Inflammatory seborrhic keratosis resolution after hyperbaric oxygen therapy: Case presentation and pathophysiology review

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Abstract

Seborrhoeic keratosis (SK) is a common epidermal tumor, consisting of a benign proliferation of immature keratinocytes. The natural history of SK is a slow progression over time and complete remission is not expected. The article presents the first case of a complete resolution of a large (2.5 cm diameter) SK lesion after hyperbaric oxygen therapy (HBOT). In addition to the case presentation, the pathophysiology of SK and the potential beneficial physiological effects of HBOT are reviewed and discussed.

Introduction

Seborrhoeic keratosis (SK) is a common epidermal tumor, consisting of a benign proliferation of immature keratinocytes. Clinically, SK manifests as solitary or multiple, well demarcated brownish papules or plaques with a verrucous surface, that predominantly localizes at the head, neck and trunk areas.1 Since the tumor slowly grows, in many cases the lesion is surgically removed due to cosmetic reasons or because the lesions are traumatized and become symptomatic.2,5

In this article we report an unexpected full resolution of a large SK lesion in a patient treated by hyperbaric oxygen therapy (HBOT) due to hemorrhagic cystitis. In addition to the case presentation, the pathophysiology of SK and the potential beneficial physiological effects of HBOT in the setting of SK will be reviewed and discussed.

Case Report

A 74 years old man was admitted for HBOT due to chronic late post radiation hemorrhagic cystitis (grade 3-4 RTOG). In addition, on his left temple, he had a slowly growing SK lesion, known for about 24 months prior to the treatment. The SK presented as a red non-tender demarcated lesion, 2.5 cm diameter in size, with clear boundaries and occasional mild oozing (Figure 1A).

The lesion was diagnosed as inflammatory seborrhoeic keratosis and was scheduled for surgical removal after the intendent HBOT.

The medical history of the patient included prostate cancer (Glisson score 6), diagnosed 13 years prior to his admission, that was treated with radiation. Since he had chronic unremitting hemorrhagic cystitis, he was referred to HBOT. In addition to the cystitis, he had type-II diabetes mellitus, hypertension, osteoporosis, gastro-esophageal reflux, hyperlipidemia, s/p resection of craniopharyngioma in 2019 and a fractionated stereotactic radiation therapy, gait disturbance with parkinsonism and a primary unspecified kidney tumor.

The HBOT protocol used for his hemorrhagic cystitis included a total of 56 hyperbaric sessions, five days per week of 90 minutes 100% oxygen at 2 ATA with five-minute air breaks every 20 minutes. The treatment went well with no significant side effects and the clinical symptoms of the hemorrhagic cystitis resolved.

Surprisingly, in addition to the resolution of the cystitis, there was a full resolution of the 2.5 cm seborrhoeic keratosis lesion (Figure 1B). During a follow-up, 16 weeks after the last hyperbaric session, the skin was still intact without any sign of recurrence.

Discussion and Conclusions

SKs are common epidermal tumors that usually develop after the age of 50. Skin aging and cumulative UV exposure are considered to play a major role in SK pathogenesis and seem to cause increased expression of amyloid precursor protein (APP),6 which is a marker of cellular senescence and chronic inflammation, particularly in human keratinocytes (Figure 2 summarizes the pathophysiology cascade of SK).7

APP expression has been evaluated in SKs versus normal skin by different methods and it was found that APP and its downstream products (i.e. amyloid-β42) are highly expressed in SK lesions as compared to the adjacent normal skin tissues.8 Similar to growth factors and tumor growth factor alpha (TGF-α), APP can induce proliferation of epidermal keratinocytes, and contribute to mitochondrial dysfunction and oxidative phosphorylation.9 In addition to the skin, APP is highly expressed and has a pathophysiology role in many of the age-related diseases such as Alzheimer’s disease, atherosclerosis, Parkinson and macular degeneration.8,10 HBOT utilizes 100% oxygen in an environmental pressure higher than one absolute atmosphere (ATA) to enhance the amount of oxygen dissolved in body’s tissues. Repeated intermittent hyperoxic exposures have been shown to induce physiological effects which normally occur during hypoxia in a hyperoxic environment.
including stem cells proliferation, generation of new blood vessels (angiogenesis) and enhanced tissue regeneration.  

The direct effect of HBOT on keratinocytes was evaluated by the use of monolayer cultures including dermal fibroblasts, keratinocytes and melanocytes. It was found that repeated HBOT sessions at a pressure of 2 atmospheres, as was used in our treatment protocol, inhibits keratinocytes proliferation. In addition, in another study it was demonstrated that HBOT ameliorates APP and amyloid beta (Aβ) plaques in the brain of Alzheimer prone mice. With regards to UV related injury, it was found that pretreatment with HBOT significantly reduces UV-A induced apoptosis and proliferation in hairless SKH1-E mice. Hence, as summarized in Figure 2, HBOT has the potential to intervene and revert the pathophysiological processes responsible for the development of SK lesions.

In our patient, the SK lesion at a diameter of 2.5 cm had a complete remission after HBOT (Figure 1).

The natural history of SK is a slow progression over time and complete remission is not expected. The only available treatments for SK include surgical removal, cryosurgery, curettage, electrodessication, shave excision or laser therapies (CO2, YAG). To the best of our knowledge, this is the first reported case of complete resolution of SK without local intervention. Had the lesion not been so large in size and not at such a notable location, we would have probably missed HBOT’s effect on the lesion. Until this case, we have not monitored SK lesions in patients treated at our center, and the skin lesions have not received the attention they deserve. By presenting this unique case, we hope that awareness of the potential beneficial effect of HBOT on SK lesions will increase and prospective clinical trials on different types of SK will be initiated. These studies may potentially shed additional light on the pathophysiology of SK and may help in developing noninvasive biological interventions, such as HBOT, to treat it.

References

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