Vitreous levels of intercellular adhesion molecule 1 (ICAM-1) as a risk indicator of proliferative vitreoretinopathy

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Abstract

Objective—To investigate whether high vitreous levels of the soluble intercellular adhesion molecule 1 (sICAM-1) may be related to clinical risk factors of proliferative vitreoretinopathy (PVR) and whether their measurement may serve as an additional risk indicator of this complication in eyes with rhegmatogenous retinal detachment (RRD).

Methods—Levels of sICAM-1 were measured by enzyme linked immunosorbent assays (ELISA) in vitreous from 36 eyes with RRD clinically considered to be at high risk of developing PVR (large retinal breaks, vitreous haemorrhage, long standing RRD, and previous vitreoretinal surgery). Levels of sICAM-1 in this group were compared with those in vitreous from 31 eyes with RRD without clinical risk factors for PVR, 32 eyes with established PVR and 10 eyes with macular holes.

Results—Vitreous from eyes with RRD at high risk of developing PVR contained significantly higher levels of sICAM-1 (range 6.1–97.7 ng/ml; Mann–Whitney test, p=0.0002) than those from eyes with RRD at low risk of developing this complication (range 4.8–17.7 ng/ml). Vitreous sICAM-1 levels in eyes with RRD at high risk of developing PVR were significantly lower than in eyes with established PVR (p=0.037), but higher than in eyes with macular holes (p <0.0001). Levels of sICAM-1 >15 ng/ml (≥ median of the levels present in control eyes) provide a useful cut off point for a highly specific test (96.7%) with high positive (91.6%) and negative (96.7%) predictive values, despite a relatively low sensitivity (30.5%).

Conclusions—The present findings suggest that laboratory measurement of sICAM-1 levels in vitreous from eyes with RRD may constitute an additional factor for identifying patients at high risk of PVR. Hence, determination of sICAM-1 levels may aid in the monitoring of patients likely to develop this complication and in the identification of patients who may benefit from adjuvant anti-inflammatory therapy.
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from patients with established PVR and other retinal proliferative disorders.15–16

Our previous observations that high levels of this molecule were present in two eyes which, at the time of vitrectomy for RRD, did not have PVR greater than grade B, but that later developed this complication,17 prompted us to investigate whether high vitreous levels of sICAM-1 are related to the known clinical risk factors for PVR, and whether their laboratory measurement could be used as an additional factor to identify those patients likely to develop this complication. On this basis, we investigated the levels of sICAM-1 in vitreous from eyes with uncomplicated RRD with well recognised features of high clinical risk of developing PVR,19,20 and compared these with the levels found in vitreous from eyes with RRD at low clinical risk of PVR, and eyes with established PVR. Vitreous samples from eyes undergoing pars plana vitrectomy for treatment of macular holes were used as controls.

Patients and methods

Vitreous and serum samples were obtained from 109 patients undergoing pars plana vitrectomy (PPV) for the treatment of uncomplicated RRD (less than grade B, 67 patients), established PVR (32 patients), or macular holes (10 patients). Of the 67 patients with uncomplicated RRD, 31 were at low clinical risk of PVR, while 36 were at high clinical risk of PVR, as judged by well recognised clinical features that predispose to this complication.19,20 These included the presence of giant or very large retinal breaks (13 patients), severe vitreous haemorrhage (six patients), long standing RRD (>3 months) with history of previous retinal surgery (17 patients). In the uncomplicated RRD group, vitrectomy was considered the best choice of operation because of either opacities in the media (obscuring the fundal view), or to the complex size, site, or distribution of retinal breaks. Eyes with established PVR were grade B+ according to the classification of Machemer et al.21 Ten vitreous specimens from eyes undergoing PPV for treatment of macular holes were used as disease controls. Clinical follow up of the patients under study revealed that none of the patients with low clinical risk of PVR developed this complication, while one of the 36 patients presenting with high clinical risk factors actually developed PVR. Vitreous specimens (approximately 0.75 ml) were centrifuged and transferred to cryotubes for storing at −70°C until use. The study was approved by the local health authority ethics committee and it was performed in accordance with the ethical standards laid down in the 1964 Declaration of Helsinki.

Levels of sICAM-1 were measured by enzyme linked immunosorbent assays (ELISA) using commercially available kits (R&D Systems, Abingdon) as follows. Microtitre well plates coated with anti-ICAM antibodies were incubated with 100 µl of a 1:10 dilution of vitreous, together with 100 µl of anti-ICAM-1-HRP conjugate for 1.5 hours, after which antibodies and test samples were removed and the plate washed six times with phosphate buffered saline (PBS) containing 0.05% Tween-20. The amount of conjugated antibodies was detected by addition of 100 µl of tetramethylbenzidine (substrate) and incubated for 30 minutes at room temperature. The reaction was stopped by addition of 100 µl of 1M H₂SO₄, and the absorbance read at 450 nm, in a Dynatech MR5000 reader. Levels of sICAM-1 present in the vitreous or serum samples were interpolated from specific calibration curves prepared with known standard solutions. The specificity of the reaction was controlled by addition of interleukin 6 and human serum albumin to normal cadaveric vitreous to a final protein concentration of 3.0 mg/ml (similar to the levels observed in pathological specimens). Addition of these proteins did not modify the levels of sICAM-1 detected in the vitreous samples, thus indicating the specificity and validity of the test.

STATISTICAL ANALYSIS OF THE RESULTS

The significance of difference between corresponding groups of observations was evaluated by the Mann–Whitney test. Acceptable significance was recorded when p values were <0.05. The sensitivity, specificity, and positive and negative predictive values of sICAM-1 measurements were evaluated according to standard methods.22 For analysis, values corresponding to 3× and 2× the median levels of sICAM-1 observed in eyes with macular holes (>15 ng/ml and >10 ng/ml, respectively) were taken as cut off points for evaluation of the test. A comparison was made between these levels and the presence or absence of clinical risk factors of PVR. Accordingly, sensitivity was defined as the prevalence of the positive test in subjects with clinical risk factors for PVR. Specificity was defined as the prevalence of the negative test in individuals without clinical risk factors for PVR. The positive predictive value was calculated as the proportion of subjects with a positive test who presented with clinical risk factors for PVR, while the negative predictive value was calculated as the proportion of subjects with negative test who did not present with clinical risk factor for PVR.22

Results

VITREOUS LEVELS OF SICAM-1 IN EYES WITH HIGH CLINICAL RISK OF PVR

Figure 1 shows that levels of sICAM-1 in vitreous from eyes with uncomplicated RRD at high clinical risk of PVR are significantly higher (range 6.1–97.8 ng/ml) than in vitreous from eyes with RRD at low clinical risk of PVR (range 4.9–17.7 ng/ml: Mann–Whitney U test, p=0.00027) and vitreous from eyes with macular holes (range 4.8–7.6 ng/ml: Mann–Whitney U test, p=0.000038). However, vitreous sICAM-1 levels in eyes with RRD at high risk of developing PVR were significantly lower than in eyes with established PVR (Mann–Whitney U test: p=0.037), and vitreous levels of sICAM-1 in eyes with RRD at low risk of PVR, were significantly higher than in eyes with macular holes (Mann–Whitney U test: p=0.00047). Only one of the 31 vitreous
vitreous sICAM-1 were present in vitreous from eyes with macular holes (15 ng/ml).

Table 1 Analysis of the sensitivity, specificity, and predictive value of vitreous sICAM-1 measurement

<table>
<thead>
<tr>
<th>Factor for positive test*</th>
<th>Sensitivity†</th>
<th>Specificity‡</th>
<th>Positive predictive value§</th>
<th>Negative predictive value¶</th>
</tr>
</thead>
<tbody>
<tr>
<td>15 ng/ml (3 × MoC)</td>
<td>11/36 (30.5%)</td>
<td>30/31 (96.7%)</td>
<td>11/12 (91.6%)</td>
<td>30/31 (96.7%)</td>
</tr>
<tr>
<td>10 ng/ml (2 × MoC)</td>
<td>21/36 (58.3%)</td>
<td>21/31 (67.7%)</td>
<td>21/31 (67.7%)</td>
<td>21/36 (58.3%)</td>
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</table>

*MoC, median of the control; †prevalence of positive test in subjects with clinical risk factors of PVR; ‡prevalence of negative test in subjects without clinical risk factors of PVR; §proportion of subjects with positive tests that presented with clinical risk of PVR; ¶proportion of subjects with negative test that did not present with clinical risk of PVR.

Figure 1 Levels of sICAM-1 in vitreous from eyes with rhegmatogenous retinal detachment (RRD) at high clinical risk of developing proliferative vitreoretinopathy (PVR). Comparison with vitreous levels of the same molecule in eyes with RRD at low risk of PVR, established PVR and eyes with macular holes. *Mann–Whitney U test, p=0.00047 (v eyes with macular holes); **Mann–Whitney U test, p=0.00027 (v RRD at low risk of PVR); ***Mann–Whitney U test, p=0.000038 (v eyes with macular holes); ****Mann–Whitney U test, p=0.037 (v RRD at high risk of PVR), p=0.000075 (v RRD at low risk of PVR). The broken line represents three times the median of the ICAM-1 levels in vitreous from eyes with macular holes.

Figure 2 Comparison between vitreous levels of sICAM-1 in eyes with various factors known to predispose to PVR. *Mann–Whitney U test, p= 0.00003 (v eyes with macular holes); **Mann–Whitney U test, p= 0.00033 (v eyes with macular holes); ***Mann–Whitney U test, p= 0.044 (v eyes with vitreous haemorrhage); ****Mann–Whitney U test, p=0.0028 (v eyes with macular holes).

Discussion

This study shows that vitreous from eyes with RRD at high risk of developing PVR contained higher levels of sICAM than those from eyes with RRD without clinically recognisable risk of PVR. Analysis of the various clinical factors known to predispose to PVR showed that there were no significant differences in sICAM-1 vitreous levels between eyes with large retinal breaks and eyes that had undergone previous operations or that presented with vitreous haemorrhage (p >0.5). However, vitreous from eyes which had been subjected to previous operations contained higher levels of sICAM-1 than eyes with severe vitreous haemorrhage (Mann–Whitney U test: p=0.044).
reinforces our view that levels of vitreous sICAM-1 depend upon the severity of cytokine mediated reactions at the blood-retinal barrier and therefore upon the severity of inflammation caused by the trauma of RRD.

Since considerable levels of sICAM-1 may be found in normal serum (102–450 ng/ml), it was important to consider the possibility that sICAM-1 found in eyes with vitreous haemorrhage may have derived from blood. As such, one should expect much higher levels of this molecule in vitreous from eyes with vitreous haemorrhage when compared with other clinical risks for PVR. However, sICAM-1 levels in eyes with vitreous haemorrhage were significantly lower than in vitreous from eyes which had undergone previous retinal surgery, strongly suggesting that this molecule is locally released by cells of the retinal microenvironment. This is supported by our previous observations that ICAM-1 expression is increased on the vascular endothelium of retinal vessels during anterior PVR and on the extracellular matrix and infiltrating cells of PVR membranes.

Vitreoretinal surgeons would prefer to prevent PVR as the surgery for PVR itself is unpredictable, both anatomically and visually. The clinical factors known to predispose RRD to PVR include the presence of vitreous haemorrhages, large retinal breaks, longstanding retinal detachments, and previous operations. However, at present there are no laboratory tests that may help to identify those at risk of PVR. The present study showed that vitreous levels of sICAM-1 > 15 ng/ml (3x median of the levels present in control eyes) provide a useful cut off point for a highly specific test (96.7%) with high positive (91.6%) and negative (96.7%) predictive values, despite a relatively low sensitivity (30.5%). Hence, we propose that assessment of the vitreous levels of sICAM-1 may constitute an additional factor for identifying patients at high risk of developing PVR.

In the present study only one of the 36 patients with high clinical risk factors for PVR developed this complication, which is in accordance with the known incidence of PVR. On this basis we could not predict PVR development by high vitreous levels of sICAM-1 for which, in our view, the significance of high vitreous levels of sICAM-1 as a risk indicator for PVR lies in the relation between the laboratory findings and the presence of clinical risk factors for PVR.

A common feature of the clinical risk factors of PVR is a high degree of inflammation, which bears the common features of cell migration and release of cytokines and growth factors within the vitreous cavity. High vitreous levels of sICAM-1 will raise the index of suspicion for the development of PVR, encourage the advisability of monitoring patients after surgery, and the targeting of those likely to benefit from adjunct anti-inflammatory therapy. Laboratory evidence of those at risk of developing PVR will advance the clinical and surgical management of this condition.

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Proliferative vitreoretinopathy (PVR) is a serious complication of retinal detachment surgery. The presence of the molecule 1 (ICAM-1) as a risk indicator of PVR has been investigated. In the study, vitreous levels of intercellular adhesion molecule 1 (ICAM-1) were measured in patients with retinal detachment and compared to controls. The results showed a significant increase in ICAM-1 levels in patients with PVR, indicating that ICAM-1 may be a useful marker for the risk indicator of PVR. These findings support the use of ICAM-1 as a potential target for the prevention and treatment of PVR.