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5 Mpox in persons with advanced HIV infection: a global case series

6

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78 **ABSTRACT 350**

79 Background. People living with HIV account for 38-50% of those affected in the 2022 multi-80 country mpox outbreak. Most reported cases had high CD4 counts and similar outcomes to 81 those without HIV. Emerging data suggest worse clinical outcomes and higher mortality in 82 more advanced HIV. We describe the clinical characteristics and outcomes of mpox in a cohort of persons with HIV and low CD4 cell counts (CD4 <350 cells/mm³). 83 Methods. A network of clinicians from 19 countries provided data from confirmed mpox 84 85 cases between May 11 and Jan 18, 2023, in persons with HIV infection and CD4 counts <350 cells/mm³. We describe their clinical presentation, complications, and causes of death. 86 Analyses were descriptive. 87 Findings. We include data of 382 cases: 367 cisgender men, 4 cisgender and 10 transgender 88 89 women with a median age of 35 years. At mpox diagnosis, 349 (91.4%) were known to be 90 living with HIV; 228/349 (65.3%) adherent to antiretroviral therapy (ART); 32/382 (8.4%) had 91 a concurrent opportunistic illness. The median CD4 count was 211 cells/mm³ (IQR 117-291); 92 with 85 (22.3%) individuals with CD4 counts <100 cells/mm³, and 94 (24.6%) 100-200 93 cells/mm³. Of those with HIV viral load data available, 193/354 (54.5%) were undetectable. 94 Severe complications were more common in persons with CD4<100 compared to those with 95 >300 cells/mm³, including necrotising skin lesions (54.1% vs. 6.7%), lung involvement (29.4% 96 vs. 0%) occasionally with nodules, and secondary infections and sepsis (43.5% vs. 9.3%). Overall, 107/382 (28.0%) were hospitalised of whom 27/107 (25.2%) died. All deaths 97 occurred in those with CD4 counts <200 cells/mm^{3.} Amongst those with CD4 counts <200 98 cells/mm³ more deaths occurred in those who also had high a HIV viral load. An immune 99 100 reconstitution inflammatory syndrome to mpox was suspected in 21/85 (24.7%) persons

- 101 initiated or re-initiated on ART of whom 12/21 (57.1%) died. Sixty-two (62/382, 16.2%)
- 102 received tecovirimat and 7 (2.2%) cidofovir or brincidofovir; three had laboratory
- 103 confirmation of tecovirimat resistance.
- 104 Interpretation. A severe necrotizing form of mpox in the context of advanced
- 105 immunosuppression appears to behave like an AIDS-defining condition, with a high
- 106 prevalence of fulminant dermatological and systemic manifestations and death.
- 107 Funding. None

109 **RESEARCH IN CONTEXT 635**

110 Evidence before this study

111 In 2022, mpox, a disease caused by an orthopox virus referred to as monkeypox virus (MPXV) 112 has caused outbreaks in 110 countries. Two distinct clades of MPXV, Clade I and Clade II, have 113 existed in different geographic regions. The subclade IIb, identified in the 2022 outbreak, 114 originates from subclade IIa mpox, which is considered a self-limiting disease. Unlike the 115 previous epidemiological descriptions in West Africa, mpox transmission in this outbreak has 116 been closely associated with the sexual networks of gay-and-bisexual men-who-have-sex-117 with-men (GBMSM), in which a high proportion of people are living with HIV. Some evidence suggests greater disease severity in people with advanced HIV. This finding warrants careful 118 119 evaluation of the interplay between HIV, immune status and clinical manifestations of mpox. 120 We searched PubMed for the terms "monkeypox, mpox AND (HIV)" from inception to Dec 31, 121 2022. Publications were predominantly letters, perspectives, case reports, and public health 122 agency reports. In the multi-country outbreak, scientific publications of case series have 123 described similar clinical outcomes in people living with HIV to those in people without HIV 124 infection. However, most cases series included people with HIV and high CD4 counts (> 500 125 cells/mm³) and suppressed HIV viral loads. In contrast, a Nigerian case series in the 2017-18 126 outbreak reported that more severe outcomes in hospitalized people were observed in 127 persons living with HIV especially those who were viraemic and immunosuppressed. In a Center for Disease Control (CDC) report on 758 mpox cases in persons living with HIV during 128 129 the multi-country outbreak (median CD4 639 cells/mm³; 3% < 200/mm³) 10% (68/758) were 130 hospitalized for a median duration of 2 days (0-7). Worse rectal symptoms were described in those with HIV. A second CDC report identified adverse outcomes in 47 people with HIV and 131

mpox who had low CD4 counts, 12(26%) died and 5 deaths were attributed to mpox. Based on these data, individuals with HIV and advanced disease have been identified as cases requiring expert clinical advice, close surveillance and prioritised for anti-viral treatments such as tecovirimat, and preventive vaccines (where available).

136

137 Added value of this study

138 This mpox case series is the largest in those with advanced HIV disease. We characterized

139 382 persons with HIV and CD4 < 350 cells/mm³. Individuals with lower CD4 counts

140 presented with widespread, large, necrotising, and coalescing skin lesions. Some individuals

also developed lung nodules without an alternative confirmed or suspected diagnosis.

142 Severe secondary bacterial infections were common. Frequent and severe oral, ano-genital,

143 and ocular presentations and complications are described. Immune reconstitution

144 inflammatory syndrome (IRIS) was suspected in a quarter of those starting or re-initiating

antiretroviral therapy (ART) after mpox diagnosis, 57% of whom died. The greatest disease

severity, hospitalisation, and mortality was observed in individuals with both low CD4 count

147 and high HIV viral load. This international case series includes 27 of the 60 persons reported

to have died of mpox in the multi-country outbreaks, all 27 are persons with HIV and CD4 <

149 200 cells/mm^{3.}

150

151 Implications of all the available evidence

Our findings support the consideration of a severe, disseminated, and necrotising form of mpox infection as an AIDS-defining condition in CDC and WHO HIV disease classifications. This is based on the observation of protracted illness with fulminant disseminated necrotising

155 cutaneous lesions, systemic complications, and mortality in those with CD4 cell counts <200 156 cells/mm³. Clinicians should also be aware that starting ART in people with advanced HIV and 157 mpox may contribute to deterioration and possible death, possibly as part of an immune 158 reconstitution syndrome. Our data reinforces the importance of HIV testing in mpox cases. 159 Our findings support the recommendations that all at-risk persons with HIV with a CD4 count 160 <200 cells/mm³ should be prioritised for preventive mpox vaccination. There should also be consideration for use of potential mpox antivirals where available despite lacking data on 161 their effectiveness and a concerted global effort to ensure access in countries without access 162 163 to antivirals and vaccines.

164 BODY TEXT

165 INTRODUCTION 313

166 Since May, 2022, around 85,000 human mpox infections have been reported in 110 countries, with transmission predominantly through sexual contact amongst GBMSM.¹ The multi-167 168 country outbreak was declared a Public Health Emergency of International Concern (PHEIC) by the World Health Organisation (WHO) in July 2022.² People with HIV, accounting for 38-169 170 50% of persons diagnosed with mpox,³ have been disproportionately affected.³ Most persons 171 living with HIV described in the 2022 case series had HIV viral suppression with median CD4 counts >500 cells/mm³ and had similar clinical presentations, time to viral clearance, and 172 outcomes to persons without HIV.⁴⁻¹³ 173

174

175 Data from Nigeria and the USA suggest worse clinical outcomes in those with more HIVrelated immunosuppression.^{4,14,15} Two reports from Nigeria during the 2017-2018 outbreak 176 177 suggested that people with advanced HIV presented with more severe or prolonged mpox. The first described that 4 of 7 deaths in 122 individuals with mpox, occurred in people with 178 179 untreated advanced HIV.¹⁴ The second report included 9 people with HIV with CD4 cell counts ranging from 20 to 357 cells/mm^{3.15} The authors described confluent rashes, higher rates of 180 secondary bacterial infections and more prolonged illness.¹⁵ More recently, a report from the 181 US-CDC during the 2022 outbreak, confirmed these finding in 47 cases of severe mpox among 182 people with advanced uncontrolled HIV infection.¹⁶ All were hospitalized, had prolonged 183 184 disease courses, or developed complications, and some had fatal outcomes (i.e., 5 deaths attributed to mpox).¹⁶ Worse rectal disease was described in another CDC series in which 185 82% of people living with HIV were on ART and 72% were virally suppressed.⁴ 186

187

Based on the existing data we hypothesized that in the current outbreak, mpox presentations and outcomes in persons living with HIV may differ by CD4 strata and HIV viral load. We leveraged global research networks to describe the characteristics, clinical course and outcomes of mpox in persons with HIV and CD4 count <350 cells/mm³.

192

193 METHODS

194 Case definition and identification

195 Participating clinicians were recruited through the international research networks of the 196 London-based Sexual Health and HIV All East Research (SHARE) Collaborative and the 197 Network of the Skin Neglected Tropical Diseases and Sexually Transmitted Infections Unit of the Hospital Germans Trias i Pujol in Spain.^{8–10} Researchers in geographic locations with 198 199 high numbers of mpox diagnoses were approached and invited to contribute mpox cases 200 diagnosed between May 11, 2022, and January 18, 2023. A confirmed case was defined as a 201 polymerase-chain-reaction (PCR)-confirmed MPXV infection in a specimen from any 202 anatomical site. We restricted this series to adults older than 18 years living with HIV and 203 CD4 <350 cells/mm³ or, in settings where a CD4 count was not always routinely available, 204 HIV infection clinically classified as CDC stage C. We included people living with HIV and CD4 205 counts <350 cells/mm³ in line with the widely accepted 2010 consensus statement which 206 defines late presentation of HIV as CD4 <350 cells/mm³ or an AIDS-defining illness.¹⁷

CD4 count was categorised as <100, 101-200, 201- 300, and 301-350 cells/mm³, because CD4 208 209 count cut-offs of 100 and 200 are associated with different risk for opportunistic infections 210 (e.g, cryptococcal meningitis is associated with CD4 < 100 cells vs. Pneumocystis jirovecci pneumonia (PJP) or toxoplamosis <200 cells).¹⁸ For strata comparison, we grouped the seven 211 212 individuals with a missing CD4 count with those who had a CD4 count < 100 cells/mm³. Three 213 of these had an AIDS-defining condition and four had a positive point-of-care qualitative CD4 214 count test [Visitect CD4 Lateral Flow assay providing a visually interpreted result of above 200 215 (negative)- or below 200 (positive)]. HIV viral load (VL) was categorised as undetectable (<50 216 copies/mL), 50-200 (low level viraemia), 200-<log4, ≥log4 log RNA copies/ml. We categorised the presence or absence of clinician reported complications by organ system. 217

218

219 Data collection

220 Each contributing centre completed a de-identified structured case-report sheet (CRS) 221 adapted from one used in our prior case series to include variables of interest relevant to 222 persons living with HIV and to capture more severe outcomes (Supplementary Figure 1). The 223 CRS used drop down-menus and free-text fields to capture routinely collected data from 224 electronic or paper medical records. The CRS focus on HIV status included CD4 cell count, HIV 225 viral load, concurrent opportunistic infections, and adherence to ART. These data were 226 included with information on mpox presentation, diagnosis, clinical features, complications, and outcome. We also considered four outcomes for management: outpatient, 227 228 hospitalization, ICU-level care, and death. Duration of hospitalization was the number of days 229 until discharge or until data collection if ongoing by the end of data collection.

230

231	Ethical	consid	erations

Participating clinicians identified individuals living with HIV and diagnosed with mpox infection at their site. Informed consent for inclusion was obtained and maintained in accordance with local standards, along with local institutional review board approval as per each site's local requirement. Image-specific consent was obtained from participants (or their families when deceased) for the use of images. De-identified data were securely transferred to the coordinating site.

238

239 Statistical analysis

All analyses were descriptive, and no hypothesis testing was conducted. Continuous variables were described as the mean and standard deviation (SD) or median and inter-quartile range (IQR). Categorical variables are described as counts and percentages over the entire sample or the corresponding subgroup. No imputation methods were applied to missing data. Data were analysed using R version 4.2.1. Aggregate or de-identified data are presented to avoid deductive disclosure of the identities of study participants.

246

247 Funding

248 There was no specific funding for this study.

249

250 **RESULTS**

We describe 382 cases of human mpox infection in persons living with HIV with CD4 < 350/mm³ from sites in 19 countries (10 in Europe, 8 in the Region of the Americas, and 1 in Africa) (Supplementary Figure 2). Most (72.5%; 277/382) were originally from the Americas

(Argentina, Brazil, Canada, Chile, Ecuador, Mexico, Peru, USA), 25.9% (99/382) were from the
European region (Austria, Belgium, France, Germany, Greece, Italy, Portugal, Spain, Sweden,
Switzerland, UK), 1.6% (6/382) from Africa (Nigeria).

257

The demographic and epidemiological characteristics of the participants are described in Table 1. The median age was 35 years (IQR 30–43). Three hundred and sixty-seven out of 382 (96.1%) identified as cisgender men, 4 cisgender women (1.0%), 10 transgender women (2.6%), and 1 non-binary individual assigned male at birth (0.3%). The ethnicity or race of participants as described by the attending clinician was Black 14.4% (55/382), Latin-American 58.9% (225/382), or White 22.3% (85/382).

264

265 Overall, 349/382 (91.4%) were known to be living with HIV prior to mpox diagnosis, and 266 33/382 (8.6%) were newly diagnosed with HIV infection at the time of the MPXV infection. Of 267 those known to be living with HIV, 84.5% (295/349) were on ART and 65.3% (228/349) were 268 reported as adherent to treatment in the preceding six months. The median CD4 cell count 269 was 211 cells/mm³ (IQR 117-291). The distribution of CD4 counts was as follows: 85 (22.3%) <100 cells/mm³, 94(24.6%) 100-200 cells/mm³, 128(33.5%) 201-300 cells/mm³, and 75 270 271 (19.6%) 301-350 cells/mm³. CD4 counts were missing in 7 individuals who were assigned to 272 the group with CD4 <100 cells/mm³as described in the methods and in Table 1,2,3. Overall, 273 193 individuals (50.5%) were virally suppressed (HIV RNA <50 copies /ml), 26 (6.8%) had low-274 level HIV viraemia (50-200 copies/ml), 30 (7.9%) had viraemia from 200 to log4 copies/ml, 275 and 105 (27.5%) had viraemia of > log4 copies/ml. HIV RNA values taken within the past 6

276 months were not available for 28 individuals (7.3%). Overall, 8.4% of patients (32/382) had a
277 concurrent opportunistic infection at the time of the mpox diagnosis.

278

279 In terms of clinical presentations (Table 2), 243/382 (63.6%) patients had fever and 364 280 (95.3%) had a skin rash, which was initially vesiculo-pustular in 297/382 (77.7%) and 281 progressed to ulcerative in 84/382 (22.0%). The median number of skin lesions was 15 (IQR 282 8-35), and the median duration to resolution was 23 (IQR 18-33) days. Among 36 patients 283 (9.4%) who had 100 or more lesions and 43 patients (11.3%) who had a duration to resolution 284 of forty days or more, the majority had CD4 counts <200 cells/mm³ and detectable HIV plasma 285 viral loads (Supplementary Figure 3 & 4). Overall, 235 individuals (61.5%) had genital, 203 286 (53.1%) anal lesions, 144 (37.7%) oral involvement and 20 (5.2%) had ocular involvement. The 287 most common organ complications were dermatological, respiratory, and secondary bacterial infection (Table 2). A total of 94/382 (24.6%) patients developed dermatological 288 289 complications; 10 (2.6%) of these were ecchymotic or haemorrhagic lesions, and 84 (22.0%) 290 were necrotising lesions, of which 55 (14.4%) were coalescing. The most common 291 presentation was multiple, large (typically greater than 2cm in diameter), rounded ulcers with 292 necrotic centres and a fresh, raised border, located close to the oro-genital regions (Figure 293 1B. Photographs B1-4) or in distant locations (Figure 1B. B6-8), while verrucous appearance 294 (Figure 1B. B5) was rare. In many instances, erythema and oedema surrounded the ulcer. 295 Lesions involving the mouth, eye, or anus resulted in functional impairment (B1-4). In the 296 anogenital region, some individuals presented with significant tissue damage and phagedenic 297 ulcerations (Figure 1C). Some persons had progression to target-shaped lesions with 298 erythema, swelling and pallor beyond the margins of the ulcer indicating severe necrosis. 299 (Figure 1C) Pseudo Koebner phenomena (i.e., the spread of the skin infection along sites with skin microtrauma or rubbing) were manifested by ulcers exhibiting a lineal distribution or
 overlying bony prominences . In cases where epithelialization had occurred, tissue
 destruction resulted in disfiguring scarring (<u>Supplementary Figure 5 &6, and Dermatology</u>
 <u>Atlas</u>).

304

305 In total, 35/382 (9.2%) people presented with respiratory complications (Supplementary 306 Table 1). Of these, eleven individuals (2.9%) presented with numerous bilateral diffuse 307 pulmonary nodules; 4/11 diagnosed with an X-ray only, and 7/11 were further characterized 308 on CT scanning. All the radiographic images were reviewed by two specialist radiologists who 309 concurred that these nodular lesions were unusual and characterised by well-defined 310 borders, absence of cavitation and no adjacent areas of ground glass shadowing; most of the 311 nodules were peri-vascular suggesting hematogenous spread and, generally, ranged in size 312 from of 5 to 20 mm. In all three patients with nodules in whom BAL or lung biopsy were 313 performed, a positive MPXV PCR result was obtained (with negative microbiological results 314 for *P. jirovecci* and *M. tuberculosis*) (Figure 1A). 8/11 had a CD4 count below 100 cells/mm³.

315

12/35 (34.3%) individuals with respiratory complications had dyspnoea of whom - two had
normal chest X-rays, and ten had no available radiology report. Additionally, 6/35 (17.1%)
individuals presented with pleural effusion (1 with a positive MPXV PCR on BAL), and 3/35
(8.6%) with ground glass changes (2 with suspected opportunistic infections; 1 with a positive
MPXV PCR on BAL).

Twelve (3.1%) individuals were reported to have neurological involvement (Supplementary 322 323 Table 2), including one (0.3%) case classified as encephalitis with orbital, frontal and temporal 324 oedema on CT scan, and with a positive MPXV PCR result, and negative HSV-1 and 2, and VZV 325 results in cerebro-spinal fluid (CSF). Of the nine (2.4%) cases with altered mental status or 326 confusion six had normal CSF and/or radiological findings and three did not undergo imaging 327 or CSF examination. In five cases confusion was attributed to sepsis, in one to respiratory 328 failure in one to hepatic encephalopathy, and the cause was undetermined in two cases. 329 Neurological symptoms were almost exclusively described in persons with HIV with CD4 less 330 than 100 cells/mm³.

331

In 76/382 (19.9%) individuals, secondary bacterial infections were diagnosed, including cellulitis, abscesses, and sepsis. Among 17 patients with sepsis, 8 had positive blood cultures and the following pathogens were identified: three *Pseudomonas aeruginosa*, two ESBL *E. coli*, two polymicrobial, and one *Shigella flexneri*. Additionally, 12 patients had a positive result from an abscess or deep wound sample culture: three *Pseudomonas aeruginosa*, two *Klebsiella pneumoniae*, two ESBL *E. coli*, three methicillin-sensitive and two methicillinresistant *Staphylococcus Aureus*.

339

All complications were more common in individuals with CD4<100 compared to individuals
with CD4>300 cells/mm³. This included dermatological (57.6% vs. 9.3%), respiratory (29.4%
vs. 0%), CNS (10.6% vs. 1.3%), bacterial infection (43.5% vs. 9.3%), ocular (15.3% vs. 1.3%),
gastrointestinal (27.1% vs. 6.7%), rectal complications(56.5% vs. 28.0%), oropharyngeal

344 (34.1% vs. 24.0%), and genito-urinary 34.1% vs. 9.3%) (Figure 2A).

345

346	Overall, 107/382 (28.0%) individuals were hospitalised; of these, 7 (1.8%) survived an
347	admission to intensive care and 27 (7.1%) died (Table 3). Among the 27 people who died,
348	the median CD4 count was 35 cells/mm ³ (IQR 24-100), and the median HIV viral load was 5
349	log copies/ml (IQR 4-5), only one patient was HIV virologically suppressed. Among those
350	who died, severe necrotising or haemorrhagic skin lesions occurred in 25/27 (92.6%),
351	bloodstream or deep tissue bacterial infections (24/27; 88.9%), respiratory symptoms and
352	respiratory failure (23/27; 85.2%), neurological (8/27; 29.6%), rectal (21/27; 77.8%), and
353	oropharyngeal (18/27; 66.7%) involvement were described (Table 3). Ocular disease
354	occurred in (13/27; 48.1%), 8 of whom had peri-orbital cellulitis. The reported cause of
355	death was septic shock and multi-organ failure in 20/27 (74.15%), respiratory failure 4/27
356	(14.8%), disseminated mpox in 2/27 (7.4%) and cardiac arrest in 1/27 (3.7%).
357	
358	Rate of hospitalisation and ICU admission increased with declining CD4 counts and rising viral
359	loads <mark>(Figure 2</mark>). No deaths occurred in individuals with CD4 counts >200cells/mm ³ . Mortality
360	was incrementally higher among people in the lowest CD4 strata (CD4<100 27.1% vs. CD4
361	100-200 4.3% vs. CD4>200 0%; <mark>Figure 2B)</mark> and amongst those with the highest viral loads (HIV
362	VL log≥4 16.2% vs. HIV VL undetectable 0.5%; <mark>Figure 2C</mark>). In those with CD4 count <100
363	cells/mm ³ (n=85) and available HIV VL, the death rate was 7.1% (1/14) for individuals with VL
364	<50 copies/ml and 29.8% (14/47) for those with HIV VL ≥4 log copies/ml.

365

366 Among 85 persons started or restarted on ART, in 21 (24.7%) the managing clinician

367 suspected IRIS as a cause for clinical deterioration (Supplementary Table 3). Of these, 6

368 (28.6%) were newly diagnosed and 15 (71.4%) were known to be living with HIV but either 369 not receiving or not adherent to ART. All had CD4 count <200 cells/mm³. The median time 370 from onset of mpox symptoms to the start of ART was 21 days (range 0-73), and from the 371 start of ART to worsening of mpox symptoms was 14 days (range 3-64). Nine of 21 (42.9%) 372 were treated for IRIS with steroids, and 10 (47.6%) received supportive care. Of those with 373 suspected IRIS, 3/21 (14.3%) were admitted to the ICU, 5 (23.8%) were hospitalised in a 374 general ward, and 12 (57.1%) died.

375

376 Forty-three (41.7%) of the 103 hospitalized patients and twenty-one (7.5%) of the 279 outpatients received antivirals to treat mpox. Sixty-two (62/382, 16.2%) individuals received 377 378 tecovirimat (5 received both oral and IV) and 7 (1.8%) cidofovir or brincidofovir. All patients 379 receiving mpox-specific antiviral therapy were treated in Europe or the USA, except two 380 who received tecovirimat in Brazil. Laboratory confirmation of tecovirimat resistance 381 (presence of FL13L mutations by sequencing) was detected in 3/5 people tested, who had 382 severe immunocompromise, disseminated and progressive mpox infection despite 383 prolonged treatment (>14 days) with tecovirimat and finally died. Sampling for resistance 384 testing was conducted after at least one course of tecovirimat had been completed. Nobody 385 who died had received mpox-vaccination prior to or during 2022.

386

387 DISCUSSION

388 Our large case series describes a severe, disseminated form of mpox infection with 15% 389 mortality in individuals with advanced HIV-related disease characterised by CD4 counts below 390 200 cells/mm³. This fulminant form of mpox is characterized by massive necrotising skin,

391 genital and non-genital cutaneous and mucosal lesions, sometimes accompanied by lung 392 involvement with multifocal nodular opacities or respiratory failure, severe cutaneous and 393 bloodstream secondary bacterial infections. The severity of oral and anogenital complications are more marked than previously described.^{4–10,19} As described in the CDC classification, 394 395 disseminated forms of coccidiodomycosis, histoplamosis and mycobacterium avium complex 396 are considered to be AIDS-defining illnesses.²⁰ Very similarly, we describe that people with the lowest CD4 counts (<100 cells/mm³) and highest HIV viral loads (> 4log c/ml) had 397 398 disseminated forms of mpox strongly suggesting that this severe necrotising form of mpox with systemic involvement is also an AIDS-defining condition (Supplement Table 4).²⁰ We 399 400 describe in detail the clinical course of 27 people with CD4 counts < 200 cells/mm³ who died, representing more than 40% of all mpox deaths reported in 2022. We also wish to raise 401 402 awareness of the 57% mortality rate in those in whom IRIS was suspected following ART 403 initiation/re-initiation.

404

This data builds on the observations of the altered natural history and course of mpox that is 405 emerging. To date most information about the intersection of HIV and mpox reports on those 406 with well-controlled HIV infection.⁴⁻¹¹ During the 2022 multi-country outbreak even the 407 largest series include very few people living with HIV and CD4 counts < 350/mm³ (12%) or 408 409 <200/mm³ (3%).^{4,21,22} We hypothesized that mpox may have a different clinical presentation in individuals with advanced immunosuppression, as can be the case with some pathogens. 410 411 Although the self-limiting clinical course in individuals with well-controlled HIV is very similar 412 to that of individuals without HIV, our series provides evidence that the disease is very 413 different in those with advanced HIV. The protracted duration and larger number of skin

414 lesions in these individuals also raises the possibility of a more prolonged period of infectivity,
415 but further studies are needed to investigate this.

416

417 Prior work has shown that people living with HIV with high CD4+ T-cell counts (>350 cells/ mm³) mount a poxvirus-specific T-cell response that is similar to those without HIV infection,²³ 418 419 but there are no data on immunological responses in those with low CD4 counts (<350 cells/ 420 mm³). In our series low CD4 cell count especially when < 200/mm³ was strongly associated with increasing severity of mpox disease relative to those with CD4 200-350/mm³ and 421 422 compared to previous reports where CD4 > 500/mm³. Data from animal model MPXV studies have shown that CD4 depletion before immunisation of non-human primates decreased the 423 development of protective B-cell responses and antibodies, and increased infection 424 425 severity after monkeypox virus challenge, which is consistent with our findings.²⁴ Moreover, 426 in our series, the effect of a low CD4 count on severity or death varied with the HIV plasma 427 viral load, with higher viral loads associated with increased frequency of severe illness in any 428 given CD4 group. Prior work has shown that replicating HIV virions target antigen-specific T 429 cells that are activated to combat other pathogens, resulting in impaired T-cell responses.^{25–} ²⁷ Thus, it is possible that a substantial fraction of MPXV-specific CD4 T cells might die or be 430 431 impaired due to either complete or abortive HIV infection. Others have shown impaired 432 immune responses to hepatitis B and other vaccines in those with low CD4 and unsuppressed HIV virus replication, providing further evidence that HIV replication may interfere with 433 immune response to other pathogens.^{27,28} Based on our findings we believe that a severe 434 435 necrotising form of mpox with systemic manifestations exists. This form of mpox affected those with CD4 counts <200 cells/mm³ - the precise CD4 threshold considered to be AIDS-436

defining in international guidelines (Supplement table 4). Given the 15% mortality in this
group, strong consideration should be given to designating this disseminated form of severe
mpox as a new AIDS-defining condition in definitions and guidelines.

440

441 Several limitations of our research need to be highlighted. Our data are derived from an 442 observational retrospective convenience case series from countries with high numbers of mpox infections. We were, therefore, unable to assess how well our cohort represents the 443 444 entire population of people living with HIV and who developed mpox infection. Study sites 445 may have been more likely to include individuals with more severe outcomes which may have 446 biased the relationships we saw between both CD4 counts and HIV viral load and clinical 447 outcomes. Individuals included in this case series had symptoms that led them to seek 448 medical care; therefore, persons who were asymptomatic, had milder symptoms, or lacked 449 access to medical care could have been missed and thus we may have overestimated illness severity.²⁹ Also outcomes may have been missed if people reattended with severe disease at 450 451 different sites that were not part of the case series. Due to data collection from multiple sites 452 some characteristics may have been collected in a heterogeneous manner, and laboratory 453 techniques also differed from site to site. Many of the cases had a concomitant opportunistic 454 infection and it may be difficult to ascertain many of the outcomes to mpox relative to the 455 other pathogens. For example, some healthcare settings included did not have access to 456 certain radiological and microbiological investigations so we cannot be certain of the role of 457 mpox as opposed to other opportunistic infections. However, we suggest that the coincident 458 perivascular non-cavitating lung nodular pattern described by two radiologists and seen in eleven patients, without a documented suspicion or microbiological evidence of a co-459

460 opportunistic pathogen, may be a manifestation of mpox disease and further research is461 warranted.

Many of the deaths were associated with multiorgan system failure, and the relative contribution of the mpox to death unclear. We have raised the possibility of IRIS reactions with the initiation of antiretroviral therapy but in the absence of a strict definition and in the absence of details on confounding conditions that may have contributed to adverse outcomes the data are uncertain. Data are currently not available from randomised controlled clinical trials on the impact of MPXV antivirals or preventive vaccines on the course of mpox, and with the limited use in this series their role cannot be evaluated.

469

Physicians caring for persons with mpox and advanced HIV disease must be made aware of 470 471 the severe outcomes and high mortality that can occur especially when cutaneous and 472 bloodstream bacterial super-infection set in. Clinical trials tailored to this group are needed 473 to evaluate the impact of antiviral agents and preventive vaccines to modify disease 474 outcomes. In the absence of these data, persons with HIV, low CD4 counts who need to be 475 hospitalised with mpox should be considered for expanded access to these therapies where available. We have raised the risk of deterioration after initiation of antiretroviral therapy 476 477 that carried a 57% mortality rate. This needs to be considered when caring for persons with 478 HIV and advanced disease with mpox who are not on therapy. Further research into the role of IRIS is necessary to better understand the role of potential interventions, such as early 479 480 versus delayed initiation of ART, the concomitant use of steroids or other immunomodulatory 481 strategies leading to a reduction in the frequency of IRIS.

482

483 Our data also reinforces the recommendation that HIV testing (in addition to other sexual 484 transmitted pathogens) be performed in every case of mpox. Further those with HIV 485 infection and high risk for mpox infection should be prioritised for preventive vaccine. Two 486 thirds of the deaths we have reported have occurred in Latin America. Our findings are 487 particularly pertinent for countries with low levels of HIV diagnosis and/or without universal free access to ART and intensive care units, where the interaction of uncontrolled HIV 488 infection and mpox is more prevalent. In these countries, a concerted effort to provide 489 490 urgent access to mpox antivirals and vaccines is of vital importance.

491

492 **CONTRIBUTORS**

493 OM and CMO conceived and designed the study. CMO, and OM co-ordinated the global 494 collaboration. AA and CGC managed the global data collection. CMO, OM, AA, MM, CGC 495 developed the case report form. MM and OM analysed and interpreted and vouch for the 496 data. All authors except MM, and CMO submitted cases. OM, CMO, AA, CGC and MM drafted 497 the first draft of the manuscript. CGC and JV prepared the image library. SW edited the final draft. All authors reviewed the manuscript and vouch for the accuracy and completeness of 498 499 the data. All authors were responsible for the final decision to submit for publication and have 500 seen and approved the manuscript. CMO, MM and OM had full access to all data.

501

502 DATA SHARING

503 De-identified participant data collected, including individual participant data, and will be 504 made available from the corresponding author on reasonable request.

505

506 **DECLARATION OF INTERESTS**

507 We declare no competing interests.

508

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594		

597

596 **FIGURE LEGENDS**

598 **FIGURE 1.** Skin presentation of mpox in advanced HIV disease

599
600 Panel A: Disease progression in a patient with CD4 count of 18 cell/mm3 and viral load log5,
601 with pcr- confirmed lung involvement, bowel perforation, IRIS, and death despite having
602 received two courses of IV tecovirimat and one course of IV cidofovir.

603

604 Panel B: Photographs of necrotizing lesions in multiple patients. Lesions of the skin and 605 mucous membranes. B1: Necrotic ulcers in the peri-labial and nasal areas. Ulcer with tissue 606 destruction on the right upper lip. B2: Umbilicated vesiculopustular-like lesions on upper 607 eyelid surrounding an extensive necrotic ulcer. Eyelids and nasal radix with oedema and erythema. B3: Mucositis, oedema and erosions of the labial mucosa and tongue. B4: Necrotic 608 609 ulcers with raised edges, some confluent, on the scrotum, dorsum of the fingers, groin, and thighs. B5: Numerous verrucous, excrescent, yellowish facial lesions. B6: Multiple 610 periumbilical target-like vesiculopustular lesions, with necrotic depressed centre and 611 612 erythematous halo. **B7**: Large, necrotic, and confluent ulcer on the elbow surrounded by small 613 numerous vesiculopustular lesions. B8: Necrotic ulcers, oedema and erythema on the left 614 hand and wrist.

615

Panel C: Before and after lesions, with progression to severe confluent target-shaped ulcers
with dark necrotic centre surrounded by a vesiculopustular halo and peripheral oedema, in
the perianal area and back.

619

620 Credits: Pictures courtesy of Dr. Judit Villar-García (Figure 1A), Dr Maria Fernanda Peña
621 Vazquez (Figure 1B 1), Dr. Rodríguez Aldama (Figures 1B 3,4,6,7; Figures 1C), Dr Cecilia Agurto
622 Lescano, Dr. Katty Chong Chinchay (Figure1B 5), Dr. Jenny Valverde (Figure 1 B2), Dr. Gonzalez
623 Rodriguez (Figures 2 B8).

624 625

FIGURE 2. Complications stratified by CD4 count (A) and outcomes stratified by CD4 count (B) and viral load (C)

630 TABLE 1. BASELINE DEMOGRAPHIC DATA

						
		TOTAL n (%)	CD4 <100*	CD4 100-200	CD4 201-300	CD4 >300
		N = 382	N = 85	N =94	N =128	N = 75
Age, m years	edian (IQR)	35 (30-43)	35 (32-43)	35 (29 –42)	34(31-42)	36 (30-44)
Gende	r					
	cisgender women	4 (1.0%)	4 (4.7%)	0	0	0
	transgender women	10 (2.6%)	4 (4.7%)	3 (3.2%)	3 (2.3%)	0
	cisgender men	367(96.1%)	77 (90.6%)	91(96.8%)	125(97.7%)	74(98.7%)
	Non-binary individual**	1 (0.3%)	0	0	0	1 (1.3%)
-	where al care was ed					
	Africa	6 (1.6%)	3 (3.5%)	1 (1.1%)	2 (1.6%)	0
	Europe	99 (25.9%)	20 (23.5%)	18 (19.1%)	39 (30.5%)	22 (29.3%)
	Latin-America	212 (54.5%)	37 (43.5%)	65 (69.1%9	67 (52.3%)	43 (57.3%)
	North- America	65 (17.0%)	22 (25.9%)	13 (13.8%)	19 (14.8%)	11 (14.7%)
Ethnici	ty					
	Asian	7 (1.8%)	1 (1.2%)	1 (1.1%)	3 (2.3%)	2 (2.7%)
	Black	55 (14.4%)	26 (30.6%)	10 (10.6%)	14 (10.9%)	5 (6.7%)
	Latin- American	225(58.9%)	44 (51.8%)	63 (67.0%)	76 (59.4%)	42(56.0%)
	Mixed	10 (2.6%)	0	1 (1.1%)	5 (3.9%)	4 (5.3%)
	White	85 (22.3%)	14 (16.5%)	19 (20.2%%)	30 (23.4%)	22 (29.3%)
HIV sta	itus					
	Previously known PLWH currently adherent to ARV	228 (59.7)	17(20%)	53 (56.4%)	100 (78.1%)	58(77.3%)
	Previously known PLWH not on ARV or non- adherent	121(31.6%)	53(62.3%)	33 (35.1%)	25(19.6%)	10 (13.3%)
	Newly diagnosed	33(8.6%)	15(17.6%)	8 (8.5%)	3 (2.3%)	7 (9.3%)

	with HIV infection					
CD4 co (cells/r (IQR)	bunt mm ³), median	211(117-291)	47(27-77)	156 (125- 184)	259 (221- 280)	326 (316- 338)
people	ount among 27 9 who died, mm ³), median	35 (IQR 24- 100)				
	al load strata opies/ml)					
	Not available	28(7.3%)	11 (12.9%)	4 (4.3%)	10 (7.8%)	3 (4%)
	<50	193 (50.5%)	14 (16.5%)	50(53.2%)	80 (62.5%)	49 (65.3%)
	50-200	26(6.8%)	3 (3.5%)	6 (6.4%)	8 (6.3%)	9 (12%)
	201-log4	30 (7.9%)	10 (11.8%)	6 (6.4%)	10 (7.8%)	4 (5.3%)
	≥log4	105 (27.5%)	47 (55.3%)	28 (29.8%)	20 (15.6%)	10 (13.3%)
History	y of mpox					
vaccina	ation					
	Vaccination before 2022	16(4.2%)	2 (2.4%)	4 (4.3%)	7 (5.7%)	3 (4%)
	Third- generation vaccine for	21(5.9%)	4 (4.7%)	3 (3.2%)	9 (7.0%)	5 (6.7%)
	preexposure Third- generation vaccine postexposure	5 (1.3%)	1 (1.2%)	0 (0%)	3 (2.3%)	1 (1.3%)
Concur opport infectio	tunistic					
	Oesophageal candidiasis	4 (1%)	3 (3.5%)	1 (1.1%)	0 (0%)	0 (0%)
	CMV end- organ disease	1 (0.3%)	1 (1.2%)	0 (0%)	0 (0%)	0 (0%)
	Disseminated herpes simplex	1 (0.3%)	1 (1.2%)	0 (0%)	0 (0%)	0 (0%)
	Histoplasmos is	2 (0.5%)	1 (1.2%)	0 (0%)	1 (0.8%)	0 (0%)
	Isosporosis	1 (0.3%)	1 (1.2%)	0 (0%)	0 (0%)	0 (0%)
	Kaposi Sarcoma	4 (1%)	2 (2.4%)	0 (0%)	2 (0.8%)	0 (0%)
	Disseminated Mycobacteriu m Avium Intracellulare	3 (0.8%)	2 (2.4%)	1 (1.1%)	0 (0%)	0 (0%)

Pneumocystis jirovecci pneumonia	6 (1.6%)	5 (5.9%)	1 (1.1%)	0 (0%)	0 (0%)
Toxoplasmosi s	2 (0.5%)	1 (1.2%)	0 (0%)	1 (0%)	0 (0%)
Tuberculosis	8 (2.1%)	5 (5.9%)	1 (1.1%)	2 (0.8%)	0 (0%)

631 PLWH = People living with HIV; ARV = Antiretroviral therapy; Third generation vaccine= MVA - BVN

^{*}For the purpose of the table, seven individuals were classified as CD4 <100 cells/mm³ despite not having formal

633 CD4 counts: three individuals from Peru did not have information on CD4 counts due to lack of testing reagents

but had CDC Stage C disease on the basis of an opportunistic infection; four patients from Nigeria were tested

635 using a qualitative CD4 count test (Visitect CD4 Lateral Flow Assay, visually interpreted result of above or below

636 200 CD4 cells/mm³) with a result of <200 CD4 cells/mm³.

637 **** This non-binary individual was assigned male at birth.

639 TABLE 2. CLINICAL DATA

	TOTAL n (%)	CD4 <100*	CD4 100- 200	CD4 201- 300	CD4 >300
	N = 382	N = 85	N =94	N =128	N = 75
Gastrointestinal					
Overall	55 (14.4%)	23 (27.1%)	14 (14.9%)	13 (6.4%)	5 (6.7%)
Diarrhoea	38 (9.9%)	15 (17.6%)	9 (9.6%)	10 (7.8%)	4 (5.3%)
Gastrointestinal Bleeding	20 (5.2%)	6 (7.1%)	7 (7.4%)	4 (3.1%)	3 (4%)
Obstruction	6 (1.6%)	1 (1.2%)	2 (2.1%)	2 (1.6%)	1 (1.3%)
Oesophagitis	11 (2.9%)	7 (8.2%)	3 (3.2%)	1 (0.8%)	0 (0%)
Highest Care-level					
Outpatient	275 (72.0%)	32(37.6%)	69 (73.4%)	111 (86.7%)	63 (84.0%)
Hospitalization in general ward	73 (19.1%)	26 (30.5%)	19 (20.2%)	16 (12.5%)	12 (16.0%)
ICU-level#	34 (8.9%)	27 (31.8%)	6 (6.4%)	1 (0.8%)	0
					0
Ultimate Outcome					
Death #	27 (7.1%)	23 (27.1%)	4 (4.3%)	0	0
Organ Support					
Need for ventilation	21(5.5%)	16 (18.8%)	4 (4.3%)	1 (0.8%)	0
Indication for ventilation					
Respiratory failure	17 (4.5%)	14 (16.5%)	2 (2.1%)	1 (0.8%)	0 (0%)
Sedation	1 (0.3%)	0 (0%)	1 (1.1%)	0 (0%)	0 (0%)
Low GCS/Coma	3 (0.8%)	2 (2.4%)	1 (1.1%)	0 (0%)	0 (0%)
Need for Inotropes	16 (4.2%)	13 (15.3%)	3 (3.2%)	0 (0%)	0 (0%)
Antimicrobial treatment					
Antibiotics	144 (37.7%)	52 (61.2%)	34 (36.2%)	38 (29.7%)	20 (26.7%)
Tecovirimat (oral)	52(13.6%)	21(24.7%)	11(11.7%)	15 (11.7%)	5 (1.5%)
Tecovirimat (intravenous)	15 (3.9%)	13 (15.3%)	1 (1.1%)	1 (0.8%)	0 (0%)
IVIG	6 (1.6%)	6 (7.1%%)	0 (0%)	0 (0%)	0 (0%)
Cidofovir/Brincidof ovir	7 (1.8%)	5 (5.9%)	2 (2.1%)	0 (0%)	0 (0%)
Genotypic resistance to Tecovirimat					
Samples sequenced	5	4	1	0	0
Presence of F13L mutations	3	3	0	0	0

conferring resistance					
restitution atory syndrome					
Antiretroviral started or restarted	85 (22.3%)	40 (47.1%)	23 (24.5%)	15 (11.70%)	7 (9.3%)
Deterioration consistent with IRIS	21 (5.5)	15(17.6%)	6 (6.4%)	0	0
IRIS treatment provided	19 (5.0%) Steroids 9 NSAIDS 1 Supportive care 9	14 (16.5%) Steroids 8 NSAIDS 1 Supportive care 5	5 (5.3%) Steroids 1 NSAIDS 0 Supportiv e care 4	NA	NA

640 *For the purpose of the table, seven individuals were classified as CD4 <100 cells/mm³ despite not having formal

641 CD4 counts: three individuals from Peru did not have information on CD4 counts due to lack of testing reagents

642 but had CDC Stage C disease on the basis of an opportunistic infection; four patients from Nigeria were tested

643 using a qualitative CD4 count test (Visitect CD4 Lateral Flow Assay, visually interpreted result of above or below

644 200 CD4 cells/mm³) with a result of <200 CD4 cells/mm³.

645 ***The categories within a group of organ involvements are not mutually exclusive; therefore, an individual may*

646 present with multiple manifestations of the group.

647 #All individuals who died received ICU-level care.

648 *‡* Among the 12 patients with dyspnoea, two had a normal chest X-ray, and ten either had no radiology

649 *examinations done or the report was unavailable.*

650

651 Further detail on respiratory and IRIS cases in supplementary tables 1 and 3.

TABLE 3. Detailed information about fatal cases

<u>, , , , , , , , , , , , , , , , , , , </u>			-																	
Patient	Age	Region where medical care was provided	CD4 (cells/mm3)	HIV Viral Load (copies/ml)	HIV and ART Status	Opportunistic Infections	Peak number of lesions	Necrotising skin lesions	Bacterial Infections (Culture result when available)	Respiratory Complications	Ventilatory Support provided *	CNS Complications	Ocular Complications	Rectal Complications	Oropharyngeal Complications	MPX Antiviral Therapy	Started ARV on admission	Suspected IRIS (treatment)	Days from symptom onset to death	Cause of Death
1	35	America s	32	Log 4	Known HIV but not adherent to ART	Oesophagea I candidiasis	300	Yes	Pyomysitis/absces s (skin biopsy: K. Pneumoniae and P. aeruginosa)	None	No	None	None	Pain	Tonsilitis	None - Not available	No	No	51	Shock and Multi-organ failure
2	31	Europe	24	Log5	New diagnosis	None	100	Yes	None	Diffuse perivascular nodules (MPXV positive BAL specimen)	IMV	None	None	None	None	Oral and IV TPOXX, and IVIG	Yes	Yes (nsaids)	196	Shock and Multi-organ failure
3	33	America s	17	Log 5	Known HIV not on ART	PJP*	100	Yes	Sepsis (blood: ESBL <i>E. Coli</i>)	Ground-glass opacification	IMV	None	Periorbi tal cellulitis	Pain	Lymphade nopathy	Oral and IV TPOXX, Cidofovir, and IVIG	Yes	Yes (steroids)	63	Shock and Multi-organ failure
4	46	America s	57	Log 4	Known HIV not on ART	None	100	Yes	Sepsis	Ground-glass opacification (MPXV positive BAL specimen)	IMV	Confu sion	None	Procti tis	Throat Pain	Oral and IV TPOXX, and IVIG	Yes	Yes (steroids)	87	Shock and Multi-organ failure
5	30	America s	121	Log 5	New diagnosis	None	250	Yes	Non-genital cellulitis and sepsis	Diffuse perivascular nodules	IMV	None	Periorbi tal cellulitis	Procti tis	Tonsilitis	None - Not available	Yes	Yes (Supportiv e care)	47	Shock and Multi-organ failure
6	44	America s	106	Log 5	Known HIV not on ART	None	100	Yes	Non-genital cellulitis and sepsis	Diffuse perivascular nodules	IMV	Encep halitis	Periorbi tal cellulitis	Procti tis	Throat Pain	None - Not available	Yes	Yes (Supportiv e care)	49	Shock and Multi-organ failure
7	37	America s	25	<50	Known HIV not on ART	Oesophagea I candidiasis	150	Yes	Necrotising Cellulitis and sepsis (Swab: K. pneumoniae, E. faecalis)	Diffuse perivascular nodules	IMV	Confu sion	None	Procti tis	Tonsilitis	None - Not available	Yes	Yes (Supportiv e care)	38	Respiratory Failure
8	34	Europe	13	Log 5	Known HIV not on ART	None	25	Yes	Perianal and rectal abscesses and sepsis (Blood: ESBL <i>E. Col</i> i)	Diffuse perivascular nodules and pleural effusion (MPXV PCR positive in transthoracic lung biopsy)	NIMV	None	Keratitis	Perfo ration	Lymphade nopathy	Oral and IV TPOXX, and Cidofovir	Yes	Yes (steroids)	117	Shock and Multi-organ failure

9	41	America s	7	Log 5	Known HIV not on ART	None	200	Yes	Sepsis	Diffuse perivascular nodules	IVM	None	None	None	None	None – Not available	No	No	15	Respiratory Failure
10	41	America s	171	Log 6	Known HIV not on ART	PJP*	30	Yes	Sepsis	Ground-glass opacification and large lung cavity	No	None	None	Procti tis	Tonsilitis	None – Not available	No	No	18	Shock and Multi-organ failure
11	32	America s	Unk now n	Log 5	Known HIV not on ART	PJP*	20	Haem orrha gic	Sepsis	Consolidation	No	Confu sion	Periorbi tal cellulitis	Procti tis	Lymphade nopathy	None – Not available	No	No	39	Shock and Multi-organ failure
12	23	America s	Unk now n	Unk now n	New diagnosis	PJP*	15	Yes	Genital cellulitis	Consolidation	No	None	None	Pain	Throat Pain	None – Not available	No	No	18	Respiratory Failure
13	32	America s	Unk now n	Unk now n	Known HIV not on ART	TB*	10	Yes	None	Diffuse perivascular nodules	IMV	None	None	Procti tis	Throat Pain	None – Not available	No	No	32	Shock and Multi-organ failure
14	35	Europe	33	Log 6	New diagnosis	Visceral leishmaniasi s	150	Yes	Necrotising Cellulitis (Blood: <i>P. aeruginosa</i>)	None	IMV	None	None	Pain	Lymphade nopathy	ΙΥ ΤΡΟΧΧ	Yes	Yes (steroids)	87	Respiratory Failure
15	46	America s	6	Log 5	Known HIV but not adherent to ART	Kaposi Sarcoma	50	Yes	Sepsis	Pleural Effusion	IMV	Confu sion	None	Procti tis	None	None - Not available	No	No	40	Shock and Multi-organ failure
16	40	Africa	99	Unk now n	Known HIV not on ART	TB (pulmonary) *	200	Yes	Sepsis	Consolidation, multilobar	No	None	None	Pain	None	None - Not available	Yes	Unknown	26	Shock and Multi-organ failure
17	47	America s	32	Log 4	New diagnosis	None	Not kno wn	Yes	Gluteal abscess & Sepsis (Blood: P. aeruginosa)	None	No	None	Keratitis	Pain	None	Oral TPOXX, Cidofovir and IVIG	Yes	No	94	Shock and Multi-organ failure
18	41	America s	10	Log 5	Known HIV not on ART	None	Not kno wn	Yes	Sepsis (Blood: P. aeruginosa, Clostridum sporogenes)	Shortness of Breath	IMV	None	Periorbi tal cellulitis	Pain	Tonsilitis	Oral and IV TPOXX, Cidofovir, and IVIG	Yes	Yes (steroids)	85	Disseminate d mpox
19	32	America s	58	Log 5	Known HIV not on ART	Oesophagea I candidiasis	Not kno wn	No	Non-genital cellulitis	Pleural effusion	No	Confu sion	Peri- orbital oedema	None	Tonsilitis	Oral and IV TPOXX, and Brincidofovir	Yes	No	78	Cardiac Arrest
20	34	America s	115	Log5	Known HIV not on ART	None	35	Yes	Sepsis	Shortness of Breath	No	None	None	Pain	None	None - Not available	No	No	26	Shock and Multi-organ failure
21	38	America s	70	Log5	Known HIV not on ART	None	32	No	Sepsis	Shortness of Breath	IMV	None	Conjunc tivitis	Pain	None	None - Not available	Yes	Yes (Supportiv e care)	71	Shock and Multi-organ failure
22	32	America s	23	Log5	Known HIV not on ART	CMV retinitis	23	Yes	Sepsis	None	No	None	None	Pain	None	None - Not available	Yes	Unknown	78	Shock and Multi-organ failure
23	48	Africa	90	Unk now n	Known HIV but not adherent to ART	TB (disseminat ed)	100 0	Yes	Sepsis (Swab: K. pneumoniae, P aeruginosa)	Shortness of breath	NIMV	None	Keratitis	None	Throat Pain	None – Not available	Yes	No	25	Shock and Multi-organ failure
24	45	Africa	110	Unk now n	Known HIV not on ART	None	100 0	Yes	Sepsis	Shortness of Breath	NIMV	None	Periorbi tal cellulitis	None	None	None – Not available	No	No	4	Shock and Multi-organ failure

25	28	Africa	99	Unk now n	Known HIV not on ART	TB (pulmonary)	100 0	Yes	None	Shortness of breath	No	Confu sion	Keratitis	None	Throat Pain	None – Not available	No	No	4	Shock and Multi-organ failure
26	29	America s	35	Log5	New diagnosis	None	50	Yes	Necrotising Cellulitis & sepsis (Blood: Polymicrobial)	Pleural effusion	IMV	Confu sion	Periorbi tal cellulitis	Procti tis	Tonsilitis	Oral and IV TPOXX, and IVIG	Yes	Yes (supportiv e care)	83	Disseminate d mpox
27	33	America s	35	Log4	Known HIV not on ART	None	50	Yes	Non-genital cellulitis	Pleural effusion & ulcerative lesions on the trachea (MPXV PCR positive on BAL specimen)	IMV	None	None	Procti tis and Bowe I obstr uctio n	Throat pain	Ιν τροχχ	Yes	Yes (steroids)	46	Shock and Multi-organ failure

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655 Legend: All deceased individuals were cis male, except for patient 16 and 24 that were cis female.

656 None had received smallpox vaccination before 2022 or had been vaccinated as pre-exposure or post-exposure since May 2022. * Since

657 ventilation was often unavailable, answering "no" does not necessarily mean it was not needed.

658 IMV – Invasive Mechanical Ventilation, NIV – Non-Invasive Ventilation

659 *Clinical suspicion, not microbiological confirmation