| 1 | Cancer in adolescents and | young adults living with HIV |
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17

18 Acknowledgements

19 The authors would like to thank Mary Mahy and Juliana Daher from Joint United Nations Programme

20 on HIV/AIDS (UNAIDS) for providing number of adolescents living with HIV stratified by age group.

21

22 Financial support and sponsorship

23 This work was supported by core funding to the UK Medical Research Council [MC_UU_12023/26],

the National Institute of Allergy and Infectious Diseases of the National Institutes of Health [5U01-

AI069924-05], and the Swiss National Science Foundation [320030_169967]. The funders had no

26 direct role in manuscript writing or the decision to submit for publication.

| 27 | Conflict | t of interest | |
|----|--|---|--|
| 28 | There a | re no conflicts of interest. | |
| 29 | | | |
| 30 | Keywo | rds | |
| 31 | Adolescents, cancer, HIV, prevention, young adults | | |
| 32 | | | |
| 33 | Key poi | ints | |
| 34 | • | AYALHIV are at increased risk of AIDS and non-AIDS defining malignancies, associated with | |
| 35 | | immune dysregulation and coinfection with oncogenic viruses. | |
| 36 | • | Non-Hodgkin lymphoma and Kaposi sarcoma are the commonest malignancies occurring in | |
| 37 | | AYALHIV globally. | |
| 38 | • | Reducing the risk of cancer in AYALHIV requires increased access to suppressive | |
| 39 | | antiretroviral therapy, HPV and HBV vaccination, screening and treatment for HBV/hepatitis | |
| 40 | | C virus coinfection, and programmatic screening for cervical and anogenital cancers. | |
| 41 | • | Improvement in cancer estimates for AYALHIV requires data disaggregated by age and route | |
| 42 | | of HIV transmission, which is currently lacking. | |
| 43 | • | Enabling long-term follow-up of children and adolescents living with HIV, including survivors | |
| 44 | | of a dual diagnosis of HIV and malignancy as they undergo transition into adult services, | |
| 45 | | requires effective linkage of pediatric and adult cohorts. | |

46 3 Tables and 1 Figure

- 47 **Table 1:** Infection-associated cancers and related clinical HIV stage
- 48 **Table 2:** Search strategy for Medline (PubMed)
- 49 **Table 3:** Kaposi sarcoma incidence rate in HIV-positive adolescents and young adults who started
- 50 antiretroviral therapy.
- 51 **Figure 1:** Estimates for number of adolescents and young adults living with HIV in 2016 stratified by
- 52 age group, sex, and UNAIDS region.
- 53

54 ABSTRACT

55 <u>Purpose of the review</u>

Adults living with HIV have an increased risk of malignancy yet there is little data for adolescents and
young adults. We reviewed recently published cancer epidemiology, treatment, and outcome data
for adolescents and young adults living with HIV (AYALHIV) aged 10 to less than 25 years between
2016 and 2017.

60

61 <u>Recent findings</u>

- 62 AYALHIV are at increased risk of developing cancer compared to their uninfected peers. Kaposi
- 63 sarcoma and non-Hodgkin lymphoma occur most frequently with variation by geographical region.
- 64 Increased cancer risk is associated with HIV-related immunosuppression and coinfection with
- 65 oncogenic viruses. Published data, particularly on posttreatment outcomes, remain limited and
- 66 analyses are hampered by lack of data disaggregation by age and route of HIV transmission.

67

68 <u>Summary</u>

- 69 Although data are sparse, the increased cancer risk for AYALHIV is the cause for concern and must be
- 70 modified by improving global access and uptake of antiretroviral therapy, human papilloma virus
- 71 (HPV) and hepatitis B virus (HBV) vaccination, screening for hepatitis B and C infection, and
- 72 optimized cancer screening programs. Education aimed at reducing traditional modifiable cancer risk
- 73 factors should be embedded within multidisciplinary services for AYALHIV.

74 INTRODUCTION

The number of adolescents and young adults living with HIV (AYALHIV) continue to rise due to high rates of new infections and increasing life expectancy on antiretroviral therapy (ART). AYALHIV (between the ages of 10 to <25 years) account for 13% of those living with HIV; the majority of them are from sub-Saharan Africa (SSA). Adolescence is the only age group with a rising AIDS-related mortality [1] (Fig. 1).

80

81 Adolescence, transition, and cancer risk

82 Historically, malignancies in people living with HIV are categorized as AIDS-defining and non-AIDS 83 defining (Table 1) [2,3]. AYALHIV are at increased risk of developing both AIDS and non-AIDS-defining 84 cancers compared to HIV-negative individuals [4]. The increased cancer risk for those living with HIV 85 is driven by interlinked immunosuppression, decreased cancer surveillance, persistent coinfection 86 with oncogenic viruses, and HIV viremia. Immediate ART initiation, before immunosuppression 87 occurs, significantly reduces risk of cancer [5,6]. However, AYALHIV have lower rates of ART uptake, 88 increased nonadherence to ART, and higher rates of loss to follow-up compared to younger children 89 and older adults resulting in poorly controlled HIV [4,7&,8]. 90 Currently, the most frequent cancers in AYALHIV are, depending on geographic region, Kaposi 91 sarcoma, non-Hodgkin lymphoma(NHL), and leiomyosarcoma [4,9,10,11]. Kaposi sarcoma is 92 associated with human herpes virus 8 (HHV-8) infection, leiomyosarcoma, and some NHL subtypes, 93 with Epstein–Barr virus (EBV) infection [12,13]. In American adolescents (15–19 years) living with 94 HIV, 49% [95% confidence interval (CI) 39–63] of all cancers were attributable to infectious 95 precipitants [9]. Higher rates of unprotected sex in behaviourally infected AYALHIV increase 96 acquisition of sexually transmitted oncogenic viruses including high-risk human papilloma viruses 97 (hrHPVs) and hepatitis B and C viruses (HBV, HCV) potentiating cervical, oropharyngeal, anogenital 98 cancers, and hepatocellular carcinomas (HCCs), respectively [13]. Perinatally infected AYALHIV may

face increasedriskof cancer compared to their behaviourally infected peers due to lifelong exposure
to HIV, immune dysregulation, and if coinfected perinatally with HBV and/or HCV.

101 Last, the period of transition of healthcare between pediatric and adult services is associated with

102 poorer health outcomes in many chronic diseases, including HIV [14]. Global models of transition

103 vary widely between countries, income settings, and individual diseases and an adolescent living

104 with HIV and a previous or current cancer diagnosis may have to negotiate two independent

transition processes [15,16]. For young people living with HIV, transition typically occurs during late

teens or early 20s, an age with peak incidence in Hodgkin lymphoma diagnoses within the general

107 population [16–18].

108

109 METHODS

110 We searched PubMedon November 2nd 2017 (search terms are shown in Table 2). We restricted the

search to January 1st 2016 to November 1st 2017. We identified 289 references, which were

reviewed by the authors. We included papers that reported cancer incidence rates, risk factors,

survival, or prevention interventions in AYALHIV aged 10 to less than 25 years. Papers reporting

incidence rate in adults without further age disaggregation for less than 25 year olds were not

115 considered. We included original articles, systematic reviews, and case reports. Expert reviews were

116 excluded. A few older important studies were used to support key statements.

117

118 CANCER EPIDEMIOLOGY IN ADOLESCENTS AND YOUNG ADULTS LIVING WITH HIV

119 In the recent literature, there were limited data on cancers disaggregated by age and virtually no

120 data disaggregated by mode of HIV transmission. It was, therefore, not possible to describe

121 differences in perinatally and horizontally infected AYALHIV. Although pediatric and adult cohorts

122 linking is being developed [19,20,21], there are only few longitudinal follow-up results for cancer risk

in children and adolescents transitioning to adult care [22].

125 Non-Hodgkin lymphoma

126 A single-center cohort study from the United Kingdom reported on the increased risk of a new 127 lymphoma diagnoses in young adults living with perinatally acquired HIV (PaHIV) following transition 128 to adult care [22]. A total of 5 out of 147 (3.4%) developed lymphoma at a median (range) age of 19 129 (18–23) years. Patients presented with advanced disease (Ann Arbor stage III/IV) mainly diffuse large 130 B-cell lymphomas, a prolonged history of nonadherence, with a life time average of 14 years with 131 detectable viraemia and a low nadir CD4 cell count [157 (90-220) cells/ml]. Small numbers 132 precluded formal risk factor analysis; however, the NHL incidence rate significantly exceeded that of 133 the age matched general UK population; incidence rate ratio 25.9 (95% CI 8.31–61.7), P < 0.0001. 134 Treatment outcomes were not reported. This study echoes a previous report from Italy, describing 135 two cases of Burkitt lymphoma in AYALHIV who were chronically exposed to high-level HIV viremia 136 [23]. These two case series support the concerns of longer term oncogenic risk for the current 137 generation of perinatally infected AYALHIV who experienced prolonged viremia due to late diagnosis 138 and have low rates of viral suppression due to previous inferior ART regimens, suboptimal dosing, 139 nonadherence and the evolution of resistance [22]. Improved HIV diagnosis and linkage to care, 140 adherence support and potent, and well-tolerated ART is required to achieve virological suppression. 141 Greater awareness and prompt investigation of symptoms is needed to diagnose NHL at early stages 142 [22].

143

144 Kaposi sarcoma

We identified one Kaposi sarcoma case series [24] and two cohort studies reporting incidence rates in AYALHIV [25,26]. A cohort study conducted in Uganda and Kenya reported crude Kaposi sarcoma incidence rates for AYALHIV (18–24 years) higher in ART nonusers than in ART users [13]. Incidence rates tended to be higher in young men than in women;[25] although, it is unclear whether this is explained by higher prevalence of HHV-8 [27], delayed access, and poorer adherence to ART in young men or additional factors. Another study from Malawi reported a steady increase in Kaposi 151 sarcoma cases in adolescents per annum despite improved ART coverage [24]. The average annual 152 number of Kaposi sarcoma diagnoses in children and adolescents from 2006 to 2010 (n = 89) was 153 17.8 cases per year, compared to 25.2 cases per year from 2011 to 2015 (n = 126) [24]. This may be 154 explained by better Kaposi sarcoma diagnosis with the improved HIV care [24&]. A third study 155 reported Kaposi sarcoma incidence rates for AYALHIV (aged 16-24 years) from Europe, South Africa, 156 North America, and Asia. In this multiregional cohort analysis adolescents in South Africa had very 157 high Kaposi sarcoma incidence rates (303, 95% CI 176–523) per 100 000 person years, followed by 158 adolescents in Latin America, North America, and Europe (Table 3).

159

160 Invasive cervical cancer

161 Cervical cancer is the fourth leading cause of cancer incidence and mortality for women globally 162 [28,29]. In AYALHIV, one cohort study reported an incidence rate for invasive cervical cancer (ICC) in 163 young women (18–25 years) of 223 (100–496) per 100 000 person years [30]. Women living with HIV 164 have higher hrHPV prevalence [31,32] and more diverse HPV subtypes than their HIV negative 165 counterparts [33,34,32]. HIV-infected young women have high incidence of cervical dysplasia 166 [35];compared to HIV uninfected peers, the rate has been with reported incidence to be three times

167 higher [36].

168

169 Hepatocellular carcinoma

HIV/HBV and HIV/HCV coinfections are associated with an increased risk of liver disease including
HCC in adults; however, there are minimal data in those coinfected either perinatally or in childhood
[37]. Two cases of HCC in adolescents are described in the literature. One male, of black African
origin, with PaHIV/HBV developed a rapidly progressive HCC aged 19 despite more than a decade of
suppressive ART for both HBV and HIV and regular HCC screening. Despite timely surgery, he died of
recurrent metastatic HCC within a year of diagnosis [37]. A second adolescent with PaHIV developed
an HCC but with no evidence of hepatitis coinfection. He had slow disease progression despite being

- severely immunocompromised, with no evidence of recurrence more than a year from surgicalresection [38].
- 179

180 Smooth muscle tumours

A recent study from South Africa reported a case series of EBV-associated smooth muscle tumors in
AYALHIV and adults [39]. Five cases occurred in adolescents (10–15 years) with median CD4 cell
count 616 (range 1–1331) cells/ml; all were female, and all but one survived [39].

184

185 TREATMENT, PROGNOSIS, AND SURVIVORSHIP

186 There were limited published data for cancer outcomes in AYALHIV; however, adult studies suggest 187 disparities in access to cancer treatment and poorer outcomes in adults living with HIV compared to 188 their uninfected peers [40]. A retrospective observational study from Malawi reported treatment 189 outcomes for 70 children and adolescents with HIV (median age 8.6 (1.7–17.9) years) diagnosed with 190 Kaposi sarcoma [41]. Local first-line chemotherapy included bleomycin and vincristine (BV). In 2012, 191 doxorubicin became available in Malawi, which was added for second-line therapy. Paclitaxel was 192 used for the third line. ART-naïve individuals started nevirapine-based ART within 2 weeks of 193 chemotherapy. Of all patients, 28% had severe immunosuppression and nearly half were on ART at 194 time of Kaposi sarcoma diagnosis. The combination of BV was well tolerated with ART, with minimal 195 severe adverse events. Over half (58%) have survived at median follow-up of 29 (15–50) months. 196 Lymphadenopathic Kaposi sarcoma, the most common clinical presentation in children in eastern 197 Africa, was associated with the best outcomes. Kaposi sarcoma with woody edema had a more 198 chronic disease course, whereas visceral disease and Kaposi sarcoma with more than 20 widespread 199 'disseminated' skin/oral lesions were independently associated with increased mortality. Identifying 200 risk factors associated with unfavorable outcomes may be critical to determining which patients will 201 require alternative therapeutic strategies [41].

202 Timely ART initiation in individuals with HIV-related malignancies reduces morbidity associated with 203 opportunistic infections and improves overall survival. However, preexisting HIV-associated organ 204 dysfunction, coexistence of opportunistic infections, compound immunosuppression caused by HIV 205 and chemotherapy, as well as drug interactions between ART and chemotherapy and overlapping 206 treatment-related toxicities make management of patients with HIV and cancer complex. A recent 207 study suggests coadministration of chemotherapy with ART based on integrase strand-transfer 208 inhibitors or nonnucleoside reverse transcriptase inhibitors but not boosted protease inhibitors 209 results in better safety profiles and higher suppressed viral replication [39]. 210 Adult survivors of childhood/adolescent cancer have a lifelong increased morbidity and mortality as 211 well as amplified risk of secondary malignancy [42]. Morbidity may be multisystem impacting on 212 cardiorespiratory, skeletal, renal, neurocognitive, endocrine, and reproductive health compounded

213 with significant psychosocial issues affecting mental health [43]. Annual reviews are recommended

for survivors of childhood cancer for screening, prevention, and treatment of late effects; however,

215 uptake following transition to adult care is poor [44]. AYALHIV who survived malignancy face similar

216 issues compounded by risk of cumulative long-term sequelea of HIV. Potentially, they have an

217 increased risk of a secondary malignancy due to their underlying immune dysregulation and require

218 enhanced support during transition to ensure retention in care and viral suppression.

219

220 PRIMARY AND SECONDARY PREVENTION

Early HIV diagnosis and timely ART may substantially reduce the risk of AIDS-defining cancers
[5,6,10,11,45]. Unlike for HPV, HBV, and HCV, there are no vaccines or specific treatment for EBV
and HHV-8, and early access to suppressive ART remains the most important preventive measure for
cancers related to these infections.

226 High-risk variants of human papilloma virus

227 The high global prevalence of persistent hrHPV infection in both female and male AYALHIV 228 [33,34,46,47,35] and high proportion of high-grade precancerous lesions [35] highlight the 229 importance of sex-neutral HPV vaccination. HPV vaccination induces good HPV-specific cell-230 mediated immune responses in AYALHIV, compared to HIV-uninfected age-matched controls; 231 although, three rather than two doses are still recommended for AYALHIV due to a data gap [48]. 232 Currently, only 11 (6%) countries vaccinate males in their national immunization programs [49]. 233 WHO and American Society of Clinical Oncology (ASCO) guidance prioritize vaccination of girls based 234 on cost-effectiveness analyses for prevention of cervical cancer; boys can be included if the vaccine 235 uptake among priority female population is less than 50% and resources are available [49,50]. A 236 relatively high proportion of hrHPV types in young women in SSA are not covered by currently 237 available HPV vaccines, [31,51] which supports early initiation of cervical screening for all sexually 238 active women living with HIV irrespective of age as recommended by WHO and ASCO [49,52]. A 239 study from Saudi Arabia showed that male circumcision may play role in reduction of HPV infection, 240 penile cancers, and cervical cancer among women with circumcised partners [53]. Screening for anal 241 cancers is not routinely recommended; although, some experts suggest that this might be effective 242 [54,55]. There is an urgent need of prospective studies validating different approaches for 243 prevention and screening of cervical and anogenital cancers.

244

245 Hepatitis B and C

Occurrence of HCC early in adulthood underlines the importance of primary prevention with HBV
 vaccination and screening for chronic HBV and HCV coinfection. Systematic screening for HBV and
 HCV infection is limited in most sub-Saharan African countries [56]. Hepatitis B vaccination from
 birth with serological monitoring and boosting when appropriate, and education around prevention
 of HCV acquisition should be embedded within the life span care of those living with HIV. There is no
 consensus on HCC screening; although, 6 monthly liver ultrasounds and alpha-fetoprotein are

supported by WHO guidance [38,57]. Reducing risk of HCC includes HBV viral suppression with
tenofovir-based regimens, avoidance of excessive alcohol and weight optimization. Increased
advocacy for rapid access to curative direct-acting antivirals for HCV for coinfected adolescents is
urgently required.

Knowledge, awareness, and uptake of sexual and reproductive health services (SRS) is insufficient
among young people [58,59]. Enhanced counselling, integration, or linkage to SRS can improve the
uptake of voluntary male circumcision and cervical cancer screening [60]. AYALHIV require access to
'youth friendly' SRS integrated within multidisciplinary HIV care that includes primary prevention
packages addressing vaccination, ART adherence, smoking [33], alcohol and substance use [61],
weight management, and where appropriate screening for HPV, HBV, and HCV-related malignancies.

262

263 CONCLUSION

264 People living with HIV, including adolescents and young adults, are at increased risk of malignancy, 265 due to immune dysregulation and the persistence of oncogenic viruses. While the excess cancer risk 266 is reduced with suppressive ART, ART coverage is still suboptimal in many settings, and AYALHIV 267 have the lowest rates of engagement with each aspect of the HIV care cascade. Improving HIV 268 diagnosis, linkage, and retention in care on sustained suppressive ART for AYALHIV remains the most 269 important cancer preventive measure. However, this must go hand in hand with integrated cancer 270 screening and education programs including prevention of traditional cancer modifiable risk factors 271 for a vulnerable population who currently face an increased life time risk of malignancy, while they 272 negotiate their transition to adulthood living with HIV. Increased awareness among healthcare 273 workers and prompt investigation of suggestive symptoms is needed to diagnose cancers at early 274 stages. In the era of effective ART, AYALHIV should have access to cancer treatment and supportive 275 care compared to their uninfected peers.

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| | |

491 **TABLES**

- 492 Table 1
- 493 Infection-associated cancers and related clinical HIV stage
- ^aThe Center for Disease Control (CDC) lists NHL, Kaposi sarcoma, and invasive cervical cancer as
- 495 Category C (AIDS-defining) illnesses and leiomyosarcoma as Category B (symptomatic HIV-infection
- 496 entities not included in Category C) illnesses [2]. The WHO lists all of these under Clinical Stage 4 and
- 497 does not classify leiomyosarcoma [3].
- ^bMany other neoplastic disorders, such as anal cancer, oral squamous carcinoma, and testicular
- 499 cancer have been linked to HIV infection and included in the group of non-AIDS-defining illnesses.
- 500 CDC, Center for Disease Control; EBV, Epstein–Barr virus; HBV, hepatitis B virus; HCV, hepatitis C
- 501 virus; HHV-8, human herpes virus-8.

| Malignancy | Association with oncogenic viruses | | | |
|--|---------------------------------------|----------------|--|--|
| AIDS-defining malignancies | | | | |
| Non-Hodgkin lymphoma | | | | |
| Burkitt's lymphoma Large cell (immunoblastic) lymphoma Primary central nervous system lymphoma | EBV EBV | CDC C WHO 4 | | |
| Kaposi sarcoma | HHV-8 | CDC C, WHO 4 | | |
| Invasive cervical carcinoma | HPV | CDC C, WHO 4 | | |
| Non-AIDS-defining malignancies ^b | | | | |
| Smooth muscle tumours | | | | |
| Leiomyoma (benign) Leiomyosarcoma (malignant | EBV) EBV | CDC B | | |
| Hodgkin lymphoma | EBV | - | | |
| Hepatocellular carcinoma | HBV, HCV | - | | |
| Anal cancer | HPV | _ | | |

502

| 504 | Table 2 |
|-----|---|
| 505 | Search strategy for Medline (PubMed) |
| 506 | (('Neoplasms'[Mesh]) OR (neoplasm*[Title/Abstract] OR cancer*[Title/Abstract] OR |
| | |
| 507 | carcinoma*[Title/Abstract] OR tumor*[Title/Abstract] OR Tumor*[Title/Abstract] OR |
| 508 | malignanc*[Title/Abstract] OR leukemic*[Title/Abstract] OR leukemic*[Title/Abstract] OR |
| 509 | hematopoietic stem cell transplantation*[Title/Abstract] OR hematopoietic stem cell |
| 510 | transplantation*[Title/Abstract] OR hematopoietic cell transplantation*[Title/Abstract] OR |
| 511 | hematopoietic cell transplantation[Title/Abstract])) AND |
| 512 | (('Adolescent'[Mesh]) OR ('Young Adult'[Mesh]) OR (adolescen*[Title/Abstract] OR |
| 513 | juvenile*[Title/Abstract] OR youth*[Title/Abstract] OR teen*[Title/Abstract] OR |
| 514 | underage*[Title/Abstract] OR underage[Title/Abstract] OR pubescen*[Title/Abstract] OR young |
| 515 | adult*[Title/Abstract])) AND (Search HIV Infections[MeSH] OR HIV[MeSH] OR hiv[tw] OR HIV-1*[tw] |
| 516 | OR HIV-2*[tw] OR HIV1[tw] OR HIV2[tw] OR HIV infect*[tw] OR human immunodeficiency virus[tw] |
| 517 | OR human immunedeficiency virus[tw] OR human immuno-deficiency virus[tw] OR human immune- |
| 518 | deficiency virus[tw] OR ((human immune*) AND (deficiency virus[tw])) OR acquired |
| 519 | immunodeficiency syndrome[tw] OR acquired immunedeficiency syndrome[tw] OR acquired |
| 520 | immuno-deficiency syndrome[tw] OR acquired immune-deficiency syndrome[tw] OR ((acquired |
| 521 | immune*) AND (deficiency syndrome[tw])) OR 'sexually transmitted diseases, viral'[MH]) |
| 522 | |
| | |

523 Table 3

524 Table 3: Kaposi sarcoma incidence rate in HIV-positive adolescents and young adults who started

525 antiretroviral therapy.

| Study | Country/region | Age group [years] | Rate (95% CI) per 100,000 person-years |
|----------------------|----------------|-------------------|--|
| Semeere et al. [25] | Uganda, Kenya | 18-19 | 245 (79–760) |
| | Uganda, Kenya | 20-24 | 323 (245–426) |
| Rohner et al. [30**] | South Africa | 16–25 | 303 (176–523) |
| | Latin America | 16–25 | 248 (141–438) |
| | North America | 16–25 | 95 (36–253) |
| | Europe | 16–25 | 115 (93–143) |

527

528 FIGURE

- 529 Figure 1
- 530 Estimates for number of adolescents and young adults living with HIV in 2016 stratified by age
- 531 group, sex, and UNAIDS region. Estimates for (i) western, central Europe and North America and (ii)
- eastern Europe and central Asia for adolescents aged 10 to <15 years are not available.
- 533 F, female; M, male. Axis Y: Number of adolescents and young adults. Axis X: UNAIDS regions. Source:
- 534 UNAIDS 2017 estimates [1].

