VOLUME 18 NO 12 PP 1424-1443 DECEMBER 2013

Systematic Review

Epidemiology of ocular surface squamous neoplasia in Africa

Stephen Gichuhi^{1,2}, Mandeep S. Sagoo^{3,4}, Helen A. Weiss² and Matthew J. Burton^{2,3}

- 1 Department of Ophthalmology, University of Nairobi, Nairobi, Kenya
- 2 London School of Hygiene and Tropical Medicine, London, UK
- 3 Moorfields Eye Hospital, London, UK
- 4 UCL Institute of Ophthalmology, University College London, UK

Abstract

OBJECTIVES To describe the epidemiology and an aetiological model of ocular surface squamous neoplasia (OSSN) in Africa.

METHODS Systematic and non-systematic review methods were used. Incidence was obtained from the International Agency for Research on Cancer. We searched PubMed, EMBASE, Web of Science and the reference lists of articles retrieved. Meta-analyses were conducted using a fixed-effects model for HIV and cigarette smoking and random effects for human papilloma virus (HPV). RESULTS The incidence of OSSN is highest in the Southern Hemisphere (16° South), with the highest age-standardised rate (ASR) reported from Zimbabwe (3.4 and 3.0 cases/year/100 000 population for males and females, respectively). The mean ASR worldwide is 0.18 and 0.08 cases/year/100 000 among males and females, respectively. The risk increases with exposure to direct daylight (2–4 h, OR = 1.7, 95% CI: 1.2–2.4 and \geq 5 h OR = 1.8, 95% CI: 1.1–3.1) and outdoor occupations (OR = 1.7, 95% CI: 1.1–2.6). Meta-analysis also shows a strong association with HIV (6 studies: OR = 6.17, 95% CI: 4.83–7.89) and HPV (7 studies: OR = 2.64, 95% CI: 1.27–5.49) but not cigarette smoking (2 studies: OR = 1.40, 95% CI: 0.94–2.09). The effect of atopy, xeroderma pigmentosa and vitamin A deficiency is unclear.

CONCLUSIONS Africa has the highest incidence of OSSN in the world, where males and females are equally affected, unlike other continents where male disease predominates. African women probably have increased risk due to their higher prevalence of HIV and HPV infections. As the survival of HIV-infected people increases, and given no evidence that anti-retroviral therapy (ART) reduces the risk of OSSN, the incidence of OSSN may increase in coming years.

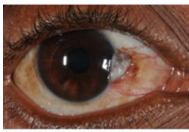
keywords ocular surface squamous neoplasia, conjunctival intraepithelial neoplasia, conjunctival intraepithelial dysplasia, ocular surface epithelial dysplasia, conjunctival squamous cell carcinoma, risk factors, incidence

Introduction

Ocular surface squamous neoplasia (OSSN) is the most common ocular surface tumour (Grossniklaus *et al.* 1987). Other synonymous terms include 'conjunctival epithelial neoplasia', 'ocular surface epithelial dysplasia' and 'conjunctival squamous cell neoplasia' (Lee & Hirst 1992; McDonnell *et al.* 1992; Tulvatana 2003). OSSN covers a spectrum of disease ranging from non-invasive intra-epithelial dysplasia of the conjunctiva and cornea (CCIN) to invasive squamous cell carcinoma (Lee & Hirst 1995).

Clinical features

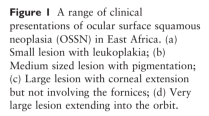
The disease may present with irritation, red eye, raised gelatinous mass and leucoplakia (Tunc *et al.* 1999). In Africans, it is often pigmented brown (Figure 1). OSSN is usually unilateral (Chisi *et al.* 2006) and arises at the limbus – the junction between the cornea and conjunctiva (Lee & Hirst 1997). Most lesions occur within the exposed part of the eyeball between the lids (Ateenyi-Agaba 1995; McKelvie 2002; Waddell *et al.* 2006). Up to 31.2% of cases seen are recurrent lesions (Chisi *et al.* 2006). Late stages present with a large fungating oculo-orbital mass (Ogun *et al.* 2009). Early lesions resemble



(a) Small lesion with leuloplakia



(b) Medium sized lesion with pigmentation





(c) Large lesion with corneal extension but not involving the fornices



(d) Very large lesion extending into the orbit

Table I Histopathological classification of ocular surface squamous neoplasia (OSSN), Basti & Macsai (2003) and American Joint Committee on Cancer (2010)

Benign

Squamous papilloma

Pseudoepitheliomatous hyperplasia

Benign hereditary intraepithelial dyskeratosis

Pre-invasive

Conjunctival intraepithelial neoplasia (CIN)

CIN I (mild dysplasia) – confined to the basal third of the conjunctival epithelium

CIN II (moderate dysplasia) – extends into the middle third of the conjunctival epithelium

CIN III (severe dysplasia) – extends into the superficial third of the conjunctival epithelium

CIS (carcinoma-in-situ) - full thickness dysplasia*

Invasive

Squamous cell carcinoma

GX - grade cannot be defined

G1 – Well differentiated

G2 - Moderately differentiated

G3 - Poorly differentiated

G4 – undifferentiated

Mucoepidermoid carcinoma

benign growths such as pterygia and pingueculae. OSSN can be the first manifestation of HIV infection in about 50% of cases in HIV-endemic settings (Porges & Groisman 2003; Spitzer *et al.* 2008).

Histopathology

Histologically, OSSN may be classified into 3 forms: benign, pre-invasive and invasive (Table 1; Basti & Macsai 2003). The term OSSN usually excludes the benign forms. The term 'invasive' indicates infiltration through the basement membrane of the conjunctival epithelium into the underlying stroma (Basti & Macsai 2003; Shields & Shields 2004).

Epidemiology overview

Two disease patterns of OSSN are recognised: older, predominantly male in temperate climates, not associated with HIV or human papilloma virus (HPV); and younger men and women, in tropical climates, associated with HIV and HPV. The latter represents a public health challenge in Africa in relation to the HIV pandemic and late presentation of large tumours (Ukponmwan et al. 2002: Chisi et al. 2006; Ogun et al. 2009), diagnostic difficulties (Furahini & Lewallen 2010), malignant transformation and high recurrence rates after treatment (1-year recurrence of 16.6% reported in Tanzania; Makupa et al. 2012). Experienced surgeons report lower recurrences (3.2%) after excision (Waddell et al. 2006). Trial data to guide management in this context are lacking (Gichuhi & Irlam 2013). For the temperate pattern of disease, one randomised controlled crossover trial in Australia compared mitomycin-C with placebo in participants

^{*}The American Joint Committee on Cancer (AJCC) staging manual 2010 classifies CIS under CIN.

whose average age was 67 years (Hirst 2007). There was a significant treatment effect on clinically assessed complete resolution of lesions (P = 0.0005), but no effect on histologically assessed complete resolution (P = 0.49).

Incidence rates and geographical variation

Incidence estimates for OSSN are difficult to ascertain and vary regionally (Table 2). The first paper to examine this used cancer registry data from International Agency for Research on Cancer (IARC; Newton et al. 1996). A subset of these data were used in a subsequent publication looking at variation in incidence across the USA (Emmanuel et al. 2012). However, published results need to be interpreted with caution - firstly, all eye cancers are classified together by the International Classification of Diseases for Oncology (ICD-O-3 C.69) while other databases classify squamous cell carcinoma of the conjunctiva (SCCC) with head and neck cancers (Lee et al. 2000; Curado et al. 2007; Parkin et al. 2010). OSSN is not recognised as a separate entity. Squamous cell carcinomas that are site-coded for the eye (C69) probably include some cancers that originate in the eyelid skin (WHO 2000, 2010; Curado et al. 2007). Secondly, the availability of histopathology services to confirm OSSN diagnosis is often limited in low- and middleincome countries (Furahini & Lewallen 2010). Thirdly, health information systems tend to capture invasive squamous cell carcinoma (SCC) but not earlier stages. Countries reporting higher rates of SCC (mostly in Africa) only started sending cancer registry data to IARC in the mid-1980s (Curado et al. 2007). Completeness of the current IARC database is hampered in that only data from 80 countries were submitted, of which 75% was of acceptable quality, and not all countries had data on

squamous cell carcinoma in the eye under code C69. Africa had the lowest level of acceptable quality of data (36%). Fourthly, crude incidence rates can be influenced by population structure, a problem often addressed by reporting age-standardised incidence rates. Finally, in areas with limited health facilities for cancer treatment where a large number of patients are treated outside the reference area, incidence may be underestimated. Moreover, in defining incidence from different sources, it may be difficult to distinguish between recurrence or extension of an existing cancer on one hand and the development of a new primary on the other. Analysis of incidence time trends is also difficult if geographical coverage, ICD revisions and disease definitions in a registry change.

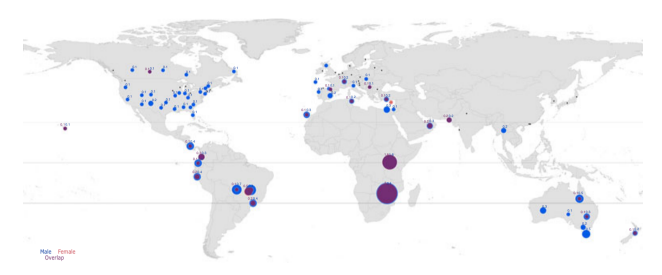
Methods for this review

Systematic and non-systematic review methods were used. No a priori systematic review protocol had been published. Incidence data were obtained from the current IARC report (9th Volume) covering the period 1998-2002. The IARC collates data from cancer registries worldwide. The report uses ICD codes to show the age-standardised incidence per 100 000 population stratified by sex and histological type. Under code C.69 where eye cancers are reported, the four main groups are retinoblastoma, malignant melanoma, carcinomas (11.4% of all eye cancers), sarcoma and other unspecified tumours. Under carcinomas, there are three subgroups - SCC (principally tumours of the conjunctiva and cornea, comprising 70% of the carcinoma subgroup), other specified carcinoma (adenocarcinomas of the lacrimal gland and lacrimal duct) and unspecified carcinomas. We extracted data from the SCC subgroup.

Table 2 Age-standardized incidence rates of squamous cell carcinoma in the eye (ICD-O-3 C.69) by continent for the period 1998–2002 (Curado *et al.* 2007)

	Age-standardized incidence rate			
Region	Males mean (95% CI)	Females mean (95% CI)	P-value	
Africa	1.38 (-1.00 to 3.75)	1.18 (-1.08 to 3.43)	0.853	
Central & South America	0.48 (0.33 to 0.62)	0.21 (0.10 to 0.33)	0.005	
Oceania	0.28 (0.14 to 0.41)	0.05 (0.01 to 0.10)	0.002	
North America	0.08 (0.06 to 0.10)	0.00 (0.00 to 0.01)	< 0.001	
Asia	0.08 (0.01 to 0.14)	0.05 (0.00 to 0.09)	0.416	
Europe	0.05 (0.02 to 0.08)	0.01 (0.00 to 0.03)	0.033	
Southern Hemisphere	0.61 (0.14 to 1.09)	0.33 (-0.12 to 0.78)	0.355	
Northern Hemisphere	0.10 (0.06 to 0.14)	0.05 (0.00 to 0.08)	0.045	
Worldwide estimate	0.18 (0.09 to 0.26)	0.08 (0.01 to 0.15)	0.091	

CI = confidence interval.



KEY: Dot size is directly proportional to incidence. Males are shown in blue and females in red. Overlaps between males and females appear purple in colour

Figure 2 Worldwide mapping of the age-standardized incidence rates (ASR) of squamous cell carcinoma of the eye (ICD-O-3 C.69) for the period 1998–2002 (Curado *et al.* 2007). Key: Dot size is directly proportional to incidence. Males are shown in blue and females in red. Overlaps between males and females appear purple in colour.

The coordinates locating each registry were obtained from http://itouchmap.com/latlong.html.

We searched PubMed, EMBASE and Web of Science for systematic reviews, meta-analysis and case—control studies using 'OSSN', 'conjunctival squamous cell carcinoma', 'risk factors' and their synonyms as key words with no language restrictions. Abstracts were assessed and studies were selected if they reported analysis of known or suspected risk factors. The search was conducted on 2 January 2013 and updated on 31 May 2013. Data were extracted from the full texts of articles and additional articles obtained from their reference lists. Meta-analyses were conducted where appropriate. A fixed-effects model was used for HIV and cigarette smoking. A random-effects model was chosen for HPV after investigation of heterogeneity.

Results and discussion

Africa has the highest age-standardised incidence rate of ocular SCC followed by Central and South America then Oceania (Australia, New Zealand and Hawaii), respectively (Table 2 and Figure 2). The rate in Africa is about 9–10 times higher than in Europe and North America. The highest incidence rate is 3.4 cases/year/100 000 among males and 3.0 cases/year/100 000 among females in Zimbabwe (Curado *et al.* 2007). Uganda follows with 1.6 cases/year/100 000 for males and females. Australia

comes third with 0.3–0.5 cases/year/100 000 in parts of that country. Other countries have rates between 0 and 0.1 cases/year/100 000. The rates have a right-skewed bell-shaped distribution peaking at latitude 16° South (Figure 3). Incidence rates are higher in the Southern Hemisphere than the Northern Hemisphere, with male ASR = 0.61 cases/year/100 000 (95% CI: 0.14–1.09) and female ASR = 0.33 (95% CI: –0.12 to 0.78) in the Southern Hemisphere, compared with male ASR = 0.10 (95% CI: 0.06–0.14) and female ASR = 0.05 (95% CI: 0.00–0.08) in the Northern Hemisphere.

The high rates in Africa are consistent with other estimates from the region. A Tanzanian study estimated the incidence of suspected OSSN from 2006 to 2008 using operating theatre records across the country. Although there was no histological confirmation in all cases, the incidence was found to be 2.2 cases/year/100 000 (Furahini & Lewallen 2010). Uganda reported a peak incidence of 3.5 cases/year/100 000 in 1992 (Ateenyi-Agaba 1995). More recent data from the Kampala Cancer Registry also show a marked increase, although it is reported as ocular cancer, rather than specifically as OSSN (Wabinga *et al.* 2000).

Cancer registry data in two African countries show that OSSN has become more prevalent with time. In Zimbabwe, the age-adjusted annual incidence rates of SCCC underwent a more than 10-fold increase from 0.17 to 1.8/100 000 between 1990 and 1999 (Masanganise *et al.* 2008) while the prevalence of OSSN among ocular

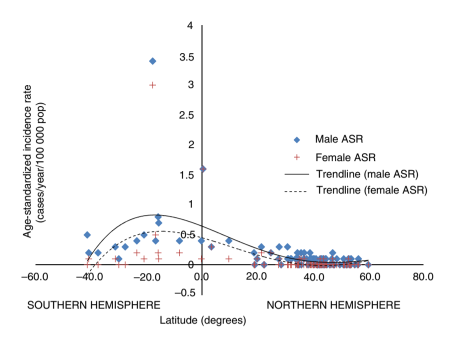


Figure 3 The age-standardized incidence rates (ASR) of squamous cell carcinoma of the eye (ICD-O-3 C.69) for the period 1998–2002 (Curado *et al.* 2007).

Table 3 The proportion of orbital exenterations performed due to ocular squamous cell carcinoma in different regions of the world

Year (ref.)	Country	No. of exenterations (N)	No. due to SCCC (n)	Proportion (n/N) (%)
2011 (Ackuaku-Dogbe 2011)	Ghana	25	19	76
2001 (Masanganise & Magava 2001)	Zimbabwe	23	13	57
2007 (Nemet et al. 2007)	Australia	38	12	32
2004 (Pushker et al. 2004)	India	26	3	15
2008 (Croce et al. 2008)	Italy*	6	1	13
2005 (Rahman et al. 2005)	UK†	69	6	9

^{*}Included children.

surface tumour biopsy specimens increased from 33% in 1996 to 58% by 2000 (Pola *et al.* 2003).

OSSN is the most common indication for orbital exenteration performed in adults in Africa (Table 3; Pola *et al* 2003). This surgical procedure to excise all the orbital tissue including stripping the periosteum from the orbital walls is performed in cases with advanced disease. More than half (≥57%) the exenterations performed in Africa are for OSSN compared with 32% in Australia and 9–15% in Europe and India. Although available data does not clearly distinguish those performed for primary eyelid disease from conjunctival disease, SCC still emerges as an important cause in Africa. Eyelid SCC is uncommon in Africa (Templeton 1967, 1973).

Incidence of OSSN by age and sex

In temperate countries, OSSN remains a rare, slow-growing tumour of elderly males (70-80% are males

with a mean age of about 60 years; Lee & Hirst 1997; Tunc *et al.* 1999). In contrast, in tropical countries, particularly in Eastern and Southern Africa, the prevalence is highest among young people in their 30s and among women (50–70%; Table 4; Poole 1999; Pola *et al.* 2003; Chisi *et al.* 2006; Furahini & Lewallen 2010). Within East Africa, the pattern of SCCC in the 1960s differed to that seen today. In 1967, the average age of affected patients was 48 years, and males were four times more frequently affected than females (Templeton 1967).

Worldwide, IARC data show that the overall incidence is higher in males than females but the difference is not statistically significant (Figure 3 and Table 2). The mean male ASR worldwide is 0.18 cases/year/100 000 (95% CI: 0.09-0.26) and 0.08 (95% CI: 0.01-0.15) among females (P = 0.09). Incidence is significantly higher in males than females except in Africa and Asia where both

[†]Mainly elderly patients.

Table 4 The age and	sex of patients affected	by ocular surface squamous	neoplasia (OSSN)
Table 4 The age and	Sex of patients affected	. Dy ocuiai surface squamous	neodiasia (Ossin)

Year (ref.)	Country	Mean age (years)	Male (%)	Female (%)	Male:Female ratio
1995 (Ateenyi-Agaba 1995)	Uganda	33	52	48	1:2.3
2008 (Spitzer et al. 2008)	Malawi	33	42	58	1:2.1
2010 (Simbiri et al. 2010)	Botswana	39	39	61	1:1.6
2003 (Pola et al. 2003)	Zimbabwe	35	30	70	1:1.4
2002 (Mahomed & Chetty 2002)	S. Africa	37	50	50	1:1.3
2006 (Chisi et al. 2006)	Kenya	38	50	50	1:1
2012 (Makupa <i>et al.</i> 2012)	Tanzania	39	32	68	1:1
2009 (Ogun et al. 2009)	Nigeria	54	43	57	1:0.9
1999 (Tunc et al. 1999)	USA	64	70	30	1:0.4
2002 (McKelvie 2002)	Australia	69	77	23	1:0.3

sexes are equally affected (Table 2). Prevalence in Africa is higher in females than males (Table 4). This may be related to Africa having the highest prevalence of both HIV and HPV, which may increase the risk of OSSN in women and gender differences in mortality of HIVinfected adults. In South Africa, HIV-infected females have a longer life expectancy than HIV-infected males (Cornell et al. 2012; Johnson et al. 2013; Maskew et al. 2013). Men present in later stages of HIV/AIDS for antiretroviral therapy (ART) and possibly have poorer adherence to ART (Taylor-Smith et al. 2010). This has also been observed in Latin America, China and Lao (Dou et al. 2011; Gonzalez et al. 2011; Bastard et al. 2013). In Europe, the response to ART and mortality is similar for both sexes (Perez-Molina et al. 2012; Thorsteinsson et al. 2012).

Variation in disease severity

There may be variation in disease stage at presentation, with more advanced disease present at time of surgery in East Africa, compared with other regions (Table 5; Chisi et al. 2006; Waddell et al. 2010; Kao et al. 2012; Makupa et al. 2012). This may reflect delayed presentation to ophthalmic services in this region, leading to more advanced pathology by the time of surgery. Histopathological reporting is also subjective, and pathologists may not always grade tumours the same way (Margo et al. 2002). Alternatively, the disease may be intrinsically more aggressive in the East African region or HIV worsens disease progression.

Risk factors

Various factors are thought to influence the causation of OSSN, but it is not clear how they interact or which is the most potent. The rising incidence of OSSN in recent decades may be driven by increased prevalence of these

factors. We found no systematic reviews of risk factors for OSSN after the literature search. Of the case–control studies found, two in Uganda and Australia examined the association with solar exposure; six in Africa examined the association with HIV; sixteen examined the association with HPV; seven in Africa, five in Asia, one in Brazil, two in USA and one in Australia. Two studies examined cigarette smoking in Uganda.

Ultraviolet solar radiation. Several cutaneous malignancies, including melanoma and SCC, have a strong association with solar radiation. It was first noted in the 1960s that SCCC was relatively common in East Africa, and this apparent excess risk was attributed to higher exposure to sunlight (Templeton 1967). There is a strong relationship between the incidence of SCCC and increasing Ultraviolet (UV) levels (Newton et al. 1996). Using IARC data and published measurements of ambient solar ultraviolet light, the incidence of SCCC was found to reduce by 49% for every 10° increase in latitude from 1.2 cases/ year/100 000 (Table 7) in Uganda (latitude 0.3°) to <0.02/year/100~000 in the UK (latitude $> 50^\circ$). More recently, the National Institutes of Health/American Association of Retired Persons (NIH-AARP) Diet and Health Study in the USA found a slightly lower risk of SCCC in those who lived >35° compared with ≤35° from the equator, although this was not statistically significant (adjusted Hazard Ratio = 0.92, 95% CI: 0.49-1.71; Emmanuel et al. 2012). The USA has comparatively lower HIV prevalence, solar irradiance and incidence of OSSN than Africa, which is bisected by the equator. The high incidence of ocular SCC near the equator may be related to high solar irradiance (the amount of solar radiant energy incident on a surface per unit area and per unit time) in the world (World Energy Council 2007).

A case-control study in Uganda adjusted for age, sex, residential district, and HIV serostatus demonstrated that the risk of OSSN was higher with increasing time spent

Table 5 Stages	of ocular surface so	uamous neoplasia (OSSN)) seen at presentation in	Africa and USA
----------------	----------------------	-------------------------	---------------------------	----------------

	Stage of O	SSN, n (%)					
Country year (ref.)	Mild dysplasia (CIN I)	Moderate dysplasia (CIN II)	Severe dysplasia (CIN III)	Carcinoma in situ (CIS)	Well differentiated SCC	Moderately differentiated SCC	Poorly differentiated SCC
Kenya 2006 (Chisi et al. 2006)	7 (21.9)				1 (3.1)	9 (28.1)	15 (46.9)
Uganda 2008 (de Koning et al. 2008)	17 (21.0)	18 (22.2)	22 (27.2)	0 (0)	24 (29.6)		
Uganda 2010	39 (29.3)				94 (70.7)		
(Ateenyi-Agaba et al. 2010)							
Uganda 2010 (Waddell et al. 2010)	48 (15.1)	66 (20.8)	81 (25.5)	0 (0)	123 (38.7)		
Tanzania 2012 (Makupa et al. 2012)	28 (21.2)		73 (55.3)	0 (0)	31 (23.5)		
Malawi 2013 (Tiong et al. 2013)	1 (2.0)	5 (10.2)	9 (18.4)	17 (34.7)	17 (34.7)		
USA 2012 (Kao et al. 2012)	48 (8.1)	98 (16.4)	59 (9.9)	322 (54.0)	69 (11.6)		

in daylight (Waddell *et al.* 2010). Compared with those who reported spending up to 1 h a day in direct sunlight, the odds ratio (OR) for those who spent 2–4 h was 1.7 (95% CI: 1.2–2.4), and for those who spent 5 or more hours a day, it was 1.8 (95% CI: 1.1–3.1). A case–control study in Australia reported that the strongest risk factor was a past history of skin cancer (OR = 15, 95% CI: 2.0–113.6), although other factors, including outdoor activity, pale skin and irides and propensity to burn, were also important (Lee *et al.* 1994).

More direct evidence for UV radiation induced damage in the pathophysiology of SCCC was described in another case—control study in Uganda in which 52% of the cases had mutations in the tumour suppressor gene TP53 compared with 14% of controls (Ateenyi-Agaba *et al.* 2004a). The mutations were mainly of the CC TT type, consistent with UV-induced mutagenesis. This gene also downregulates the replication of HPV type 16 via the viral E2 protein, suggesting that its mutation may allow replication of HPV particles (Brown *et al.* 2008). Further, exposure to UV radiation is associated with altered expression of matrix metalloproteinases (MMPs) and the tissue inhibitors of these metalloproteinases (TIMPs), molecules that may be responsible for tissue invasion and metastasis of tumours (Ng *et al.* 2008).

In addition, OSSN lesions occur more often at the limbus. A study in Uganda demonstrated that tumours almost always occur in sun-exposed areas of the eye (Waddell *et al.* 2006). It is thought that the human eye is more exposed laterally, making this a large collecting zone of peripheral sunlight, which, depending on the incident angle and radius of curvature of the cornea, is focused on the limbus, lens and lid margin, which are the main foci of sun-related eye diseases such as pterygium, OSSN, cataract and lid malignancies (Maloof *et al.* 1994). Low doses of ambient sunlight received on every

day exposure inhibit immunity in the skin and internal organs (Halliday et al. 2012).

HIV. There is strong evidence that HIV is a major risk factor for OSSN. Uganda, which had a cancer registry since 1951, was the first country to report a dramatic increase in the annual incidence of SCCC shortly after the outbreak of HIV/AIDS. There was a sixfold increase from 0.6 cases/year/100 000 between 1970 and 1988 to 3.5/ year/100 000 by 1992 (Figure 4; Ateenyi-Agaba 1995). A marked rise was also observed in the USA with the onset of the HIV pandemic (Guech-Ongey et al. 2008). At the same time, a US study observed a strong association in an HIV-infected cohort (OR = 13.0, 95% CI: 4-34; Goedert & Cote 1995). In Tanzania, regional incidence rates were significantly correlated with regional HIV prevalence (Pearson's r = 0.53, P = 0.03; Furahini & Lewallen 2010). The majority of patients (60–77%) with OSSN seen in Africa are HIV-infected (Table 6). A meta-analysis of 6 case-control studies (Table 7) in Uganda, Rwanda and Zimbabwe shows a strong association with HIV infection (pooled OR = 6.17, 95% CI: 4.83–7.89; Figure 5).

The association with HIV suggests that immunosuppression plays a role in OSSN; however, a linear association between the CD4 lymphocyte count and OSSN has not been confirmed. A cross-sectional study conducted in Tanzania found a median CD4 cell count of 71 cells/ μ l among HIV-infected individuals with OSSN (Makupa *et al.* 2012). HIV-infected cases tended to have larger lesions: 71% had lesions >5 mm in diameter *vs.* 27% among HIV-negative individuals (OR = 3.13, 95% CI: 1.5–6.5). HIV-infected cases were also more likely to develop recurrent tumours within a year of excision (82% *vs.* 18%; OR = 3.54, 95% CI: 1.12–11.2). However, there was no significant trend found between CD4 count and the grade of OSSN (P = 0.94). In a Ugandan study,

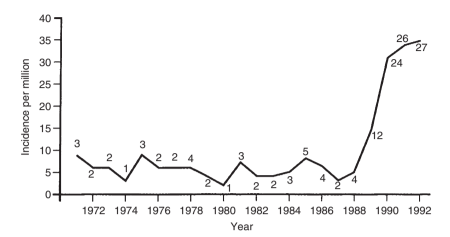


Figure 4 Sudden rise in the annual incidence rates of conjunctival SCCC in Kampala with the onset of the HIV pandemic – number of cases shown (Ateenyi-Agaba 1995).

Table 6 Prevalence of HIV infection in cases of squamous cell carcinoma of the conjunctiva in Africa

Year (ref.)	Country	Study period	HIV prevalence in SCCC cases (%)
2012 (Makupa <i>et al.</i> 2012)	Tanzania	2005–2008	60
2011 (Osahon et al. 2011)	Nigeria	1999–2009	75
2002 (Mahomed & Chetty 2002)	South Africa	1995–1997	71
1995 (Ateenyi-Agaba 1995)	Uganda	1990–1991	75
1996 (Waddell et al. 1996)	Uganda	1993-1994	71
2003 (Porges & Groisman 2003)	Zimbabwe	1993–1995	91
2001 (Newton et al. 2001)	Uganda	1994–1998	77

among 112 HIV-infected cases of CIN and invasive SCC, the median CD4 count at diagnosis was 111 cells/ μ L (IQR; 62–221; Waddell *et al.* 2006). Excess risks standardised incidence ratio (SIR = 19.5, 95% CI: 6.3–45.5) have also been observed among a cohort of kidney transplant recipients in Australia suggesting that immune suppression from other causes may play a role (Vajdic *et al.* 2007).

HAART does not reduce the incidence of SCCC according to data from the US HIV/AIDS Cancer Match (HACM) Study (Guech-Ongey *et al.* 2008) which compared SIRs in the pre-HAART and HAART eras among 491 048 adults aged \geq 15 years with HIV/AIDS diagnosed from 1980 to 2004. The SIRs here estimate the excess risk of SCCC attributable to HIV/AIDS compared with a population with negligible HIV/AIDS prevalence and were similar at 12.0 (95% CI: 5.5–22.8) and 12.6 (95% CI: 4.6–27.4) in the pre- and post-HAART eras, respectively (P = 0.79). There is, however, a case report of ART causing tumour regression in an otherwise inoperable case (Holkar *et al.* 2005).

Human papilloma virus. The relationship between human papilloma virus (HPV) and OSSN is rather controversial with variable results. (Tulvatana 2003;

Moubayed et al. 2004; Sen et al. 2007; de Koning et al. 2008; Guthoff et al. 2009; Simbiri et al. 2010; Yu et al. 2010). A review of 12 case series and 17 case-control studies concluded that there was no causal association between mucosal HPV types and OSSN while the role of cutaneous types was uncertain (de Koning et al. 2008). The studies included used different methods for testing of HPV (including PCR and serology), and different HPV types were examined. Conversely, a random-effects meta-analysis of various case-control studies shows that OSSN is associated with HPV infection in sub-Saharan Africa (pooled OR = 2.64, 95% CI: 1.27-5.49) and worldwide (pooled OR = 4.00, 95% CI: 2.11-7.57; Figure 6). The prevalence of HPV in OSSN ranges from 0% to 100% depending on geographical region with subtypes HPV18 and HPV16 being the most common (Table 8; di Girolamo 2012). Most African studies report prevalence of 75-85% (Ateenyi-Agaba et al. 2004b; Simbiri et al. 2010; Yu et al. 2010). HPV is more commonly isolated in OSSN than pterygium - on average, considering studies from different regions of the world, 33.8% of OSSN lesions and 18.6% of pterygia are HPV positive (di Girolamo 2012). There may be a true geographical variation in the prevalence of HPV in OSSN.

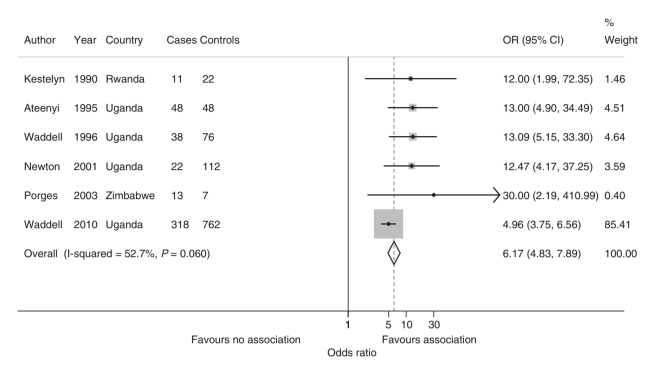


Figure 5 Meta-analysis of case-control studies of HIV infection in ocular surface squamous neoplasia (OSSN) in Africa (fixed effect).

Differences in HPV prevalence in OSSN may be influenced by patient selection, sample handling in the operating theatre, preparation, storage, overseas shipping and the detection method. Variations may also be due to different testing methodology and the specific HPV types tested for. Most existing molecular diagnostic tests applied in OSSN testing for HPV were developed for cervical tissue testing. The sensitivity and specificity of various polymerase chain reaction (PCR) tests varies and may be influenced by various factors including the PCR design (nested, broad spectrum or type-specific), size of amplified product and choice of polymerase used (Munoz et al. 2012; Mesher et al. 2013). Detection of E6/E7 mRNA transcripts by quantitative reverse transcriptase-PCR (qRT-PCR) has been proposed as the gold standard for HPV testing (Smeets et al. 2007). However, RNA is unstable limiting this test to fresh frozen tissue (Kim et al. 2013). Testing for HPV DNA by PCR from paraffin-embedded archived tumour blocks may be complicated by contamination between samples at the time of initial tissue sectioning for DNA harvest (Boyd et al. 1996; Iftner & Villa 2003).

Generally, only a limited subset of HPV types has been investigated among OSSN cases. There are 170 genotypes of HPV described to date, which are broadly subdivided into cutaneous and mucosal types (de Villiers 2013).

There are conflicting reports on which of these two are more commonly associated with OSSN. One study conducted in Uganda reported that among OSSN cases, the prevalence of mucosal types was higher than cutaneous types (38% vs. 22%) while from another study in the same population, the prevalence of cutaneous types was higher than mucosal types (43.6% vs. 6.8%; Table 8; de Koning et al. 2008; Ateenyi-Agaba et al. 2010). Multiple HPV types have been found in individual patients with OSSN tumours. One Ugandan study reported multiple HPV types in 57.1% of SCCC and 75% of dysplasia cases by PCR (Ateenyi-Agaba et al. 2010). In Botswana, multiple HPV types were identified in all OSSN and all pterygium specimens by DNA sequencing (Simbiri et al. 2010). The HPV types found by sequencing ranged from 4 to 21 types per sample. The same study also described co-infection with multiple other viral types per individual in 17 of 18 (94%) histologically proven OSSN specimens by PCR; 83% were positive for Epstein-Barr virus (EBV), 72% were HPV positive, 67% were Kaposi's sarcoma-associated herpesvirus (KSHV) positive, 67% were herpes simplex virus (HSV-1/2) positive and 56% were cytomegalovirus (CMV) positive. All the pterygium specimens from that study similarly had multiple viruses; 75% were positive for each of EBV, KSHV, CMV and HSV while 50% were

Table 7 Characteristics of case–control studies included in the meta-analysis of HIV as a risk factor of ocular surface squamous neoplasia (OSSN)

Study period (ref.), Country	Cases	Controls
1989–1990 (Kestelyn <i>et al.</i> 1990), Rwanda	11 patients with clinical evidence of conjunctival dysplasia or malignancy seen at Centre Hospitalier de Kigali	22 controls. 2 controls per case from the same area matched for age and sex within 5 years. Referrals from elsewhere were excluded
1990–1991 (Ateenyi-Agaba 1995), Uganda	48 patients with conjunctival growths who presented to the eye clinic at Mulago Hospital, Kampala	48 patients matched for age and sex attending the same eye clinic with other eye diseases
1993–1994 (Waddell <i>et al.</i> 1996), Uganda	38 patients in seven countrywide eye clinics including New Mulago Hospital, Kampala who had suspicious conjunctival lesions had excision biopsy of the lesion	76 controls. 2 controls per case matched for age and sex. 16 Controls were patients in the eye clinic without neoplasia or clinical features of HIV disease; the remainder were general (non-eye clinic) anonymous outpatients at the same health units
1993–1995 (Porges & Groisman 2003), Zimbabwe 1994–1998 (Newton <i>et al.</i> 2001),	13 cases from patients who underwent excisional biopsy for conjunctival lesions at Bindura Provincial Hospital (Mashonaland Central, Zimbabwe) 22 cases. Patients aged >15 years with a provisional diagnosis of cancer from all wards and out-patient	7 controls. Patients were from the same group as cases but had benign lesions on histology 112 controls. 93 patients with tumours not suspected to be of infectious
Uganda	clinics of the 4 main hospitals in Kampala: Mulago, Nsambya, Mengo and Rubaga	aetiology and 19 with non-malignant conditions
2001–2005 (Waddell <i>et al.</i> 2010), Uganda	318 cases recruited from country-wide ophthalmology clinics in Uganda. Anyone with a suspected OSSN was offered surgical treatment and histology, together with enrolment into a case-control study	762 controls were recruited from 2 sources. The first group comprised patients attending the ophthalmology clinics with concerns or conditions other than OSSN. This group also included those individuals who were originally recruited as cases, but where histology subsequently revealed another diagnosis. The second group comprised people who were recruited through the voluntary HIV counselling and testing (VCT) service

HPV positive. The proportion of HPV infection in this series was much higher than any other studies in the region have reported raising the question whether this could be due to the methodology used.

The mechanism by which HPV is associated with OSSN is unknown. HPV is associated with causation of metaplasia in squamocolumnar epithelial transition zones such as the corneoscleral limbus and eyelid skin of the eye, the cervix and anus where there is active cell turnover and continuous cell division to replace desquamated cells (Chow *et al.* 2010). HPV also promotes degradation of the p53 gene (Scheffner *et al.* 1990).

The epidemiology of OSSN is closely related to that of cervical cancer with respect to high incidence in Africa and the association with HIV and HPV mainly types 18 and 16 (Sun *et al.* 1997; Clifford *et al.* 2003; Stanley 2010). A meta-analysis of HPV prevalence reports worldwide shows that Africa has the highest adjusted preva-

lence (22.1%; 95% CI: 20.9–23.4%) among women with cytologically normal cervical pap smears using PCR-based or high-risk Hybrid Capture 2 (HC-2) technology to detect HPV DNA (de Sanjose *et al.* 2007). Whether vaccination against HPV may help to reduce the incidence of OSSN remains to be seen (Hughes *et al.* 2008).

Occupation. Outdoor occupations have been associated with OSSN, probably related to UV solar radiation exposure. In Uganda, those with outdoor occupations had an OR of 1.7 (95% CI: 1.1–2.6) compared to those with indoor occupations (Waddell *et al.* 2010). Another in Uganda reported that 74% of 133 patients with SCCC or dysplasia had outdoor occupations (Ateenyi-Agaba *et al.* 2010). In Japan, exposure to petroleum products was also described as a risk factor for conjunctival intraepithelial neoplasia (synonym of OSSN) in a

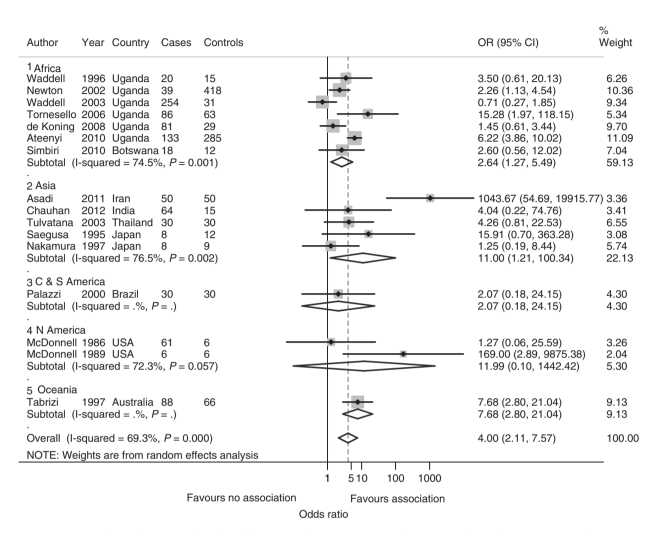


Figure 6 Meta-analysis of case-control studies of human papilloma virus (HPV) infection in ocular surface squamous neoplasia (OSSN) (random effects).

small age–sex-matched case–control study (Napora *et al.* 1990). Exposure to smoke from burning wood in the kitchen was described as a risk factor for cervical cancer among HPV-infected women in Honduras (Velema *et al.* 2002).

Cigarette smoking. Cigarette smoking is implicated in other squamous cell cancers (Haverkos 2004). There is, however, evidence of no effect from smoking on OSSN in Africa. In Uganda, two case–control studies showed that current smokers were not at a significantly higher risk for OSSN than non-smokers (Waddell *et al.* 2010; Ateenyi-Agaba *et al.* 2010; pooled OR = 1.40; 95% CI: 0.94–2.09; Figure 7). In a Nigerian series of 37 SCCC cases, only two patients (5.4%) had a history of cigarette smok-

ing (Ogun *et al.* 2009) while in a series from Australia, 5 of 11 cases of SCCC (45%) were smokers (McKelvie 2002).

Allergy. There is little evidence that allergic conjunctivitis is a risk factor. Among 215 SCCC cases in Tanzania, 1.9% had allergic conjunctivitis (Poole 1999). In Rwanda, allergic conjunctivitis was found in 4% of children and was responsible for 3–6% of hospital visits of all ages (de Smedt *et al.* 2013). In a case—control study in Uganda, none of the cases of OSSN had a history of allergic eye disease (Waddell *et al.* 2010). However, a case series of SCCC from Germany reported that 6/10 cases had atopic eczema, so this may be of more importance in temperate climates (Heinz *et al.* 2003).

Table 8 Studies on the prevalence and subtypes of human papilloma virus (HPV) in ocular surface squamous neoplasia (OSSN)

shipped to the Netherlands Fresh frozen tissue shipped paraffin-embedded tissue paraffin-embedded tissue paraffin-embedded tissue paraffin-embedded tissue paraffin-embedded tissue paraffin-embedded tissue tissue transport medium Plasma shipped in dry ice Serum shipped in dry ice Fresh tissue shipped in Fresh frozen biopsies shipped overseas Formalin-fixed Formalin-fixed Formalin-fixed Formalin-fixed Formalin-fixed Fresh tissue Tissue used to France to France to France to USA 6.4% muc, 44.7% cut 7.7% muc, 41% cut 20, CJ198, indeterm 18, 38, 100, DL473, 50% gen, 28% cut 27% gen, 23% cut 42% gen, 13% cut 35% gen, 29% cut 6, 11, 16, 18, 31, 6, 11, 16, 18, 33, HPV subtypes found 14, 27, 37, 38 14, 20,CJ198 21 subtypes* Nil detected PPHLIFR 16, 18, 45 18, 100 58, 72 16, 18 16, 18 16, 18 16, 18 16 91 prevalence HPV (% 98 7 00 72 61 15 36 47 56 45 22 45 41 31 33 13 00 000 08 00 98 50 sequencing antibodies antibodies Diagnostic anti-HPV anti-HPV PCR/IHC PCR/DB method DNA IHC PCR PCR PCR PCR PCR [SH Sample size 30 254 39 17 18 22 24 24 39 39 118 23 10 3 v ∞ 6 45 ∞ 21 CIN I-III Dysplasia Dysplasia Dysplasia included CIS/SCC CIN III CIN III CIN II CIN II CIN I CINI OSSN OSSN OSSN OSSN OSSN SCC SCC SCC SCC SCC SCC Saudi Arabia Botswana Country Uganda Uganda Uganda Uganda Taiwan Uganda Uganda Japan USAUSA USA USA USA 2010 2002 2010 2006 1991 1992 Year 2004 2003 2008 2002 1990 6661 2006 1997 1997 McDonnell (McDonnell et al. 1992) Nakamura (Nakamura et al. 1997) Karcioglu (Karcioglu & Issa 1997) Waddell (Waddell et al. 2003) (Ateenyi-Agaba et al. 2004a) Newton (Newton et al. 2002) Dushku (Dushku et al. 1999) (Ateenyi-Agaba et al. 2010) Simbiri (Simbiri et al. 2010) Odrich (Odrich et al. 1991) Lauer (Lauer et al. 1990) Scott (Scott et al. 2002) Tornesello (Tornesello de Koning (de Koning Kuo (Kuo et al. 2006) Lead author (ref.) North America Ateenyi-Agaba Ateenyi-Agaba et al. 2006) et al. 2008)

(continued)

supplemented with 7 fresh paraffin-embedded tissue Freshly prepared Formalin-fixed Formalin-fixed, formalin-fixed Formalin-fixed Formalin-fixed Formalin-fixed Formalin-fixed Formalin-fixed Formalin-fixed Formalin-fixed Tissue used tissues 6, 11, 13, 16, 18 HPV subtypes Nil detected Nil detected Nil detected Nil detected Nil detected Nil detected 16, 18 16 16 16 18 prevalence HPV (%) 0 0 0 0 1760 0 38 25 39 4 PCR/ISH-CARD Diagnostic PCR/ISH PCR/IHC PCR/DB method PCR PCR IHC PCR PCR PCR Sample ∞ 16 30 4 size 48 20 30 12 11 31 88 Dysplasia included CIS/SCC Disease OSSN OSSN OSSN OSSN OSSN OSSN OSSN SCCSCC SCC Saudi Arabia Hungary Thailand Australia Germany Germany Country Poland Taiwan Finland Japan India India 2000 2000 Year 1995 2009 2002 2003 2007 1997 2008 2005 1992 Reszec(Reszec & Sulkowski 2005) Guthoff (Guthoff et al. 2009) Saegusa (Saegusa et al. 1995) (Auw-Haedrich et al. 2006) Fulvatana (Tulvatana 2003) Fabrizi (Tabrizi et al. 1997) (Tuppurainen et al. 1992) Manderwad (Manderwad Toth (Toth et al. 2000) Foth (Toth et al. 2000) Eng (Eng et al. 2002) Sen (Sen et al. 2007) Lead author (ref.) Auw-Haedrich Tuppurainen et al. 2009) Oceania Europe

*The 21 subtypes were HPV types 1, 3, 7, 11, 13, 16, 18, 29, 39, 40, 43, 45, 59, 61, 68, 70, 77, 85, 89, 91, 97. ? - means unclear or not mentioned.

Table 8 (Continued)

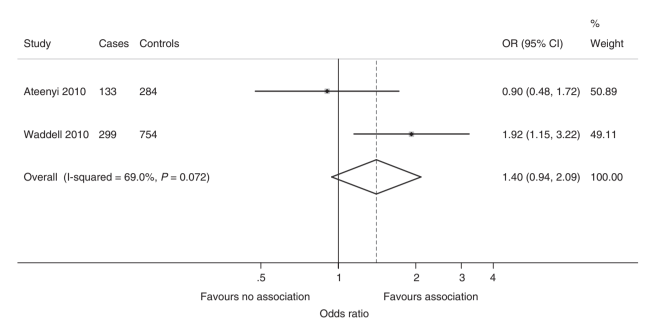


Figure 7 Meta-analysis of case-control studies in Uganda on cigarette smoking and ocular surface squamous neoplasia (OSSN) in Africa (fixed effect).

Xeroderma pigmentosum. Xeroderma pigmentosum (XP), a rare, inherited skin disease characterised by high sensitivity to UV damage is associated with a high prevalence (40%) of specific mutations of the TP53 tumour suppressor gene (Dumaz et al. 1993). Over a 25-year period in Zimbabwe, in a series of 12 cases, 2 had SCCC while the rest had SCC of the skin, lip or tongue (Chidzonga et al. 2009). From a series of 7 XP cases in India, 6 of the 14 eyes (42.9%) had invasive SCC and eight eyes (57.1%) had CIN (Gupta et al. 2011). A larger series of 32 cases in France found that 59% of them had ocular and periocular malignancies (Touzri et al. 2008).

Vitamin A deficiency. The importance of vitamin A in maintaining the health of the ocular surface is well known, but the role of vitamin A deficiency in OSSN has not been established. Deficiency of vitamin A induces keratinisation of the ocular surface (Beitch 1970; Pfister & Renner 1978). Keratinisation is commonly observed as leucoplakia in OSSN lesions (Figure 1). There is a synergistic interaction between vitamin A and zinc in maintenance of the corneal and conjunctival epithelium (Kanazawa *et al.* 2002). In South Africa, it was shown that 54% of HIV-infected adults are deficient in vitamin A (plasma retinol <1.05 μm) and 33% deficient in zinc (<10.7 μm; Visser *et al.* 2003). In Ethiopia, 53% of HIV-infected adults were deficient in vitamin A (Fufa

et al. 2009). As most patients with OSSN are also HIV-infected, it is plausible that vitamin A deficiency contributes to the aetiology.

Other risk factors. There is limited evidence of a role for exposure to dust, ocular trauma and pre-existing benign conjunctival lesions such as pterygia and pingueculae (Templeton 1967; Margo & Groden 1986; Waddell *et al.* 2010).

Protective factors. One of the Ugandan case–control studies found that some factors are associated with a lower risk for SCCC such as higher personal income (adjusted OR = 0.4, 95% CI: 0.3–0.7) and decreasing age at leaving home (P = 0.05), perhaps reflecting less exposure to sunlight consequent to rural-to-urban migration (Newton *et al.* 2002).

Aetiological model of OSSN

Various models have been proposed to simultaneously address the role of two or more risk factors in cancer causation within hierarchical levels (Victora *et al.* 1997). Most such models focus on social and environmental hypothesis but do not incorporate biological factors. A recently proposed framework called Multi-level Biological And Social Integrative Construct (MBASIC) includes

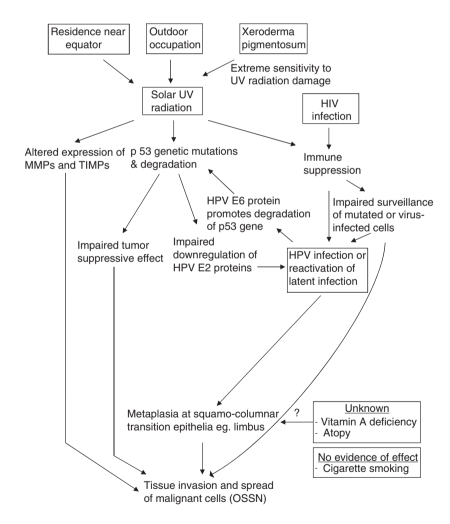
biological factors together with macro-environmental and individual level factors (Lynch & Rebbeck 2013). Using the existing evidence reviewed in this article, we propose an aetiological model that might explain how the risk factors discussed may be involved development of OSSN (Figure 8).

Conclusions

OSSN is a disease of increasing importance in Africa. A triad of ultraviolet solar radiation, HIV and HPV form the major risk factors and this may explain the high incidence rates in Africa. There is evidence from case—control studies that exposure to UV radiation, outdoor occupations – perhaps due to exposure to sunlight, HIV and

HPV infection are associated with a higher risk for OSSN. These studies also show no evidence of effect of cigarette smoking. Dust, ocular trauma and pre-existing benign conjunctival tumours may play a role. Although mentioned in the literature, the effect of atopy and xero-derma pigmentosa is unclear. The effect of vitamin A deficiency has not been examined in case—control studies.

The highest incidence of OSSN is found in Africa, where males and females are equally affected, unlike other continents where male disease predominates. This probably reflects that African women have increased risk due to their higher prevalence of HIV and HPV infections. As people with HIV are living longer, and given no evidence that ART reduces risk of OSSN, one could expect incidence of OSSN to increase in Africa in coming years.



MMPs - matrix metalloproteinases

TIMPs - tissue inhibitors of metalloproteinases

Figure 8 An aetiological model illustrating how ocular surface squamous neoplasia (OSSN) might develop. MMPs, matrix metalloproteinases; TIMPs, tissue inhibitors of metalloproteinases.

Currently, the best available options for OSSN control remain early detection and effective treatment. However, there are no early non-invasive diagnostic methods in use and no trial evidence to guide treatment. OSSN is currently largely neglected by both eye and HIV care programmes. Eye care programmes prioritise preventable blindness while OSSN often in early stages does not affect vision. OSSN may, however, lead to facial disfigurement and death in late stages. In Africa, a key research question is whether scale-up of ART and HPV vaccination will impact on OSSN.

Acknowledgements

SG received funding from the British Council for Prevention of Blindness (BCPB) to conduct this study. MJB is supported by The Wellcome Trust (Grant Number 098481/Z/12/Z). We acknowledge Benjamin D. Hennig from the University of Oxford (http://www.view softheworld.net) for help with preparing the incidence map (Figure 2).

References

- Ackuaku-Dogbe E (2011) Review of orbital exenterations in Korle-Bu teaching hospital. *Ghana Medical Journal* **45**, 45–49. American Joint Committee on Cancer 2010. Carcinoma of the conjunctiva. In: *Cancer Staging Handbook*, 7th edn (eds SB Edge, DR Byrd, CC Compton, AG Fritz, FL Greene & A
- Ateenyi-Agaba C (1995) Conjunctival squamous-cell carcinoma associated with HIV infection in Kampala, Uganda. *Lancet* **345**, 695–696.

Trotti) Springer, New York, 597.

- Ateenyi-Agaba C, Dai M, le Calvez F *et al.* (2004a) TP53 mutations in squamous-cell carcinomas of the conjunctiva: evidence for UV-induced mutagenesis. *Mutagenesis* **19**, 399–401.
- Ateenyi-Agaba C, Weiderpass E, Smet A *et al.* (2004b) Epidermodysplasia verruciformis human papillomavirus types and carcinoma of the conjunctiva: a pilot study. *British Journal of Cancer* **90**, 1777–1779.
- Ateenyi-Agaba C, Franceschi S, Wabwire-Mangen F et al. (2010) Human papillomavirus infection and squamous cell carcinoma of the conjunctiva. British Journal of Cancer, 102, 262–267.
- Auw-Haedrich C, Sundmacher R, Freudenberg N et al. (2006) Expression of p63 in conjunctival intraepithelial neoplasia and squamous cell carcinoma. Graefes Archive for Clinical and Experimental Ophthalmology 244, 96–103.
- Bastard M, Soulinphumy K, Phimmasone P et al. (2013) Women experience a better long- term immune recovery and a better survival on HAART in Lao People's Democratic Republic. BMC Infectious Diseases 13, 27.
- Basti S & Macsai MS (2003) Ocular surface squamous neoplasia: a review. *Cornea* **22**, 687–704.

- Beitch I (1970) The induction of keratinization in the corneal epithelium. A comparison of the "dry" and vitamin A-deficient eyes. *Investigative Ophthalmology* 9, 827–843.
- Boyd AS, Annarella M, Rapini RP, Adler-Storthz K & Duvic M (1996) False-positive polymerase chain reaction results for human papillomavirus in lichen planus. Potential laboratory pitfalls of this procedure. *Journal of the American Academy of Dermatology* 35, 42–46.
- Brown C, Kowalczyk AM, Taylor ER, Morgan IM & Gaston K (2008) P53 represses human papillomavirus type 16 DNA replication via the viral E2 protein. *Virology Journal* 5, 5.
- Chidzonga MM, Mahomva L, Makunike-Mutasa R & Masanganise R (2009) Xeroderma pigmentosum: a retrospective case series in Zimbabwe. *Journal of Oral and Maxillofacial Surgery* 67, 22–31.
- Chisi SK, Kollmann MK & Karimurio J (2006) Conjunctival squamous cell carcinoma in patients with human immunodeficiency virus infection seen at two hospitals in Kenya. *East African Medical Journal* 83, 267–270.
- Chow LT, Broker TR & Steinberg BM (2010) The natural history of human papillomavirus infections of the mucosal epithelia. *APMIS: Acta Pathologica, Microbiologica, et Immunologica Scandinavica* 118, 422–449.
- Clifford GM, Smith JS, Plummer M, Munoz N & Franceschi S (2003) Human papillomavirus types in invasive cervical cancer worldwide: a meta-analysis. *British Journal of Cancer* 88, 63–73.
- Cornell M, Schomaker M, Garone DB *et al.* (2012) Gender differences in survival among adult patients starting antiretroviral therapy in South Africa: a multicentre cohort study. *PLoS Medicine* 9, e1001304.
- Croce A, Moretti A, D'agostino L & Zingariello P (2008) Orbital exenteration in elderly patients: personal experience. Acta Otorhinolaryngologica Italica 28, 193–199.
- Curado MP, Edwards B, Shin HR et al. (eds.) (2007). Cancer incidence in Five Continents. International Agency for Research on Cancer, Lyon, France.
- de Koning MN, Waddell K, Magyezi J et al. (2008) Genital and cutaneous human papillomavirus (HPV) types in relation to conjunctival squamous cell neoplasia: a case- control study in Uganda. Infectious Agents and Cancer 3, 12.
- de Sanjose S, Diaz M, Castellsague X et al. (2007) Worldwide prevalence and genotype distribution of cervical human papillomavirus DNA in women with normal cytology: a meta-analysis. The Lancet Infectious Diseases 7, 453–459.
- de Smedt S, Wildner G & Kestelyn P (2013) Vernal keratoconjunctivitis: an update. *British Journal of Ophthalmology* 97, 9–14.
- de Villiers EM (2013). Cross-roads in the classification of papillomaviruses. Virology.
- di Girolamo N (2012) Association of human papilloma virus with pterygia and ocular-surface squamous neoplasia. *Eye* (London, England) 26, 202–211.
- Dou Z, Xu J, Jiao JH *et al.* (2011) Gender difference in 2-year mortality and immunological response to ART in an HIV-infected Chinese population 2006–2008. *PLoS One* 6, e22707.
- Dumaz N, Drougard C, Sarasin A & Daya-Grosjean L (1993) Specific UV-induced mutation spectrum in the p53 gene of

- skin tumors from DNA-repair-deficient xeroderma pigmentosum patients. *Proceedings of the National Academy of Sciences, USA* 90, 10529–10533.
- Dushku N, Hatcher SL, Albert DM & Reid TW (1999) p53 expression and relation to human papillomavirus infection in pingueculae, pterygia, and limbal tumors. *Archives of Ophthalmology* 117, 1593–1599.
- Emmanuel B, Ruder E, Lin SW, Abnet C, Hollenbeck A & Mbulaiteye S (2012) Incidence of squamous-cell carcinoma of the conjunctiva and other eye cancers in the NIH-AARP Diet and Health Study. *Ecancermedicalscience* 6, 254.
- Eng HL, Lin TM, Chen SY, Wu SM & Chen WJ (2002)
 Failure to detect human papillomavirus DNA in malignant epithelial neoplasms of conjunctiva by polymerase chain reaction. American Journal of Clinical Pathology 117, 429–436.
- Fufa H, Umeta M, Taffesse S, Mokhtar N & Aguenaou H (2009) Nutritional and immunological status and their associations among HIV-infected adults in Addis Ababa, Ethiopia. Food and Nutrition Bulletin 30, 227–232.
- Furahini G & Lewallen S (2010) Epidemiology and management of ocular surface squamous neoplasia in Tanzania. *Ophthalmic Epidemiology* 17, 171–176.
- Gichuhi S & Irlam JH (2013) Interventions for squamous cell carcinoma of the conjunctiva in HIV-infected individuals. Cochrane Database Systematic Review, 2, CD005643.
- Goedert JJ & Cote TR (1995) Conjunctival malignant disease with AIDS in USA. Lancet 346, 257–258.
- Gonzalez MA, Martin L, Munoz S & Jacobson JO (2011) Patterns, trends and sex differences in HIV/AIDS reported mortality in Latin American countries: 1996–2007. BMC Public Health 11, 605.
- Grossniklaus HE, Green WR, Luckenbach M & Chan CC (1987) Conjunctival lesions in adults. A clinical and histopathologic review. Cornea 6, 78–116.
- Guech-Ongey M, Engels EA, Goedert JJ, Biggar RJ & Mbulaiteye SM (2008) Elevated risk for squamous cell carcinoma of the conjunctiva among adults with AIDS in the United States. *International Journal of Cancer* 122, 2590–2593.
- Gupta N, Sachdev R & Tandon R (2011) Ocular surface squamous neoplasia in xeroderma pigmentosum: clinical spectrum and outcome. *Graefes Archive for Clinical and Experimental Ophthalmology* **249**, 1217–1221.
- Guthoff R, Marx A & Stroebel P (2009) No evidence for a pathogenic role of human papillomavirus infection in ocular surface squamous neoplasia in Germany. *Current Eye Research* **34**, 666–671.
- Halliday GM, Damian DL, Rana S & Byrne SN (2012) The suppressive effects of ultraviolet radiation on immunity in the skin and internal organs: implications for autoimmunity. *Journal of Dermatological Science* 66, 176–182.
- Haverkos HW (2004) Viruses, chemicals and co-carcinogenesis. Oncogene 23, 6492–6499.
- Heinz C, Fanihagh F & Steuhl KP (2003) Squamous cell carcinoma of the conjunctiva in patients with atopic eczema. *Cornea* 22, 135–137.

- Hirst LW (2007) Randomized controlled trial of topical mitomycin C for ocular surface squamous neoplasia: early resolution. *Ophthalmology* **114**, 976–982.
- Holkar S, Mudhar HS, Jain A et al. (2005) Regression of invasive conjunctival squamous carcinoma in an HIV-positive patient on antiretroviral therapy. *International Journal of STD and AIDS* 16, 782–783.
- Hughes DS, Powell N & Fiander AN (2008) Will vaccination against human papillomavirus prevent eye disease? A review of the evidence. *British Journal of Ophthalmology* 92, 460–465.
- Iftner T & Villa LL (2003) Chapter 12: Human papillomavirus technologies. Journal of the National Cancer Institute, Monographs 31, 80–88.
- Johnson LF, Mossong J, Dorrington RE et al. (2013) Life expectancies of South African adults starting antiretroviral treatment: collaborative analysis of cohort studies. PLoS Medicine 10, e1001418.
- Kanazawa S, Kitaoka T, Ueda Y, Gong H & Amemiya T (2002) Interaction of zinc and vitamin A on the ocular surface. Graefes Archive for Clinical and Experimental Ophthalmology 240, 1011–1021.
- Kao AA, Galor A, Karp CL, Abdelaziz A, Feuer WJ & Dubovy SR (2012) Clinicopathologic correlation of ocular surface squamous neoplasms at bascom palmer eye institute: 2001 to 2010. Ophthalmology 119, 1773–1776.
- Karcioglu ZA & Issa TM (1997) Human papilloma virus in neoplastic and non-neoplastic conditions of the external eye. *British Journal of Ophthalmology* 81, 595–598.
- Kestelyn P, Stevens AM, Ndayambaje A, Hanssens M & van de Perre P (1990) HIV and conjunctival malignancies. *Lancet* **336**, 51–52.
- Kim Y, Choi KR, Chae MJ et al. (2013) Stability of DNA, RNA, cytomorphology, and immunoantigenicity in Residual ThinPrep Specimens. APMIS: Acta Pathologica, Microbiologica, et Immunologica Scandinavica doi: 10.1111/apm.12082 [Epub ahead of print].
- Kuo KT, Chang HC, Hsiao CH & Lin MC (2006) Increased Ki-67 proliferative index and absence of P16INK4 in CIN-HPV related pathogenic pathways different from cervical squamous intraepithelial lesion. *British Journal of Ophthalmology* 90, 894–899.
- Lauer SA, Malter JS & Meier JR (1990) Human papillomavirus type 18 in conjunctival intraepithelial neoplasia. *American Journal of Ophthalmology* 110, 23–27.
- Lee GA & Hirst LW (1992) Incidence of ocular surface epithelial dysplasia in metropolitan Brisbane. A 10-year survey. *Archives of Ophthalmology* 110, 525–527.
- Lee GA & Hirst LW (1995) Ocular surface squamous neoplasia. Survey of Ophthalmology 39, 429–450.
- Lee GA & Hirst LW (1997) Retrospective study of ocular surface squamous neoplasia. *Australian and New Zealand Journal of Ophthalmology* 25, 269–276.
- Lee GA, Williams G, Hirst LW & Green AC (1994) Risk factors in the development of ocular surface epithelial dysplasia. *Ophthalmology* **101**, 360–364.

- Lee SB, Au Eong KG, Saw SM, Chan TK & Lee HP (2000) Eye cancer incidence in Singapore. *British Journal of Ophthalmology*, **84**, 767–770.
- Lynch SM & Rebbeck TR (2013) Bridging the gap between biologic, individual, and macroenvironmental factors in cancer: a multilevel approach. *Cancer Epidemiology Biomarkers & Prevention* 22, 485–495.
- Mahomed A & Chetty R (2002) Human immunodeficiency virus infection, Bcl-2, p53 protein, and Ki-67 analysis in ocular surface squamous neoplasia. *Archives of Ophthalmology* **120**, 554–558.
- Makupa II, Swai B, Makupa WU, White VA & Lewallen S (2012) Clinical factors associated with malignancy and HIV status in patients with ocular surface squamous neoplasia at Kilimanjaro Christian Medical Centre, Tanzania. *British Journal of Ophthalmology*, 96, 482–484.
- Maloof AJ, Ho A & Coroneo MT (1994) Influence of corneal shape on limbal light focusing. *Investigative Ophthalmology* & Visual Science 35, 2592–2598.
- Manderwad GP, Kannabiran C, Honavar SG & Vemuganti GK (2009) Lack of association of high-risk human papillomavirus in ocular surface squamous neoplasia in India. *Archives of Pathology and Laboratory Medicine* 133, 1246–1250.
- Margo CE & Groden LR (1986) Squamous cell carcinoma of the cornea and conjunctiva following a thermal burn of the eve. Cornea 5, 185–188.
- Margo CE, Harman LE & Mulla ZD (2002) The reliability of clinical methods in ophthalmology. *Survey of Ophthalmology* 47, 375–386.
- Masanganise R & Magava A (2001) Orbital exenterations and squamous cell carcinoma of the conjunctiva at Sekuru Kaguvi Eye Unit, Zimbabwe. *Central African Journal of Medicine* 47, 194 199
- Masanganise R, Rusakaniko S, Makunike R et al. (2008) A historical perspective of registered cases of malignant ocular tumors in Zimbabwe (1990 to 1999). Is HIV infection a factor? Central African Journal of Medicine 54, 28–32.
- Maskew M, Brennan AT, Westreich D, McNamara L, Macphail AP & Fox MP (2013) Gender differences in mortality and CD4 count response among virally suppressed HIV- positive patients. *Journal of Women's Health* (2002) 22, 113–120.
- McDonnell JM, McDonnell PJ & Sun YY (1992) Human papillomavirus DNA in tissues and ocular surface swabs of patients with conjunctival epithelial neoplasia. *Investigative Ophthalmology & Visual Science* 33, 184–189.
- McKelvie PA (2002) Squamous cell carcinoma of the conjunctiva: a series of 26 cases. *British Journal of Ophthalmology* 86, 168–173.
- Mesher D, Szarewski A, Cadman L *et al.* (2013) Comparison of human papillomavirus testing strategies for triage of women referred with low-grade cytological abnormalities. *European Journal of Cancer* 49, 2179–2186.
- Moubayed P, Mwakyoma H & Schneider DT (2004) High Frequency of Human Papillomavirus 6/11, 16, and 18 Infections in Precancerous Lesions and Squamous Cell Carcinoma of the Conjunctiva in Subtropical Tanzania. American Journal of Clinical Pathology 122, 938–943.

- Munoz M, Camargo M, Soto-De Leon SC *et al.* (2012) The diagnostic performance of classical molecular tests used for detecting human papillomavirus. *Journal of Virological Methods*, 185, 32–38.
- Nakamura Y, Mashima Y, Kameyama K, Mukai M & Oguchi Y (1997) Detection of human papillomavirus infection in squamous tumours of the conjunctiva and lacrimal sac by immunohistochemistry, in situ hybridisation, and polymerase chain reaction. British Journal of Ophthalmology 81, 308–313.
- Napora C, Cohen EJ, Genvert GI *et al.* (1990) Factors associated with conjunctival intraepithelial neoplasia: a case control study. *Ophthalmic Surgery* **21**, 27–30.
- Nemet AY, Martin P, Benger R et al. (2007) Orbital exenteration: a 15-year study of 38 cases. Ophthalmic Plastic and Reconstructive Surgery 23, 468–472.
- Newton R, Ferlay J, Reeves G, Beral V & Parkin DM (1996) Effect of ambient solar ultraviolet radiation on incidence of squamous-cell carcinoma of the eye. *Lancet* 347, 1450–1451.
- Newton R, Ziegler J, Beral V et al. (2001) A case-control study of human immunodeficiency virus infection and cancer in adults and children residing in Kampala, Uganda. *Interna*tional Journal of Cancer 92, 622–627.
- Newton R, Ziegler J, Ateenyi-Agaba C et al. (2002) The epidemiology of conjunctival squamous cell carcinoma in Uganda. British Journal of Cancer 87, 301–308.
- Ng J, Coroneo MT, Wakefield D & di Girolamo N (2008)
 Ultraviolet radiation and the role of matrix metalloproteinases in the pathogenesis of ocular surface squamous neoplasia.

 Investigative Ophthalmology & Visual Science 49, 5295–5306.
- Odrich MG, Jakobiec FA, Lancaster WD *et al.* (1991) A spectrum of bilateral squamous conjunctival tumors associated with human papillomavirus type 16. *Ophthalmology*, **98**, 628–635.
- Ogun GO, Ogun OA, Bekibele CO & Akang EE (2009) Intraepithelial and invasive squamous neoplasms of the conjunctiva in Ibadan, Nigeria: a clinicopathological study of 46 cases. *International Ophthalmology* **29**, 401–409.
- Osahon AI, Ukponmwan CU & Uhunmwangho OM (2011) Prevalence of HIV seropositivity among patients with squamous cell carcinoma of the conjunctiva. *Asian Pacific Journal of Tropical Biomedicine* 1, 150–153.
- Parkin DM, Nambooze S, Wabwire-Mangen F & Wabinga HR (2010) Changing cancer incidence in Kampala, Uganda 1991–2006. *International Journal of Cancer* 126, 1187–1195.
- Perez-Molina JA, Mora Rillo M, Suarez-Lozano I *et al.* (2012) Response to combined antiretroviral therapy according to gender and origin in a cohort of naive HIV-infected patients: GESIDA- 5808 study. *HIV Clinical Trials*, 13, 131–141.
- Pfister RR & Renner ME (1978) The corneal and conjunctival surface in vitamin A deficiency: a scanning electron microscopy study. *Investigative Ophthalmology & Visual Science* 17, 874–883.
- Pola EC, Masanganise R & Rusakaniko S (2003) The trend of ocular surface squamous neoplasia among ocular surface tumour biopsies submitted for histology from Sekuru Kaguvi Eye Unit, Harare between 1996 and 2000. *Central African Journal of Medicine* 49, 1–4.

- Poole TR (1999) Conjunctival squamous cell carcinoma in Tanzania. British Journal of Ophthalmology 83, 177–179.
- Porges Y & Groisman GM (2003) Prevalence of HIV with conjunctival squamous cell neoplasia in an African provincial hospital. *Cornea* 22, 1–4.
- Pushker N, Kashyap S, Balasubramanya R et al. (2004) Pattern of orbital exenteration in a tertiary eye care centre in India. Clinical & Experimental Ophthalmology 32, 51–54.
- Rahman I, Cook AE & Leatherbarrow B (2005) Orbital exenteration: a 13 year Manchester experience. British Journal of Ophthalmology 89, 1335–1340.
- Reszec J & Sulkowski S (2005) The expression of P53 protein and infection of human papilloma virus in conjunctival and eyelid neoplasms. *International Journal of Molecular Medicine* 16, 559–564.
- Saegusa M, Takano Y, Hashimura M, Okayasu I & Shiga J (1995) HPV type 16 in conjunctival and junctional papilloma, dysplasia, and squamous cell carcinoma. *Journal of Clinical Pathology* 48, 1106–1110.
- Scheffner M, Werness BA, Huibregtse JM, Levine AJ & Howley PM (1990) The E6 oncoprotein encoded by human papillomavirus types 16 and 18 promotes the degradation of p53. *Cell* 63, 1129–1136.
- Scott IU, Karp CL & Nuovo GJ (2002) Human papillomavirus 16 and 18 expression in conjunctival intraepithelial neoplasia. *Ophthalmology* **109**(542–7), 39.
- Sen S, Sharma A & Panda A (2007) Immunohistochemical localization of human papilloma virus in conjunctival neoplasias: a retrospective study. *Indian Journal of Ophthalmology* 55, 361–363.
- Shields CL & Shields JA (2004) Tumors of the conjunctiva and cornea. Survey of Ophthalmology 49, 3-24.
- Simbiri KO, Murakami M, Feldman M et al. (2010) Multiple oncogenic viruses identified in Ocular surface squamous neoplasia in HIV-1 patients. Infectious Agents and Cancer 5, 6.
- Smeets SJ, Hesselink AT, Speel EJ *et al.* (2007) A novel algorithm for reliable detection of human papillomavirus in paraffin embedded head and neck cancer specimen. *International Journal of Cancer* **121**, 2465–2472.
- Spitzer MS, Batumba NH, Chirambo T et al. (2008) Ocular surface squamous neoplasia as the first apparent manifestation of HIV infection in Malawi. Clinical & Experimental Ophthalmology 36, 422–425.
- Stanley M (2010) Pathology and epidemiology of HPV infection in females. *Gynecologic Oncology* 117, S5–10.
- Sun EC, Fears TR & Goedert JJ (1997) Epidemiology of squamous cell conjunctival cancer. Cancer Epidemiology Biomarkers & Prevention 6, 73–77.
- Tabrizi SN, McCurrach FE, Drewe RH, Borg AJ, Garland SM & Taylor HR (1997) Human papillomavirus in corneal and conjunctival carcinoma. Australian and New Zealand Journal of Ophthalmology 25, 211–215.
- Taylor-Smith K, Tweya H, Harries A, Schoutene E & Jahn A (2010) Gender differences in retention and survival on antiretroviral therapy of HIV-1 infected adults in Malawi. *Malawi Medical Journal* 22, 49–56.

- Templeton AC (1967) Tumors of the eye and adnexa in Africans of Uganda. *Cancer* 20, 1689–1698.
- Templeton AC (1973) Tumours of the eye and adnexa. *Recent Results in Cancer Research*, 41(20), 3–14.
- Thorsteinsson K, Ladelund S, Jensen-Fangel S et al. (2012) Impact of gender on the risk of AIDS-defining illnesses and mortality in Danish HIV-1-infected patients: a nationwide cohort study. Scandinavian Journal of Infectious Diseases 44, 766–775.
- Tiong T, Borooah S, Msosa J et al. (2013) Clinicopathological review of ocular surface squamous neoplasia in Malawi. British Journal of Ophthalmology 97, 961–964.
- Tornesello ML, Duraturo ML, Waddell KM *et al.* (2006) Evaluating the role of human papillomaviruses in conjunctival neoplasia. *British Journal of Cancer* **94**, 446–449.
- Toth J, Karcioglu ZA, Moshfeghi AA, Issa TM, Al-Ma'ani JR & Patel KV (2000) The relationship between human papillomavirus and p53 gene in conjunctival squamous cell carcinoma. *Cornea* 19, 159–162.
- Touzri RA, Mohamed Z, Khalil E et al. (2008) Ocular malignancies of xeroderma pigmentosum: clinical and therapeutic features. Annales de Dermatologie et de Venereologie 135, 99–104.
- Tulvatana W (2003) Risk factors for conjunctival squamous cell neoplasia: a matched case-control study. *British Journal of Ophthalmology* 87, 396–398.
- Tunc M, Char DH, Crawford B & Miller T (1999) Intraepithelial and invasive squamous cell carcinoma of the conjunctiva: analysis of 60 cases. *British Journal of Ophthalmology* 83, 98–103.
- Tuppurainen K, Raninen A, Kosunen O *et al.* (1992) Squamous cell carcinoma of the conjunctiva. Failure to demonstrate HPV DNA by *in situ* hybridization and polymerase chain reaction. *Acta Ophthalmologica* 70, 248–254.
- Ukponmwan CU, Igbokwe UO & Aligbe JU (2002) Squamous cell carcinoma of the conjunctiva in Benin City Nigeria. *Nigerian Journal of Medical Practice* 5, 143–147.
- Vajdic CM, van Leeuwen MT, McDonald SP et al. (2007) Increased incidence of squamous cell carcinoma of eye after kidney transplantation. Journal of the National Cancer Institute 99, 1340–1342.
- Velema JP, Ferrera A, Figueroa M et al. (2002) Burning wood in the kitchen increases the risk of cervical neoplasia in HPVinfected women in Honduras. *International Journal of Cancer* 97, 536–541.
- Victora CG, Huttly SR, Fuchs SC & Olinto MT (1997) The role of conceptual frameworks in epidemiological analysis: a hierarchical approach. *International Journal of Epidemiology* 26, 224–227.
- Visser ME, Maartens G, Kossew G & Hussey GD (2003) Plasma vitamin A and zinc levels in HIV-infected adults in Cape Town, South Africa. *British Journal of Nutrition* 89, 475–482.
- Wabinga HR, Parkin DM, Wabwire-Mangen F & Nambooze S (2000) Trends in cancer incidence in Kyadondo County, Uganda 1960–1997. British Journal of Cancer 82, 1585–1592.

- Waddell KM, Lewallen S, Lucas SB, Atenyi-Agaba C, Herrington CS & Liomba G (1996) Carcinoma of the conjunctiva and HIV infection in Uganda and Malawi. *British Journal of Ophthalmology* **80**, 503–508.
- Waddell K, Magyezi J, Bousarghin L et al. (2003) Antibodies against human papillomavirus type 16 (HPV-16) and conjunctival squamous cell neoplasia in Uganda. British Journal of Cancer 88, 2002–2003.
- Waddell KM, Downing RG, Lucas SB & Newton R (2006) Corneo-conjunctival carcinoma in Uganda. Eye (London, England) 20, 893–899.
- Waddell K, Kwehangana J, Johnston WT, Lucas S & Newton R (2010) A case-control study of ocular surface squamous neoplasia (OSSN) in Uganda. *International Journal of Cancer* 127, 427–432.
- WHO (2000) International Classification of Diseases for Oncology (ICD-O-3) [Online]. Available: http://www.who.int/classifications/icd/adaptations/oncology/en/index.html [Accessed 14th March 2013.
- WHO (2010) International Statistical Classification of Diseases and Related Health Problems 10th Revision (ICD-10) [Online]. Available: http://apps.who.int/classifications/icd10/browse/2010/en [Accessed 4th January 2013].
- World Energy Council (2007) *Survey of Energy Resources* [Online]. Available: http://www.worldenergy.org/documents/fig_solar_10_2.gif [Accessed 4th March 2013].
- Yu JJ, Fu P, Pink JJ et al. (2010) HPV infection and EGFR activation/alteration in HIV-infected East African patients with conjunctival carcinoma. PLoS One 5, e10477.

Corresponding Author Stephen Gichuhi, Department of Ophthalmology, University of Nairobi, Nairobi, Kenya. E-mail: sgichuhi@uonbi.ac.ke