Ocular lesions in sickle cell disease patients from Bahia, Brazil

Lesões oculares em pacientes com doença falciforme da Bahia, Brasil

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ABSTRACT

Objective: The present study aims to describe ocular alterations in sickle cell disease patients in Bahia, a Northeast state, with the highest prevalence of the disease in Brazil.

Methods: We carried out a cross-sectional study in a group of 146 (292 eyes) sickle cell disease patients (90 HBSS and 56 HBSC). Ophthalmologic examination including indirect binocular ophthalmoscopy was performed. Examination was completed by fluorescein angiography to detect retinal lesions.

Results: The most frequent ocular lesions identified were “vascular tortuosity” and “black sunburst”. Proliferative retinopathy was found in 22 (12.2%) eyes of HBSS patients and 25 (22.3%) eyes of HBSC patients (OR=2.06; CI95%: 1.5-4.06, p=0.022). Its frequency was higher among HBSS patients aged 20-39 years, while in HBSC patients, it peaked after 40 years (35.7% and 42.8%) and dropped sharply afterwards.

Conclusion: Proliferative retinopathy was described as early as 10 years of age in both patients groups. Proliferative sickle retinopathy can result in blindness and the knowledge of the most prevalent ocular alterations and age risk will be important to establish a protocol of ophthalmologic follow-up, in order to prevent a severe visual loss and increase patient’s life quality.

Keywords: Anemia, sickle cell; Hemoglobin SC disease; Eye injuries; Retinal diseases

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INTRODUCTION

The sickle cell hemoglobin (HbS) is characterized by a single nucleotide change (GAG→GTG) in the sixth codon of the β-globin gene, which leads to a valine instead of glutamine in the sixth position of the β-globin chain. (1)

Because of its decreased solubility under deoxygenated conditions, hemoglobin S polymerizes in red blood cells, forming a highly ordered fiber aggregates that distort the cells' shape into an elongated forms. The polymer makes reversible and irreversible changes in sickle cell erythrocytes and is responsible for the sickle cell phenotype, including the homozygous state of HBSS and heterozygous combinations such as HBSC and HBS-β thalassemia that have a less severe manifestations than those found in sickle cell anaemia patients. The sickle cell trait (HBAS) is an exception and is not considered a sickle cell disease since the carriers do not have any clinical manifestations. (1)

The overall frequency of HbS is high worldwide, and in Brazil its distribution is heterogeneous. In Northeastern Brazil, mainly in Bahia, the population is a tri-racial mixture of Europeans (mostly Portuguese), Africans and indigenous. (2) The frequency of HbS in the state of Bahia is the highest in Brazil, varying from 4.5 to 14.7% in several population groups studied. (3)

The sickle cell disease is characterized by a variety of clinical abnormalities frequently linked to hemolytic anaemia and vaso-occlusive processes, which are responsible for causing pain and other clinical features such as retinopathy. (1,4)

Ocular lesions results from stasis and occlusion of the small eye vessels by sickled erythrocytes. Transient dark red spots, representing plugs of sickled erythrocytes within superficial capillaries can be seen on the surface of the optic disc and conjunctiva. Vaso-occlusive disease of the retina can be responsible for nonproliferative and proliferative ocular changes. The nonproliferative lesions consist of ocular lesions such as a “salmon patches”, vessel tortuosity, “black sunbursts”, iridescent spots and angioid streaks that characterize hemorrhagic, infarctive, and resolving lesions of sickle retinopathy. (5) Proliferative ocular lesions can result in partial or total loss of vision and are classified into different clinical stages. Stage I is characterized by arteriolar obstruction; stage II, by arteriovenous anastomoses; stage III, by neovascularization; stage IV, by vitreous hemorrhage and stage V, by retinal detachment. (5)

Despite the high prevalence of sickle cell disease in Bahia and the high frequency of ocular changes in sickle cell disease patients, there have been no studies of these alterations with sickle cell retinopathy in Bahia, a state with African ethnic characteristics. (2) We therefore consider important to characterize the ocular lesions found among sickle cell patients from northeast Brazil.

METHODS

This study was approved by the Oswaldo Cruz (Research Foundation's Human Research Ethics Committee), and informed consent was obtained in accordance with ethical principles and the Helsinki Declaration of 1975, and by the Brazilian resolution 196/96, of the Ministry of Health, Law 6,638/79 and Normative Resolution 04/97. Ophthalmologic examinations were carried out, and peripheral blood samples were collected only after signed informed consent was obtained.

The cross-sectional study involved a group of 146 (292 eyes) sickle cell patients (90 HBSS and 56 HBSC) from the State of Bahia in northeast Brazil and was carried out between July, 2002, and April, 2006.

The patients were selected among those attending the hematologic ambulatory at the Bahia Foundation of Hematology and Hemotherapy (HEMOBA), a reference center attending to sickle cell disease patients who are seen in routinely visits. Patients were then sent to the Brazilian Institute of Ophthalmology and Blindness Prevention (IBOPC) for an ocular examination, including fundus biomicroscopy, indirect binocular ophthalmoscopy and fluorescein angiography when the retinopathy could not be adequately characterized solely by means of a fundoscopic examination. We observed the following in the fundoscopic examination: vascular tortuosity, alterations of papila, “salmon patches”, angiod streaks and iridescent spots. The pathologic classification of fundoscopic alterations as proliferative was based on Goldberg’s five-stage groups; stage I - peripheral arteriolar occlusions, stage II - peripheral arteriovenous anastomoses, stage III - preretinal neovascularization, stage IV - vitreous hemorrhage and stage V - retinal detachment.

Patients presenting with proliferative retinopathy were separated by age and classified as severe when displaying stages III to V when retinal neovascularization was seen. The hemoglobin pattern was confirmed at the Pathology and Molecular Biology Laboratory of Fiocruz and UFBA using High Performance Liquid Chromatography (HPLC-VARIANT I/BIO-RAD, CA, USA).

EpiInfo software for Windows, version 3.3.2 was used...
to store and analyze data. The Chi-squared Pearson’s test or the Fisher’s exact test was used when necessary to compare both groups of hemoglobinopathy carriers. A p-value of less than 0.05 was considered statistically significant.

**Results**

The number of sickle cell disease patients enrolled in the study was 146 (292 eyes): 90 (61.6%) with sickle cell anemia or HBSS and 56 (38.4%) with HBSC disease. The majority of these patients (58.9%) were from Salvador, and the others came from other cities in the state. Overall, the patients had an average age of 26.7 (+11.6) years. The HBSS patients group had an average age of 26.7 (+10.1) years and the HBSC group had an average age of 26.9 (+13.9) years. Eighty-four (57.5%) were women, and 62 (42.5%) were men. Among the HBSS patient group, there were 39 (43.3%) men and 51 (56.7%) women, while among the HBSC group, there were 23 (41.1%) men and 33 (58.9%) women. Statistically non-significant differences were found when comparing the age (p=0.625) of the HBSS and HBSC patients. There were no significant differences in gender (p=0.923) when comparing both patient groups studied.

Two HBSC disease patients had an intraocular pressure ≥ 20mmHg; however, no overall change in intraocular pressure was observed among the different patient groups. Visual acuity was 20/20 or 20/25 for the best eye in 133 patients, corresponding to 91.1% of the total number of cases.

Age-related ocular lesions, including nonproliferative and proliferative lesions, were very frequent among both groups of patients (HBSS and HBSC). Table 1 shows the ocular lesion distribution across six age groups. The HBSS group had more ocular changes in the age range of 20-39. In the HBSC group of patients these changes became more frequent in the age range of 40 to more than 50 years.

Vascular tortuosity and “black sunbursts” were the most frequent fundoscopic ocular lesions found in both sickle cell disease groups (HBSS and HBSC). Table 2 shows that the vascular tortuosity percentage was similar in both sickle cell disease patients’ groups. The “black sunburst” was more frequent in the HBSC disease group. The frequency of “salmon patches” and angiod streaks, however, was low in both groups, and the disc-shaped signal and iridescent spots were only found in the HBSS group.

In both the HBSS and HBSC disease groups, there were patients with proliferative ocular lesions; however, the HBSC group had more proliferative ocular lesions than the HBSS group.

The HBSS group had more severe ocular disease in the age range of 30-39 than the HBSC group. Table 3 shows the distribution of proliferative ocular lesions across six age groups related to stages I to V and III to V. The HBSS group had the highest frequency of proliferative changes, mainly in the age range of 20-39. The HBSC patient group had more ocular changes in the age range of 40-49. Table 4 shows that the HBSC group had more cases of proliferative ocular lesions per eye than the HBSS group. Figure 1 shows fluorescein angiography of peripheral arteriolar occlusions, peripheral arteriovenous anastomoses, preretinal neovascularization and extensive capillary non-perfusion from a 31-year-old woman with sickle cell anemia (HBSS).

**Discussion**

Several ocular changes were observed in the studied patient groups; however, no change in visual acuity was found, as previously reported. (5-10)

With regard to the fundoscopic lesions, vascular tortuosity and “black sunbursts” were the most frequent changes identified, corroborating previous Brazilian studies. (7-10) The overall frequency of vascular tortuosity was similar in both patient groups (HBSS and HBSC), as previously described. (10,11) The “black sunburst” lesion was seen in both groups, although its frequency in HBSC patients was much lower than that described in other studies. (10,12,13) The low frequency of other ocular lesions such as the disc sign and iridescent spots could be attributed to their transient status and to the fact that they do not produce clinically visual impairment that requires serial follow-up testing. (14)

Proliferative retinopathies were significantly more frequent in the HBSS group than in HBSS group, as described previously. (1,7,15-17) When the HBSC patients were stratified according to age, however, we observed two peaks of ocular changes, one between 10 and 19 years of age and another between 40 and 49 years of age that subsequently dropped sharply. This finding is in contrast to previous studies. (15,18,19) Indeed, Fox et al. (18) studied Jamaican patients and observed that ocular proliferative lesions increased with age in both genotypes. Interestingly, the HBSS patients had more severe proliferative changes (stages III to V) between the ages of 10 to 19 years and 30 to 39 years when compared to the HBSC group, characterizing an elevated frequency when compared with others studies. (20,21) On the basis of
our results, additional studies are warranted to determine if these findings are related to an increase of vascular-occlusive crisis among these specific ages or with the presence of others biomarkers for assessing relative risk in these group of SCA patients.

This is the first study of ocular lesions among HBSS and HBSC patients from Bahia. The lower percentage of severe proliferative retinopathy found among HBSS patients when compared to HBSC patients could be attributed to auto infarction or a spontaneous regression of proliferative retinopathies, a phenomenon described in sickle cell disease. (23) A milder disease phenotype was previously described among these patients, (22,23) protecting against the early vessel occlusion described in HBSS and contributing to proliferative lesion development among the older patients. In addition, the age-related variation in proliferative lesions observed in HBSC patients could be related to more severe anemia, resulting in a decrease in blood viscosity. The results presented here show that sickle cell disease patients from Bahia could have specific risk factors for retinal vessel changes other that those described before. (22,23)

**Conclusion**

In conclusion, the ocular lesions described here could help to define the clinical and ophthalmologic protocols of HBSS and HBSC patient’s follow-up in specific ages. The presence of proliferative ocular alterations are a well documented cause of blindness and the knowledge of the most prevalent ocular alterations and age risk of these in HBSS and HBSC patients will be important to establish ophthalmologic follow-up protocols, preventing a severe visual loss. Further studies, including the association between ocular lesions and hematological, environmental and genetic factors - in a larger number of HBSS and HBSC patients in Brazil should contribute to improving patient quality of life, bringing new knowledge of ophthalmologic alterations among these patients.

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**References**


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