Risk factors for proliferative sickle retinopathy

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Abstract
The prevalence, incidence, and risk factors associated with proliferative sickle retinopathy (PSR) were investigated in 786 patients with homozygous sickle cell (SS) disease and 533 patients with sickle cell haemoglobin C (SC) disease. PSR was more common in SC disease, in which there was a significant predominance of males, and it increased with age in both genotypes. In SC disease the risk of developing PSR was highest between 15 and 24 years in males, between 20 and 39 years in females, and in SS disease between 25 and 39 years in both sexes. PSR tended to be bilateral, especially in SC disease. There was no evidence of familial clustering of PSR in SC siblings, and insufficient numbers of SS siblings were available to test for clustering. Haematological risk factors associated with PSR in SS disease were a high haemoglobin in males and a low fetal haemoglobin in both sexes and in SC disease, a high mean cell volume, and a low fetal haemoglobin in females.

Proliferative sickle retinopathy (PSR) is the major sight threatening complication in sickle cell eye disease, and may reach frequencies as high as 70% in patients with sickle cell haemoglobin C (SC) disease. The risk factors for its development are still largely unknown, partly because detailed ocular information has not been available in sickle cell populations large enough to allow comprehensive statistical analysis. A study of 261 patients with homozygous sickle cell (SS) disease including 29 with PSR noted that male patients with PSR had significantly higher total haemoglobin (Hb) and lower fetal haemoglobin (HbF) but that no correlations occurred in females.1 Another study of 243 patients with SC disease including 90 with PSR found significant correlations of PSR with low HbF in both sexes and with a high mean cell volume (MCV) in males.2 Since those earlier Jamaican reports 333 cases of PSR including 112 incident cases have been observed. This larger group has enabled more detailed analyses on prevalence, incidence, familial clustering, and haematological risk factors. The results of these analyses are now presented.

Patients and methods
The patients attended the Sickle Cell Clinic at the University of the West Indies. All SS and SC patients examined between August 1970 and October 1988 were included in this study, although children in a cohort study from birth were excluded. Ocular examination included visual acuities, fundal examination through dilated pupils, and fluorescein angiography where indicated. Patients were classified as having PSR regardless of whether or not the lesions were perfused when examined.

The patient's age and PSR status for the prevalence analysis was defined at the first eye examination. As patients with proliferative disease were reviewed more frequently, using age at last visit would have exaggerated the prevalence of PSR. Therefore all cases of PSR that developed during the study were excluded from the prevalence analysis. The exact age at onset of PSR was rarely available, and for the incidence analysis it was assumed that PSR developed at the mid point between the last negative and the first positive examination. Patients were divided into the following age groups for analysis: 0–9 years, 10–14, 15–19, 20–24, 25–29, 30–39, and 40 years and above.

The diagnosis of SS and SC disease was based on standard criteria. Haemoglobin, red cell indices, and platelets were measured in electronic counters (Coulter ZBI6 or S plus 4, Coulter Electronics), fetal haemoglobin (HbF) by an alkali denaturation method, bilirubin by the method of Lathe and Ruthven, and serum iron by the method of Beale et al. Mean cell haemoglobin concentration (MCHC) was derived from the spin microhaematocrit. α Thalassaemia status was determined by restriction endonuclease analysis of DNA obtained from the buffy coat.3

For comparison of haematological indices in patients with and without PSR at their last examination, levels were derived from the mean of multiple steady state observations. This study was confined to a subset of 1181 patients (740 SS, 441 SC) for whom complete haematological data were available. Indices with skewed distributions (HbF, bilirubin) were logarithmically transformed (log10 [HbF + 1], log10 bilirubin) to give more normal distributions prior to statistical analysis. Simple comparison of the distribution of haematological indices between patients with and without PSR can be misleading, as certain indices are interdependent and age related, so multiple logistic regression analysis' was also used, separately for both sexes and genotypes. This approach assessed the effect of a factor assuming all other factors in the model were fixed and allowed for age related trends. Haematological indices that appeared to influence the simple comparison by Student's t test were included in the logistic regression. Comparison of proportions across age groups was performed by the Mantel-Haenszel test, and a log linear regression model was used to analyse age related trends in incidence.

Results
A total of 786 SS and 533 SC patients were examined, representing 29% of all SS patients and 55% of all SC patients registered at the clinic.
The excess of females examined (58% SS, 54% SC) reflects the attendance pattern of the clinic. Proportionately more of the SC patients were examined because of the known higher prevalence of PSR in SC disease and because they comprised a greater proportion of the adult clinic.

**Prevalence**

The prevalence of PSR at the initial eye examination by age group in SS and SC disease is shown in Tables I and II respectively. Proliferative retinopathy was more common in SC than SS disease (<0.001) and increased with age in both genotypes. In SS disease PSR was first observed in the 15–19 years age group, and the prevalence increased gradually to peak in patients aged 30 years and over. The apparent fall in frequency of PSR in male SS patients from the 30–39 year age group to the 40 years and over age group was not significant. In SC disease PSR was first seen in the 10–14-year age group, and the prevalence of PSR increased steeply with age, reaching maximum frequencies in SC males in the 25–29-year age group and in SC females aged 40 and above. As to the effect of sex, males with SC disease were significantly more likely to have PSR than females (p<0.001), but a similar trend in SS disease was not significant (p=0.14). Symptoms attributable to PSR prior to first eye examination occurred in 10% SC patients and 1% SS patients, but symptomatic referral did not significantly affect the prevalence data.

**Incidence**

The development of PSR for the first time was observed in 41 patients with SS disease (Table III) and in 71 patients with SC disease (Table IV). In SS disease incident cases were first observed in the 15–19 year age group, and the incidence rate appeared to increase to those aged 25 and over, though this trend was not significant. In SC disease incident cases first occurred in the 10–14 year age group, and incidence rates peaked at age 20–24 in males and at age 25–29 in females. However, neither the age related trend nor the apparent sex difference in SC disease reached statistical significance.

**Bilaterality**

The frequency of bilateral involvement was examined, since this may distinguish the role of systemic factors (favouring bilateral involvement) from local factors (favouring unilateral involvement). Bilateral PSR occurred in 44/90 (49%) of patients with SS disease and in 170/243 (70%) of patients with SC disease, being significantly more common (p<0.001) in SC disease. In both genotypes bilateral PSR was more common than would be expected if PSR had developed independently in the second eye (p<0.001).

The development of bilateral PSR was prospectively observed in the study in 59 patients (18 SS, 41 SC). The interval between onset in the first and second eye was examined, since simultaneous onset might reflect some acute haemato logical change or clinical event relevant to the pathogenesis of PSR. In only three patients (1 SS, 2 SC) was PSR observed to develop in both eyes within one year, and in none did this occur simultaneously. In 14 patients (2 SS, 12 SC) the interval between onset in the first and second eye was at least one year. In the remaining 42 patients the intervals between visits were too irregular to provide useful data.

**Familial Clustering**

The clustering of PSR affected patients within families was examined, since this might identify other genetic factors relevant to the development of PSR. In SS disease there were insufficient numbers of PSR affected patients to conduct this.
TABLE V

<table>
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<th>Number of siblings examined</th>
<th>Sibling with PSR</th>
<th>Observed</th>
<th>Expected</th>
<th>O-E</th>
<th>O-E</th>
<th>Sibling without PSR</th>
<th>Observed</th>
<th>Expected</th>
<th>O-E</th>
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HAEMATOLOGICAL VARIABLES

Comparison of haematological variables in patients with and without PSR showed significant associations of PSR with high Hb in SS males and a low HbF in SS patients of both sexes (Table VI). In SC disease there were associations with high Hb and high MCHC in males and with high MCV and low HbF in both sexes (Table VII).

Further examination of these relationships by multiple logistic regression analysis, which allows for effects of age and for interrelationships between haematological variables, showed that in SS disease both the high Hb for males and low HbF in both sexes remained significant (Table VIII). In SC disease all the significant relationships in males apparent on simple regression were no longer significant, though a low HbF and a high MCV remained significant in females. The effects of these factors can be illustrated by using the fitted logistic regression models to predict the probability of PSR. For example, a 35-year-old SS male with Hb 7 g/dl and HbF 6% would have a 6% chance of having PSR, whereas with Hb 10 g/dl and HbF 1.5% this would rise to 53%. A 35-year-old SC female with Hb 12 g/dl, mean corpuscular haemoglobin concentration (MCHC) 35 g/dl, mean cell volume (MCV) 75 fl, and HbF 2% would have a 29% chance of having PSR, though changing the latter two indices to 90 fl and 0.5% would increase this chance to 92%.

The α globin gene number was known in 351/786 (45%) SS patients. PSR was observed at the first visit in 2/42 (5%) SS patients homozygous for α thalassaemia, 8/116 (7%) SS patients heterozygous for α thalassaemia, and 15/192 (8%) SS patients with a normal α globin gene complement, with no significant difference between groups. Among SS patients with a normal α globin gene complement PSR occurred in 10/82 (12%) males compared with 5/110 (5%) females, though this difference just failed to reach statistical significance (z=1.95, p=0.051). No α globin gene number data were available in SC disease.

Discussion

The observation that proliferative sickle retinopathy occurs at such high frequency in a generally benign genotype of sickle cell disease is of considerable interest. Most patients with sickle cell haemoglobin C disease run a mild clinical...
course with near normal haematology and infrequent vaso-occlusive episodes. Patients with homozygous sickle cell disease generally have frequent vaso-occlusion and a severe clinical course, yet proliferative sickle retinopathy is significantly less frequent. This enigma suggests that the two genotypes may be useful models for understanding the risk factors for PSR.

One hypothesis attempting to reconcile this unexpected observation envisaged three models with different vaso-occlusive tendencies. Patients with low vaso-occlusive indices would be unlikely to develop retinal ischaemia and therefore have no stimulus to develop PSR. Patients with moderate vaso-occlusive indices would develop peripheral retinal closure and proceed to the development of preproliferative or proliferative disease. Patients with high vaso-occlusive indices would develop extensive peripheral retinal vascular closure, with a stimulus to PSR formation, but would proceed to occlude preproliferative arteriovenous anastomoses or nascent PSR. It was proposed that SC disease represented the intermediate model, with sufficient vaso-occlusion to produce retinal ischaemia but insufficient to occlude the lesions of developing PSR. An analogous situation would be the persistence of splenomegaly in SS patients with high levels of HbF, representing the survival of a capillary bed not damaged by vaso-occlusion and hence an expression of pathological mildness. This hypothesis implies that the risk factors for retinal vaso-occlusion may differ from those for PSR. Only limited data are available on the risk factors for peripheral retinal vaso-occlusion. Observations in the Jamaican cohort study confirmed that closure generally increased progressively with age, was more common in males than females at an early age, and was not significantly more frequent in SS disease. Analysis of haematological risk factors in SS disease showed that retinal closure was significantly associated with low total haemoglobin, low fetal haemoglobin, a high reticulocyte count, and high counts of irreversibly sickled cells. The effect of total haemoglobin disappeared on multiple logistic regression, suggesting that its effect was entirely secondary to its relationship with HbF, but a later matched pairs analysis of patients with minimal or complete closure showed an independent effect of haemoglobin level. In SS disease, therefore, closure was associated with low Hb, low HbF, high reticulocyte, and high ISC counts whereas PSR was associated with a high Hb (males only) and low HbF. In SC disease closure was associated with a high reticulocyte count, though this was not apparent on a later matched pairs analysis, which showed effects of high MCV and low platelet counts. By contrast haematological risk factors for PSR in SC disease were apparent only in females and included a high MCV and low HbF.

The results of analysis for haematological risk factors for PSR in the present study are broadly similar to previous observations. In SS disease observations in males were identical to those of Hayes et al., whereas in females a previously noted and unexplained contribution of low serum iron was lost, and an effect of low HbF has emerged in the present study. In SC disease the results on simple regression in males were again identical to those of an earlier study, though logistic regression in the currently available larger data set shows that none of the apparent contributions of high Hb, low HbF, high MCHC, and high MCV remain significant. In females only a low HbF was significant in the earlier study, but a high MCV also appears to be a risk factor.

These haematological data may contribute to the understanding of the mechanisms of PSR formation, since different indices are believed to affect flow in different sized vessels. Thus flow in large vessels is affected predominantly by viscosity (haemoglobin, haematocrit) or number of particles (red cell count), whereas flow in the capillary bed is affected by the characteristics of the individual red cells such as membrane deformability and intracellular viscosity. In sickle cell disease the latter is particularly affected not only by the MCHC but also by HbF, which affects the intracellular polymerisation of HbS. The haematological data are therefore more consistent with a capillary site of obstruction. Furthermore the lack of demonstrable differences in blood viscosity between groups with and without PSR in SS and SC disease is less consistent with a role of larger vessel flow. The elegant angiographic studies of Galinos et al. showing non-perfusion at or near Y shaped branches of the precapillary arterioles would be consistent with either interpretation.

Other observations in the present study have contributed to the understanding of PSR. Both the prevalence and incidence data show that PSR is rare in the first decade, suggesting that extensive retinal damage must have accumulated before the PSR develops. The high frequency of bilateralism suggest that systemic factors contribute to the development of PSR, though the lack of simultaneous development argues against a role of acute haematological change or a severe clinical event. The analysis of familial clustering was limited by the relatively small numbers available, and, though showing a trend in that there was a slight excess of families in which all or none of the siblings had PSR, the differences were far short of significance, contrary to our clinical impression.

Do the available data fit the hypothesis? The first model of low vaso-occlusive indices is self-evident, since PSR does not develop in the absence of retinal ischaemia. However, this is likely to be a small group, since retinal closure occurred in approximately 90% children by 12 years of age, the factors protecting against closure being either homozygous α thalassaemia or very high levels of HbF. The middle group...
with moderate vaso-occlusive indices should include the majority of SC patients and a smaller proportion of those with SS disease. The group with high vaso-occlusive indices should contain predominantly patients with SS disease. This genotype is already recognised to be associated with increased automaticity of established PSR, but, if the hypothesis is correct, patients with the greatest vaso-occlusion should be experiencing preproliferative changes and not proceeding to the development of PSR.

Testing the hypothesis requires an accurate indicator of vaso-occlusive tendency. Using HbF level as such an indicator, we could not demonstrate a lower frequency of PSR among those with the lowest HbF levels. Although inconsistent with the hypothesis, this observation casts doubt on the validity of HbF as an indicator of vaso-occlusion and does not necessarily refute the hypothesis.

We thank other ophthalmologists associated with the Jamaican ophthalmic studies, especially Mr Patrick Condon, Robert Acheson, and Brendan Moriarty, who have all contributed to the data. We thank Dr Douglas Higgs for the studies on o globin gene number.