HIV-associated malignancies in children

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Purpose of review
HIV-infected children are at an increased risk of developing cancer. Many of the cancers in HIV-infected children are linked to immunosuppression and oncogenic coinfections. Worldwide most HIV-infected children live in sub-Saharan Africa, but cancer data for this population are scarce. In this article, we review the current literature on the epidemiology and prevention of cancer in HIV-infected children.

Recent findings
Combined antiretroviral therapy (cART) reduces the risk of developing cancer in HIV-infected children. Cancer risk remains increased in children who start cART at older ages or more advanced immunosuppression as compared with children who start cART at younger age and with mild immunosuppression. Starting cART before severe immunosuppression develops is key to prevent cancer in HIV-infected children but most children in low-income countries start cART at severe immunosuppression levels. Vaccination against high-risk variants of human papillomavirus may protect against human papillomavirus-associated cancer later in life. However, tailoring of human papillomavirus vaccination guidelines for HIV-infected children and young women awaits answers to determine the best vaccination strategies.

Summary
Better data on the short-term and long-term risks of developing cancer and the effects of preventive measures in HIV-infected children from regions with high burden of HIV/AIDS are urgently needed.

Keywords
cancer, children, epidemiology, HIV, prevention

INTRODUCTION
HIV-infected children have an increased cancer risk due to a compromised immune system and an increased susceptibility to oncogenic viruses \cite{1,2}. Worldwide 1.8 million children aged 0–14 years are HIV-infected and at increased risk of developing cancer. Most of these children live in Africa: 1.04 million (58\%) in Eastern and Southern Africa and 0.51 million (29\%) in Western and Central Africa. In Asia-Pacific, the total number of HIV-infected children is 193,000 (11\% of HIV-infected children worldwide) and 32,200 (2\%) in Latin America and the Caribbean’s (Fig. 1). Cancer mortality in HIV-infected children remains high in resource-limited countries. Median survival in HIV-infected children and adolescents with Kaposi sarcoma was less than 6 months in a recent trial from Malawi \cite{3}. In Uganda, median survival in HIV-infected children with Burkitt lymphoma was 11.8 months \cite{4}. In a study from South Africa \cite{5}, 10\% of all HIV-infected children with cancer died of treatment-related complications and severe infections. Reliable estimates on the burden of cancer and measures to prevent cancer in HIV-infected children are warranted, especially in resource-limited countries with large burdens of HIV/AIDS. There is a paucity of cancer incidence studies in HIV-infected children from sub-Saharan Africa (SSA), wherein the majority of HIV-infected children live \cite{6}. In this article, we review the current literature on the epidemiology and prevention of cancer in HIV-infected children.

METHODS
We searched Medline for the period August 2011 to August 2016 using the following search terms: (((HIV[Title/Abstract] OR AIDS[Title/Abstract])

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AND (cancer[Title/Abstract] OR neoplasm[Title/Abstract] OR tumor[Title/Abstract] OR tumour[Title/Abstract]) AND (infant OR children).

Additional searches were done for specific topics of interest, such as African countries and prevention. We included articles that were published in the last 5 years and highlighted those of special interest from the last 18 months. Cancer treatment and outcome are not covered in this article.

**KEY POINTS**

- HIV-infected children are at increased risk of developing cancer due to immunosuppression and most cancers are associated with oncogenic coinfections.
- Most HIV-infected children live in SSA, but cancer studies in HIV-infected children from this region are scarce.
- Starting cART before severe immunosuppression develops is key to prevent cancer in HIV-infected children but most children in low-income countries start cART at severe immunosuppression levels.
- Vaccination against high-risk variants of HPV may protect against cancer later in life, however, tailoring of HPV vaccination guidelines for HIV-infected children and young women awaits answers to determine the best vaccination strategies.

**Epidemiology of Cancer in HIV-Infected Children**

Producing reliable cancer incidence estimates in HIV-infected pediatric populations is a challenge: HIV cohorts may not record cancer cases and cancer registries may not record HIV infection status. Record linkages between data of HIV-treatment programmes or HIV registries and cancer registries have been identified as a potential solution to this problem [7,8]. Three recent studies [9,10–11] from the USA, Taiwan and South Africa have used this method to estimate cancer incidence rates in HIV-infected children. The South Africa study [9] linked data at five pediatric combined antiretroviral therapy (cART) programmes using probabilistic record linkage methods with four referral pediatric oncology units. Amongst a total of 11 707 children, 47 prevalent and 24 incident cancer cases were identified. The overall cancer incidence rate was 82/100 000 person-years. The most frequent cancers were Kaposi sarcoma and non-Hodgkin lymphoma (NHL) with incidence rates of 34 and 31/100 000 person-years, respectively. The incidence rate for all non-AIDS defining cancers (NADCs) combined was 17/100 000 person-years. Risk factors identified were older age at starting cART [10 years versus <3 years, adjusted hazard ratio (HR) 7.3, 95% confidence interval (CI) 2.2–24.6] and severe and advanced immunodeficiency as compared with mild or no immunodeficiency at enrolment into programme (adjusted HR 3.5, 95% CI 1.1–12). The risk of

**FIGURE 1.** Joint United Nations Programme on HIV/AIDS (UNAIDS) estimates for number of children (in thousands) living with HIV stratified by age group and region in 2015. Latin America and the Caribbean: 7200 children aged 0–4 years; 11 000 children aged 5–9 years; 14 000 children aged 10–14 years. Middle East and North Africa: 5200 children aged 0–4 years; 3400 children aged 5–9 years; 2300 children aged 10–14 years. Estimates for Western, Central and Eastern Europe, Central Asia and North America are not available (NA). Source: unpublished UNAIDS 2016 estimates.
developing cancer was reduced by 70% in children receiving cART compared with children not receiving cART (adjusted HR 0.29, 95% CI 0.09–0.86). There were insufficient incident cancers to identify risk factors for specific cancers. This was one of the first studies to show a protective effect of cART on the risk of developing cancer at patient level in children. Although the study used data from referral pediatric oncology units to improve cancer ascertainment, under-ascertainment cannot be excluded given that children diagnosed with cancer may not reach specialized services. The study highlighted the need for early HIV diagnosis and cART initiation before advanced immunodeficiency develops to further reduce the burden of cancer in HIV-infected children. A recent multiregional analysis has shown that many children in low-income countries start cART at severe immunosuppression levels (62 and 65% in girls and boys) compared with high-income countries (21 and 28% in girls and boys) [12**]. Reasons for the late start are manifold and several studies evaluated acceptance of HIV testing, linkage to care and cART initiation [13]. However, few reported long-term outcomes [14] and none provided cancer data.

Simard et al. [10] investigated the long-term cancer risks in young adults diagnosed with AIDS during childhood. The study was based on the United States HIV/AIDS Cancer Match Study using record linkage between HIV and cancer registries. Participants were followed up for up to 10 years. Compared to the general population, children diagnosed with AIDS had an increased risk of developing Kaposi sarcoma, NHL and NADCs, with leiomyosarcoma being the most frequent NADC. Comparing cancer incidence in the pre-cART with the cART era the risk of developing Kaposi sarcoma was reduced by almost 90% [relative risk (RR) 0.13, 95% CI 0.02–0.74] and the NHL risk by 60% (RR 0.40, 95% CI 0.21–0.75). The risk of developing NADCs did not decrease with the advent of cART (RR 0.98, 95% CI 0.33–2.86). The study highlighted that the risk of developing certain cancers remains increased in persons diagnosed with AIDS during childhood and the need for continued cancer monitoring in this population even with the initiation of cART. With improved access to care in low-income and middle-income countries there are now growing numbers of HIV-infected children who start cART earlier and who live into adolescence and adulthood. Studies looking at long-term cancer risk in persons with long-term exposure to HIV infection, immunosuppression and cART are urgently needed for African and other affected countries.

Lastly, the study [11] from Taiwan used record linkage methods to estimate cancer incidence in HIV-infected children using the Nationwide Health Insurance database. However, with 207 included children the study was rather small. Incidence rates for Kaposi sarcoma, NHL and NADC differ substantially between these three studies from different geographic areas (Table 1). It is also difficult to compare these studies given the differences in study design, inclusion criteria and analysis methods. This problem was addressed in a recent multiregional cohort study that directly compared the risk of developing Kaposi sarcoma in children starting cART in different geographic regions. Of the 24 991 HIV-infected children from Eastern Africa, Southern Africa and Europe starting cART, 26 children developed incident Kaposi sarcoma [15**]. The study showed that Kaposi sarcoma [caused by an infection with human herpesvirus 8 (HHV-8)], is a frequent cancer in SSA and in SSA migrants in Europe, but cases are rare in children outside of this region. Incidence rates per 100 000 person-years were 86 in Eastern Africa, 11 in

Table 1. Literature review 2011–2016: cancer incidence in HIV-infected children in the era of combination antiretroviral therapy

<table>
<thead>
<tr>
<th>Study</th>
<th>Countries</th>
<th>Calendar years</th>
<th>Children included</th>
<th>Total follow-up time (years)</th>
<th>Cancer cases identified</th>
<th>Incidence rate per 100 000 person-years</th>
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*Included children and adolescents diagnosed with AIDS only.

Incident defined as: after cohort enrolment.
Southern Africa and 81 in children of SSA origin living in Europe, but zero in Asia-Pacific and European children of non-SSA origin (Fig. 2). Higher age at starting cART and advanced HIV/AIDS stage were risk factors for developing Kaposi sarcoma. Regional differences persisted after adjusting for potential confounders and may point to higher HHV-8 coinfection rates in HIV-infected children from SSA. However, data for HHV-8 seroprevalence were not available in that study. In conclusion, the study underscores the need to start cART early in HIV-infected children at high risk for Kaposi sarcoma before advanced immunosuppression develops.

Early cART initiation may also help to reduce the risk of new HHV-8 infections in children living in endemic areas. Recent studies [16,17] demonstrated that HIV-infected children have a higher risk of coinfection with HHV-8 compared with HIV-uninfected children. An observational study from Zambia suggested that in HIV-infected children receiving cART, the risk of HHV-8 infection was similar to HIV-uninfected children [17]. In contrast, in HIV-infected children not receiving cART the risk of acquiring HHV-8 was six times higher compared with HIV-uninfected children (incidence rate ratio 5.97, 95% CI 3.13–11.41) [17].

PREVENTION OF CANCER IN HIV-INFECTED AND EXPOSED CHILDREN

In this section, we review the latest World Health Organization (WHO) guidelines for the early testing and treatment of HIV-infected children and its implications, including cancer risks in HIV and antiretroviral drugs (ARV) exposed but uninfected children. Lastly, we discuss vaccination against human papillomavirus (HPV) in HIV-infected children to prevent cervical cancer later in life.

EARLY TESTING AND TREATMENT

In the WHO ‘Consolidated guidelines on HIV testing services’ (2015), special mention is made of HIV testing for infants/children and adolescents [18]. Scaling up of early infant diagnosis (EID) is thought to be improved by task shifting to trained and supervised lay providers. The use of point-of-care HIV testing is carefully considered with rapid diagnostic tests for HIV serology recommended to assess HIV exposure in infants less than 4 months, to rule out HIV in asymptomatic HIV-exposed infants at 9 months and to diagnose HIV in children older than 18 months. The development of point-of-care virological assays could further improve EID access [18]. WHO suggests governments reconsider the age of consent to allow greater autonomy for HIV testing among adolescents [18]. However, to optimally prevent malignancy and achieve mortality and morbidity benefits, HIV diagnosis needs to be linked to prompt cART initiation and long-term retention on treatment, which could prove challenging in resource-limited settings.

Updated treatment recommendations support the initiation of cART for HIV-infected adolescents and children regardless of WHO clinical stage and CD4+ cell count, particularly infants diagnosed within the first year of life. Priority is given to children 2 years and younger, children between 2 and 5 years with WHO clinical stage 3 or 4 or CD4+ count ≥750 cells/mm³ or CD4+ percentage less than 25%, and children older than 5 years (including adolescents) with WHO stage 3/4 or CD4+ count ≥350 cell/mm³ [19].

The new guidelines, however, introduce the concern of cumulative effects of ARV therapy on a growing child as a result of early cART initiation with life-long duration. Possible strategies to reduce overall lifetime exposure and mitigate cumulative side-effects could include early time-limited cART. The CHER study (Children with HIV Early Antiretroviral randomized trial) investigated the effects of early time-limited cART initiation compared with deferred cART in infants [20]. HIV-infected infants 6–12 weeks of age with CD4+ percentage less than 25% were randomized to deferred therapy (ART-Def), early cART restricted to 40 weeks (ART-40W) or early cART restricted to 96 weeks (ART-96W) with 3.5 years follow-up. Hazard ratios for death for
cART-40W and cART-96W were 0.40 ($P = 0.02$) and 0.45 ($P = 0.36$), respectively compared with cART-Def. Cumulative probability of clinical disease progression or death by 3.5 years was 41% in cART-Def, 28% in the cART-40W and 21% in the cART-96W groups. The CHER trial reported that primary clinical, immunological and virological endpoints for early-time limited cART were superior to deferred cART in asymptomatic HIV-infected infants. However, there was no comparison between time-limited and continuous early cART, follow-up was too short to fully assess the effects of the different strategies on incident malignancy and the strategy may not be feasible in lower-resource settings because of its complexity and the need for close clinical and CD4+ monitoring. Overall, the idea of time-limited cART in infants is appealing but data regarding its efficacy compared with continuous cART is lacking.

**CANCER RISK IN HIV AND ANTIRETROVIRAL EXPOSED BUT UNINFECTED CHILDREN**

Although ARV prophylaxis for the prevention of mother-to-child HIV transmission has been tremendously successful, new challenges regarding the health outcomes of HIV and ARV exposed, uninfected infants are anticipated. Amongst these is the risk of cancer amongst infants exposed to HIV and ARV perinatally. Few studies [21–24,25] have comprehensively investigated this risk with most studies hampered by short-term follow-up periods. In the most recent study with a 16-year follow-up period in New Jersey, USA, cancer registry data were linked to HIV case surveillance data to determine the risk of cancer amongst HIV-exposed, uninfected infants perinatally exposed to ARV prophylaxis [26*]. For 3805 HIV-exposed uninfected children the incidence of all cancer types was 13.7/100,000 person-years (95% CI 3.7–35.2). Cancer incidence was not significantly different between HIV-exposed children unexposed to ARV prophylaxis (22.5/100,000 person-years) and children exposed to ARV prophylaxis (14.3/100,000 person-years). Another study [25*] showed an increased risk of developing cancer in children exposed to didanosine in-utero as compared with the other ARVs. The authors concluded that didanosine should be strictly contraindicated in pregnancy. Continued vigilance of the health effects of ARV prophylaxis in infants is required as the prevention of mother-to-child transmission programme gains momentum and success globally in heterogeneous settings.

**HPV VACCINATION FOR THE PRIMARY PREVENTION OF INVASIVE CERVICAL CANCER**

HIV-infected individuals are at higher risk for all types of HPV-related cancer compared with the HIV-uninfected individuals [27]. Bivalent, quadrivalent and the newly introduced nanovalent HPV vaccine offer effective protection against infection with high-risk HPV variants and have acceptable safety profiles in the general population [28**]. Vaccinations are recommended for girls and women aged 9 years up to 25 years old prior to HPV exposure [29]. There are few studies investigating the immunogenicity and safety of the HPV vaccination in HIV-infected populations [30]. However, from the available evidence, both the bivalent and quadrivalent HPV vaccines have demonstrated high seroconversion rates and acceptable safety profiles in various HIV-infected populations (children, female adolescents and adults). The nanovalent vaccine is still untested in the HIV-infected population [31*]. Seroconversion rates for the quadrivalent vaccine of more than 92% were reported in HIV-infected boys and girls aged 7–12 years [32] and HIV-infected young women aged 16–23 years (ART and non-ART groups) [33]. However Levin et al. [32] reported lower antibody titres to HPV-6 and HPV-18 in HIV-infected children (7–12 years) compared with historical controls. Kahn et al. [33] also reported lower antibody titres in young HIV-infected women (16–23 years) not on cART, but similar antibody titres for women receiving cART compared with the HIV-uninfected comparison group.

In a study [34] of the bivalent vaccine in HIV-infected women in South Africa (18–25 years), all study participants remained seropositive for HPV-16 and HPV-18 at 12-month follow-up. Antibody titres were 50% and 70% lower at 7 and 12 months in the HIV-infected group compared with the HIV-uninfected group [30,34]. However, HPV-16 and HPV-18 geometric mean titres in the HIV-infected group remained 26-fold and 16-fold higher at 1 year than those reported in healthy women (15–25 years) who cleared a natural infection. The bivalent vaccine was reported to have an acceptable safety and reactogenicity profile in HIV-infected young women in this trial.

Although the immunogenicity and safety of HPV vaccination in HIV-infected children and young women has been demonstrated [30,32–34], no long-term follow-up of HIV cohorts receiving HPV vaccination have been reported. Therefore, the duration of vaccine-induced immunogenicity and the clinical significance of the lower antibody titre in HIV-infected children and young women receiving the bivalent or quadrivalent HPV vaccination
remain unknown. Tailoring of HPV vaccination guidelines for HIV-infected children and young women awaits answers to pertinent questions such as the adequacy of a two-dose quadrivalent vaccination schedule, the evaluation of the newly available nonvalent HPV vaccine, and the requirement/timing of a booster dose in this group.

CONCLUSION

Many of the cancers occurring in HIV-infected children are linked to immunosuppression and oncogenic coinfections. Starting cART before severe immunosuppression develops and vaccination against oncogenic viruses appears key to prevent cancer in HIV-infected children. Better data on the short-term and long-term risks of developing cancer and the effects of preventive measures in HIV-infected children from regions with high burden of HIV/AIDS are urgently needed.

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Conflicts of interest

There are no conflicts of interest.

REFERENCES AND RECOMMENDED READING

Papers of particular interest, published within the annual period of review, have been highlighted as:
|| of special interest
|| of outstanding interest


Observational study demonstrating a beneficial effect of antiretroviral therapy on the risk of acquiring human-herpesvirus 8, the underlying cause of Kaposi sarcoma.

16. International multiregional study comparing directly the risk of developing Kaposi sarcoma in HIV-infected children starting cART in Asia, Africa and Europe.


19. WHO. Consolidated guidelines on HIV testing services 2015. World Health Organization. Comprehensive guidelines on HIV testing services (HST). The document consolidates previous and new HST recommendations that address the existing gaps and limitations to current approaches. It highlights tailored approaches to HST for key populations. Essential reading for programme managers, healthcare workers and other stakeholders in HIV care.

20. WHO. Consolidated guidelines on the use of antiretroviral drugs for treating and preventing HIV infection. 2015. Following an extensive review of evidence in 2015, the WHO has updated its 2013 guidelines on the use of ARVs including 10 new recommendations to improve the quality and efficiency of HIV services provided to the HIV-infected population with an emphasis on earlier initiation of ART for improved clinical outcomes.


Evaluation of cancer incidence amongst HIV-infected children exposed to nucleos(t)ide reverse transcriptase inhibitors in utero. Didanosine, still in use in some African countries, accounted for one-third of cancers in exposed children and was associated with a higher cancer risk in multivariate analysis.


28. Van Damme P, Olsson SE, Block S, et al. Immunogenicity and safety of a 9-valent HPV vaccine. Pediatrics 2015; 136:e28–39. Multicentre study (72 sites) investigating the immunogenicity of nanovalent HPV vaccine in girls and boys aged 9–15 years compared with young women aged 16–26 years. Nanovalent HPV was well tolerated in girls and boys with immune responses noninferior to those of young women, supporting that efficacy findings in young women could be bridged to younger girls and boys (9–15 years).


