Sickle cell retinopathy: diagnosis and treatment

Retinopatia falciforme: diagnóstico e tratamento

M ARIA T E RESA B RIZZI C HI ZOTTI B ONANOMI 1, M ARC ELO M ENDES L AVEZZO 2

ABSTRACT

Hemoglobinopathies are a group of inherited disorders characterized by quantitative or qualitative malformations of hemoglobin (Hb). Some of these diseases present vaso-occlusive phenomena that are responsible for high morbidity in clinical and/or ophthalmologic terms. Diagnosis of hemoglobinopathies is performed exclusively through hemoglobin electrophoresis. From the ophthalmologic perspective, the most important representative of this group of diseases is sickle cell retinopathy, which presents a wide spectrum of fundus manifestations and may even lead to irreversible vision loss if not properly diagnosed and treated. The aim of this review is to present the classification of sickle cell retinopathy and to describe current management and future perspectives for its treatment, taking into consideration the clinical management of these patients.

Keywords: Retinal diseases/diagnosis; Retinal diseases/therapy; Hemoglobin, sickle; Sickle cell trait; Hemoglobin SC disease; Vitreoretinal surgery

INTRODUCTION

The hemoglobinopathies are a group of hereditary disorders characterized by abnormal hemoglobin (Hb) formation. The changes in hemoglobin caused by these disorders may be quantitative or qualitative. Qualitative changes occur when the number of hemoglobin chains present is normal, but their chemical composition is changed. Sickle hemoglobinopathies are part of this group.

The most common genotypes associated with hemoglobinopathies depend on the population studied. In the Brazilian population, the abnormal hemoglobins S and C are the most frequent. Using high-performance liquid chromatography analysis, the abnormal hemoglobins most commonly found is HbS, in one study, the most common abnormal genotype was HbAS, which occurred in 9.11% of the samples. Other genotypes represented included a genotype associated with beta-thalassemia in 5.50% of the samples, HbAC in 2.47%, HbSS in 0.76%, and HbSC in 0.39% (1). If the numbers of individuals with HbS and HbC undergo a conformational change under conditions of hypoxia, acidosis, oxidative stress and infection, changing the erythrocytes into rigid sickle-shaped (sickled) cells called drepanocytes. Drepanocytes are less flexible and more prone to hemolysis than normal red blood cells, and their formation leads to various clinical manifestations of anemia and vaso-occlusive events.

The aim of this review is to describe the vitreoretinal findings related to sickle cell disease and to discuss the complications of the disease that can affect vision, as well as their treatment.

EPIDEMIOLOGY AND PATHOGENESIS

The pathophysiology of sickle cell disease is not limited to abnormalities of erythrocytes; its clinical manifestations are related to processes and complex metabolic pathways that include endothelial activation, inflammation, nitric oxide bioavailability, oxidative stress and regulation of the adhesiveness of several types of blood cells (2).

The Hb molecule is composed of two alpha and two beta polypeptide chains. In hemoglobin S and C, an amino acid substitution occurs at position 6 of the beta chain. In HbS, glutamic acid is replaced by valine, while in HbC it is replaced by lysine. The diagnosis of hemoglobinopathies is performed exclusively through hemoglobin electrophoresis (3). The sickling test cannot be used for this purpose because it is nonspecific (4).

The sickling process is influenced by infections, dehydration and acidosis. The blood flow in terminal retinal arterioles may be decreased by the aggregation of drepanocytes, which form small plugs. Under these conditions, the available Hb is deoxygenated and the oxygen demand of the tissue is not supplied, resulting in tissue acidosis. This causes greater sickling, increased blood viscosity and decreased blood flow. This sequence of events has been called the “vicious cycle of erythrostasis”.

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1 Physician, School of Medicine, Universidade de São Paulo - São Paulo (SP), Brazil.

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Correspondence address: Marcelo Mendes Lavezzo. Rua Raul Pompeia, 229 - Apt. 102, São Paulo (SP) - 05025-010 - Brazil - E-mail: mmlavezzo@gmail.com

The sickle cell hemoglobinopathy with the greatest number of clinical manifestations is the SS type, which leads to hemolysis and sickling crises. Patients of the SC genotype, who present rare systemic changes, are usually those with the most severe retinal manifestations.

The frequency of retinopathy is greatest in adulthood, but retinopathy has also been described in children. For patients bearing the SC genotype, the risk of developing proliferative sickle-cell retinopathy (PSR) is highest between the ages of 15-24 for males and 25-39 for females; for those with the SS genotype, it is higher between the ages of 25-39 for both sexes.

Individuals with sickle-cell trait comprise 8% of the African-Caribbean and over 10% of the Brazilian population. Although these individuals often show no relevant clinical manifestations, they may present various retinal changes such as hemorrhages, exudates, angioid streaks, acute chorioretinal infarction, choroidal lesions, vitreous hemorrhage, retinal vascular abnormalities such as tortuous/dilated retinal veins, microaneurysms, central retinal artery occlusion and retinal proliferation.

**DIAGNOSIS**

Because the early stages of sickle-cell disease often remain undetected, a meticulous eye examination is necessary. This assessment should be periodic and should include measurement of visual acuity, intraocular pressure and evaluation of the anterior/posterior segment structures by fluorescein angiography.

Ocular findings are represented by orbital, conjunctival, uveal, papillary and retinal alterations. The most relevant anterior segment abnormalities are the conjunctival sign and iris atrophy, both of which are diagnostic indicators of sickle-cell disease, and hyphema, for which there exists a particular therapeutic approach in patients.

In this review, we will focus on various aspects of sickle cell retinopathy and its complications.

**RETINAL ALTERATIONS**

The retinal manifestations of sickle-cell disease can be non-proliferative (yielding diagnostic value) or proliferative; in the latter case, there is a real risk of ocular morbidity. Both non-proliferative and proliferative manifestations are caused by erythroblastosis secondary to sickling.

**Non-proliferative retinopathy**

Venous tortuosity

Vascular tortuosity is a sign described in anemia; its frequency in homozygous sickle-cell disorder reaches 50% (22,23). However, vascular tortuosity is a nonspecific sign that is common to several diseases associated with blood hyperviscosity.

Hemorrhage, iridescent bodies and black sunburst

When an arteriole of intermediate size is clogged (occluded) by a plug of sickled erythrocytes, hemorrhage may occur, presumably by ischemic necrosis of the vessel wall. This type of hemorrhage is round or oval-shaped, bright red and measures ¼-1 disc diameter. In days or weeks, a modification of the orange-red color occurs that contrasts with the dark color of the retinal pigment epithelium (RPE); the area of the hemorrhage then appears to be salmon-colored. This is known as salmon patch (24-29). Over time, hemoglobin degradation occurs. The hemorrhagic defect then appears as bright yellow dots at several levels of the sensory retina; these are known as iridescent bodies.

If the hemorrhage occurs in the outer retinal layers, it stimulates the proliferation of RPE, leading to the formation of dark, oval or round, ¼-2 disc-diameter chorioretinal lesions. These lesions, which are similar in appearance to choriotetic scars, are known as black sunbursts (Figure 1).

**Figure 1. Patient with sickle-cell trait. A and B) Black sunbursts in the temporal and nasal peripheries of the right eye, respectively. C) Angiographic aspect of the black sunburst (window defect due to the atrophy of the RPE at the edges of the lesion and blockage by RPE mobilization to the center of the lesion). D) Temporal periphery showing the boundary between the vascularized and non-vascularized zones of the retina.**

The hemorrhages are temporary, and those that occur in the posterior part of the eye are difficult to observe. Their late signs, such as iridescent bodies and black sunbursts, can be observed in approximately 25-40% of cases and are more frequent in SC hemoglobinopathy.

**Macular changes**

Sickle-cell maculopathy occurs as a result of chronic changes in the perifoveal capillary network. It may be present in 10-40% of patients (26-29). The SS genotype is the most affected.

Ischemic changes are chronic and insidious, and the patient may display no symptoms. There is usually some disparity between the angiographic findings and the visual symptoms. Normal visual acuity can coexist with a very enlarged foveal avascular zone, and the presence of ischemic regions would be detected only by central visual field testing (30). However, some patients display low visual acuity and a well-perfused macular capillary network. In these cases, an old ischemic lesion of the photoreceptors and recanalization of the affected capillaries may have occurred (28).

In addition to the presence of an enlarged foveal avascular zone showing perifoveal capillary loss and nerve fiber layer infarcts, another finding in sickle-cell maculopathy is the formation of microaneurysm-like dots and vascular loops. These changes occur primarily in the non-perfused areas and do not cause leaking of fluorescein (26,28).

The incidence of acute symptoms caused by the presence of an average-sized arteriole occlusion in the macula is low. When this occurs, a thin zone of the retina, the retinal depression sign, which is linked to a scotoma or to reduced visual acuity, is formed (31). In any case, the ischemic macular changes associated with sickle hemoglobinopathy are due to the occlusion of arteriolar circulation around the foveal avascular zone and branches that supply the temporal horizontal raphe (32). Those changes may or may not be associated with peripheral vascular occlusion and can occur in children (32).

Ischemic macular changes are uncommon in angiograms of patients with sickle hemoglobinopathy. With the introduction of optical coherence tomography (OCT), atrophy of the inner retinal...
layers could be demonstrated in patients with macular infarction(31); even without angiographic changes, this type of atrophy was demonstrated with spectral domain OCT (SD-OCT)(32). These findings are consistent with the atrophy of ganglion cells, of the inner nuclear layer and of Müller cells that has been demonstrated histologically in patients suffering from chronic ischemia(25). SD-OCT can document this atrophy even in patients with 20/20 vision (Figures 2 and 3). An initial study of patients of several genotypes showed a significantly thinner retina using SD-OCT in all macular sectors in approximately 50% of the eyes of patients of the SS (16 patients), SC (2 patients) and STahl (2 patients) genotypes compared with normal subjects of the same age group (20-60 years old)(33). Patients with sickle-cell disease may also have significantly reduced retinal sensitivity to microperimetry compared to those without focal thinning or to age- and visual acuity-matched controls(33).

Optic nerve sign

Vascular changes in the optic nerve are transient and consist of dark, small and red dilated capillaries that show occlusion in fluorescein angiography(22,42), without any impact on acuity(17).

Proliferative retinopathy

The involvement of vision in any stage of life of a patient with sickle-cell disease is reported to be 10-20%(43). Although PSR is the primary cause of visual acuity reduction in these patients(36), the risk of this event is unknown. In a 20-year longitudinal study in which 307 SS patients and 166 SC patients were recruited at 5 years of age, it was established that the incidence of PSR is 0.5 cases/year per 100 SS patients and 2.5 cases/year per 100 SC patients(44). The emergence and development of PSR are usually insidious, beginning in the first decade of life(45). The condition remains asymptomatic until complications such as vitreous hemorrhage (VH) or retinal detachment (RD) occur typically between 20-30 years of age(46). The significance of PSR as a cause of reduced visual acuity is confounded by the spontaneous regression of the neovascular complexes, which has been described in 20-60% of the cases(46) and often occurs approximately two years after the development of proliferative lesions.

The peripheral retina, especially the temporal portion, is highly susceptible to the effects of intravascular sickling. Because it exhibits vascular network architecture of the terminal type with very thin vascular diameter, it is predisposed to occlusion. The events that lead to the occurrence of PSR typically begin in this region. Due to the insufficient number of capillaries, the collateral circulation is not developed when a vascular segment is occluded. The sequence of events was described by Goldberg(47). The recently blocked vessel

Figure 2. A 23-year-old patient with sickle cell anemia (HbA1 - 0%, HbA2 - 2.7%, Fetal Hb - 25.4% and HbS - 71.9%) with macular atrophy with 20/20 visual acuity in both eyes. A and B) Retinography. C and D) Fluorescein angiography. E and F) Decrease of the central thickness with enlargement of the foveal depression demonstrated by spectral domain OCT. Courtesy of Dr. Aline Sokolowski da Silva.

Figure 3. A 52-year-old male patient with sickle-cell trait was hospitalized for 10 days in the Intensive Care Unit due to severe pneumonia and anemia and remained on mechanical ventilation and sedation. On day 11 of hospitalization, the patient presented a sudden decrease in the visual acuity in both eyes (20/80 and 20/100). A and B) Retinography showing perifoveal retinal pallor in both eyes and sparse cotton-wool exudates. C) Arteriolar branch near to the fovea showing “string of pearls” obstruction in its pathway. D and E) Diffused decrease of the thickness of the inner layers of the retina, visible in the temporal region of both eyes, as demonstrated by spectral domain OCT; there is some irregularity in the RPE level, however without compromising the outer layers of the retina. F and G) After blood transfusion, there was an improvement in visual acuity (20/20 and 20/25) but maintenance of the structural changes in the macula of both eyes. Courtesy of Dr. Carlos Alexandre Garcia.
can be occluded permanently and recanalized with or without the occurrence of salmon patch hemorrhage. The plugs of erythrocytes may leave the site in up to 12 hours, but remodeling of the vascular network takes months or years. If this remodeling process is efficient and forms new vascular arcades similar to normal ones, it eventually leads to the posterior formation of a peripheral capillary network (Figure 1D) without risk of sea fan development. If the remodeling is partial, an irregular vascular pattern that is more prone to neovascularization develops\(^{(48)}\). The irreversible vascular occlusion in the peripheral retina characterizes Stage I of the process. The occluded segment can display a “silver wire” appearance, but the ischemic phenomenon is more easily observed by fluorescein angiography (Figure 4). Following these events, there is a tortuous elongation of the arteriole end and dilation of capillaries that form arteriovenous communications in the boundary between the vascular and ischemic retinas, characterizing Stage II. The neovessels are formed from these anastomoses and are derived from the venous side of the circulation\(^{(48)}\). The presence of neovessels defines Stage III. Characteristically, the neovascular tuft occurs at the boundary between the non-vascularized and vascularized zones of the retinal periphery (Figures 4 and 5); the tuft is initially flat, with thin walls and located in the extreme periphery. The neovascular tuft is, therefore, difficult to diagnose ophthalmologically without a careful search, and angiography of the suspected sites can help in this process. The well-developed neovascular tissue exhibits a typical fan shape. This morphology was compared by Welch & Goldberg\(^{(23)}\) to that of the marine invertebrate *Gorgonia flabelum* or sea fan (SF). Subsequently, a fibroglial tissue membrane develops on the SF, obscuring the blood column (Figures 4-6). At the same time, the SF tends to rise into the vitreous and form anastomoses, which leads to the emergence of large fibrous bands in the retinal periphery. The degree of perfusion of these lesions can be demonstrated by fluorescein angiography\(^{(48)}\).

Because sickle-cell anemia is a thromboembolic disease, spontaneous regression of SF by auto-infarction may occur\(^{(49,50)}\) (Figure 5).

The progression is more intense in the 20-40 year age group. The regression occurs throughout life, and it is unusual for an SF to remain perfused in patients over 40 years old who bear the SS genotype\(^{(48,51)}\).

The term “proliferative sickle-cell retinopathy” is often used to characterize Stage III retinopathy. For patients with SS and SC hemoglobinopathies, Stage III retinopathy has a prevalence of 26% and 63%\(^{(52)}\), 14.6%, and 54.5%\(^{(53)}\), 2% and 24%\(^{(48)}\) and 14% and 43%\(^{(46)}\), respectively. In addition to being more common in the SC genotype, the retinopathy is also more severe in this genotype\(^{(39,40)}\). The tendency to spontaneous regression occurs in both groups, and the difference between them is controversial\(^{(8,44,46)}\). Within the SC genotype, PSR is more common when the blood levels of hemoglobin is above 12.5 g (73%) compared with lower levels of Hb (15%)\(^{(49)}\).

Although most SF are small, occupying generally less than one clock hour of the retinal periphery\(^{(55,52)}\), they can still cause VH. The presence of VH characterizes Stage IV. The higher the circumferential involvement of the eye by the SF, the greater the tendency for VH to occur\(^{(53)}\) (Figure 4). The frequency of stage IV is difficult to establish because small VH near the SFs may go unnoticed; in fact, these are described in approximately 5% of cases\(^{(56)}\). Their occurrence is pathologically relevant because it can worsen the retinal traction, which greatly contributes to the development of RD. The abnormal traction of the SFs is often accompanied by elevation of the adjacent retina in the form of tractional retinoschisis or RD. The association between retinal atrophy (post-ischemic) and vitreous traction can lead to the progression of the disease.

**Figure 4.** Proliferative sickle retinopathy at various stages in the same eye of a 47-year-old patient (HbA - 0%, HbS - 50.3%, Hbc - 48.9%, Hdf - 0.9%). A) Fluorescein angiography of the inferior temporal periphery showing the abrupt interruption of both the arterioles and venules of the blood column. Partial vascular remodeling has moved the peripheral vascular arcade to a more posterior position toward the equator of the eye, clearly delimiting the vascularized and non-vascularized zones of the retina (stages I and II). B) Neovessel formation or sea fan (SF) on the boundary of the perfused zone (stage III). C) Retinographical appearance of the SF, which is out of the retinal plane towards the vitreous with fibroglial proliferation on its surface and localized vitreous hemorrhage (stage IV). D) In the angiogram, this SF was intensely vascularized. In all photos of the angiogram, the wall of the retinal vessels shows no fluorescein staining; staining is only observed in the SF (a purely obstructive phenomenon). Courtesy of Dr. Gustavo Miki.

**Figure 5.** A) Retinography of the posterior pole of the right eye in a SC patient showing the streaks of vitreous organization. B) The temporal periphery shows three contiguous SFs with fibrous proliferation and localized vitreous hemorrhage. C) The middle SF is not perfused by the vascular loop (arrow) and suffered an auto-infarction.

**Figure 6.** Complications of proliferative sickle retinopathy in a patient of SC genotype. A) Macular epiretinal membrane. B) Traction on the peripheral retina with rupture. C and D) The contralateral eye of the patient, showing typical peripheral proliferative retinopathy. The tortuous vessel is the retinal arteriole that feeds the SF.
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formation of retinal tears. These are typically followed by RD, which is usually mixed and characterizes Stage V; it occurs together with VH in 1.2% of SS patients and in 13.4% of SC patients\([5]\) (Figure 6). Blindness originating from this untreated pathological process is rare\([46,49,50]\); however, it has been described in 12% of cases, half of which occurred between 22-28 years of age\([46]\) justifying the use of preventive treatment.

The occurrence of neovascularization of the optic disc is rare\([22,46,48,53,54]\).

DIFFERENTIAL DIAGNOSIS

Other diseases that occur with macular ischemia or neovascularization should be considered in the differential diagnosis of sickle-cell retinopathy. We must remember that in sickle-cell retinopathy the neovessels are always outside the vascularized retina, i.e., situated anterior to the boundary between the vascularized and non-vascularized retina; in addition, when fluorescein angiography is used, there is typically no dye in the retinal vessels in the perfused retina. This means that there is no inflammation of the vascular wall; the disease process represents a purely occlusive phenomenon (Figure 4).

The diseases that occur concomitantly with macular ischemia or neovascularization and should be considered in the differential diagnosis are\([56]\):

I. Other causes of macular ischemia:

- Diabetic retinopathy
- Retinal vascular occlusion
- Embolic phenomena (e.g., talc retinopathy)
- Infectious diseases (dengue)\([56]\)

II. Other causes of peripheral retinal neovascularization, VH and RD:

- Ischemic vascular disease:
  - Proliferative diabetic retinopathy
  - Branch retinal vein occlusion
  - Ocular ischemic syndrome
  - Retinopathy of prematurity
- Eales’ disease (Figure 7)
- Familial exudative vitreoretinopathy
- Chronic myelogenous leukemia
- Scleral buckle
- i. Hyperviscosity syndrome\([51]\) (Figure 8)
- Inflammatory disease with possible ischemia:
  - Sarcoidosis
  - Retinal vasculitis
  - Intermediate uveitis
  - Acute retinal necrosis
- Miscellaneous:
  - Incontinentia pigmenti
  - Autosomal dominant vitreoretinochoroidopathy
  - Old RD

TREATMENT

In sickle-cell retinopathy, VH resulting from PSR is the primary cause of reduced acuity. The treatment strategy is based on closing the SF\([58,66]\). Patients in stages I and II are not treated because treatment of the ischemic retina does not prevent the formation of SF, and the majority of patients do not develop SF or its complications\([49,50]\). The treatment is still controversial because many SF regress spontaneously, especially in older SS patients\([50]\), in whom the incidence of blindness is very small\([5,58,47,60,62]\). Therefore, the recommended approach is to treat SF in all SC patients and SS patients below 40 years of age with sectorial technique.

Patients with recent vitreous hemorrhage and visual acuity impairment are usually followed for at least six months to allow spontaneous clearing. Photocoagulation or cryotherapy can be performed when the visibility permits it.

Figure 7. Differential diagnosis of sickle cell retinopathy: Eales’ Disease, showing zones of hypofluorescence caused by blockage (hemorrhages). In the angiogram, note that the retinal vessel wall shows fluorescein staining that is suggestive of an inflammatory component (vasculitis). Courtesy of Dr. Polo Eduardo San Martin González.

The photoocoagulation technique currently used consists of sectoral treatment of the SF. This type of treatment has been effective and safe in the treatment of PSR in Stage III and prevents subsequent complications.

Surgical treatment

Surgical treatment is indicated in complications of PSR (VH and RD). Because patients with sickle-cell disease are susceptible to intra- and postoperative complications, some general precautions must be taken.

The presence of VH (Stage IV) causes the first visual symptoms. Due to the high ocular morbidity that these patients exhibit during surgery, there is a 6-month waiting period for reabsorption of the VH, after which the SF is treated with laser or cryotherapy.

The subsequent procedures must be performed with ultrasound and intraocular pressure (IOP) measurement because of the risks of concomitant RD and secondary glaucoma. Pars plana vitrectomy (PPV) is considered when the VH is massive and long in duration or if the complications listed above occur. SF treatment is performed during vitrectomy after relieving the traction on these structures. Reports of surgical treatment for VH are rare.

The major technical problem is the manipulation of the extreme retinal periphery, where iatrogenic breaks and cataract may occur. The presence of hyphema, residual VH and secondary glaucoma are complications that are difficult to treat.

The use of scleral buckle surgery to treat RD is challenging because there is a great chance that these patients will develop vascular occlusion secondary to the procedure that may progress to phthisis bulb.

In a retrospective study of 27 patients, 2 of whom were SS and the remaining were SC, 10 eyes were observed with VH (3 patients), tractional RD (4 patients) and rhegmatogenous RD (3 patients). PPV was performed in 18 eyes of 17 patients, including VH (7 eyes), tractional RD and rhegmatogenous RD (3 eyes each), epiretinal membrane (3 eyes of 2 patients) and macular hole (2 eyes). Of the 27 patients, 15 (83%) experienced improved visual acuity, including two patients (3 eyes of 2 patients) and macular hole (2 eyes). Of the 27 patients, 15 (83%) experienced improved visual acuity, including two patients with spontaneous resolution of RD (1 tractional and 1 rhegmatogenous). It was reported that iatrogenic breaks occurred as a result of the delamination technique used in the vitrectomy. Thus, segmentation rather than removal of traction streaks should be performed, and increased IOP should be avoided to control intraoperative bleeding and prevent intense photoocoagulation. Erythrocytapheresis should be considered in SC patients.

General care related to surgery

A cooperative study of sickle-cell disease in the United States reported that total mortality was 1.1% within 30 days after any surgical procedure and stated that preoperative transfusion could reduce the risk of complications. Thus, some general preoperative precautions should be taken in addition to proper hydration, acidosis correction, and the use of saline solution and a properly heated room. The following points should be considered:

1. Blood transfusion - To reduce immediate postoperative morbidity in patients with sickle-cell disease, the Department of Apheresis of the Fundação Pró-Sangue Hemoctroto de São Paulo (FPS/HSP) and the Hematology service of the Faculdade de Medicina da Universidade de São Paulo (FMUSP) have designated as standard protocol performing erythrocytapheresis 48-72 hours prior to surgery. During the apheresis session, an exchange of the erythrocyte volume of the patient, calculated according to volemia and hematocrit, is performed. The blood replacement is isovolumetric, and concentrated erythrocytes that have been previously washed or filtered are used. The replacement of this volume is sufficient to reduce the abnormal hemoglobin levels of the patient by 40 to 50%.

2. Use of supplemental oxygen.

3. Local anesthesia should be used whenever possible to avoid hypotension with poor perfusion caused by general anesthesia.

4. Dilution of pupils - Phenylephrine drops should be avoided; parasympatholytics are preferred because they do not cause vasoconstriction in the anterior segment.

5. Decreased IOP - An IOP as low as possible should be used to improve tissue perfusion. Single doses of carbonic anhydrase inhibitors or intravenous mannitol can be used to achieve this goal. Repeated use of these drugs can cause hemococoncentration or acidosis, leading to sickling.

Alternative treatments/future perspectives

Siqueira et al. reported a case of a 36-year-old man with PSR in the temporal peripheral retina of both eyes and VH in the right eye. Intravitreal injection of bevacizumab (1.5mg/0.06ml) was selected because the patient refused to undergo vitrectomy. There was a reduction of fluorescein leakage in the angiography one week later; four weeks later, he presented better visual acuity and regression of the neovascularization. At that time, photoocoagulation of the non-perfused area of the temporal periphery of both eyes was performed. No adverse effects were described. The authors state that the results require further investigation to verify the safety of bevacizumab as well as to determine whether the neovascularization effectively regressed. This type of therapy could be useful as an adjunct to photoocoagulation in the management of PSR and could obviate the need for PPV in some patients. However, it is important to emphasize that for many patients there is a risk of worsening the tractional component; this risk is very common in patients with sickle-cell disease due to the use of anti-angiogenic drugs and has been well described in diabetic retinopathy.

Finally, we cannot forget that the ocular changes that compromise the vision of patients with sickle-cell disease are the consequences of a complex systemic pathophysiological process. The prevention of these ocular complications can be achieved through the use of new drugs that focus on the physiopathology of the disease. The multifaceted nature of sickle-cell disease allows the development of vascular occlusion to be arrested at multiple points, including HbS polymerization, erythrocyte density and cell-cell interactions. The emerging treatment options that can be employed to improve tissue perfusion in these patients include:


II. Erythrocyte hydration: clotrimazole, magnesium pilolate.


IV. Antioxidant therapy: glutamine, deferiprone.

V. Antithrombotic agents: heparin, ticlopidine, warfarin.

VI. Vasodilation: nitric oxide, arginine, Flocor.

VII. Decrease in HbS cells: transfusion, apheresis.

VIII. Hematopoietic stem cell transplantation and gene therapy.

References


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