Ocular surface squamous neoplasia in HIV-infected patients: current perspectives

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Abstract: Ocular surface squamous neoplasia (OSSN) refers to a spectrum of conjunctival and corneal epithelial tumors including dysplasia, carcinoma-in-situ, and invasive carcinoma. In this article, we discuss the current perspectives of OSSN associated with HIV infection, focusing mainly on the epidemiology, pathophysiology, clinical manifestations, diagnosis, and treatment of these tumors in patients with HIV. Upsurge in the incidence of OSSN with the HIV pandemic most severely affected sub-Saharan Africa, due to associated risk factors, such as human papilloma virus and solar ultraviolet exposure. OSSN has been reported as the first presenting sign of HIV/AIDS in 26%–86% cases, and seropositivity is noted in 38%–92% OSSN patients. Mean age at presentation of OSSN has dropped to the third to fourth decade in HIV-positive patients in developing countries. HIV-infected patients reveal large aggressive tumors, higher-grade malignancy, higher incidence of corneal, scleral, and orbital invasion, advanced-stage T4 tumors, higher need for extended enucleation/exenteration, and increased risk of tumor recurrence. Current management of OSSN in HIV-positive individuals is based on standard treatment guidelines described for OSSN in the general population, as there is little information available about various treatment modalities or their outcomes in patients with HIV. OSSN can occur at any time in the disease course of HIV/AIDS, and no significant trend has been discovered between CD4 count and grade of OSSN. Furthermore, the effect of highly active antiretroviral therapy on OSSN is controversial. The current recommendation is to conduct HIV screening in all cases presenting with OSSN to rule out undiagnosed HIV infection. Patient counseling is crucial, with emphasis on regular follow-up to address high recurrence rates and early presentation to an ophthalmologist for any symptoms in the unaffected eye. Effective evidence-based interventions are needed to allow early diagnosis and treatment, as well as prevention of the disease.

Keywords: eye, conjunctiva, OSSN, ocular surface squamous neoplasia, HIV, human immunodeficiency virus

Introduction

Ocular surface squamous neoplasia (OSSN) refers to a spectrum of conjunctival and corneal epithelial tumors including dysplasia, carcinoma-in-situ and invasive carcinoma (squamous cell carcinoma) which may or may not be associated with intraocular or orbital extension.1,2 OS malignancies, such as OSSN, Kaposi’s sarcoma, and non-Hodgkin’s lymphoma, are notably expressed in people with HIV/AIDS, among which OSSN is known to occur in 4%–8% of patients.3,4
Although there has been a decline in the incidence of HIV in recent years, there has been a surge in the incidence of OSSN in the population with HIV, due to its linear relationship with the HIV pandemic in the last few decades.\(^5\)\(^\text{-}^\text{18}\)

This dramatic rise in HIV infection during the pandemic has resulted in gradual drift in the age of presentation, clinical course, and management of patients with OSSN.\(^5\)\(^\text{-}^\text{10}\)

HIV not only increases risk of OSSN but also influences the severity of disease and its prognosis.

### Incidence and risk of OSSN with HIV infection

According to a World Health Organization report, the prevalence of HIV/AIDS worldwide was 36.7 million at the end of 2016. While the burden of the epidemic continues to vary substantially between countries and regions, sub-Saharan Africa remains most severely affected and accounts for almost two-thirds of the total new HIV infections globally.\(^20\)

The geographical distribution of OSSN all over the world is also affected by the HIV pandemic.\(^10\)\(^\text{-}^\text{21}\)

HIV infection is recognized as a risk factor for the development of OSSN in various studies from sub-Saharan Africa.\(^13\)\(^\text{-}^\text{22}\)\(^\text{-}^\text{26}\) The Kampala Cancer Registry in Uganda recorded a sixfold increase in the incidence of conjunctival squamous-cell carcinoma: from an average of six per million per year in 1988 to 35 per million per year in 1992.\(^10\) Various studies have observed a three- to 30-fold increased risk of OSSN developing in HIV infected individuals.\(^7\)\(^\text{-}^\text{27}\)\(^\text{-}^\text{29}\)

Although a strong association between HIV and OSSN has been established by numerous studies, some patients with OSSN are not aware of their HIV status until HIV screening confirms the seropositivity. In studies from sub-Saharan Africa where HIV screening was done in all patients with OSSN, seropositivity was detected in 49%–92% cases, indicating a high association between OSSN and HIV status in African countries.\(^5\)\(^\text{-}^\text{7}\)\(^\text{-}^\text{12}\)\(^\text{-}^\text{14}\)\(^\text{-}^\text{15}\)\(^\text{-}^\text{30}\) In Africa, OSSN has been reported as the first presenting sign of HIV/AIDS in 50%–86% cases.\(^8\)\(^\text{-}^\text{12}\)\(^\text{-}^\text{31}\) This trend is now well documented in many African countries, and with increasing migration it is appreciated in other developing and developed countries as well. In studies from India by Kaliki et al and Kamal et al, HIV positivity was noticed in 38%–41% of diagnosed OSSN patients, of which 70% were unaware of their status prior to screening.\(^32\)\(^\text{-}^\text{33}\) In 26% of patients, OSSN was the first and only evident manifestation of HIV.\(^32\)

There are limited data from the US, which shows a higher incidence of OSSN among people with HIV than the general population. Guech-Ongey et al noted a significant 12-fold risk of OSSN in people with HIV/AIDS.\(^6\) Frish et al also noticed similar results while studying relative risk of human papilloma virus (HPV)-associated cancers in HIV-infected individuals.\(^34\)

Goedert and Coté observed that the risk of OSSN among people with HIV changed with duration from diagnosis of AIDS, with an exponential rise in squamous-cell carcinoma in HIV patients 2 years after the diagnosis of AIDS.\(^35\) In a study by Karp et al among younger patients (<50 years) with OSSN, 50% were HIV-positive, 33% of whom were unaware of their HIV infection.\(^14\) In a systematic review and meta-analysis of 12 studies, it was revealed that HIV infection augmented the risk of OSSN, with an overall relative risk estimate of 8.06 (95% CI 5.29–12.3).\(^18\)

### Epidemiology

#### Mean age at presentation

The epidemiology of OSSN has changed over the last few decades, especially in developing countries. Although once considered an uncommon tumor occurring in the elderly,\(^36\) it is becoming more common and more likely to affect young populations.\(^12\)\(^\text{-}^\text{14}\)\(^\text{-}^\text{32}\)\(^\text{-}^\text{37}\)\(^\text{-}^\text{40}\) It has been observed that the mean age at presentation of OSSN has dropped to the third to fourth decade in HIV-positive patients from the sixth decade in HIV-negative patients.\(^7\)\(^\text{-}^\text{11}\)\(^\text{-}^\text{12}\)\(^\text{-}^\text{39}\)\(^\text{-}^\text{41}\)\(^\text{-}^\text{42}\)

The mean age at presentation of OSSN in people with HIV/AIDS in various studies is approximately 35–41 years.\(^7\)\(^\text{-}^\text{8}\)\(^\text{-}^\text{11}\)\(^\text{-}^\text{12}\)\(^\text{-}^\text{14}\)\(^\text{-}^\text{15}\)\(^\text{-}^\text{32}\)\(^\text{-}^\text{39}\)\(^\text{-}^\text{39}\)\(^\text{-}^\text{41}\)\(^\text{-}^\text{41}\)\(^\text{-}^\text{41}\) In developed countries, patients with OSSN have a different disease profile and continue to present in elderly in both the general population\(^36\)\(^\text{-}^\text{38}\)\(^\text{-}^\text{43}\)\(^\text{-}^\text{44}\) and in HIV-infected individuals.\(^5\)

#### Sex predisposition

Studies from Africa have reported a striking feature of either a female preponderance\(^1\)\(^\text{-}^\text{10}\)\(^\text{-}^\text{14}\)\(^\text{-}^\text{39}\)\(^\text{-}^\text{41}\)\(^\text{-}^\text{44}\) or no sex difference\(^3\) in HIV-positive OSSN patients, while others have noted dominance by elderly males.\(^8\)\(^\text{-}^\text{15}\)\(^\text{-}^\text{45}\) This predilection may result from the presence of the HIV/AIDS pandemic, high HPV exposure, and solar radiation in the region.\(^45\) In developed countries, males are more commonly affected than females in the general population.\(^36\)\(^\text{-}^\text{38}\)\(^\text{-}^\text{43}\)\(^\text{-}^\text{44}\) as well as in people with HIV.\(^6\) Studies from Australia, Britain, and San Francisco have found that 70%–80% of patients with OSSN are males,\(^36\)\(^\text{-}^\text{38}\)\(^\text{-}^\text{46}\) with rare occurrence of HIV seropositivity.

#### Etiology

The cause of OSSN is multifactorial, but the precise etiopathogenesis is unknown. HIV,\(^10\)\(^\text{-}^\text{11}\) HPV,\(^11\)\(^\text{-}^\text{13}\)\(^\text{-}^\text{47}\) and solar ultraviolet exposure\(^14\)\(^\text{-}^\text{48}\) are postulated risk factors for the development of OSSN.\(^5\)\(^\text{-}^\text{17}\) An association between OSSN
and cigarette smoking, vitamin A deficiency, and allergic eye disease has also been described. It is suggested that the increase in incidence of OSSN in sub-Saharan Africa is related to the coexistence of the HIV/AIDS pandemic, infection with HPV, and exposure to solar radiation in the region. The overlap of these factors in parts of the world with rapidly rising incidence of OSSN indicates that they possibly interact with one another. Therefore, it is difficult to describe a single one as the causative factor.

It is considered that HIV infection causes the breakdown of immunosurveillance of malignant cells as a result of suppression of cell-mediated immunoreponse. The relationship between HPV and OSSN is controversial, with variable results about the two types of HPV. Studies have established no association between mucosal HPV types and OSSN, while others found an uncertain role of cutaneous HPV type in OSSN. A systematic review by Carreira et al suggested that only the cutaneous HPV subtypes were associated with an increased risk of OSSN. Studies suggest that HPV alone may be a contributing factor to OSSN, but is unlikely to cause it. Karp et al hypothesized that HIV predisposes to OSSN by creating a “permissive environment” for activation of oncogenic HPV, which subsequently act as a cofactor in the development of neoplasia. This also explains the increased prevalence of HPV in immunosuppressed individuals with HIV disease.

**Clinical features**

**Symptoms**

Symptoms in patients with OSSN may range from none at all to severe pain and/or visual loss. The most common presentations are a red eye, ocular irritation, or appearance of a mass lesion in the eye. OSSN presents as a slowly growing lesion in the general population, while it behaves aggressively in HIV-infected individuals in both developing and developed countries. It often presents with large and unsightly disfiguring lesions in late disease. In HIV-infected individuals, longer duration between onset of symptoms and diagnosis of tumor is noted in a few studies from Africa, with a mean history of 3 months at presentation. Such longer duration and largeness of the lesion imply that many patients either do not seek medical care early or receive long-term conservative treatment for other ocular conditions due to misdiagnosis. Satisfactory training of eye-health care professionals is necessary for active participation to rule out malignancy clinically, to consider incisional biopsy for histopathological confirmation in doubtful cases, and to follow up closely.

**Signs**

OSSN commonly affects the interpalpebral conjunctiva, and frequently arises from the nasal limbus. It can present either as a solitary growth or with diffuse involvement of the OS. Solitary tumors can be nodular, noduloulcerative, gelatinous, leukoplakic, placoid, or papillary in morphology (Figure 1). Makupa et al noted higher incidence of leukoplakia and feeder vessels, while Kabra et al mentioned large lesions with fornical extension at the time of presentation in HIV-infected patients.

The majority of cases are unilateral, however, 15% of cases present with bilateral involvement, and multifocal lesions are noted in 3% of cases. Bilateral presentation can be simultaneous or sequential, as highlighted in a series of four patients in seropositive patients by Masanganise et al. Finances and fear of losing the only eye were the possible reasons for delayed follow-up and advanced disease requiring enucleation or orbital exenteration.

Studies from the sub-Saharan region have established that HIV-infected patients have larger tumors, higher-grade malignancy, and increased risk of tumor recurrence. Kaliki et al found significant differences between HIV-positive and HIV-negative patients, including larger tumors, higher incidence of corneal, scleral, and orbital invasion, advanced stage T4 American Joint Committee on Cancer tumors, and

![Figure 1](https://www.dovepress.com/)

**Figure 1** Clinical presentation of ocular surface squamous neoplasia in human immunodeficiency virus-infected patients.

**Notes:** (A) Leukoplakic lesion in the temporal limbus and bulbar conjunctiva; (B) gelatinous lesion involving inferonasal limbus and peripheral cornea with diffuse limbal thickening extending from 9 o’clock to 6 o’clock hour positions; (C) nodular lesion involving temporal bulbar conjunctiva; (D) nodular lesion in the inferior peripheral cornea and bulbar conjunctiva covered with extensive keratin; (E) diffuse papillary lesion involving superior, temporal, and inferior quadrants of bulbar, fornical, and tarsal conjunctiva; (F) fungating mass involving the entire anterior ocular surface with anterior orbital extension.
higher need for extended enucleation/orbital exenteration in HIV-positive cases. Similar findings of aggressive and invasive OSSN in individuals with HIV were described in a study by Shields et al. The invasive nature of the tumor may be correlated with longer duration of symptoms.

Rarely, this tumor can present as necrotizing scleritis with scleral perforation and prolapse of uveal tissue as a manifestation of intraocular extension. Early identification of corneal, scleral invasion, intraocular or orbital extension clinically or with ancillary investigations is crucial. This will reduce morbidity and will result in better anatomical outcomes in terms of globe salvage and cosmesis. An unusual case in a seropositive male has been described who presented with widespread metastatic squamous-cell carcinoma after the primary OSSN in the lower palpebral conjunctiva had been misdiagnosed and treated conservatively as chalazion. OSSN can often extend to regional lymph nodes, surrounding paranasal sinuses and the brain in HIV-positive individuals. Death may result from regional or distant metastases, as well as intracranial spread. Therefore, antiretroviral therapy (ART)-center referral is mandatory in every single OSSN patient with positive serology for a complete systemic examination, counseling, and management. The published literature on OSSN and HIV is summarized in Table 1.

### Diagnosis

Although OSSN is predominantly a clinical diagnosis, overlap between the clinical features of OSSN and benign lesions sometimes causes difficulty in differentiating between the two. Additionally, the occurrence of malignant features on histopathology in pterygia and other benign lesions shows that histopathological confirmation is indispensable for a definitive diagnosis of OSSN (Figure 2). Investigative modalities, such as ultrasound biomicroscopy, anterior-segment optical coherence tomography, computed tomography, or magnetic resonance imaging, of the orbit are necessary to rule out stromal invasion, corneal/scleral involvement, and intraocular/orbital extension of the tumor.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Location</th>
<th>Type of study</th>
<th>OSSN patients with HIV/total OSSN patients</th>
<th>Mean (range) age at OSSN presentation, years</th>
<th>Sex, male: female</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kestelyn et al (1990)</td>
<td>Rwanda, Africa</td>
<td>Case–control</td>
<td>9/11 (82%)</td>
<td>37 (26–51)</td>
<td>1.75:1</td>
</tr>
<tr>
<td>Ateenyi-Agaba (1995)</td>
<td>Kampala, Uganda, Africa</td>
<td>Case–control</td>
<td>36/48 (75%)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Karp et al (1996)</td>
<td>Miami, USA</td>
<td>Retrospective</td>
<td>3/6 (50%)</td>
<td>43 (38–49)</td>
<td>All male</td>
</tr>
<tr>
<td>Malawi, Africa</td>
<td>Case series</td>
<td>25/29 (86%)</td>
<td>31 (22–50)</td>
<td>1:1.1</td>
<td></td>
</tr>
<tr>
<td>Porges and Groisman (2003)</td>
<td>Zimbabwe, Africa</td>
<td>Case–control</td>
<td>12/13 (92%)</td>
<td>37 (21–60)</td>
<td>1:1.8</td>
</tr>
</tbody>
</table>
Management
Although the management of OSSN in the general population has been well elaborated in numerous studies, current clinical practice in treatment of OSSN in HIV-positive individuals is based on limited medical literature. There is little information about various treatment modalities currently used against this tumor or their outcomes in patients with HIV. The treatment of OSSN in patients with HIV/AIDS depends on the tumor laterality, extent, invasion into adjacent structures, and overall systemic status. The most commonly performed treatment modality for resectable tumors (fewer than two quadrants) with well-defined lesions is surgical management by wide excision. The main objective of surgical excision is complete removal of the tumor to minimize recurrence.70 Wide excision biopsy is executed by a “no-touch” technique with 4 mm tumor-free margins for conjunctival component and alcohol keratoepitheliektomy with 2 mm tumor-free margins for corneal component, followed by adjunct double-freeze–thaw cryotherapy to the surrounding resected conjunctival margins and OS reconstruction by direct closure or amniotic membrane graft, based on the size of the surgical defect.32,70,71

Partial lamellar sclerectomy is needed in cases with clinical evidence of scleral invasion. Adjunctive cryotherapy is applied to the tumor bed after tumor excision in cases with episcleral fixity, scleral invasion on imaging, or adherence of lesion to the base during the surgery.72 Concomitant primary simple limbal epithelial transplantation after wide excisional biopsy for OSSN involving >3 clock hours prevents LSCD in cases requiring extensive corneoscleral limbal dissection ≥6 clock hours during surgery.72

Diffuse tumors (2 three quadrants of the OS) and large annular lesions can be managed by chemoreduction/chemotherapy with topical agents, such as mitomycin-C, cidofovir, and 5-fluorouracil,74–77 or immunoreduction/immunotherapy with topical and periocular injection IFNα.73,77,78 Chemoprevention with topical chemotherapeutic agents or immunoprevention with topical IFNα can be considered in patients with microscopic tumor residue.72,74

<table>
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<th>Treatment/tumor recurrence</th>
<th>Remarks</th>
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<tr>
<td>Mean duration of symptoms</td>
<td>Tumor characteristics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>6 months</td>
<td>–</td>
<td>CIN 45%, SCC 55%</td>
<td>Prompt and complete surgical excision</td>
</tr>
<tr>
<td>&lt;3 months</td>
<td>Limbal mass, aggressive</td>
<td>–</td>
<td>Excision biopsy</td>
</tr>
<tr>
<td>3 weeks, 2 months, 2 months</td>
<td>Orbital extension 1, nasal mass 2</td>
<td>SCC 100%</td>
<td>Extended enucleation/subtotal orbital exenteration</td>
</tr>
<tr>
<td>2 weeks, 2 months, 1 year</td>
<td>Masses in superior, temporal quadrants</td>
<td>CIN 100%</td>
<td>Excision biopsy (n=2), topical IFNα (n=1)</td>
</tr>
<tr>
<td>1 week–2 years</td>
<td>Base 3 mm to 3 cm, all lesions at 3:00 or 9:00, melanin positive in few, fungating rough surface positive</td>
<td>SCC 27%, CIN 11%</td>
<td>Excision biopsy + superficial keratectomy, tumor recurrence (n=1)</td>
</tr>
<tr>
<td>3–&lt;12 months</td>
<td>MC corneal overriding 92%, alteration of conjunctival color 66.7%, nasal 66.7%</td>
<td>CIS 16%, SCC 84%, pterygium harboring carcinoma 16.7%</td>
<td>–</td>
</tr>
</tbody>
</table>

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### Table 1 (Continued)

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<tbody>
<tr>
<td>Guech- Ongey et al (2008)*</td>
<td>Nine states, USA</td>
<td>Cancer registry-record review</td>
<td>15/491,048 patients with AIDS</td>
<td>15–29 (n=1) 30–39 (n=5) 40–49 (n=4) ≥50 (n=5)</td>
<td>14:1</td>
</tr>
<tr>
<td>Spitzer et al (2008)*</td>
<td>Malawi, Africa</td>
<td>Case series</td>
<td>30/38 (79%)</td>
<td>33 (19–60)</td>
<td>1.4:1</td>
</tr>
<tr>
<td>Shields et al (2011)*</td>
<td>Philadelphia, USA</td>
<td>Retrospective case series</td>
<td>4</td>
<td>54 (42–65)</td>
<td>All 4 male</td>
</tr>
<tr>
<td>Pradeep et al (2012)*</td>
<td>Karnataka, India</td>
<td>Consecutive case series</td>
<td>6/21 (29%)</td>
<td>36 (31–40)</td>
<td>4.2:1</td>
</tr>
<tr>
<td>Makupa et al (2012)*</td>
<td>Tanzania, Africa</td>
<td>Prospective</td>
<td>79/132 (60%)</td>
<td>38 (18–86)</td>
<td>1.2:1</td>
</tr>
<tr>
<td>Gichuhi et al (2015)*</td>
<td>Kenya, Africa</td>
<td>Prospective</td>
<td>98/133 (74%)</td>
<td>39</td>
<td>1:1.8</td>
</tr>
<tr>
<td>Kamal et al (2015)*</td>
<td>Hyderabad, India</td>
<td>Retrospective, cross sectional</td>
<td>83/200 (41%)</td>
<td>40</td>
<td>2.8:1</td>
</tr>
<tr>
<td>Kabra and Khaitan (2015)*</td>
<td>Ahmedabad, India</td>
<td>Retrospective</td>
<td>11/48 (22%)</td>
<td>33 (14–66)</td>
<td>10:1</td>
</tr>
<tr>
<td>Kaliki et al (2016)*</td>
<td>Hyderabad, India</td>
<td>Retrospective</td>
<td>86/228 (38%)</td>
<td>41 (24–26)</td>
<td>3:1</td>
</tr>
</tbody>
</table>

**Abbreviations:** AKE, alcohol keratoepitheliectomy; CIN, conjunctival intraepithelial neoplasia; CIS, carcinoma in situ; MMC, mitomycin C; OSSN, ocular surface squamous neoplasia; PCR, polymerase chain reaction; SCC, squamous-cell carcinoma.
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<td></td>
<td></td>
</tr>
<tr>
<td>1 month–2 years</td>
<td>Tumor size 2 mm to several cm, orbital involvement 6%, nasal predominance</td>
<td>SCC 100%</td>
<td>–</td>
</tr>
<tr>
<td>Leukoplakic 100%, mean tumor basal dimension 13 mm</td>
<td>–</td>
<td>Excision biopsy + AKE + margin cryotherapy, tumor recurrence (n=1)</td>
<td>Mean duration of immunosuppression 5 years.</td>
</tr>
<tr>
<td>1–4 months</td>
<td>All tumors at presentation &gt;8 mm or &gt;3 clock hours, all cases in temporal quadrant</td>
<td>SCC 100%</td>
<td>Excision biopsy + margin cryotherapy ± topical MMC</td>
</tr>
<tr>
<td>≥6 months in 69%</td>
<td>Size of lesion &gt;5 mm 72%, leukoplakia 64%, feeder vessels 67%</td>
<td>Mild–moderate dysplasia 15.4%, severe dysplasia 53.8%, SCC 30.8%</td>
<td>Tumor recurrence in 82%</td>
</tr>
<tr>
<td>8 months</td>
<td>Mean diameter of lesion 6.8 mm, MC-quadrant nasal limbus 61%, leukoplakia 72%, corneal involvement 65%, severe inflammation associated</td>
<td>Moderately differentiated squamous-cell carcinoma 46%, mild–moderate dysplasia 18%, severe dysplasia 21%</td>
<td>Excision biopsy (cryotherapy not done)</td>
</tr>
<tr>
<td>15 months</td>
<td>MC-quadrant temporal 45%, mean basal dimension 12 mm, mean thickness 3 mm, tumor epicenter limbus 80%, corneal involvement 82%</td>
<td>Mild–moderate dysplasia 6%, severe dysplasia 1%, CIS 33%, stromal invasion 60%, scleral invasion 19%, corneal invasion 18%, surgical margins positive 23%, base positive 20%</td>
<td>Excision biopsy + AKE + margin cryotherapy 69%, topical MMC 12%, extended exenteration 4%, orbital exenteration, 16%, tumor recurrence in 30%</td>
</tr>
<tr>
<td>16 months</td>
<td>Bilateral 15%, mean basal dimension 11 mm, orbital involvement 9%, intraocular extension 1%</td>
<td>SCC 55%</td>
<td>Excision biopsy + AKE + margin cryotherapy 75%, topical MMC 7%, extended exenteration 6%, orbital exenteration, 12%, tumor recurrence in 23%</td>
</tr>
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natural killer cells, which further recognize and destroy tumor cells. An intact immune system could be an important connection between IFNα-2b therapy and tumor resolution. Therefore, topical IFNα-2b may not be an ideal choice for patients with underlying immunosuppression, and it is preferred to switch the treatment to a nonimmunomodulating agent, such as 5-fluorouracil or mitomycin C in this patient population.

Moreover, in patients with HIV, treatment response could be paradoxical, and lesions may increase in size following the use of IFNα-2b. Therefore, serology for HIV is recommended in patients with OSSN before commencing treatment with topical IFNα-2b.

**Histopathology**

Higher-grade malignancy in the form of stromal, corneal, and scleral invasion has been noted in HIV-positive OSSN patients. Invasive squamous-cell carcinoma has been noted in 55%–80% of patients with OSSN and HIV, and it is associated with reduced prognosis. It is noted that OSSN in HIV-positive patients has higher-grade malignancy with increased tumor invasion, irrespective of the size of lesion or age at presentation.

**Recurrence of OSSN in HIV population**

Recurrence rates after complete surgical excision of OSSN in the general population vary between 5% and 33% in various studies from Australia, the US, and Canada. In a study from the US, a recurrence rate of 28.5% was noted with simple excisional biopsy, which decreased to 7.7% when combined with cryotherapy. About 43% of patients experienced recurrence after treatment with topical medications.

Furthermore, HIV-positive OSSN patients have also shown high recurrence after surgical excision. Recurrence rates of 3%–43% were noted postsurgery in several studies from Africa, frequently occurring during 6 months of follow-up. High recurrence rates in Africa may be due to late presentation, poor surgery with incomplete or simple excision, and unavailability of adjunctive therapies, such as cryotherapy and chemotherapy, and amniotic membranes. However, a study from India also documented high recurrence of 30% in HIV-positive patients compared to 20% in HIV-negative patients, despite adjunctive cryotherapy following wide surgical excision of OSSN.

The standardized surgical procedure for conjunctival tumors described by Shields et al is fundamental in minimizing recurrences, irrespective of HIV status. Combined treatment with surgical excision and topical IFNα-2b might reduce the risk of recurrence or new tumors in eyes with

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**Immunotherapy in OSSN and related concerns in HIV patients**

IFNα-2b is used in medical treatment of OSSN due to its antiviral and oncostatic effects, as well as its property to activate

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**Figure 2** Clinicopathological correlation of noduloulcerative variant of ocular surface squamous neoplasia in human immunodeficiency virus-infected patients.

**Notes:** (A) A 36-year-old female presented with noduloulcerative lesion involving the nasal quadrant, with scleral thinning, thickening of surrounding conjunctiva with overlying keratin, peripheral corneal opacity, and extension into the anterior orbit. Histopathology of the anterior orbital exenteration specimen revealed (B) invasive squamous-cell carcinoma in the bulbar conjunctiva (hematoxylin and eosin (H&E) stain, magnification 4×), with tumor infiltration into the (C) sclera (H&E stain, magnification 10×). (D) Iris (H&E stain, magnification 10×). (E) Ciliary body (H&E stain, magnification 10×). (F) Choroid (H&E stain, magnification 10×), and (G) anterior orbit (H&E stain, magnification 4×). (H) Fine-needle aspiration biopsy from the preauricular lymph nodes revealed tumor extension into the regional lymph nodes (H&E stain, magnification 10×).
advanced disease. It is not clear whether these interventions have different efficacy in people with HIV infection. However, postoperative topical 5-FU drops have been shown to reduce the recurrence of OSSN in HIV patients.

Moreover, accurate documentation of histopathology reports, including interpretations about the base and margins of the tumor, is mandatory. Tabin et al validated the importance of excision margins at the time of surgery in predicting recurrence. They observed that recurrence was 33% in completely excised tumors, but 56% in incompletely excised lesions with mild–severe dysplasia in surgical margins. They described that in intraepithelial lesions, the most important factor for tumor recurrence was incomplete tumor excision, not histologic depth of the lesion. It has been noted that disease prognosis depends on the grades and types of OSSN, with worse prognosis in mucoepidermoid and spindle-cell carcinoma, but recurrence rates have not been compared with them.

Cases with poor response or tumor recurrence after topical mitomycin C treatment and residual/recurrent tumors after excisional biopsy require appropriate secondary treatment. Protocol for positive margins includes repeated surgical resection or topical IFN-α2b (1 million IU/mL). Despite aggressive treatment, patients with advanced disease or multiple recurrences are prone to develop intraocular invasion, with subsequent need for enucleation or orbital exenteration.

Posttreatment follow-up
As the majority of tumors recur within 6 months of treatment, frequent follow-up is recommended post-operatively at 1 week, 6 weeks, 3 months, and 6 months, and subsequently at 6-monthly intervals till 2 years after treatment. Recurrent lesions have slow growth and malignant potential, and thus lifelong annual follow-up is mandatory for all patients with a history of OSSN. Patient counseling is crucial, with an emphasis on regular follow-up and early presentation to an ophthalmologist for any symptoms in the unaffected eye.

Highly active antiretroviral therapy and OSSN
The effect of highly active ART (HAART) on OSSN is controversial. According to a study by Guech-Ongey et al, HAART does not reduce the incidence of OSSN. However, complete regression of invasive OSSN with HAART alone has been reported in a patient 6 months after initiation of treatment, though anecdotal evidence suggests that in patients who develop the malignancy prior to commencing HAART, HAART does not interrupt tumor growth or recurrence.

With the introduction of HAART, HIV/AIDS has gradually transformed from an acute and fatal disease into a treatable chronic condition with increased survival. This may lead to a possible increase in the disease burden of OSSN, as well as prevalence of bilateral OSSN. Strategies for the long-term management of ophthalmic disorders for HAART responders are important. In developed regions, especially the US and Europe, most patients experience a rise in CD4 lymphocytes during the first few months of HAART. This is followed by reduction in opportunistic infections involving the posterior segment, which frequently occur in advanced HIV/AIDS.

On the other hand, the majority of HIV-infected people have no access to HAART in developing countries, and hence develop other HIV-related ocular complications before reaching levels of immunosuppression so as to cause Cytomegalovirus retinitis. Therefore, anterior-segment involvement by HIV appears to predominate in the HAART era, and thus attention should be focused on these disorders for early diagnosis and treatment.

Association with CD4 counts
OSSN can occur at any time in the disease course of HIV/AIDS. It is hypothesized that altered tumor surveillance on the OS in addition to decreased circulating CD4 T cells can aid in the development of conjunctival and corneal lesions. However, no significant trend has been discovered between CD4 count and grade of OSSN. It is speculated that immunosuppression resulting from HIV plays a role in the etiopathogenesis of OSSN. Therefore, CD4 monitoring is recommended in all OSSN patients affected by HIV. A CD4 lymphocyte count <200 cells/mm3 has been noted in 85%–100% patients at the time of OSSN detection. These cell counts indicate that a majority of HIV-positive patients with OSSN are significantly immunosuppressed at presentation. However, a linear relationship between OSSN presentation and CD4 lymphocyte count or initiation of HAART has not been confirmed. It is suggested that OSSN be considered a criterion to commence AR treatment in HIV-positive patients in African countries with financial and technological restraints when CD4 counts are not accessible.

Serotesting for HIV in OSSN patients
Previously, HIV screening was advised in patients presenting with OSSN at a younger age (<40 years) and with a history of rapid tumor growth (<6 months). The current
recommendation is to conduct HIV screening in all cases presenting with OSSN to rule out undiagnosed HIV infection, especially in those <60 years, atypical conjunctival lesions at presentation, large lesions, bilateral or multifocal tumors, and history of aggressive tumor growth. In an era of increasing global travel, young patients from countries with high HIV prevalence and high ultraviolet B-radiation exposure should be given a low threshold for serotesting and excision biopsy.

Comments and recommendations

Africa is the global epicenter of the HIV/AIDS pandemic, a region with the highest prevalence of HIV in the world and high solar ultraviolet exposure year-round. In the presence of this triad of risk factors, the number of cases of OSSN is expected to stay high. After the introduction of HAART, HIV/AIDS has progressively changed into a nonfatal chronic disease with increased life expectancy. This may lead to a possible increase in the burden of patients living with HIV and OSSN prevalence subsequently in countries with limited resources.

Large unsightly lesions in the eye expose affected persons to the stigma and discrimination associated with HIV infection, and untreated tumors threaten survival. Without an orbital prosthesis, the cosmetic outcome of advanced disease after orbital exenteration is very poor, which leads to facial disfigurement and limited social interaction subsequently. Orbital prostheses and facilities for periocular reconstruction are often unavailable in low-income countries.

Treatment of opportunistic tumors in HIV/AIDS is a very important component in HIV care. HIV/AIDS research should focus on treatment of this tumor. Further understanding of the correlation between HIV and OSSN would be of great significance, because of the potential to reduce the spread of HIV with early diagnosis and to reduce morbidity in already immunosuppressed individuals. It is imperative to improve our understanding of this condition so that we can identify and manage it better as it becomes more common.

Effective evidence-based interventions are needed to allow early diagnosis and treatment. Active search for early manifestations of OSSN, early and complete surgical excision, and close follow-up to address current high recurrence rates is recommended. No randomized controlled trials on the effectiveness of interventions against OSSN in HIV-infected individuals were identified in this review. The majority of studies from the developed world have small samples, due to the rarity of the condition. More studies need to be conducted in regions with high disease prevalence so that results can be extrapolated. There is a need to explore the alternative methods like the use of topical antimetabolites alone or in combination with surgery in early OSSN. Efficacy of new techniques used in the developed world, such as amniotic membrane transplant and photodynamic therapy, need to be assessed.

Practical and inexpensive ways of preventing OSSN in the HIV population should be recognized in view of the gradual rise in incidence of the disease. Interventions to reduce the mutagenic effects of solar ultraviolet B radiation, such as protective sunglasses, and the scope for development of HPV vaccination for oncogenic viruses associated with this disease need to be identified.

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Disclosure

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References


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