Update on pharmacotherapy for ocular surface squamous neoplasia

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Abstract
The most frequently encountered non-pigmented tumor of the ocular surface is ocular surface squamous neoplasia (OSSN). Over the past two decades, the pharmacological management of OSSN has grown, with topical 5-fluorouracil, mitomycin, and interferon alpha 2b all being successfully used to treat this disease. Other agents, such as anti-vascular endothelial growth factor (VEGF), retinoic acid, cidofovir and Aloe vera, have less frequently been used in the treatment of OSSN. This review will discuss these pharmacologic agents, summarizing available data and presenting the approach to the treatment of these tumors.

Keywords: Ocular surface squamous neoplasia (OSSN), Mitomycin-C, Interferon alpha 2b, 5-fluorouracil, Conjunctival neoplasia, Corneal neoplasia

Background
Ocular surface squamous neoplasia (OSSN) encompasses a spectrum of corneal as well as conjunctival lesions ranging in severity from epithelial dysplasia to conjunctival intraepithelial neoplasia in situ to invasive squamous cell carcinoma (SCC). Although OSSN is typically a localized and slow growing tumor, it can rarely invade the sinuses, orbit or the globe [1]. OSSN most often originates from the limbus and lesions are often localized within the interpalpebral fissure, corresponding to areas with sun exposure [2–4]. While the majority of OSSN lesions are non-pigmented, pigmented variants do exist, especially in Africa [5].

Epidemiology
OSSN incidence has been reported to range from 0.13 to 1.9 per 100,000 individuals, and varies with geography, with higher incidences in areas of greater sun exposure (such as Africa) as compared with areas with less sun (such as Denmark) [6,7]. Men in their 6th decade of life are commonly affected, but these lesions can also occur in younger individuals, especially when associated with xeroderma pigmentosum (XP) or human immunodeficiency virus (HIV) [8–11]. Studies in Africa report an increased incidence of OSSN in individuals with HIV compared to non-infected individuals [9,12]. Furthermore, the biology of OSSN is more aggressive in African American individuals with HIV, with more frequent invasion into the orbit than is seen in individuals with HIV in western countries [13].

Risk factors
Risk factors for OSSN include environmental and genetic factors, with the strongest risk factor being exposure to ultraviolet (UV) radiation. As with cutaneous malignancies, UV can damage DNA and lead to the development of cancer promoting mutations [14]. Individuals with XP are at an increased risk of UV-induced cell damage due to an inability to repair mutations in the DNA [10,15,16]. Another possible risk factor is the human papilloma virus (HPV) which has been found more frequently in OSSN specimens compared to healthy conjunctiva [18], although the frequency of detection varies with region (low in countries such as India, Germany and Taiwan but high in cities such as Miami, Florida). HPV-induced carcinogenesis has been attributed to the ability of its oncoproteins, specifically E6 and E7, to target and interact with host cellular proteins, such as p53, and to enhance degradation of normal proteins [19]. Finally, immune-
suppressive and immune-deviated states, such as in HIV and atopy/allergy, have been associated with OSSN, likely due to impairment in tumor recognition by immune cells [9, 15, 20–22].

**Main text**

**Clinical presentation**

OSSN typically presents unilaterally, but it can also be bilateral, especially in individuals with XP, HIV, and atopic states/allergy [23]. OSSN has several classic features, which can be present in isolation or in combination, including leukoplakia, opalescence, papillomatous or gelatinous features, abnormal vascularity (presence of hairpin vessels), and irregular corneal borders [10, 24, 25]. OSSN most frequently originates at the limbus and can extend into the cornea or conjunctiva but can involve either of these structures in isolation [24, 25]. Due to the shared risk factor of UV exposure, OSSN can be associated with other ocular surface lesions such as pterygia and pinguecula, and often these lesions can co-exist [26]. Individuals with OSSN can be asymptomatic and their tumors are often identified identically on a routine eye examination. Others can present with bothersome symptoms, including persistent tearing, redness, grittiness or pain.

**Diagnostic modalities**

The diagnosis of OSSN is usually made clinically. Vital dye staining with rose bengal, toluidine blue, or lissamine green can help determine the diagnosis as well as to outline the borders of the tumors [27–29]. Confirmation by pathology after incisional or excisional biopsy is considered the gold standard for diagnosis. Other less invasive modalities including fine needle aspiration biopsy, impression cytology, or exfoliative cytology have been used to sample tissue [24]. Imaging modalities such as high resolution segment optical coherence tomography (HR-OCT), confocal microscopy and fluorescent microscopy have more recently been reported to aid in the diagnosis of OSSN [24, 30–37]. HR-OCT is the imaging modality most frequently used in our institution for the diagnosis and monitoring of OSSN [31–36, 38, 39]. OSSN has a classic appearance on OCT with three characteristic features: (1) hyperreflective, (2) thickened epithelium with (3) an abrupt transition point between normal and abnormal epithelium (Fig. 1). A study of 34 individuals with OSSN or with pterygium found a 94% sensitivity and a 100% specificity of a ultra-high resolution custom OCT device in differentiating OSSN from pterygium, using an epithelial thickness cut off of approximately 140 μm on OCT images [34]. Similarly, another study of 21 lesions of OSSN or of pterygium found a 100% sensitivity and specificity of the RT Vue (Optovue, Fremont, CA) in differentiating these conditions with an epithelial thickness cut off of 120 μm on OCT images [33]. Other studies have shown that distinctive features on HR-OCT have been able to rule-in and rule-out OSSN in the setting of co-existing ocular surface diseases (such as rosacea, limbal stem cell deficiency, mucous membrane pemphigoid, pterygia, pinguecula or Salzmann’s nodular degeneration) that can make the identification of neoplasia challenging [33, 35]. Advantages of OCT are that images are easily and quickly obtained in a non-contact manner. However, limitations of the technology are that the region of scanning is manually determined by the technician and as such, without automated scanning technology of the entire ocular surface, areas of subclinical disease may not be captured. Furthermore, shadowing can obscure the depth of penetration especially in thick tumors and the tumor invasion cannot be detected [24, 25, 32–36]. HR-OCT can also play a role in the ongoing surveillance of OSSN, especially in identifying residual sub-clinical disease despite apparent clinical resolution.

Confocal microscopy has been used less frequently for the diagnosis and monitoring of OSSN, but has
shown some utility [30, 37]. On confocal microscopy, large areas of superficial cell and keratin debris can be seen with loss of the normal structure of the conjunctival epithelium and corneal basal epithelium, papillomatous organization and large fibrovascular cores. In vivo cyto-morphologic studies using confocal microscopy can help elucidate abnormalities at the cellular level including high nuclear to cytoplasmic ratios, pleomorphism and hyperreflective and indistinctly bordered cytoplasm [37, 40]. Advantages of confocal microscopy include its ability to examine neoplastic tissue in 5 to 20 μm sections as well as at the cytologic level. Disadvantages include the small area of visualized tissue, the lack of cross-sectional visualization, and the inability to effectively illuminate and reflect through keratinized and necrotic neoplastic tissue. Thus, interpretation of confocal images can be more challenging in tumors than when imaging optically transparent healthy corneal and conjunctival tissue [37, 40].

Differential diagnosis
Several other lesions must be considered in the differential diagnosis of a corneal or conjunctival OSSN including pterygium, pinguecula, corneal pannus, Salzmann’s nodular degeneration, pyogenic granuloma, as well as amelanotic melanoma [24].

Treatment
Surgical removal has been considered the gold standard of care for OSSN [41–43]. Surgery has the advantage of serving as both a diagnostic and therapeutic procedure, providing both an accurate histological diagnosis and rapid tumor resolution. However, surgery has potential disadvantages including limbal stem cell deficiency or conjunctival scarring and leaving residual disease that can lead to subsequent tumor recurrence. Given these potential disadvantages, topical chemotherapy has become a popular alternative to surgery [41, 42, 44]. Advantages of topical therapy include treatment of the entire ocular surface which can address areas of subclinical disease, as well as a lower conjunctival scarring risk [45].

Surgical treatment
Surgical removal is still a widely used approach for the treatment of OSSN, especially in lesions that affect < 4 clock hours of limbus [43]. Excision is generally accomplished with wide margins, typically 4 mm, using a no-touch technique. Cryotherapy of the limbus and conjunctival borders in a double freeze/slow thaw manner and absolute alcohol placement on the limbus are usually performed as adjuvant therapies. If pathological evaluation of the specimen reveals tumor cells at the conjunctival margin, additional surgery or postoperative adjunctive chemotherapeutic drugs may be used after surgery, including interferon alpha 2b (IFNα-2b), 5-fluorouracil (5-FU) 1%, and mitomycin C (MMC) 0.02–0.04%. MMC has also been used as an adjuvant agent intraoperatively [15, 46, 47]. Adjuvant therapies are used both pre- and postoperatively as recurrence rates are higher when margins have residual disease (> 50% in some studies [48]), as compared to when margins are negative for tumor (5 to 33% recurrence frequency with follow-up of 15 years [48]). Table 1 outlines all topical chemotherapy drops and associated side effects.

Interferon alpha 2b (IFNα-2b)
Pharmacology
Interferons are leukocyte-derived proteins that can enhance phagocytic and cytotoxic mechanisms, inhibit biosynthetic enzymes, decrease blood vessel proliferation, induce apoptosis and inactivate viral RNA. Interferon α-2b (IFNα-2b) specifically is a cytokine containing 165 amino acid residues with immunomodulatory effects. Intralesional injections of IFNα-2b enhance the production of IL-2 and IFN-γ mRNA by the immune system and lower the production of IL-10. These mechanisms help in the recognition and targeting of neoplastic cells [49].

Table 1 Topical chemotherapy for ocular surface squamous neoplasia (OSSN)

| Treatment options | Formulation | Dosage | Side effects |
|-------------------|-------------|--------|--------------|
| **IFN-α2b**      | Topical: 1 MIU/ml (Alternative: 2–3 MIU/ml), Subconjunctival injections: 3 million IU/0.5 ml (Alternative: 10 MIU/month) | Topical: 4 times a day drops continuously, Subconjunctival: Weekly injections until resolution (typically 4–5 weeks) | Minimal side effects for drops, flu-like malaise with injections |
| **5-FU**         | Topical: 1% drops | 4 times a day for 1 week with 3 weeks off (Alternative: 4 times daily for 2 days to 4 weeks) | Mild pain, lid edema, epitheliopathy |
| **MMC**          | Topical: 0.02–0.04% drops | 4 times a day for 1 week followed by 2–3 weeks off until the eye is quiet. Usually 3–4 cycles until resolution (Alternative: 7–14 day cycles) | Pain, keratopathy, punctal stenosis, LSCD |

*LSCD*= limbal stem cell deficiency; *MIU*= million international units; *LSCD*= limbal stem cell deficiency; *MIU*= million international units; *IFN*-α2b= interferon alpha 2b; *5-FU*= 5-fluorouracil; *MMC*= mitomycin C
Treatment of OSSN

When used to treat OSSN, IFNα-2b for OSSN can be used as topical eye drops, subconjunctival injections, or a combination of both [50–54]. Topical and intraliesional IFNα-2b can be used as primary or adjuvant therapies (Fig. 2a and b).

Dosage

Topically, the most frequently prescribed IFNα-2b dose is 1 million IU/ml. Typically, the medication is used four times a day without interruption until one or two more months after clinical resolution of the lesion. The average time for clinical resolution is about 4 months [10, 25, 50, 53]. Other doses, including 2 and 3 million IU/ml, have also been reported and are used in the same manner. In one study, a similar efficacy was found between the 1 million IU/ml and 3 million IU/ml doses, with no statistically significant differences in clinical resolution and duration of treatment [53]. When used as a post-surgical adjuvant in individuals with positive margins, topical IFNα-2b 1 million IU/ml is administered four times daily for a duration of 2 months post-operatively [3]. Subconjunctival IFNα-2b injections (3 million IU/0.5 ml) are administered weekly until OSSN resolution (typically requiring 4 to 5 injections to achieve clinical resolution) [50, 52, 54]. Higher concentrations have also been reported, the highest being 10 million IU given monthly [25, 54]. Topical eye drops have the advantage of ease of use and a minimal side effect profile. Subconjunctival injections have more side effects than the drops (e.g. flu-like symptoms) but have the benefits of decreased out-of-pocket costs (because the medication is usually covered by insurers), quicker resolution, no need for compounding, and assured compliance [50].

Side effects

When administered topically, IFNα-2b eye drops generally have minimal to no side effects. However, hyperemia and follicular conjunctivitis have infrequently been reported along with formation of epithelial microcysts [55]. The main side-effect associated with subconjunctival injections is a flu-like syndrome that lasts for approximately 48 h after administration of the injection. To lessen the severity of the side effects, 1 g of oral acetaminophen is administered at the time of the injection and then as needed until symptoms resolve [10, 25, 50, 52, 56].

Efficacy and cost

Many studies have looked at the use of topical interferon as primary treatment for OSSN. In a study of 24 eyes using topical IFNα-2b (1 million IU/mL) 4 times daily, 22 patients (92%) had complete resolution with treatment and 2 patients (8%) (2/22) did not respond to treatment. The mean time to resolution was 3.25 months and the group was followed for a mean of 18 months [57]. In another case series of 22 patients (17 on IFNα-2b eye drops (1 million IU/mL), 1 receiving subconjunctival injections (10 million IU/ml), and 4 on a combination of therapy, 18 (82%) of tumors resolved completely with treatment and 4 (18%) showed a partial response [54]. In a study of 98 eyes, topical IFNα-2b (n = 49) was compared to surgical excision (n = 49). Of the 49 eyes receiving chemotherapy, 40 eyes were treated with topical IFNα-2b (1 million IU/ml or 3 million IU/ml), 1 received subconjunctival injections (3 million IU in 0.5 ml), and 8 eyes received both modalities. Recurrence rates were similar between the groups with a 1 year OSSN recurrence rate of 3% in the IFNα-2b group and 5% in the surgery group. In both treatment groups, side effects were mild [58]. Downsides of topical IFNα-2b include the need for refrigeration and the high costs to compound (approximately $600 for a month supply in the USA) [25]. In other countries, IFNα-2b may be quite inexpensive. In the USA, these compounded medications are generally out of pocket costs for the patient. Subconjunctival injections are usually covered by insurers and they are available commercially without the need for compounding.
Alternative uses

IFN-α-2b has been used to treat other types of tumors including cervical intraepithelial neoplasia, actinic keratosis and metastatic malignant melanoma [59, 60]. Interferon has also been used to treat infections such as chronic hepatitis B and C [61, 62]. In the field of ophthalmology, interferons have also been used to treat conjunctival papillomas, lymphoma, and conjunctival Kaposi sarcoma [63–67].

5-fluorouracil (5-FU)

Pharmacology

5-FU is a pyrimidine analog that blocks thymidine synthase, which inhibits DNA formation. This leads to a reduction in RNA, which causes poor cell growth and cell death [68].

Treatment of OSSN

5-FU is a well-tolerated and effective agent in the primary treatment of OSSN as well as an adjuvant after surgical excision [69–71] (Fig. 3a and b).

Dosage

5-FU is compounded as a 1% solution and is typically used in a cyclical pattern four times daily for a week followed by a 3-week holiday [69, 71]. This pattern makes one cycle, which is repeated on average of 4 to 6 cycles based on clinical response (noted by both slit lamp examination and HR-OCT imaging). Alternative regimens have been reported in the literature, including continuous administration of 5-FU 3 to 4 times a day for 4 weeks or administration of 5-FU for only 2 to 4 days with a 30 to 45 day drug-holiday. [45, 47, 70, 72–74]. 5-FU is preferably kept in refrigeration [69].

Side effects

When comparing side effects of topical 5-FU with IFNα-2b, 5-FU has more side effects, most frequently pain and redness at the instillation side, but 5-FU has fewer side effects than MMC (discussed below). Other side effects of 5FU include eyelid swelling, conjunctival congestion, filamentary keratitis and rarely superficial stromal melting [25, 69, 71, 72]. To reduce symptoms, topical corticosteroids are used in conjunction with preservative free artificial tears. Punctal or canalicual stenosis can occur with systemic 5-FU treatment but not with topical 5-FU use [25].

Efficacy and cost

In a study of 44 eyes, 5-FU 1% was used as a primary therapy for the treatment of OSSN, administered 4 times daily for a week followed by 3 weeks of no treatment for a mean of 4 cycles [69]. With this therapy, 36 eyes (82%) had complete tumor resolution, 4 (9%) had partial resolution, and 4 (9%) had no response. In those with a complete resolution, the recurrence rate was 11% (4/36) at 2 years with a mean follow up time of 10 months [69]. In another study of 22 eyes, 5-FU 1% was used four times a day continuously for 4 weeks as primary therapy of OSSN [70] and found that all eyes had complete tumor resolution. However, 3 eyes (14%) experienced a recurrence and were re-treated with additional doses of chemotherapy over a follow up time of 63 to 122 months [70]. A large study compared topical 5-FU 1% to topical IFNα-2b for the primary treatment of OSSN in over 100 patients [71] and found comparable results between the two drugs. 52 of 54 eyes (96%) had complete resolution with 5-FU 1% compared to 39 of 48 eyes (81%) with IFNα-2b [71]. A 1 month treatment of 5-FU costs approximately $38 in the US [25, 71].

Alternative uses

5-FU can be used as treatment for colorectal cancer, actinic keratosis, as well as head and neck solid tumors [75–77]. 5-FU also has others uses in the field of ophthalmology, namely as an adjuvant agent in glaucoma, pterygium, and vitreoretinal surgery. It is also is used in

Fig. 3 Slit-lamp photo of OSSN before and after 5-FU treatment. a. Slit-lamp picture of right eye with a large papillary OSSN located temporally abutting the limbus with extension onto the cornea. b. Slit-lamp picture after 4 cycles of 5-fluorouracil (5-FU) 1% shows the clinical regression of the papillary OSSN. There is a small underlying remaining pterygium.
the post-operative period to revive failing glaucoma blebs [78].

**Mitomycin C (MMC) Pharmacology**

Mitomycin is an alkylating agent with antineoplastic/antibiotic properties. It binds to DNA during DNA synthesis and causes inhibition of its synthesis and function [79]. At higher concentrations, MMC inhibits nucleotide synthesis which interferes with RNA transcription and protein synthesis. MMC is toxic to proliferating and non-proliferating cells by inducing apoptosis and inhibition of fibroblast migration [80, 81].

**Treatment of OSSN**

MMC can be used as a primary treatment agent for OSSN [82–85] but can also be used intraoperatively, as an adjuvant to excisional biopsy, and post-operatively in patients with positive conjunctival or deep margins [82, 86] (Fig. 4a and b).

**Dosage**

When used as a primary therapy for OSSN, MMC is generally compounded into a concentration between 0.02 and 0.04% [25, 82–85, 87] although one group used a lower concentration (0.002%) [88]. MMC is used 4 times daily for 1 week followed by 2 to 3 weeks of no treatment. The length of the holiday depends on how long it takes the eye to recover from the 1 week of treatment. Other regimens include 1 week on/1 week off, 2 weeks on with variable duration between cycles [10, 83, 89], and 3 weeks of continuous therapy [87]. It is not known which of these concentrations and/or regimens is optimal in maximizing efficacy while minimizing toxicity [85]. MMC can also be used intraoperatively during tumor excision, typically soaked on a sponge at a concentration of 0.02% and applied to the conjunctival margins for a duration of 1–3 min [84, 90, 91]. Postoperatively, in patients with positive margins, MMC drops can be used after the surface heals. The dose and protocol during post-operative use is similar to that of primary use [92]. MMC refrigerated at a temperature of (4 °C) [93].

**Side effects**

MMC has more frequent and severe adverse effects when compared to IFNa-2b or 5-FU, including frequent pain and epitheliopathy. In addition, allergic conjunctivitis, hyperemia, ectropion [10, 94], and punctal stenosis have all been described [95]. Due to the risk of punctal stenosis, punctal plugs should be inserted prior to initiating topical MMC [95]. The 2 to 3 week medication holiday in each cycle helps with patient comfort and compliance.

**Efficacy and cost**

A large randomized controlled trial of 48 patients with OSSN compared MMC use as primary therapy to placebo. MMC 0.04% (0.4 mg/ml) was administered four times a day for 3 weeks. In the 26 patients treated with MMC, 24 (92%) had complete clinical resolution and 2 (8%) had no response. In the placebo group, none of the 22 individuals responded to therapy over the 8 week study period [87]. In 18 eyes, MMC 0.02% was used four times a day continuously for 28 days as primary treatment for OSSN with complete resolution in all patients. One recurrence was noted after 24 months [89]. In 20 eyes with OSSN (3 primary lesions and 17 recurrent lesions) treated with MMC 0.02% (n = 6) or 0.04% (n = 14) four times a day for an average of 2 cycles (1 week on/1 week off), the mean time to resolution was 4.5 weeks in the 18 (90%) eyes with complete resolution. Overall, four recurrences were noted over an average follow-up period of 57 weeks, 2 of which were successfully retreated with MMC and 2 with alternative therapies [96]. In 7 eyes, MMC 0.04% was used for primary OSSN, given four times a day in cycles of 1 week on/1 week of until tumor resolution (average 2 cycles) [97]. Six eyes (85%) had complete resolution and 1 partial resolution with a mean follow up time of 9 months [97].
In a study comparing MMC 0.04 mg/ml to IFN 1 million IU/ml, similar rates of clinical resolution were found (92% with MMC with 89% with IFN); however, statistically significant shorter resolution times were noted with MMC (median 1.5 months) as compared with IFN (mean 3.5 months), \( p < 0.0005 \). Unfortunately, patients treated with MMC (88%) experienced significantly more adverse effects compared to those patients treated with IFN (12%), \( p < 0.0005 \) \[98\].

Similarly, in a study that used either 5-FU 1% or MMC 0.04% for the primary treatment of diffuse OSSN, both 5-FU and MMC caused drug-related complications in approximately 58% of treated cases (7 of 12 cases with 5-FU and 23 of 39 cases with MMC). This study also showed that diffuse disease (involving > 5 limbal clock hours or extensive corneal spread) was recalcitrant to only one treatment cycle of either 5-FU or MMC and required additional treatment cycles \[81\]. As seen from these studies, MMC while effective has an unfavorable side effect profile; as such, authors of this paper prefer to use IFN or 5-FU as initial treatment modalities, whenever possible. MMC is often reserved for more recalcitrant cases that have failed prior therapy with alternative agents. MMC costs about $100–190/cycle in the United States but is inexpensive outside of the US.

**Alternative uses**

MMC can be used for the treatment of squamous cell carcinoma of the bladder, gastric, pancreatic and colorectal carcinoma \[99–102\]. MMC has other uses in the field of ophthalmology including as an adjuvant agent in pterygium, glaucoma, refractive, and oculoplastic (dacryocystorhinostomy) procedures \[80\].

**Retinoic acid**

**Pharmacology**

Vitamin A (retinol), and retinoic acid are involved in various cellular processes such as differentiation and growth. Retinoids have potent antineoplastic effects, including regulation of cellular growth by blocking cell maturation and modulating apoptosis \[103\].

**Treatment of OSSN**

In limited studies, retinoic acid has been used alone or in conjunction with IFNα-2b for OSSN treatment \[104, 105\]. Given that most studies used it in combination with IFNα-2b, it is difficult to elucidate the independent effect of retinoic acid on OSSN \[104–107\].

**Dosage**

In one case report, topical retinoic acid 0.01% was applied once every other day for 9 months \[107\]. In another case, topical retinoic acid 0.01% was applied once every other day in conjunction with topical IFNα-2b until clinical resolution and then continued for a duration of up to 12 months \[105\]. Another study reported on retinoic acid ointment 0.05% used 1–3 times a day for 1 to 2 weeks \[104\].

**Side effects**

Retinoic acid has been reported to cause eye lid irritation \[104\], epithelial microcysts and marginal keratitis \[105\]; however, side effects are overall mild.

**Efficacy and cost**

In the largest series to date, retinoic acid 0.01% was used in 89 patients once every other day in combination with topical IFNα-2b 1 million IU/ml drops 4 times daily. Complete clinical resolution was noted in 98% (87/89) and partial response was noted in 2% (2/89). Four tumor recurrences were noted over a mean follow up of 4 years \[105\]. Another study reported two patients who used retinoic acid ointment 0.05%, applied 3 times a day for 10 weeks. Both eyes experienced complete tumor response within 4 weeks \[104\]. Each 5 ml of 0.01% retinoic acid costs approximately $75 USD.

**Alternative uses**

Retinoic acid has been used in the dermatologic field both topically and systemically to treat moderate to severe acne and psoriasis \[108\]. It can be used systemically in acute promyelocytic leukemia \[109\]. In the eye, it has been used to treat ocular consequences of vitamin A deficiency (e.g. keratomalacia) \[110\].

**Aloe Vera**

**Pharmacology**

Aloe plants demonstrate antiproliferative, immunostimulatory and antioxidant properties. Aloe consists of the molecules aloin and aloe emodin which mediate the anti-proliferative activity \[111, 112\]. Studies have shown that aloe emodin can induce apoptosis in vitro \[112\]. Acemannan is a D-isomer mucopolysaccharide in *Aloe vera* with immunostimulatory effects, likely inducing an anti-neoplastic outcome \[111, 112\] .

**Treatment of OSSN**

There has been one report on the use of skin *Aloe vera* drop as a primary treatment modality of OSSN \[112\].

**Dosage**

*Aloe vera* oil drops were used three times daily for 3 months \[112\]. In this case report, a commercially available formulation of *Aloe vera* was used: Aloe Fleur de Jouvence (Forever Living Products; Scottsdale, AZ).
Side effects
No side effects were reported with the topical use of Aloe vera [112].

Efficacy and cost
In this case, Aloe vera drops used three times daily led to a complete clinical resolution after 3 months. No recurrence was noted for 6 years [112]. One bottle of Aloe vera oil costs approximately $25.

Alternative uses
Aloe vera has been used orally and topically to treat different dermatological skin conditions (e.g., burns, frostbite, psoriasis, sunburn), osteoarthritis and other inflammatory processes [113].

Cidofovir
Pharmacology
Cidofovir is a monophosphate nucleotide analogue that demonstrates in vitro activity against a number of DNA viruses [114]. Its clinical activity has been widely demonstrated against cytomegalovirus infection. Since HPV has been implicated in the etiology of OSSN, the anti-viral effects of cidofovir may underlie the efficacy seen in the one published case in OSSN [115].

Treatment of OSSN
One case report demonstrated the use of topical cidofovir as primary treatment for OSSN [115].

Dosage
Topical cidofovir eye drops (2.5 mg/ml) were administered six times a day for 2 weeks, then four times daily for 2 weeks, and then three times daily for 2 weeks to treat diffuse superior tarsal OSSN. The majority of the lesion resolved and the remaining focal area of disease at the nasal edge of the tarsus was surgically excised with adjuvant cryotherapy. No recurrences were noted after 2 years of follow up [115].

Side effects
In this one case, cicatrization of the inferior punctum was reported [115]. Ocular surface irritation, conjunctival erythema, and pain are the main side effects of topical cidofovir, which typically resolve after drop cessation.

Efficacy and cost
Cidofovir is a compounded medication and costs approximately $300–400 USD for one vial (75 mg/ml, 5 ml) in the US.

Alternative uses
Cidofovir has been used as a treatment for various DNA viruses. Topical cidofovir is used to treat herpes viruses, pox viruses, and papilloma viruses of the skin that do not respond to conventional therapies [116]. Cidofovir has also been applied to molluscum contagiosum and condylomata acuminata lesions. Oral cidofovir is used to treat mucosal neoplasms thought to have a viral etiology, such as squamous papilloma of the pharynx and esophagus and Kaposi’s sarcoma [117]. In the eye, cidofovir has been used to treat CMV retinitis through intravitreal injections [117, 118].

Anti-vascular endothelial growth factor (VEGF)
Pharmacology
Anti-VEGF agents are monoclonal antibodies that block the interaction of vascular endothelial growth factor (VEGF) and its receptor, interfering with the growth of blood vessels [119]. Bevacizumab is a complete anti-VEGF antibody while ranibizumab is a fragment of the antibody [119].

Treatment of OSSN
There have been a few case reports on the use of anti-VEGF agents as primary therapy in OSSN and as adjuvants after surgical excision [120–123]. Therefore, their role in treating OSSN remains uncertain.

Dosage
Dosages of anti-VEGF therapy vary between studies. As primary therapy in OSSN, topical bevacizumab (5 mg/ml) was given 4 times a day for up to 8 weeks [120] and subconjunctival/perilesional bevacizumab (2.5 mg in 0.1 ml) was given weekly for 2 weeks [121]. Lower concentrations of subconjunctival bevacizumab (1.25 mg in 0.05 ml) have also been reported [122]. In another study, subconjunctival ranibizumab (0.05 mg of 10 mg/ml) was injected once or twice a month for a period ranging from 6 to 25 months [123].

Side effects
No side effects have been reported with the use of anti-VEGF agents in OSSN. However, bevacizumab may have inhibitory effects on corneal healing, especially when used in patients with epithelial defects [120, 124].

Efficacy and cost
A few studies have looked at the role of anti-VEGF agents in the primary treatment of OSSN and as an adjuvant to surgery. In one study, ten patients with primary OSSN diagnosed by impression cytology received 2 perilesional/subconjunctival injections of bevacizumab (2.5 mg/0.1 ml), one week apart [121]. Of these ten patients, two eyes with conjunctival OSSN had complete
tumor resolution while eight eyes with corneal OSSN had no significant change to the lesion. In another study of six eyes, topical bevacizumab (5 mg/ml) was given 4 times daily as eye drops for a period of 8 weeks [120]. Two patients (34%) had complete OSSN resolution while 4 (66%) had partial resolution and underwent excisional biopsy. No recurrences were noted after 6 months of follow up [120]. Another case series of five patients were treated with perilesional ranibizumab (0.05 mg of 10 mg/ml) once or twice a month for an average of 19 months (ranging from 6 to 25 months) [123]. Three (60%) had complete resolution and 2 (40%) had partial tumor resolution [123]. Intralesional bevacizumab (1.25 mg/0.05 ml) was also used in a patient with OSSN that failed treatment with MMC, 5-FU and IFNα-2b eye drops [122]. No change in OSSN was noted after the intrallesional injection of bevacizumab [122].

**Alternative uses**

Bevacizumab is approved by the FDA for the treatment of colorectal cancer [125]. In ophthalmology, it is used off-label in age-related choroidal neovascularization, proliferative and non-proliferative diabetic retinopathy, neovascular glaucoma, and anterior segment neovascularization [126–128]. Similarly, ranibizumab is FDA approved for the treatment of patients with wet age-related macular degeneration, macular edema following retinal vein occlusion, diabetic macular edema, diabetic retinopathy, and myopic choroidal neovascularization.

**Conclusion**

In summary, great advances in the management of OSSN have occurred with the development of numerous topical therapeutics, and the ability to diagnose and follow the disease with HR-OCT. Fortunately, OSSN is a neoplastic disorder amenable to treatment with topical chemotherapeutic agents. The most commonly used agents in OSSN are IFNα-2b, 5-FU, and MMC, with robust evidence in the literature on their efficacy; less information is available on other agents such as cidofovir, anti-VEGF and retinoic acid. In the authors’ clinical practice, 5-FU is used as first line therapy due to great efficacy, mild side effects and affordable costs. In the rare occasions that 5-FU does not work and causes side effects due to a fragile ocular surface, IFNα-2b is a great alternative due to a similar efficacy profile and milder side effects. Its use is often limited by cost and access to the drug. Interestingly, IFNα-2b may have decreased efficacy compared with 5-FU in patients with underlying immunosuppression [129]. In our center, MMC is considered a second line agent for OSSN when the others fail because of its unfavorable side effect profile, notably corneal toxicity, limbal stem cell deficiency, and punctal stenosis. Surgery is still an excellent option in patients with small lesions (< 4 clock hours) or with rapidly growing lesions and allows for rapid resolution of disease. Topical chemotherapy however, is preferred in large and multifocal lesions, allowing for complete treatment of the ocular surface and a decreased risk for subsequent limbal stem cell deficiency. With topical treatment, the duration is longer and requires compliance. Overall, eye care providers need to consider patient preferences, cost, and access to care when deciding on a treatment approach; however, the pendulum is swinging in favor of the use of topical drugs in the management of ocular surface squamous neoplasia.

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**Authors’ contributions**

GAB was involved in the scientific literature research, writing and editing of the manuscript. DA was involved in the scientific literature research, writing and editing of the manuscript. NV was involved the writing and editing of the manuscript. AG was involved in the conception, design, and editing of the manuscript. CLK was involved in the conception, design, and editing of the manuscript. All authors read and approved the final manuscript.

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**Competing interests**

The authors declare that they have no competing interests.

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**References**

1. Santoni A, Thariat J, Maschi C, Heraud J, Baillif S, Lassalle S, et al. Management of invasive squamous cell carcinomas of the conjunctiva. Am J Ophthalmol. 2019;2001–9.
2. Murillo JC, Galor A, Wu MC, Kye NK, Wong J, Ahmed IO, et al. Intraocular and intraocular invasion of ocular surface squamous neoplasia after intraocular surgery: report of two cases and review of the literature. Ocul Oncol Pathol. 2017;3(1):66–72.
3. Kao AA, Galor A, Karp CL, Abdelaziz A, Feuer WJ, Dubovy SR. Clinicopathologic correlation of ocular surface squamous neoplasms at Bascom Palmer Eye Institute: 2001 to 2010. Ophthalmology. 2012;119(9):1773–6.
4. Pe'er J. Ocular surface squamous neoplasia. Ophthalmol Clin North Am. 2005;18(1):1–13, vi.
5. Shields CL, Manchandia A, Subbiah R, Eagle RC Jr, Shields JA. Pigmented squamous cell carcinoma in situ of the conjunctiva in 5 cases. Ophthalmology. 2008;115(10):1673–8.
6. Lee GA, Hirst LW. Ocular surface squamous neoplasia. Surv Ophthalmol. 1995;39(6):429–50.
7. Shields CL, Chien JL, Surakiatchanukul T, Soufik K, Lally SE, Shields JA. Conjunctival tumors: review of clinical features, risks, biomarkers, and
83. Shields CL, Naseripour M, Shields JA. Topical mitomycin C for extensive, squamous neoplasia of the conjunctiva. J Dermatol Surg Oncol. 1996;16(5):446–9.

82. Worschler WP. Considerations for use of fluorouracil cream 0.5% for the treatment of conjunctival squamous neoplasia. Cornea. 2005;24(4):443–8.

81. Bayliss PL, Everette CM, Martinis WC, Lowen MS, Barros JN. Long-term results of topical mitomycin C 0.02% for primary and recurrent conjunctival-squamous neoplasia. Ophthalmol Plast Reconstr Surg. 2009;25(4):296–9.

80. Siganos CS, Kozobolis VP, Christodoulakis EV. The intraoperative use of mitomycin-C in excision of ocular surface neoplasia with or without limbal autograft transplantation. Cornea. 2002;21(1):12–6.

79. Kemp EG, Hamnett AN, Chatterjee S, Preoperative topical and intraoperative mitomycin C adjuvant therapy in the management of ocular surface neoplasias. Br J Ophthalmol. 2002;86(1):31–4.

78. Chen C, Louis D, Dodd T, Muecke J, Mitomycin C as an adjuvant in the treatment of squamous cell carcinoma of the conjunctiva. Br J Ophthalmol. 2004;88(1):17–8.

77. Quebbemann EJ, Hoffman NE, Asuman RK, Hamid AA. Stability of mitomycin C admixtures. Am J Hosp Pharm. 1985;42(8):1750–4.

76. Khong JJ, Muecke J. Complications of mitomycin C therapy in 100 eyes with ocular surface neoplasia. Br J Ophthalmol. 2006;90(7):819–22.

75. Billings K, Karagianis A, Selva D. Punctal-canalicular stenosis associated with mitomycin-C for corneal epithelial dysplasia. Am J Ophthalmol. 2003;136(4):746–7.

74. Yousef YA, Finger PT. Squamous carcinoma and dysplasia of the conjunctiva and cornea: a review of an 101 cases. Ophthalmology. 2012;119(2):233–40.

73. Yeatts PR, Engelbrecht NE, Curry CD, Ford JG, Walter KA. 5-fluorouracil for the treatment of conjunctival intraepithelial neoplasia. Cornea. 2003;22(1):116–11.

72. Yeatts PR, Engelbrecht NE, Curry CD, Ford JG, Walter KA. 5-fluorouracil for the treatment of conjunctival intraepithelial neoplasia. Cornea. 2003;22(1):116–11.

71. Venkateswaran N, Mercado C, Galor A, Karp CL. Comparison of topical 5-fluorouracil and 5-fluorouracil plus interferon alfa-2b for the treatment of conjunctival squamous neoplasia: a long-term safety study. Br J Ophthalmol. 2011;95(3):355–9.

70. Schechter BA, Schrier A, Nagler RS, Smith EF, Velasquez GE. Regression of presumed primary conjunctival and corneal intraepithelial neoplasia with topical interferon alpha-2b. Cornea. 2002;21(1):6–10.

69. Joag MG, Sise A, Murillo JC, Sayed-Ahmed IO, Wong JR, Mercado C, et al. Regression of presumed primary conjunctival and corneal intraepithelial neoplasia with topical interferon alpha-2b. Cornea. 2002;21(1):6–11.

68. Longley DB, Harkin DP, Johnston PG. S-Fluorouracil: mechanisms of action and clinical strategies. Nat Rev Cancer. 2003;3(5):330–8.

67. Schechter BA, Schrier A, Nagler RS, Smith EF, Velasquez GE. Regression of presumed primary conjunctival and corneal intraepithelial neoplasia with topical interferon alpha-2b. Cornea. 2002;21(1):6–11.

66. Edwards L, Levine N, Smiles KA. The effect of topical interferon alpha 2b on immune cell profile in head and neck squamous cell carcinoma: a phase II trial. Am J Hosp Pharm. 1985;42(8):1750–4.

65. Venkateswaran N, Mercado C, Galor A, Karp CL. Comparison of topical 5-fluorouracil and 5-fluorouracil plus interferon alfa-2b for the treatment of conjunctival squamous neoplasia: a long-term safety study. Br J Ophthalmol. 2011;95(3):355–9.

64. Kitano S. Clinical trials of interferon in ophthalmology. Gan To Kagaku Ryoho. 1994;1(2):212–20.

63. Giaccone G, Pinto HM, Mitomycin C: mechanism of action, usefulness and limitations. Anti-Cancer Drugs. 1990;1(5):13–6.

62. Singh P. Mitomycin-C use in ophthalmology. IOSR J Pharmacy. 2013;3(1):12–4.

61. Rudkin AK, Dempster L, Muecke JS. Management of diffuse ocular surface squamous neoplasia: efficacy and complications of topical chemotherapy. Clin Exp Ophthalmol. 2015;43(1):20–5.

60. Edwards L, Levine N, Smiles KA. The effect of topical interferon alpha 2b on immune cell profile in head and neck squamous cell carcinoma: a phase II trial. Am J Hosp Pharm. 1985;42(8):1750–4.

59. Bracarda S, Eggemort AM, Samuelsson J. Redefining the role of interferon in the treatment of malignant diseases. Eur J Cancer. 2010;46(2):284–97.
110. Sommer A. Effects of vitamin A deficiency on the ocular surface. Ophthalmology. 1983;90(6):592–600.
111. Harlev E, Nevo E, Lansky EP, Offir R, Bishayee A. Anticancer potential of aloes: antioxidant, antiproliferative, and immunostimulatory attributes. Planta Med. 2012;78(9):843–52.
112. Damani NR, Shah AR, Karp CL, Orlin SE. Treatment of ocular surface squamous neoplasia with topical Aloe vera drops. Cornea. 2015;34(1):87–9.
113. Davis RH, Donato JJ, Hartman GM, Haas RC. Anti-inflammatory and wound healing activity of a growth substance in Aloe vera. J Am Podiatr Med Assoc. 1994;4(2):77–81.
114. Zabawski EJ Jr, Coderre CJ. Topical and intralungal cidofovir: a review of pharmacology and therapeutic effects. J Am Acad Dermatol. 1998;39(5 Pt 1):741–5.
115. Sherman MD, Feldman KA, Farahmand SM, Margolis TP. Treatment of conjunctival squamous cell carcinoma with topical cidofovir. Am J Ophthalmol. 2002;134(3):432–3.
116. Toro JR, Sanchez S, Turiansky G, Blauvelt A. Topical cidofovir for the treatment of dermatologic conditions: verruca, condyloma, intraepithelial neoplasia, herpes simplex and its potential use in smallpox. Dermatol Clin. 2003;21(2):301–9.
117. Vossen MG, Gattringer KB, Jäger W, Kralf S, Thalhammer F. Single-dose pharmacokinetics of cidofovir in continuous venovenous hemofiltration. Antimicrob Agents Chemother. 2013;58(4):1592–5.
118. Rahhal FM, Arevalo JF, Munigua D, Taskintuna I, Chavez de la Paz E, Azen SP, et al. Intravitreal cidofovir for the maintenance treatment of cytomegalovirus retinitis. Ophthalmology. 1996;103(7):1078–83.
119. Kim LA, D’Amore PA. A brief history of anti-VEGF for the treatment of ocular angiogenesis. Am J Pathol. 2012;181(2):376–9.
120. Asera L, Dursun Atkınoğlu S. Topical bevacizumab for the treatment of ocular surface squamous neoplasia. J Ocul Pharmacol Ther. 2015;31(8):847–90.
121. Faramarzi A, Feizi S. Subconjunctival bevacizumab injection for ocular surface squamous neoplasia. Cornea. 2013;32(7):998–1001.
122. Paul S, Stone DU. Intralunal bevacizumab use for invasive ocular surface squamous neoplasia. J Ocul Pharmacol Ther. 2012;28(6):647–9.
123. Finger PT, Chin KJ. Refractory squamous cell carcinoma of the conjunctiva treated with subconjunctival ranibizumab (Lucentis): a two-year study. Ophthalmic Plast Reconstr Surg. 2011;28(2):85–9.
124. Koenig Y, Bock F, Horn F, Kruse F, Straub K, Cursiefen C. Short- and long-term safety profile and efficacy of topical bevacizumab (Avastin) eye drops against corneal neovascularization. Graefes Arch Clin Exp Ophthalmol. 2009;247(10):1375–82.
125. Hunwitz H, Saini S. Bevacizumab in the treatment of metastatic colorectal cancer: safety profile and management of adverse events. Semin Oncol. 2006;33(Suppl 5):S26–34.
126. Rosenfeld PJ, Brown DM, Heier JS, Boyer DS, Kaiser PK, Chung CY, et al. Ranibizumab for neovascular age-related macular degeneration. N Engl J Med. 2006;355(14):1419–31.
127. Sinawat S, Rattanapakorn T, Sanguansak T, Yospaiboon Y, Sinawat S. Intravitreal bevacizumab for proliferative diabetic retinopathy with new dense vitreous hemorrhage after full panretinal photocoagulation. Eye (Lond). 2013;27(12):1391–6.
128. Goyal S, Lavallee M, Subramanian ML. Meta-analysis and review on the effect of bevacizumab in diabetic macular edema. Graefes Arch Clin Exp Ophthalmol. 2011;249(1):15–27.
129. Ashkenazy N, Karp CL, Wang G, Acosta CM, Galor A. Immunosuppression as a possible risk factor for interferon nonresponse in ocular surface squamous neoplasia. Cornea. 2017;36(4):506–10.