Purpose

First retinal hemorrhages related with high altitude were detected in 1968 in Mount Logan in Canada [1]. Their relation with high altitude, were first described in 1970 by Frayser et al. and first optic disc changes associated with high altitude were described by Singh et al. [1,2].

The mountain sickness syndrome includes “acute mountain syndrome” (AMS), “high altitude retinal hemorrhage” (HARH) and the severe forms “high-altitude pulmonary oedema” (HAPE) and cerebral oedema (HACE) [2]. These diseases are different facets of failure to acclimatize at high altitude [3,4]. These conditions are associated with the rapid ascent to above 3000 m.

Till date, many lesions in the eyes have been detected in relation with high altitude such ascornal edema, tear film alterations, pupil defects, changes in the IOP [5,6] but there's no clear relation between them and the severe forms of mountain sickness [7]. The purpose of this article is to present a case of spontaneous resolution of this disease.

Case Report

A 36-year-old man came to the institute in January 2013 with chief complaints of diminished vision in the right eye after an episode of loss of consciousness while climbing a mountain 10 days ago, (no data is recorded about meters climbed). He refers he was taken to emergency and recovery after 5 days in the hospital. He refers he didn't need to be taken to intensive care unit. By the time he arrived to the hospital, blood pressure and vital signs were within normal limits and brain and pulmonary disease were ruled out there. He noted while being at the hospital that he saw a red spot in the right eye and after returning to his hometown he decided to consult with an ophthalmologist and control clinically with a physician as being told.

On examination, he presented an UCVA of 20/100 the right eye and 20/20 in the left eye. IOP was within normal limits. External examination was normal. In the right eye fundus examination, we found a macular hemorrhage, CDR (Cup Disc Ratio) 0.2:1, two small diffuse paramacular hemorrhages were also found and the retina was attached, the exam in the left eye showed a few diffuse hemorrhages in both vascular arcades. The lesions were recorded by an Optical Coherence Tomography (OCT) image of both eyes (Figure 1).

The patient lost follow up and return to the institute a year later in January 2014. At that moment his UCVA was 20/20 in both eyes. The rest of the exam was within normal limits including IOP. The posterior segment OCT showed a complete resolution of the hemorrhages in both eyes and no evidence of any scar or intraretinal lesion were seen as shown in Figure 2.
Conclusions

While climbing a mountain and the increase of the altitude, there is a reduction in the atmospheric pressure leading to a reduction in the pressure of O$_2$ (Atmosphere O$_2$ pressure=Atmospheric pressure × % O$_2$ in the atmosphere). This lead to a reduction in the blood partial pressure of oxygen causing hypoxia if there is no time for acclimatization. This hypoxia generates a reduction in the levels of oxygen that go to the different tissues and cells [8]. To compensate this, the body reacts with tachycardia, hyperventilation, hypocapnia and metabolic. It is extremely important to know the rate of ascent suggested for each pick. A few reports mention that over 3000 mts only 300 mts should be climbed per day [9,10].

Changes on the posterior pole are produced by physiological changes and pathological ones. The physiological changes are due to acclimatization, and are mainly vasodilation and tortuosity. The pathological changes are due to hypoxia. This is thought to be the producer of tissue edema causing optic disc edema as the swollen of the disc in HACE is not related to elevation of the intracranial pressure (ICP). Hypoxia may be the cause of vessel filtration in the periphery of the retina [11]. Also this may be the causative agent of damage of the wall cells of the vessels leading to the production of hemorrhages [11]. Daniel Barthelmes et al. mentioned that higher hematocrit, may contribute to the damage of the endothelial cells [12].
Cerebral outflow limitation capacity may cause headache and it is suggested that this could be the cause of the retinal venous diameter increase [11]. Advanced technology used to detect hemorrhages in the retina helped to show that there is no correlation between HARH and AMS or HACE [13]. It is known that optic disc swollen related to high altitude exposure may have different origin if it is not related to increased intracranial pressure [11]. Although there can be a few associations between retinal disorders and HACE, everything is questionable. As mentioned in the update by Willmann et al. the retinal vein engorgement as indicator of outflow limitation must be assessed [11].

To conclude the pathophysiology of HARH remain still obscure and even the retina and the optic nerve are closely related to the brain and cranial structures there is no clear relation between HARH and HACE or even HAPE. Prevention to avoid lesions is mandatory even though retinal lesions could resolve spontaneously. Most of the posterior pole lesions resolve spontaneously in a few days or weeks, but a few of them can leave sequelae such as scotomas of visual field [7]. More research need to be done to rule out the relation between retinal and cerebral lesions to avoid severe and irreversible problems.

References


