Sickle cell retinopathy: A literature review

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**SUMMARY**

Hemoglobinopathies are a group of hereditary diseases that cause quantitative or qualitative changes in the shape, function or synthesis of hemoglobin. One of the most common is sickle cell anemia, which, due to sickling of erythrocytes, causes vaso-occlusive phenomena. Among the possible ocular manifestations, the most representative is retinopathy, which can lead to blindness if left untreated. Therefore, periodic ophthalmologic monitoring of these patients is important for early diagnosis and adequate therapeutic management, which can be done locally by treating the lesions in the eyes, or systemically.

**Keywords:** retinal diseases, anemia, sickle cell, review, hemoglobinopathies.

**INTRODUCTION**

Sickle cell disease is the most frequent and disabling chronic hemolytic anemia in our country. Sickle cell anemia is characterized by the production of abnormal hemoglobins that deform and stiffen red blood cells, causing increased blood viscosity and microcirculation occlusion to varying degrees.1 These hemoglobins are insoluble at low oxygen concentrations, and tend to crystallize.2 Sickle hemoglobin (HbS) is characterized by a mutation in the β-globin gene involving a single nucleotide (GAG → GTG) which replaces glutamine with valine in sixth amino acid position.3,4 The most common genotypes are homozygous (SS), heterozygous or sickle cell trait (SA), hemoglobin C trait (SC), hemoglobin D trait (SD), and thalassemia genotype (S-Thal).5-8

The World Health Organization (WHO) estimates that more than 5% of the world population has some type of hemoglobinopathy. The estimate of new cases in Brazil is 700 to 1,000 per year, with a prevalence of over 2 million carriers of the HbS gene.9 Sickle cell retinopathy, the subject of this study, develops in up to 42% of sickle cell individuals in the second decade of life.10

The systemic manifestations of sickle cell disease may be neurological, ophthalmologic, cardiac, pulmonary, gastrointestinal/hepatobiliary, renal/genitourinary, splenic, muscular/skeletal, and growth and developmental disorders.11 They are more severe in homozygotes for cell disease (SS) than in heterozygotes with sickle hemoglobin C (SC), and yet visual loss due to proliferative retinopathy is more common in the latter.6

Sickle cell retinopathy is not frequently reported in the literature, and studies in this regard are very old. This was one of the motivations for our study, which seeks to gather the information known and thereby clarify the progression of the disease, its diagnosis and treatment.

**METHOD**

We searched the PubMed (US National Library of Medicine – National Institutes of Health) database using the following keywords: “retinopathy,” “sickle cell,” “sickle cell anemia”.

**OCULAR MANIFESTATIONS**

Ocular manifestations of sickle cell anemia include orbital, conjunctival, uveal, papillary, and especially retinal changes.5,7,12 Retinal changes characterize sickle cell retinopathy, which may be non-proliferative or proliferative and is divided into five stages.1,10,12,13

Among the forms of sickle cell anemia, SS patients present a more severe systemic clinical picture than those with type SC. On the other hand, occlusive ocular effects are more predominant in SC patients, who present only moderate anemia and higher blood viscosity.5 These vaso-occlusions occur primarily in younger people, and are first
Sickle cell retinopathy develops in up to 42% of sickle cell individuals in the second decade of life. Vascular tortuosity is the most common finding (Figure 1), reported by the authors in about 30-50% of cases. Cury et al. found a prevalence of 19.6%, a result that may be justified by the fact that the study was conducted in children only. In addition, about 10-20% of patients will develop proliferative retinopathy, mainly in the fourth and fifth decades of life.

**Diagnosis**
In the early stages, the disease is asymptomatic, and meticulous ophthalmologic monitoring should be performed. Diagnosis is made by retinography and fluorescein angiography in cases with fundoscopic alterations, as well as measurement of visual acuity and intraocular pressure.

**Treatment**
Treatment is performed in different ways, including diathermy, cryotherapy and argon or xenon photocoagulation.
The latter is widely used to treat the typical stage III lesion (sea fan) of proliferative retinopathy. However, patients in stages I and II are not treated, since the treatment of ischemic lesions in these stages does not prevent the formation of sea fan (Figure 3), and most patients do not develop the complication. There is also surgical treatment, which is indicated for complications of proliferative retinopathy, such as retinal detachment and vitreous hemorrhage.

According to Clarkson, prophylactic photocoagulation may play a role in the treatment of selected patients with SC proliferative sickle cell retinopathy, but none of the studies reported to date have demonstrated that this treatment improves long-term visual outcome compared to natural progression, as documented in the present study. The similar visual results in the eyes analyzed in our study during the natural progression of the disease compared to those treated with photocoagulation should not be unexpected, because there is a greater predilection for spontaneous involution or neovascular tissue infarction in SC disease as opposed to neovascularization that develops in other vascular diseases of the retina. Clarkson suggests that a multicenter controlled clinical trial designed to study eyes at greater risk should be considered. There is, however, no clear definition of the risk factors leading to these advanced stages, and the value of treatment is uncertain.

In addition, since ocular disorders of retinopathy result from a systemic pathological process, prevention can be done with appropriate treatment of anemia using several emerging approaches, such as: increase in fetal hemoglobin using hydroxyurea, omega-3 and erythropoietin, 2-deoxy-5-azacytidine; erythrocyte hydration (clotrimazole, magnesium pidolate); anti-inflammatory and anti-adhesive drugs (anti-adhesive antibodies, anti-integrin antibodies, anti-Willebrand factor, sulfasalazine, statins); antioxidant therapy (glutamine, deferiprone); antithrombotic agents (heparin, ticlopidine, warfarin); vasodilatation (nitric oxide, arginine, Flocor); decrease in hemoglobin S by transfusion and apheresis; transplantation of hematopoietic cells and gene therapy.

CONCLUSION
Considering that sickle cell retinopathy is a complication that causes 42% of blindness in the affected patients, we point out the importance of new studies on the subject, since there is a gap especially in randomized clinical trials. The importance of our review is to draw attention to the need for periodic ophthalmologic monitoring in patients with anemia since childhood, aiming at prevention, diagnosis and early treatment of the disease.

RESUMO
Retinopatia da doença falciforme: revisão da literatura
As hemoglobinopatias são um grupo de doenças hereditárias que causam alterações quantitativas ou qualitativas no formato, na função ou na síntese de hemoglobinas. Uma das mais comuns é a anemia falciforme, cuja patogenia é a foicização das hemácias, causando fenômenos vaso-oclusivos. Dentre as manifestações oculares possíveis, a mais representativa é a retinopatia, que pode levar à cegueira caso não seja tratada. Por isso, é importante que haja o acompanhamento oftalmológico periô-

FIGURE 3 A. “Sea fan.” B. Angiographic appearance of the “sea fan.”
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dico desses pacientes, a fim de obter diagnóstico precoce e abordagem terapêutica adequada. Esta última pode ser de maneira direta, com tratamento das lesões oculares, ou de forma sistêmica.

Palavras-chave: doenças retinianas, anemia falciforme, revisão, hemoglobinopatias.

REFERENCES


