</td>
<td>23 (31.3)</td>
<td>18 (24.3)</td>
<td>74</td>
</tr>
<tr>
<td>55 (83.3)</td>
<td>2 (3.0)</td>
<td>1 (1.5)</td>
<td>8 (12.2)</td>
<td>66</td>
</tr>
<tr>
<td>PDR</td>
<td>0 (0)</td>
<td>0 (0)</td>
<td>30 (9.0)</td>
<td>33</td>
</tr>
<tr>
<td>BRVO</td>
<td>8 (29.6)</td>
<td>1 (3.7)</td>
<td>10 (37.1)</td>
<td>27</td>
</tr>
<tr>
<td>NPDR</td>
<td>11 (47.8)</td>
<td>2 (8.7)</td>
<td>2 (8.7)</td>
<td>23</td>
</tr>
<tr>
<td>CRVO</td>
<td>6 (30.0)</td>
<td>8 (40.0)</td>
<td>2 (10.0)</td>
<td>20</td>
</tr>
</tbody>
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*Age indicates age related change without evidence of vitreoretinal disease.

CPVD = complete posterior vitreous detachment; PPVD = partial posterior vitreous detachment; TPVC = thickened posterior vitreous cortex; PDR = proliferative diabetic retinopathy; NPDR = non-proliferative diabetic retinopathy; BRVO = branch retinal vein occlusion; CRVO = central retinal vein occlusion.

p<0.0001; aged – uveitis aged – high myopia; aged – PDR; aged – CRVO; uveitis – high myopia; uveitis – PDR; high myopia – PDR; high myopia – BRVO; high myopia – CRVO; PDR – NPDR; PDR – CRVO.
p<0.001; aged – BRVO.
p<0.005; aged – NPDR; aged – high myopia; aged – PDR; aged – BRVO; PDR – BRVO.
p<0.01; aged – BRVO; high myopia – NPDR; BRVO – CRVO.
p<0.05; uveitis – BRVO; uveitis – NPDR.
p<0.1; BRVO – NPDR; NPDR – CRVO.
p>0.5; uveitis – CRVO.

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