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Citation: Trickey A, May MT, Vehreschild J, Obel N, Gill MJ, Crane H, et al. (2016) Cause-Specific Mortality in HIV-Positive Patients Who Survived Ten Years after Starting Antiretroviral Therapy. PLoS ONE 11(8): e0160460. doi:10.1371/journal.pone.0160460

Editor: Roberto F. Speck, University Hospital Zurich, SWITZERLAND

Received: April 28, 2016

Accepted: July 19, 2016

Published: August 15, 2016

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Data Availability Statement: Data are from various cohort studies whose authors may be contacted using the email addresses from the list below: the French Hospital Database on HIV (FHDH); Dominique Costagliola—dcostagliola@ccde.chups.jussieu.fr; the Italian Cohort of Antiretroviral-naïve patients (ICONA); Antonella d'Arminio Monforte—antonella.darminio@unimi.it; the Swiss HIV Cohort Study (SHCS); Matthias Cavassini—Matthias.Cavassini@chuv.ch; the AIDS Therapy Evaluation project, Netherlands (ATHENA); Peter Reiss—p.reiss@amc.uva.nl; The Multicenter Study Group on EuroSIDA; Leah Shepherd—

RESEARCH ARTICLE

Cause-Specific Mortality in HIV-Positive Patients Who Survived Ten Years after Starting Antiretroviral Therapy

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¶ Membership of the ART-CC steering group and coordinating committee is provided in the Acknowledgments.

Abstract

Objectives

To estimate mortality rates and prognostic factors in HIV-positive patients who started combination antiretroviral therapy between 1996–1999 and survived for more than ten years.

Methods

We used data from 18 European and North American HIV cohort studies contributing to the Antiretroviral Therapy Cohort Collaboration. We followed up patients from ten years after

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Funding: This work was supported by the UK Medical Research Council (MRC) MR/J002380/1 http://www.mrc.ac.uk/--and the UK Department for International Development (DFID) under the MRC/ DFID Concordat agreement and is also part of the EDCTP2 programme supported by the European Union. Jonathan Sterne is funded by National Institute for Health Research Senior Investigator award NF-SI-0611-10168. Sources of funding of individual cohorts include the Agence Nationale de Recherche sur le SIDA et les hépatites virales (ANRS), the Institut National de la Santé et de la Recherche Médicale (INSERM), the French, Italian and Spanish Ministries of Health, the Swiss National Science Foundation (grant 33CS30_134277), the Ministry of Science and Innovation and the "Spanish Network for AIDS Research (RIS; ISCIII-RETIC RD06/006), the Stichting HIV Monitoring, the European Commission (EuroCoord grant 260694), the British Columbia and Alberta Governments, the National Institutes of Health (NIH) [UW Center for AIDS Research (CFAR) (NIH grant P30 AI027757), UAB CFAR (NIH grant P30 Al027767), The Vanderbilt-Meharry CFAR (NIH grant P30 Al54999), National Institute on Alcohol Abuse and Alcoholism (U10-AA13566, U24-AA020794), the US Department of Veterans Affairs, the Michael Smith Foundation for Health Research, the Canadian Institutes of Health Research, the VHA Office of Research and Development and unrestricted grants from Abbott, Gilead, Tibotec-Upjohn, ViiV Healthcare, MSD,

start of combination antiretroviral therapy. We estimated overall and cause-specific mortality rate ratios for age, sex, transmission through injection drug use, AIDS, CD4 count and HIV-1 RNA.

Results

During 50,593 person years 656/13,011 (5%) patients died. Older age, male sex, injecting drug use transmission, AIDS, and low CD4 count and detectable viral replication ten years after starting combination antiretroviral therapy were associated with higher subsequent mortality. CD4 count at ART start did not predict mortality in models adjusted for patient characteristics ten years after start of antiretroviral therapy. The most frequent causes of death (among 340 classified) were non-AIDS cancer, AIDS, cardiovascular, and liver-related disease. Older age was strongly associated with cardiovascular mortality, injecting drug use transmission with non-AIDS infection and liver-related mortality, and low CD4 and detectable viral replication ten years after starting antiretroviral therapy with AIDS mortality. Five-year mortality risk was <5% in 60% of all patients, and in 30% of those aged over 60 years.

Conclusions

Viral replication, lower CD4 count, prior AIDS, and transmission via injecting drug use continue to predict higher all-cause and AIDS-related mortality in patients treated with combination antiretroviral therapy for over a decade. Deaths from AIDS and non-AIDS infection are less frequent than deaths from other non-AIDS causes.

Introduction

HIV-positive patients who started combination antiretroviral therapy (ART) soon after it became widely available in Europe and North America in 1996 have been treated for up to 20 years[1]. Current mortality rates in these patients are of great interest: they started ART regimens that are less tolerable and have lower antiviral potency than those now available[2] and are likely to have switched ART regimen repeatedly as better drugs became available[3, 4]. As they age these patients are at increased risk of age-related comorbidities such as cardiovascular disease, cancer, liver and renal disease[5, 6]. The proportion of deaths classified as AIDS has decreased over this period[7, 8]. Prognosis beyond the first decade of ART is thus of interest to patients, their treating physicians, mathematical modellers, and those planning health resources.

In the early ART era, mortality was high in the first year of ART and decreased thereafter[9, 10]. For patients who started treatment with immune suppression, AIDS-related causes of death dominate soon after starting ART, but later decline relative to non-AIDS causes[5, 11, 12]. For some successfully treated patient groups mortality rates may be similar to those of the background population[13, 14].

Several factors predict short-term mortality in HIV positive patients starting ART[9, 10], but the prognostic value of some of these factors (for example CD4 count at ART start) diminishes with time[15]. We studied factors that are prognostic for mortality in a large patient cohort who started combination ART without previous exposure to antiretroviral drugs and were treated for a decade. We determined all-cause and cause-specific mortality rates and



GlaxoSmithKline, Pfizer, Bristol Myers Squibb, Roche and Boehringer-Ingelheim. The Danish HIV Cohort Study is founded by Preben and Anne Simonsens Foundation. Data from 11 European cohorts were pooled in June 2014 within COHERE in EuroCoord (www.cohere.org and www.EuroCoord.net).

COHERE receives funding from the European Union Seventh Framework Programme (FP7/2007-2013) under EuroCoord grant agreement n° 260694. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: Abbott, Gilead, Tibotec-Upjohn, ViiV Healthcare, MSD, GlaxoSmithKline, Pfizer, Bristol Myers Squibb, Roche and Boehringer-Ingelheim are commercial organisations that have funded cohorts that provide data to the ART-CC. This does not alter the authors' adherence to PLOS ONE policies on sharing data and materials. identified clinical and demographic risk factors for all-cause and cause-specific mortality present ten years after starting ART.

Methods

The Antiretroviral Therapy Cohort Collaboration (ART-CC) combined data from HIV cohorts in North America and Europe on HIV-1 positive patients aged at least 16 years who started ART with at least three drugs[16] without prior exposure to antiretroviral medications. Further details on the dataset can be found elsewhere (www.art-cohort-collaboration.org). Cohorts were approved by ethics committees or institutional review boards, used standardized methods of data collection, and scheduled follow-up visits at least every six months. Data were analysed anonymously. Eligible patients started ART during 1996–9, remained alive and in follow-up for at least ten years after ART start, and had at least one CD4 count and viral load measurement between nine and ten years after ART start. We used the CD4 count measured before and closest to ART start date plus 10 years. Patients who had stopped or interrupted ART were included. Data analysed were compiled from 18 cohorts (Appendix 1) with follow up until 31st Iuly 2013.

Information relevant to assigning causes of death was obtained either through linkage with Vital Statistics agencies and hospitals or through physician report and active follow-up. We adapted the Cause of Death (CoDe) project protocol[17] (www.cphiv.dk/CoDe.aspx) to classify causes of death. If ICD-10 codes were available, causes of death were classified by a clinician and a computer algorithm[18]. When ICD-10 codes were not available, two clinicians independently classified each death. Disagreements between clinicians and/or computer-assigned codes were resolved via panel discussion[5]. Deaths were coded as AIDS-related if there was a serious AIDS defining condition prior to death and/or a low CD4 count ($<100/\mu$ L) within a year (18 months if off treatment) of death, and a diagnosis compatible with AIDS as cause of death[6]. All other deaths, including those of unknown cause, were considered non-AIDS related. Cause of death information was analysed for six cohorts for which causes had been classified for over 70% of deaths.

Statistical Methods

Follow-up started ten years after ART start and ended at the earliest of death, loss to follow-up (LTFU) or administrative censoring (cohort-specific database close date). Patients were considered LTFU at their last clinical observation if this was over a year before the close date. AIDS was categorised as: no recorded AIDS diagnosis; first AIDS diagnosis before starting ART; and first AIDS diagnosis between starting ART and start of follow-up. Drug regimens were derived at ART start and start of follow-up and were classified as NNRTI-based, PI-based, other regimen, or no drugs at start of follow-up.

We compared characteristics of eligible patients with those no longer followed-up at ten years after start of ART because they died, were LTFU or transferred to another treatment centre. We estimated unadjusted mortality rates assuming constant rates during follow-up. Five-year cumulative mortality was estimated using the Kaplan-Meier method. We used Cox models to estimate unadjusted and adjusted hazard ratios (HR): adjustments were for sex, regimen, age $(16-39, 40-49, 50-59, 60-69, \ge 70 \text{ years})$, mode of transmission (injection drug use [IDU] or not), and HIV-1 RNA $(0-50, 51-1000, \ge 1001 \text{copies/mL})$, CD4 count $(0-99, 100-199, 200-349, 350-499, 500-749, \ge 750 \text{ cells/µL})$ and AIDS status at start of follow-up, chosen due to prior literature[19]. We investigated whether CD4 count at ART start was associated with mortality after accounting for the association of CD4 count 10 years after ART start.



We tabulated frequencies of AIDS, non-AIDS, and specific causes of death. We estimated adjusted HR for the more frequent ($n \ge 10$) specific causes of death (AIDS, cardiovascular, liver-related, malignancies (non-AIDS and not hepatitis-related), suicide/accidents) stratified by age (\geq 60 vs. <60 years), sex, IDU status, CD4 count (<200 vs. \geq 200 cells/ μ L), viral suppression (HIV-1 RNA >50 vs. ≤50 copies/mL), and AIDS diagnosis during or before the first decade of ART (vs. no AIDS). Five-year cumulative risks of death, according to age, risk group and CD4 count, viral load and AIDS status recorded ten years after starting ART, were estimated using cumulative incidence functions from a flexible Weibull model [20], stratified by IDU and CD4 count. We estimated five-year mortality risk for each patient and tabulated the frequency distribution of patients grouped by five-year mortality risk (0-1.99, 2-4.99, 5-9.99, 10-24.99 and $\geq 25\%$) by age (<60 and ≥ 60 years). We compared five-year mortality risk in HIV-positive individuals with that of age-matched French general population in 2013 using mortality.org data. We chose the French population as comparator because the largest proportion of patients were treated in France. As a sensitivity analysis we used competing risks regression with LTFU and death as the competing outcomes[21]. Analyses were done using Stata 13 (StataCorp, Texas, USA).

Results

24,445 patients started ART between 1996–1999. Of these, 3,577 (15%) died, 5,963 (24%) were LTFU during the first decade of ART, and 1,894 (8%) were excluded because CD4 count or viral load were not available at start of follow-up leaving 13,011 (53%) patients eligible for analyses. Table 1 compares characteristics at ART start of eligible patients with excluded patients who died, were LTFU, or lacked a measurement of CD4 and viral load between nine and ten years after ART start. The two groups had similar age distributions, but eligible patients were less likely to be IDU (1,636 [13%] vs 2,623 [23%]), had higher CD4 count and lower viral load at ART start.

Among 13,011 patients during follow-up from 10 years after starting ART, there were 656 deaths during 50,593 person years giving a crude mortality rate of 12.9 [95% CI 12.0,14.0] per 1,000 person years, and estimated five year survival 93.5% (93.1–94.2%). Median (IQR) follow-up was 4.0 (3.1–5.0) years and maximum follow-up time varied between cohorts from 4.5 to 7.1 years. The rate of LTFU was 5.5 (95% CI 5.3–5.7) per 100 person years (N = 5,963). Patients LTFU after 10 years were more likely to be female, IDU and younger (Table 2). Median (IQR) CD4 count at ten years after starting ART was lower in those who died, but was similar in those LTFU or who remained in care: 383 (185–554), 535 (350–740), and 566 (390–774) cells/ μ L, respectively. The proportions of patients with HIV-RNA \leq 50 copies/mL at ten years after starting ART was lower in those who died, but similar in those LTFU or who remained in care: 56%, 73%, and 77%, respectively (Table 2).

Patient demographics and clinical characteristics at start of follow-up (ten years after starting ART) are shown in Table 3. Median (IQR) CD4 count was 250 (100–401) cells/ μ L at ART start and increased to 550 (371–760) cells/ μ L at ten years. The correlation between CD4 count at ART start and at ten years was 0.27 (p<0.001). Most patients (11,504; 88%) had HIV-1 RNA \leq 200 copies/mL ten years after ART start. At ART start most patients (10,266; 79%) were on PI-based regimens with 2,229 (17%) on NNRTI-based regimens. By ten years 4,555 (35%) and 5,002 (39%) of patients were on PI- and NNRTI-based regimens, respectively, whilst 1,072 (8%) were off ART. Of the remaining patients, 1,853 (14%) were only receiving NRTIs, 108 (1%) and 9 (0.1%) were on CCR5 receptor antagonist- and integrase inhibitor-based regimens respectively and 3% were on other regimens. Of 400 (3%) patients with CD4 count <100 cells/ μ L ten years after ART start, 71 (18%) were not on treatment, 1,001 (8%) of those with CD4



Table 1. Characteristics (at start of ART) of patients who were and were not eligible for analyses of prognosis from 10 years after start of ART.

	Patients who started AF excluded fro		Patients included in analyses of prognosis from 10 years after start of ART			
Characteristics	Number (%) of patients	Number (%) of deaths	Numbers (%) of patients			
Total	11,434 (100%)		13,011 (100%)			
IDU	2,623 (23%)	820 (31%)	1,636 (13%)			
Female	2,215 (19%)	472 (21%)	2,701 (21%)			
AIDS diagnosis	2,613 (23%)	1,250 (48%)	2,714 (21%)			
Age (years)						
Median (IQR)	36 (31, 43)		36 (31,43)			
16–29	2,127 (19%)	330 (16%)	2,330 (18%)			
30–39	5,147 (45%)	1,335 (26%)	6,029 (46%)			
40–49	2,600 (23%)	1,148 (44%)	3,129 (24%)			
50–59	1,089 (10%)	614 (56%)	1,181 (9%)			
≥60	471 (4%)	296 (63%)	342 (3%)			
CD4 count (cell/µL)						
Median (IQR)	224 (85, 335)		250 (100, 401)			
≥750	381 (3%)	70 (19%)	423 (3%)			
500–749	1,166 (10%)	251 (22%)	1,483 (11%)			
350–499	1,868 (16%)	408 (22%)	2,390 (18%)			
200–349	2,799 (24%)	776 (28%)	3,372 (26%)			
100–199	2,092 (18%)	775 (37%)	2,120 (16%)			
50–99	1,103 (10%)	469 (43%)	1,229 (9%)			
25–49	760 (7%)	340 (45%)	748 (6%)			
0–24	1,265 (11%)	634 (50%)	1,246 (10%)			
HIV 1 RNA cell count						
Median (IQR)	7.4×10 ⁴ (1.7×10 ⁴ , 2.4×10 ⁵)		7.1×10 ⁴ (1.7×10 ⁴ , 2.3×10 ⁵)			
0–50	104 (1%)	11 (11%)	88 (1%)			
51–1000	580 (5%)	106 (18%)	669 (5%)			
1001–100000	5,766 (50%)	1,645 (29%)	6,794 (52%)			
>100000	4,984 (44%)	1,961 (39%)	5,460 (42%)			

LTFU: lost to follow-up; IDU: injection drug use; IQR: inter-quartile range

doi:10.1371/journal.pone.0160460.t001

count $\geq\!100$ cells/µL ten years after ART start were not on treatment. The proportions of virally suppressed patients (HIV-1 RNA <50 copies/mL) were 30% and 77% for those with CD4 counts <100 and $\geq\!100$ cells/µL, respectively. There were 6,287 (48%) patients with HIV-1 RNA <50 copies/mL and with CD4 count $\geq\!500$ cells/µL.

Mortality from ten years after start of ART

Mortality was substantially higher in patients aged 60–69 and >70 years from ten years after starting ART (adjusted HR 4.29 [95% CI 3.06–6.02] and 8.59 [5.82–12.7] respectively compared with age 16–39 (<u>Table 3</u>)). Mortality was also substantially higher in patients with CD4 count <100 cells/µL from ten years after ART start (adjusted HR 6.17 [4.46–8.53] compared with \geq 750 cells/µL). Mortality rates were similar in patients with CD4 count 500–749 and \geq 750 cells/µL. Mortality was higher in IDU compared with non-IDU (adjusted HR 2.58 [2.04–3.26]). Viral load >1000 copies/mL at ten years and an AIDS diagnosis in the first ten years



Table 2. Characteristics of patients followed from 10 years after start of ART who died, were lost to follow up (LTFU) or remained in the study until end of follow up.

	Number (%) of patients				
	Died (N = 656)	LTFU (N = 2,708)	Remained in the study (N = 9,647)		
IDU	121 (18%)	474 (18%)	1,041 (11%)		
Female	81 (12%)	669 (25%)	1,951 (20%)		
No AIDS	302 (46%)	1,867 (69%)	6,492 (67%)		
AIDS before ART start	175 (27%)	537 (20%)	2,002 (21%)		
AIDS between ART start and 10 years afterwards	179 (27%)	304 (11%)	1,153 (12%)		
Age (years)					
Median (IQR)	52 (45, 60)	45 (40, 51)	46 (41, 53)		
16–39	53 (8%)	606 (22%)	1,671 (17%)		
40–49	225 (34%)	1,287 (48%)	4,517 (47%)		
50–59	200 (30%)	570 (21%)	2,359 (24%)		
≥60	178 (27%)	245 (9%)	1,100 (11%)		
CD4 count (cells/µL)					
Median (IQR)	383 (185, 554)	535 (350, 740)	566 (390, 774)		
0–99	98 (15%)	97 (4%)	205 (2%)		
100–199	81 (12%)	176 (7%)	371 (4%)		
200–349	131 (20%)	398 (15%)	1,321 (14%)		
350–499	128 (20%)	532 (20%)	1,984 (21%)		
500–749	126 (19%)	856 (32%)	3,111 (32%)		
≥750	92 (14%)	649 (24%)	2655 (28%)		
Viral load (HIV-1 RNA copies/mL)					
% with RNA≤50	56%	73%	77%		
0–50	370 (56%)	1,977 (73%)	7,467 (77%)		
51–1000	119 (18%)	365 (13%)	1,206 (13%)		
>1000	167 (25%)	366 (14%)	974 (10%)		

IDU: injection drug use; IQR: inter-quartile range

doi:10.1371/journal.pone.0160460.t002

after starting ART were associated with higher subsequent mortality (adjusted HR 1.74 [1.40–2.15] and 1.68 [1.37–2.06] respectively). There was no difference in subsequent mortality between the patients on PI- (adjusted HR 1.17 [0.95–1.43] compared with NNRTI-) and NNRTI-based regimens at 10 years after ART start.

Lower CD4 count at ART start was weakly associated with higher mortality from ten years (Table 4). This was attenuated by adjustment for CD4 count at ten years after starting ART and other covariates, with a suggestion of lower subsequent mortality for patients with very low CD4 count at ART start after adjustment for ten year CD4 count [HR = 0.62 (0.46–0.83) for CD4 <50 vs. \ge 350 cells/ μ L].

Results were similar to those in the main analysis when LTFU was considered a competing risk to death in the sensitivity analysis.

Five-year mortality risk

Table 5 shows estimated five-year mortality risk from ten years after ART start in groups defined by combinations of age, IDU risk group, AIDS status, CD4 count and viral suppression. For patients aged 40–49 years at ten years after starting ART the subsequent five-year mortality risk ranged from 1.9% (1.5–2.4%) in those who were non-IDU and virally suppressed with CD4 count >500 cells/µL and no AIDS, to 48% (40–58%) in those who were IDU and not



Table 3. Characteristics of eligible patients 10 years after start of ART, together with unadjusted and adjusted hazard ratios (HR).

Characteristics	Number (%) of patients	Number (%) of deaths	Unadjusted HR	Adjusted HR (95% CI)*	
			(95% CI)		
Total	13,011 (100%)	<u>656</u> (100%)			
Risk group					
IDU (vs non-IDU)	1,636 (13%)	121 (7%)	2.54 (2.03, 3.18)	2.58 (2.04, 3.26)	
Sex					
Female (vs male)	2,701 (21%)	81 (3%)	0.76 (0.60, 0.96)	0.95 (0.74, 1.21)	
AIDS diagnosis					
No AIDS	8,661 (67%)	302 (3%)	1	1	
AIDS before ART start	2,714 (21%)	175 (6%)	1.73 (1.43, 2.09)	1.48 (1.23, 1.80)	
AIDS between ART start and 10 years afterwards	1,636 (13%)	179 (11%)	2.14 (1.75, 2.62)	1.68 (1.37, 2.06)	
Age (years)					
Median (IQR)	46 (41, 53)				
16–39	2,330 (18%)	53 (2%)	1	1	
40–49	6,029 (46%)	225 (4%)	1.50 (1.11, 2.03)	1.40 (1.04, 1.90)	
50–59	3,129 (24%)	200 (6%)	2.15 (1.58, 2.92)	2.14 (1.56, 2.92)	
60–69	1,181 (9%)	123 (10%)	3.39 (2.44, 4.71)	4.29 (3.06, 6.02)	
≥70	342 (3%)	55 (16%)	6.18 (4.22, 9.04)	8.59 (5.82, 12.7)	
CD4 count (cells/µL)					
Median (IQR)	550 (372, 760)				
≥750	3,396 (26%)	92 (3%)	1	1	
500–749	4,093 (31%)	126 (3%)	1.12 (0.85, 1.46)	1.01 (0.78, 1.33)	
350–499	2,644 (20%)	128 (5%)	1.72 (1.32, 2.25)	1.41 (1.07, 1.84)	
200–349	1,850 (14%)	131 (7%)	2.37 (1.81, 3.10)	1.80 (1.37, 2.37)	
100–199	628 (5%)	81 (13%)	4.28 (3.16, 5.78)	3.07 (2.24, 4.19)	
0–99	400 (3%)	98 (25%)	9.25 (6.90, 12.4)	6.17 (4.46, 8.53)	
Viral load (HIV-1 RNA copies/mL)					
Median (% with RNA≤50)	50 (75%)				
0–50	9,814 (75%)	370 (4%)	1	1	
51–1000	1,690 (13%)	119 (7%)	1.25 (1.00, 1.56)	1.09 (0.87, 1.36)	
>1000	1,507 (12%)	167 (11%)	2.41 (1.99, 2.91)	1.74 (1.40, 2.15)	

IDU: injection drug use; IQR: inter-quartile range; CI: confidence interval

doi:10.1371/journal.pone.0160460.t003

virally suppressed with CD4 count <100 cells/ μ L and AIDS during the first decade of ART. In comparison, the 5-year mortality risk in the French general population of the same age was 1%. Fig 1 shows the proportion of patients in five-year mortality risk strata, for younger (aged <60 years) and older (\geq 60) patients. Five-year mortality risk was below 25% in 12,649 (97%) patients, below 10% in 10,715 (82%) patients and below 5% in 7,829 (60%). Mortality risk was strongly age-related: only 384 (29%) patients aged \geq 60 years had five-year mortality risk less than 10%.

Cause-specific mortality from ten years after starting ART

There were 218 deaths among 7,178 patients in the eight cohorts with \geq 70% of cause of death information coded. It was possible to classify a cause of death in 180 (83%): the others were coded unknown/missing. The most frequent cause of death was non-AIDS non-liver cancer (41 deaths [25% of those classified]) followed by AIDS (35 [19%]), cardiovascular (22 [12%])

^{*}Models adjusted for all variables in table, stratified by cohort.



Table 4. Unadjusted and adjusted hazard ratios (HR) according to CD4 count at start of ART and at 10 years after start of ART.

	Cells/µL	N (%)	Unadjusted HR (95% CI)	Adjusted HR (95% CI)*
CD4 count at start of ART (cells/µL)	≥350	4,296 (33%)	1	1
	200–349	3,372 (26%)	1.09 (0.88, 1.35)	0.96 (0.77, 1.19)
	100–199	2,120 (16%)	1.38 (1.09, 1.73)	0.94 (0.74, 1.20)
	50–99	1,229 (9%)	1.60 (1.24, 2.08)	0.88 (0.67, 1.16)
	0–49	1,994 (15%)	1.19 (0.94, 1.52)	0.69 (0.52, 0.90)
CD4 10 years after start of ART (cells/µL)	≥750	3,396 (26%)	1	1
	500–749	4,093 (31%)	1.11 (0.85, 1.46)	1.04 (0.80, 1.37)
	350–499	2,644 (20%)	1.72 (1.32, 2.25)	1.47 (1.12, 1.93)
	200–349	1,850 (14%)	2.37 (1.81, 3.10)	1.92 (1.45, 2.54)
	100–199	628 (5%)	4.28 (3.16, 5.78)	3.33 (2.42, 4.60)
	0–99	400 (3%)	9.25 (6.90, 12.40)	6.85 (4.89, 9.60)

CI: Confidence interval

doi:10.1371/journal.pone.0160460.t004

and liver-related (18 [10%]). Adjusted HRs for specific causes of death stratified by age, sex, CD4 count, viral suppression and AIDS are shown in Table 6. AIDS mortality was strongly related to lower CD4 count and viral replication at ten years after starting ART. Cardiovascular mortality was substantially higher in older patients (adjusted HR 8.95 [95% CI 3.63–22.1] comparing patients aged \geq 60 with <60 years at ten years after ART start). Transmission via IDU and lower CD4 count at ten years after starting ART were strongly associated with mortality from liver-disease. Non-AIDS mortality was associated with both older age, adjusted HR compared with those aged <60 years 4.35 [95% CI 2.97–6.39], IDU status (3.40 [2.26–5.12]), lower CD4 (2.42 [1.49–3.94]) and prior AIDS diagnosis (1.58 [1.27–1.95]). Deaths from suicide/accident were higher in those with a prior AIDS diagnosis. Other causes of death (combining all causes with fewer than ten deaths) were strongly associated with older age (adjusted HR 5.71 [2.84,11.5]).

Discussion

Summary

Among HIV positive patients who started combination ART 1996–1999 without prior exposure to antiretroviral drugs and survived for at least ten years, the majority were virally suppressed and had CD4 counts ≥ 500 cells/µL. Nonetheless lower CD4 count and lack of HIV-1 viral suppression at ten years after starting ART, and AIDS before or during the first decade of ART, were strong predictors of death during the second decade of ART. Low CD4 count at ART start was no longer associated with increased mortality beyond ten years after accounting for CD4 count measured at ten years after starting ART. Compared with patients from other risk groups, those who reported IDU as route of HIV transmission experienced higher mortality from ten years after ART start. The most frequently occurring causes of death were non-AIDS-defining malignancy, followed by AIDS, cardiovascular, and liver-related causes. Older age was strongly associated with cardiovascular mortality and with non-AIDS non-liver malignancies, IDU with non-AIDS infection and liver-related mortality, and low CD4 count and detectable viral replication at ten years after starting ART with AIDS mortality. The five-year mortality risk was <5% in 60% of patients and was strongly age-related: fewer than 30% of patients aged over 60 had five-year mortality risk less than 10%

^{*}Adjusted for sex, regimen, age (16–39, 40–49, 50–59, 60–69, \geq 70 years), mode of transmission (IDU or not), HIV-1 RNA levels 10 years after start of ART (0–50, 51–1000, >1000 copies/ml), AIDS status and mutually adjusted for both CD4 measurements and stratified by cohort.



Table 5. 5-year percentage risk of death (95% CI) from 10 years after start of ART, according to age, IDU risk group, AIDS status, CD4 count and viral suppression (HIV RNA <50 copies/mL)*.

	CD4 count 10 years after start of ART (cells/µL)							
	0–99	100–199	200-349	350–499	500-749	>750		
Non-IDU, No	AIDS, Virologically su	ppressed	-					
Age 16–39	6.9 (4.9–9.8)	3.9 (2.7–5.5)	2.2 (1.6–3)	1.7 (1.2–2.3)	1.2 (0.9–1.6)	1.2 (0.8–1.6)		
Age 40–49	10.9 (8.2–14.3)	6.1 (4.6–8.1)	3.5 (2.7–4.4)	2.7 (2.1–3.4)	1.9 (1.5–2.4)	1.9 (1.5–2.4)		
Age 50–59	18.8 (14.3–24.4)	10.8 (8.2–14.2)	6.1 (4.8–7.8)	4.8 (3.8–6)	3.4 (2.7-4.3)	3.3 (2.6–4.3)		
Age 60+	36.4 (28.2–46.2)	22.1 (16.9–28.5)	12.9 (10.3–16.2)	10.1 (8.1–12.7)	7.2 (5.7–9.1)	7.1 (5.5–9.1)		
Non-IDU, No	AIDS, Not virologically	suppressed						
Age 16–39	11.3 (8.2–15.4)	6.4 (4.6–8.8)	3.6 (2.6–4.9)	2.8 (2–3.8)	2 (1.4–2.7)	1.9 (1.4–2.7)		
Age 40–49	17.5 (13.7–22.1)	10 (7.7–13.1)	5.7 (4.5–7.2)	4.4 (3.5–5.6)	3.1 (2.4–4)	3.1 (2.4–4)		
Age 50–59	29.3 (23.3–36.4)	17.4 (13.4–22.3)	10 (7.9–12.6)	7.8 (6.1–9.9)	5.6 (4.4–7.1)	5.5 (4.2–7.2)		
Age 60+	53 (43.1–63.6)	34 (26.6–42.8)	20.6 (16.4–25.7)	16.3 (12.8–20.6)	11.8 (9.1–15.1)	11.6 (8.8–15.2)		
Non-IDU, AID	S, Virologically suppr	essed						
Age 16–39	11.3 (8.1–15.8)	6.4 (4.6–9)	3.6 (2.6–4.9)	2.8 (2–3.8)	2 (1.4–2.7)	1.9 (1.4–2.7)		
Age 40–49	17.6 (13.7–22.4)	10.1 (7.7–13.1)	5.7 (4.5–7.2)	4.4 (3.5–5.6)	3.2 (2.5–4)	3.1 (2.4–4)		
Age 50–59	29.4 (23.3–36.8)	17.5 (13.6–22.3)	10.1 (8.1–12.6)	7.9 (6.3–9.8)	5.6 (4.5–7)	5.5 (4.3–7.1)		
Age 60+	53.2 (43.4–63.8)	34.2 (27.1–42.6)	20.7 (16.8–25.5)	16.4 (13.2–20.2)	11.8 (9.5–14.8)	11.7 (9.1–14.9)		
Non-IDU, AID	S, Not virologically su	ppressed						
Age 16–39	18.2 (13.4–24.3)	10.5 (7.6–14.4)	5.9 (4.3–8.1)	4.6 (3.3–6.3)	3.3 (2.4–4.5)	3.2 (2.3–4.6)		
Age 40–49	27.6 (22.3–33.7)	16.3 (12.7–20.7)	9.4 (7.4–11.8)	7.3 (5.7–9.3)	5.2 (4–6.7)	5.1 (3.9–6.8)		
Age 50–59	44.1 (36.7–52.2)	27.4 (21.9–33.9)	16.3 (13.1–20.1)	12.8 (10.2–16)	9.2 (7.2–11.7)	9 (6.9–11.8)		
Age 60+	71.8 (62.1–80.9)	50.2 (41–60.2)	32.1 (26.1–39.1)	25.8 (20.7–31.9)	19 (14.9–24)	18.7 (14.2–24.3)		
IDU, No AIDS	Virologically suppres	ssed						
Age 16–39	13.5 (9.5–19.2)	7.7 (5.4–11)	4.3 (3.1–6.1)	3.4 (2.4–4.7)	2.4 (1.7–3.4)	2.3 (1.6–3.3)		
Age 40–49	20.9 (15.8–27.2)	12.1 (9–16.1)	6.9 (5.3–8.9)	5.4 (4.1–7)	3.8 (2.9–5)	3.7 (2.8–5)		
Age 50–59	34.4 (26–44.6)	20.7 (15.4–27.6)	12.1 (9.1–15.9)	9.5 (7.1–12.5)	6.8 (5–9)	6.6 (4.9–9)		
Age 60+	60.1 (47.3–73.3)	39.7 (30–51.3)	24.5 (18.5–31.9)	19.5 (14.6–25.7)	14.1 (10.5–18.9)	13.9 (10.2–19)		
IDU, No AIDS	Not virologically sup	pressed						
Age 16–39	21.5 (15.6–29.3)	12.5 (8.9–17.5)	7.1 (5.1–9.9)	5.6 (3.9–7.8)	3.9 (2.8–5.6)	3.9 (2.7–5.6)		
Age 40–49	32.3 (25.5–40.4)	19.3 (14.7–25.2)	11.2 (8.6–14.6)	8.8 (6.6–11.5)	6.3 (4.7–8.3)	6.2 (4.5–8.4)		
Age 50–59	50.5 (40.2–61.8)	32.1 (24.4–41.5)	19.3 (14.7–25.2)	15.3 (11.4–20.3)	11 (8.1–14.9)	10.8 (7.8–14.9)		
Age 60+	78.4 (66.2–88.6)	57 (44.7–70)	37.4 (28.7–47.8)	30.3 (22.8–39.6)	22.5 (16.5–30.2)	22.1 (15.9–30.3)		
IDU, AIDS, Vir	ologically suppresse	d						
Age 16–39	21.7 (15.4–29.9)	12.6 (8.8–17.7)	7.2 (5.1–10.1)	5.6 (3.9–7.9)	4 (2.8–5.6)	3.9 (2.7–5.6)		
Age 40–49	32.5 (25.4–40.9)	19.4 (14.8–25.3)	11.3 (8.7–14.6)	8.8 (6.8–11.5)	6.3 (4.8–8.3)	6.2 (4.6–8.3)		
Age 50–59	50.7 (40.3–62.2)	32.3 (24.7–41.5)	19.4 (14.9–25.2)	15.3 (11.6–20.1)	11.1 (8.3–14.7)	10.9 (8–14.8)		
Age 60+	78.6 (66.5–88.7)	57.3 (45.4–69.7)	37.6 (29.2–47.4)	30.5 (23.3–39.2)	22.6 (16.9–29.8)	22.3 (16.3–29.9)		
IDU, AIDS, No	t virologically suppre	ssed						
Age 16–39	33.4 (24.9–44)	20.1 (14.4–27.6)	11.7 (8.4–16.2)	9.1 (6.4–12.9)	6.5 (4.5–9.3)	6.4 (4.4–9.4)		
Age 40–49	48 (39.4–57.5)	30.3 (23.5–38.4)	18.1 (14–23.3)	14.3 (10.8–18.7)	10.3 (7.7–13.7)	10.1 (7.4–13.8)		
Age 50–59	69.3 (58.5–79.5)	47.8 (37.8–58.9)	30.3 (23.4–38.5)	24.3 (18.4–31.6)	17.8 (13.2–23.7)	17.5 (12.7–23.9)		
Age 60+	92.4 (84.2–97.2)	75.8 (63.4–86.4)	54.4 (43.4–66.3)	45.4 (35.2–57.1)	34.7 (26–45.4)	34.3 (25.1–45.7)		

^{*}estimated from Weibull model

doi:10.1371/journal.pone.0160460.t005

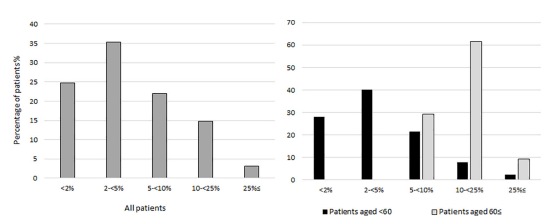


Fig 1. Frequency distribution of patients grouped by five-year mortality risk, overall and by age-group.

doi:10.1371/journal.pone.0160460.g001

Strengths and limitations

Strengths of this study include its large sample size, which meant we could investigate at risk subgroups such as the IDU risk transmission group and older patients, and the inclusion of patients from geographically diverse clinical cohorts in Europe and North America. As with any observational study our results may have been affected by unmeasured confounders. We could only include a limited number of prognostic factors in our study. The VACS Index which includes hemoglobin, composite markers of liver and renal injury [FIB-4 and estimated glomerular filtration rate (eGFR)] and hepatitis C status is a better predictor of mortality than age, CD4 count and viral load alone, but many of the cohorts in ART-CC did not collect these data [22]. Our results may have been affected by exclusion of patients without CD4 count and

Table 6. Adjusted hazard ratios (HR) for specific causes of death, adjusted for all variables in the table and stratified by cohort, deaths = 218.

		Adjusted HR (95% CI)						
Cause of death	N (%)	Age ≥60 vs. <60 years	Female vs. Male	IDU vs. non-IDU	CD4 <200 vs. ≥200 cells/μL	HIV-1 RNA >50 vs. ≤50 copies/mL	Prior AIDS diagnosis 10 years after ART start vs. no AIDS	
AIDS	35 (16%)	1.08 (0.33, 3.58)	0.38 (0.13, 1.09)	1.24 (0.52, 2.96)	5.32 (2.49, 11.4)	4.90 (2.30, 10.4)	1.75 (1.16, 2.65)	
Non-AIDS	145 (67%)	4.35 (2.97, 6.39)	0.70 (0.45, 1.06)	3.40 (2.26, 5.12)	2.42 (1.49, 3.94)	0.95 (0.62, 1.45)	1.58 (1.27, 1.95)	
Cardiovascular disease	22 (10%)	8.95 (3.63, 22.1)	0.54 (0.16, 1.84)	2.20 (0.59, 8.23)	2.70 (0.71, 10.3)	1.30 (0.44, 3.80)	1.49 (0.84, 2.63)	
Liver related	18 (8%)	1.65 (0.36, 7.50)	NA	3.84 (1.31, 11.2)	3.37 (1.11, 10.2)	2.37 (0.87, 6.40)	1.85 (1.02, 3.36)	
Non-AIDS, non-liver malignancies	45 (21%)	3.39 (1.71, 6.70)	0.81 (0.39, 1.68)	1.79 (0.77, 4.20)	1.62 (0.55, 4.80)	0.58 (0.24, 1.42)	0.88, 1.97)	
Suicide/accident	13 (6%)	2.55 (0.65, 10.0)	1.28 (0.39, 4.24)	2.45 (0.62, 9.69)	2.19 (0.42, 11.34)	0.65 (0.13, 3.17)	2.45 (1.25, 4.80)	
Other*	47 (22%)	5.71 (2.84, 11.5)	0.85 (0.42, 1.73)	6.21 (3.14, 12.3)	2.57 (1.11, 5.95)	0.93 (0.44, 1.95)	1.52 (1.04, 2.21)	
Unknown	38 (17%)	4.72 (2.28, 9.77)	0.50 (0.19, 1.27)	3.33 (1.48, 7.53)	3.38 (1.31, 8.74)	0.61 (0.24, 1.55)	1.15 (0.73, 1.83)	

IDU: injection drug use, CI: confidence interval

doi:10.1371/journal.pone.0160460.t006

^{*} Causes with fewer than 10 deaths: these were non-AIDS Infection (8); heart or vascular (6); respiratory disease (5); chronic obstructive pulmonary disease (4); renal failure (4); substance abuse (4); digestive system (4); central nervous system (3); lung embolus (1); other causes not included in CoDe classification system (15)



viral load measured during the 10th year of ART: such patients may attend care irregularly or may have been receiving care temporarily in a clinic not contributing data to ART-CC. Our results may only apply to patients who have survived 10 years on ART and continue to receive regular care. Outcomes are unknown in patients who were LTFU but CD4 counts and the proportion of patients virally suppressed were similar among those LTFU during the second decade of ART and those remaining alive and in care. Most cohorts link to death registries and so mortality rates among patients followed up in these cohorts should be reliably estimated [23], but deaths may have been under-ascertained in cohorts without links to death registries. Causes of death were classified according to a common protocol, but not all cohorts contributed data. However, the included deaths are likely representative of all deaths and we preferred not to increase the proportion of unknown or unclassifiable deaths by including cohorts with less complete data.

Context of research

Even in this cohort of patients who have survived at least ten years on ART, traditional markers of HIV progression such as CD4 count, viral load, and AIDS events remain important risk factors warranting clinical attention. CD4 measurements at start of ART were unimportant compared to those at ten years which is consistent with previous findings that recent or current values of biomarkers best predict survival [13, 24] and that the prognostic importance of values measured at start of ART diminish with time[15, 25]. Indeed, we found that for patients with the same CD4 count at ten years after ART start, subsequent mortality was lower in patients whose CD4 count had risen more over the first decade of ART, which may represent a selection effect ("survival of the fittest") as those who started ART with very low CD4 count were less likely to have survived 10 years and be included in this analysis. In this unselected treatment experienced population mortality rates remained substantially higher than in the general population. By contrast, standardised mortality rates in highly selected groups on long term ART who reached and maintained a CD4 count above 500 cells/ml have been found to approach that of the general population [26-28]. However, a recent study from Denmark found that even among well-treated HIV-infected individuals >50 years without comorbidity or AIDS-defining events the estimated median survival time remains lower than in the general population [29].

Older age was strongly associated with mortality, in particular that due to non-AIDS related morbidity. This suggests that provision of both preventive and therapeutic health care in older HIV-infected patients treated for many years will become increasingly important as the number of patients aged \geq 60 years old increases. The most common cause of death was non-AIDS cancer, implying a need for preventive and screening measures adapted for use with HIV-positive patients who have survived long-term treatment with ART to be incorporated into their routine health care[30].

Only a fifth of classifiable causes of death were due to AIDS, consistent with previous studies that found low proportions of AIDS-related deaths in those treated for many years [26, 31, 32]. Although a substantial proportion of deaths soon after start of ART are AIDS-related, [5, 6, 8, 18, 24, 33–35] this proportion decreases with duration of ART. Correspondingly, the proportion (though not the rate), of non-AIDS related deaths increases with time on ART [26, 31, 32]. Consistent with our study, the Data Collection on Adverse events of Anti-HIV Drugs (D:A:D) study found that most frequent causes of death in the 2009–11 follow-up period were due to non-AIDS cancers, AIDS, cardiovascular disease, and liver disease [8].

It is only possible to study the long-term effects of antiretroviral therapy in patients who started ART many years ago. Patients in our study will have been treated with the less potent and more toxic drugs (compared with those currently available) available during 1996–99.



Their management will have evolved as treatment guidelines changed and as new ART drugs and drug combinations were approved [2-4, 36]. Therefore long-term mortality of patients starting ART today can be expected to be lower than those found in our study. There is heterogeneity in mortality rates across cohorts [23] so that our estimates represent an average over included cohorts.

Implications for health care

Given that a quarter of patients had detectable viral load at ten years after starting ART, careful monitoring of patients treated for many years remains essential. The high burden of cancer found in this and other HIV-infected populations[37] indicates the need to optimise and improve adherence to screening guidelines[38, 39]. Higher mortality among patients diagnosed with AIDS before ART start or within the first decade of treatment may arise from chronic or late-onset sequelae of AIDS-defining illnesses, including relapse, secondary malignancy, heart disease after anti-cancer chemotherapy[40] or chronic lung injury following infection with *Mycobacterium tuberculosis*[41] or *Pneumocystis jirovecii*[42].

Patients reporting HIV transmission via IDU remained at considerably higher risk of mortality, particularly death from liver-disease, beyond ten years after starting ART. This higher mortality could arise from continued drug-use, co-infection with hepatitis C virus, higher rates of smoking and alcohol abuse, or poorer adherence to treatment [43, 44]. Several participating cohorts only assess IDU at enrolment, therefore we could not differentiate whether transmission risk of IDU remained a risk factor because of persisting drug use or because of direct sequelae or adverse socioeconomic factors associated with historic drug use. It is likely that the association of ongoing drug use with deaths from non-AIDS infection and substance use is much greater than that estimated for the transmission risk group designated IDU in this study. Current, but not former, drug use has been found to be associated with lack of adherence to ART which in turn correlates with virological outcomes [45]. It is therefore important to address continuing misuse of drugs and alcohol which may require opium substitution therapy or counselling. IDU should be screened for hepatitis C and offered treatment if warranted. Smokers should be offered nicotine substitution therapy and smoking cessation programs. Reducing mortality in IDU may also require interventions to address depression and social deprivation.

Conclusion

This study of patients who have survived ten years after starting ART found that CD4 count and viral load remain important for prognosis. However, deaths were mostly non-AIDS-related, and care for these patients should focus on improving management of non-HIV morbidity, in particular risk factors and screening for non-AIDS cancer. Whilst the majority of patients have low 5-year mortality risk, older patients have much higher risk. Patients with presumed transmission via IDU also have worse prognosis and likely require intensive management with a variety of interventions that aim to reduce their excess mortality. Our estimates of prognosis beyond ten years of ART could inform mathematical models [46] used to predict outcomes and allow comparisons to be made against the observed data to test the validity of such models. Our findings could also inform estimates of the future cost of treatment of people living with HIV.

Appendix 1

Cohorts included in this paper were the French Hospital Database on HIV (FHDH); the Italian Cohort of Antiretroviral-naïve patients (ICONA); the Swiss HIV Cohort Study (SHCS); the



AIDS Therapy Evaluation project, Netherlands (ATHENA); The Multicenter Study Group on EuroSIDA; the Aquitaine Cohort; the Royal Free Hospital Cohort, UK; the South Alberta Clinic Cohort; Cohorte de la Red de Investigación en Sida (CoRIS), Spain; The Danish HIV Cohort Study, Denmark; HAART Observational Medical Evaluation and Research (HOMER), Canada; HIV Atlanta Veterans Affairs Cohort Study (HAVACS), USA; Osterreichische HIV-Kohortenstudie (OEHIVKOS), Austria; Proyecto para la Informatizacion del Seguimiento Clinico-epidemiologico de la Infeccion por HIV y SIDA (PISCIS), Spain; VACH, Spain; Veterans Aging Cohort Study (VACS), USA; Vanderbilt, USA; University of Washington HIV Cohort, USA; and the Koln/Bonn Cohort.

Acknowledgments

We thank all patients, doctors, and study nurses associated with the participating cohort studies.

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Andrew Boulle (IeDEA Southern Africa), Christoph Stephan (Frankfurt), Jose M. Miro (PIS-CIS), Matthias Cavassini (SHCS), Geneviève Chêne (Aquitaine), Dominique Costagliola (FHDH), François Dabis (Aquitaine), Antonella D'Arminio Monforte (ICONA), Julia del Amo (CoRIS-MD), Ard Van Sighem (ATHENA), Jorg-Janne Vehreschild (Koln/Bonn), John Gill (South Alberta Clinic), Jodie Guest (HAVACS), David Hans-Ulrich Haerry (EATG), Robert Hogg (HOMER), Amy Justice (VACS), Leah Shepherd (EuroSIDA), Niels Obel (Denmark), Heidi M Crane (Washington), Colette Smith (Royal Free), Peter Reiss (ATHENA), Michael Saag (Alabama), Tim Sterling (Vanderbilt-Meherry), Ramon Teira (VACH), Matthew Williams (UK-CAB), Robert Zangerle (Austria)

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