

Unexpected increase of myocardial extracellular volume fraction in low cardiovascular risk HIV patients

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Research

Keywords: ECV, CV risk, CCT

Posted Date: July 16th, 2020

DOI: <https://doi.org/10.21203/rs.3.rs-41937/v1>

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Abstract

Background: People living with HIV (PLWH) are prone to develop sub-clinical Cardiovascular (CV) disease, despite the effectiveness of combined Antiretroviral Therapy (cART). Algorithms developed to predict CV risk in the general population could be inaccurate when applied to PLWH. Myocardial Extra-Cellular Matrix (ECM) expansion, measured by computed tomography, has been associated with an increased CV vulnerability in HIV-negative population. Measurement of Myocardial Extra-Cellular Volume (ECV) by computed tomography or magnetic resonance, is considered a useful surrogate for clinical evaluation of ECM expansion. In the present study, we aimed to determine the extent of cardiovascular involvement in asymptomatic HIV-infected patients with the use of a comprehensive cardiac computed tomography (CCT) approach.

Materials and methods: In the present study, ECV in low atherosclerotic CV risk PLWH was compared with ECV of age and gender matched HIV- individuals. 53 asymptomatic HIV+ individuals (45 males, age 48 (42.5-48) years) on effective cART (CD4+ cell count: 450 cells/ μ L (IQR: 328-750); plasma HIV RNA: <37 copies/ml in all subjects) and 18 age and gender matched controls (14 males, age 55 (44.5-56) years) were retrospectively enrolled. All participants underwent CCT protocol to obtain native and postcontrast Hounsfield unit values of blood and myocardium, ECM was calculated accordingly.

Results: The ECV was significantly higher in HIV+ patients than in the control group (ECV: 31% (IQR: 28%-31%) vs 27.4% (IQR: 25%-28%), $p < 0.001$). The duration of cART (standardized $\beta = 0.56$ (0.33-0.95), $p = 0.014$) and the years of exposure to HIV infection (standardized $\beta = 0.53$ (0.4-0.92), $p < 0.001$), were positively and strongly associated with ECV values. Differences in ECV ($p < 0.001$) were also observed regarding the duration of cART exposure (<5 years, 5-10 years and >10 years). Moreover, ECV was independently associated with age of participants (standardized $\beta = 0.42$ (0.33-0.89), $p = 0.084$).

Conclusions: HIV infection and exposure to antiretrovirals play a detrimental role on ECV expansion. An increase in ECV indicates ECM expansion, which has been associated to a higher CV risk in the general population. The non-invasive evaluation of ECM through ECV could represent an important tool to further understand the relationship between HIV infection, cardiac pathophysiology and the increased CV risk observed in PLWH.

Introduction

Myocardial Extra-Cellular Matrix (ECM) is a complex biological network consisting of a wide variety of proteins (principally glycoproteins, proteoglycans, and glycosaminoglycans) in which cardiac myocytes, fibroblasts, vascular cells, and leukocytes dwell. ECM proteins participate to myocardial strength and plasticity and play a fundamental role in cardiac development, homeostasis, and remodelling (1). ECM alterations may indicate a loss of cardiac morphological homeostasis and may represent an indicator of Cardiovascular Diseases (CVD). ECM expansion has been linked to electrical, mechanical and vasomotor dysfunctions, cardiac fibrosis and, eventually, increased mortality (2, 3). Contrast enhanced Cardiac

Computed Tomography (CCT) and Cardiovascular Magnetic Resonance (CMR) use extracellular extravascular contrast agents to non-invasively quantify the myocardial Extra-Cellular Volume (ECV) (3–6). In particular, CCT is increasingly used in experimental and clinical setting, due to its opportunity to assess lots of cardiac features other than ECV and fibrosis (such as coronary anatomy, calcification etc) in a very shorter time and with a higher spatial resolution than MRI (4, 5).

Although combined Antiretroviral Therapy (cART) has improved both the length and quality of life of people living with HIV (PLWH), cardiovascular diseases represent a challenge for clinicians and are still a leading cause of death in this population. Several studies have shown an increased frequency of coronary atherosclerotic disease, myocardial infarction, arrhythmias and sudden cardiac death (SCD) in PLWH (7). Interestingly, recent studies have shown that severe CV diseases are more frequent than expected in PLWH with a low Atherosclerotic Cardiovascular Disease (ASCVD) score, possibly as a result of premature aging due to inflammation, immune-activation, cART side effects and direct viral effect (8). Therefore, algorithms developed to predict CV risk in the general population could be inaccurate when applied to PLWH (9). In this scenario, the aim of our study was to assess ECV in PLWH with low-ASCVD score and to compare ECV of HIV + subjects to that of a control group.

Methods

Patient population

PLWH with low-ASCVD score and healthy HIV- controls were enrolled in this retrospective, monocentric study. 53 PLWH with low-ASCVD score and 20 healthy controls were consecutively and retrospectively enrolled. Clinical history, anthropometric data, as well as cardiovascular and metabolic data were collected.

All HIV+ participants underwent contrast enhanced CCT for clinical purposes. Criteria for patient's inclusion were: i) preserved glomerular filtration rate, ii) absence of contraindications to perform CCT and iii) completion of an iodine enhanced CCT scan after patient written consent acquisition. Exclusion criteria were: (i) known or suspected cardiac amyloidosis (causing interstitium expansion independently of myocardial fibrosis), (ii) hypertrophic cardiomyopathy, (iii) adult congenital heart disease.

Demographics and anamnestic data, including information on HIV status and major cardiovascular risk factors were collected and registered. Vital status was ascertained by Social Security Death Index queries and medical record review. A previous history of heart failure (HF), defined according to current guidelines, was considered in presence of physician documentation: (i) documented symptoms (e.g. shortness of breath, fatigue, orthopnoea) and physical signs (e.g. edema, rales) consistent with HF; (ii) supporting clinical findings (e.g. pulmonary edema on chest X-ray), or (iii) therapy for HF (including diuretics, digitalis, angiotensin-converting enzyme inhibitors, or beta blockers).

The 18 control subjects enrolled consisted of outpatients referred to CCT for nonspecific thoracic pain, with no history of previous cardiovascular diseases, non-smokers nor under vasoactive therapy and with a previous documented negative ECG. Selection on individuals in the control group was conducted with

age and gender matching criteria. All clinical investigations and laboratory exams were performed no more than 7 days before CCT. Review board and the local ethics committee provided approval for this investigation and informed consent was obtained from all patients.

CT image analysis

Pre- and post-contrast Hounsfield Unit (HU) were measured on CT images by the Picture Archiving and Communication System (PACS). The regions of interest (ROIs) were drawn firstly on the contrast image at the myocardial septum, in the equilibrium phase, and within the left ventricular chamber, the mean area was about 3 cm² (range: 1.5–5 cm²). ROIs were then copied to the pre-contrast image. Mean attenuation at the ROI was recorded in HU and Myocardial ECV fraction was calculated using the following equation (1, 2) (see Equations in the Supplementary Files)

where the contrast agent (CA) partition coefficient (I) represents the ratio of the change (DHU) between myocardium and blood, and HCT is the hematocrit level. DHU was determined as $DHU = HU_{\text{post}} - HU_{\text{pre}}$, where HU_{post} and HU_{pre} are attenuation after and before administration of iodinated contrast material, respectively. CT examinations were performed with coronary angiography protocol with a 64-detector Dual-source CT scanner (Somatom Definition Siemens Medical Solution, Forchheim, Germany). Cardiac scans (tube voltage, 120 kV; tube current time product, 190 mAs; section collimation, 64 detector rows, 1.2-mm section thickness; gantry rotation time, 330 ms) were acquired with prospective gating (65%-75% of R-R interval) and reconstructed into 3 mm-thick axial sections, using a B20f kernel. All patients underwent equilibrium CT, with a mean calculated effective dose of 2.18 ± 0.26 mSv and a mean iohexol volume of 150.4 ± 20.4 mL.

Virological analysis

HIV-1 RNA copy numbers were evaluated in plasma samples collected from whole blood obtained in EDTA-containing tubes and stored at -80°C. Levels of HIV-RNA were measured with Versant kPCR (Siemens Healthcare Diagnostic Inc., Tarrytown, NY) with a detection limit of 37 copies/mL.

Statistical analysis

Categorical variables were summarized as absolute frequency (and/or percentage), and continuous variables were summarized as mean and Standard Deviation (\pm SD) or median and inter-quartile range (IQR: 25th and 75th percentile). Chi-square test (with a cell count are greater than 5) or Fisher exact test (with a cell count are equal to or less than 5) compared categorical variables. Mann-Whitney test compared continuous variables, since some continuous variables exhibited skewed distributions on visual inspection. Instead, the independent Student t-test were used to compare normally distributed variables. Besides, linear regression models were used for univariate and multivariate analysis to evaluate the correlation between quantitative ECV and other study parameters. Lastly, One-way analysis of variance (ANOVA) was used for multiple-group comparisons according to four categories, followed by Bonferroni test. When interpreting the results, a p-value of less than 0.05 was considered indicative of a

significant difference. Statistical analyses were performed using statistical Program for the Social Sciences (SPSS, version 20) software package.

Results

General Characteristics

A total of 71 subjects were included in the present study (18 controls and 53 asymptomatic HIV-positive patients). The main characteristics of participants are summarised in Table 1 (anthropometrics, CV risk factors, metabolic status, immune-virologic status of HIV + subjects). No significant differences were found between the two groups in terms of gender distribution, age and other major risk factors ($p > 0.05$). Severity of HIV disease was categorized following the Center for Diseases Control and Prevention classification: 20 (38%) patients were in category A2, 10 (19%) patients in category A3, 2 (4%) patients in category B3, 3 (6%) patients in category C2 and 18 (34%) patients in category C3. Median CD4 + nadir before initiation of cART was 181 (IQR: 94–300) cells/ μ L. During their therapeutic history, 38 of the 53 HIV + participants received at least one protease inhibitor (PI) for a median of 3 (IQR: 0–14) years, while 23 HIV + received a nonnucleoside reverse transcriptase inhibitor (NNRTI) for a median of 1 (IQR: 0–9) years. With cART, plasma viremia was undetectable (lower limit of detection of 37 copies/mL) in all HIV + subjects. The hematocrit value was $35 \pm 4\%$ in HIV + participants and $41 \pm 3\%$ in controls ($p = 0.122$). No significant differences were observed regarding Heart Rate (HR) during CCT examination ($p = 0.211$).

Table 1

Baseline characteristic of the study cohort. Median and interquartile range (IQR: 25%-75%) or mean \pm SD are reported for continuous variables. Simple frequencies (n) and percentages are reported for categorical variables. Abbreviations: BMI^a, Body mass Index; HR^b, Heart Rate; HDL^c, High Density Lipoprotein; PLWH^d, People living with HIV; NA^e, not applicable.

Parameters	HIV-infected (n = 53)	Control group (n = 18)	p-value (*)
General characteristics			
Male, n (%)	45 (85)	14 (77)	0.129
Age, Years	48 (42.5–48)	55 (44.5–56)	0.122
BMI ^a , kg/m ²	22 (20.3–22)	27 (24–32)	0.116
HR ^b , bpm	67 (\pm 9.1)	65 (\pm 11.3)	0.211
Systolic blood pressure, mmHg	119 (\pm 10)	114 (\pm 11)	0.131
Diastolic blood pressure, mmHg	75 (\pm 9)	67 (\pm 6)	0.061
Laboratory results			
Total cholesterol, (mg/dl)	189 (177–189)	188 (154–201)	0.557
HDL ^c , (mg/dl)	39 (35–39)	54 (46–66)	0.046
Triglycerides, (mg/dl)	101 (78–170)	124 (80–177)	0.077
Other factors			
Diabetes, n (%)	0 (0)	0 (0)	1.000
Smoking, n (%)	9 (17)	4 (22)	0.686
Framingham Risk (%)	4 (1-6.5)	3 (1–5)	0.182
ASCVD 10 years, calculated risk	4.6 (2.7–4.6)	4.6 (1.7–3.5)	0.286
Immune-virologic status			
Years of therapy, Years	9 (6–9)	-	NA ^a
Years of exposure to HIV infection, Years	10 (1–23)	-	NA ^a
CD4 enrolment (cell/ μ l)	450 (328–750)	-	NA ^a
CD4 nadir (cell/ μ l)	181 (94–300)	-	NA ^a
% of PLWH ^e naïve for cART	12.7	-	NA ^a
HIV-RNA copies/ml in cART treated PLWH	< 37	-	NA ^a

Parameters	HIV-infected (n = 53)	Control group (n = 18)	p-value (*)
HIV-RNA copies/ml in cART naive PLWH	55.000	-	NA ^a

Measurements of Myocardial Extracellular volume fraction

The median ECV value was 27.4% (IQR: 25%-28%) in healthy controls and 31% (IQR: 28%-31%) in HIV + participants, respectively. In our study population, ECV differed significantly between the two groups ($p < 0.001$), thus indicating the presence of ECV expansion in the myocardium of HIV + subjects (Fig. 1a and b).

ECV values were not affected by patient's age neither in the HIV + group ($r = 0.12$, $p = 0.145$. Figure 2a) or in controls ($r = 0.11$, $p = 0.102$). On the contrary, ECV was found to significantly correlate with the number of years of exposure to HIV infection ($r = 0.54$, $p < 0.001$. Figure 2b), and the years on cART ($r = 0.62$, $p < 0.001$. Figure 2c), showing a trend toward ECV expansion over time in HIV infected patients.

Based on the years on cART and years of exposure to HIV, all HIV + patients were divided into three groups: (A) < 5 years, (B) 5 to 10 years and (C) > 10 years. (There was a significant difference in ECV (ANOVA, $p < 0.001$) among controls and groups (see Table 2).

The multivariable stepwise linear regression showed that the duration of therapy (β was 0.56 (0.33–0.95), $p = 0.014$) and years of HIV infection (β was 0.53(0.4–0.92), $p < 0.001$) were independent predictors of ECV expansion (Table 3).

Table 2

Myocardial ECV comparisons between control group and HIV-positive patients selected by (A) less than 5 years, (B) from 5 to 10 years and (C) greater than 10 years. Data are shown as Median and interquartile range (IQR: 25%-75%).

Parameters	Controls (n = 18)	<5 y (A)	5–10 y (B)	>10 y (C)	p-value (*)
Myocardial ECV (%)	27.4 (25–28)	-	-	-	-
Years of therapy	-	29 (26–30)	30 (27–32)	37 (33–39)	< 0.001
Years of exposure to HIV	-	29 (26–30)	31 (28–34)	35 (31–39)	< 0.001

Table 3

(A) Univariate correlation analyses between ECV and characteristics of the HIV + population. (B) Multivariate regression models for predictors of ECV expansion. Age, Heart Rate (HR), systolic and diastolic blood pressure, years of therapy and years of exposure to HIV infection were entered in this model. P-values as assessed by Pearson's correlation for normal distributions, Spearman's correlation for non-normal distributions as well as β estimates and p-values from multivariate regression models are reported.

Variable	Univariate		Multivariate	
	R (95% CI)	p-value	β (95% CI)	p-value
Intercept	25.5 (21-29.5)	0.210	25.7 (20.2–30)	0.062
Age	0.12 (0.02–0.35)	0.805	0.42 (0.33–0.89)	0.084
Gender	0.29 (0.13–0.42)	0.149	0.19 (0.01–0.34)	0.122
Heart Rate (HR)	0.15 (0.01–0.33)	0.301	1.29 (1.01–1.5)	0.062
BMI (kg/m ²)	-0.26 (-0.01-0.2)	0.068	-	-
Systolic blood pressure	-0.02 (-0.01-0.2)	0.278	-	-
Diastolic blood pressure	-0.05 (-0.02-0.1)	0.191	-	-
Years of Therapy	0.62 (0.44–0.96)	< 0.001	0.56 (0.33–0.95)	0.014
Years of exposure to HIV infection	0.54 (0.36–0.90)	< 0.001	0.53 (0.4–0.92)	< 0.001

Discussion

Given the efficacy of newer, effective antiretroviral therapies, life expectancy of PLWH has shown a widespread improvement; on the other hand, following the introduction of cART, CVD and cardiovascular deaths have increased dramatically among this population (7). Several studies conducted in PLWH with CCT angiography, showed an increased presence and extent of coronary artery disease, including non-calcified and high-risk coronary plaques, that were associated with an increase in indicators of systemic immune activation (10, 11).

In a previous work, increased myocardial inflammation has been described in HIV + patients (15). Specifically in an interesting prospective study, Luetkens et al that compared the results of comprehensive CMR studies in HIV + asymptomatic patients and controls and observed an increase in CMR markers of myocardial inflammation in the HIV + group, thus suggesting the presence of subclinical myocardial inflammation in this population, which may in the end lead to the impairment of left ventricular myocardial function. The major limitation of the cited study, as well as of our present study, is that no histological assessment has been carried out. Luetkens et al. suggested that, although direct HIV infection of myocytes is rare, it is possible that HIV itself together with other cardiotropic viral infections may lead to myocarditis and to a subsequent left ventricular dysfunction; this may also explain the high rate, observed among this population, of resting ECG abnormalities that are consistent with myocardial

damage. Authors suggested that inflammatory condition and subsequent fibrosis may be responsible of the observed CMR findings. On the other hand, our results show relevant differences in respect to those of Luetkens et al. Firstly, the statistical evaluation carried out by our colleagues is mainly descriptive and no regression analysis was performed due of the limited sample size. Moreover, since the study was conceived for preliminary investigation purposes, no correction analysis was performed, possibly leading to inflated type I errors. This assumption in our work is supported by a significant difference in ECV values between the different patient groups, which are known to represent an indirect measure of diffuse interstitial myocardial fibrosis or most serious heart diseases (17). In addition, the ECV values observed in our study were consistent with previous studies conducted in the HIV- population (18). Lastly, in our study, due to a larger sample size, we were able to perform a multivariate regression analysis.

It has been described previously that amyloid precursor protein (APP) may be an innate antiviral defence factor in macrophages and microglia that restricts HIV-1 release in the neurological environment. Highly expressed in macrophages and microglia, the membrane-associated APP binds the HIV-1 Gag polyprotein and blocks HIV-1 production and spread. To escape this restriction, Gag promotes secretase-dependent cleavage of APP, resulting in the overproduction of toxic A β isoforms. From a neuro-physiopathological point of view, this Gag-mediated A β production results in increased degeneration of primary cortical neurons. One interesting question is: could this mechanism be involved in the physiopathology of our findings of an increase in ECV? Could amyloid formation as a result of immunological response to virus in cardiovascular tissue explain cardiomyopathy and peripheral obstructive vascular lesions observed in HIV + patients? Urgent studies are clearly needed to better understand the complex dynamic that underlies the multiple peculiar features which are increasingly being observed in people chronically living with HIV, in order to design effective prevention strategies to counteract the increasing burden of cardiovascular events in our patients.

Overall, these data suggest that an increase in systemic inflammation and immune activation may manifest as a premature CVD phenotype, including disease in large and small vascular beds in patients with HIV. Among PLWH, a relevant concern is whether scores that rely on traditional CV risk factors can accurately identify those individuals with a higher probability to develop CVD (9). In this study, we highlight that PLWH with low-ASCVD risk score are likely to suffer from a silent cardiac impairment. In fact, we found a significant difference in ECV levels between PLWH and healthy subjects ($p < 0.001$), indicating the presence of ECM expansion in patients with HIV infection. More importantly, years of exposure to HIV, and years of antiretroviral therapy significantly correlated with ECV expansion.

Quantification of ECM is an indirect measure of the expansion of extracellular matrix since, in absence of confounding conditions (e.g. oedema or amyloid), it is related to collagen excess in the interstitium. ECM expansion may therefore indicate vulnerable myocardium in PLWH and represent a potential therapeutic target (7). Indeed, abnormal fibrosis may be the underlying pathological finding in many cardiovascular disorders, particularly in sudden cardiac deaths (SCD), an emerging issue in PLWH. Frieberg et al. report that PLWH had a 14% higher risk of SCD, with uncontrolled HIV infection representing a risk factor (13). Recently, a review of the health records from San Francisco between 2011 and 2016 outlined that the

cause of death for autopsy-defined sudden arrhythmic death syndrome (SADS) and SCD were 83% and 82% higher respectively in the PLWH than in general population. Interestingly, the histologic exam identified higher rates of cardiac fibrosis in PLWH underling the excess risk of SADS (14). Indeed, specific tools are sought to measure in vivo the CV risk excess of PLWH which cannot be determined by common predictors of CV scores. Quantifying ECM expansion through ECV assessment may ultimately provide a basis to improve care in HIV individuals through a better risk stratification and the possibility of targeted treatment.

Conclusion

This study investigated the role of ECV measurements in the phenotypic characterization of CV risk, demonstrating that ECV can be easily and non-invasively assessed by CT-scan and used to characterize myocardial stiffness, a clear advantage with respect to alternative methodologies (19). ECV could provide an important imaging tool to further understand the relationship between cardiac anatomy and heart function.

HIV-positive subjects present ECV expansion in the myocardium. The duration of therapy and years of HIV infection are predictors of myocardial ECV expansion. Imaging technique in PLWH may be critical to identify patients with subclinical CVD and at risk for increased SCD and SAD events, where traditional risk scores may be inaccurate. Our data show that the measurement of ECV is a promising approach to identify asymptomatic patients with high risk of CV diseases despite a low-ASCVD traditional score risk prediction.

Our findings, therefore, may help to explain the increased cardiac morbidity and mortality observed in patients with chronic HIV infection.

List Of Abbreviations

ASCVD: Atherosclerotic Cardiovascular Disease

BMI: Body mass Index

CCT: Cardiac computed tomography

CV: Cardiovascular

cART: combined Antiretroviral Therapy

CA: contrast agent

ECM: Extra-Cellular Matrix

ECV: Extra-Cellular Volume

HDL: High Density Lipoprotein

HR: Heart Rate

HU: Hounsfield Unit

NNRTI: nonnucleoside reverse transcriptase inhibitor

PLWH: People living with HIV

PACS: Picture Archiving and Communication System

PI: protease inhibitor

ROIs: Regions of interest

SD: Standard Deviation

SADS: sudden arrhythmic death syndrome

SCD: sudden cardiac death

Declarations

Ethics approval and consent to participate

The study was approved by the institutional review board (Ethics Committee of Umberto I General Hospital, Rome), and all participants signed written informed consent.

Consent for publication

Not applicable

Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Competing interests

All the authors declare that they have no competing interests.

Funding

Not applicable

Authors' contributions

C. Borrazzo participated in writing the paper, analyzed the data and performed statistical analysis; G. d'Ettoire and G. Ceccarelli participated in writing the paper, were responsible for study concept and design, and for the supervision and critical revision of manuscript for intellectual content; M. Pacilio and L. Santinelli analyzed the data; E. N. Cavallari, O. Spagnolello, V. Silvestri, and P. Vassalini were responsible for clinical evaluation of the HIV-1 positive patients and performed critical revision of manuscript for intellectual content; C. Scagnolari performed critical revision of manuscript for intellectual content; M. Francone was responsible for the clinical evaluation of the HIV-1 positive patients and performed critical revision of manuscript for intellectual content; C.M Mastroianni and I. Carbone were responsible for study concept and design, and for the supervision and critical revision of manuscript for intellectual content.

Acknowledgements

Not applicable

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Figures

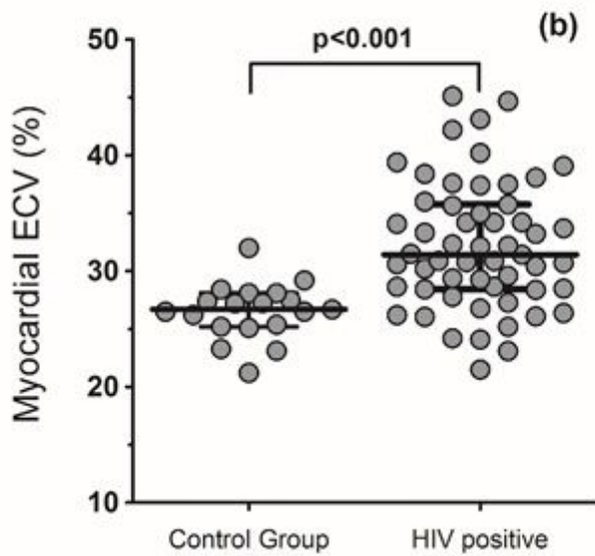
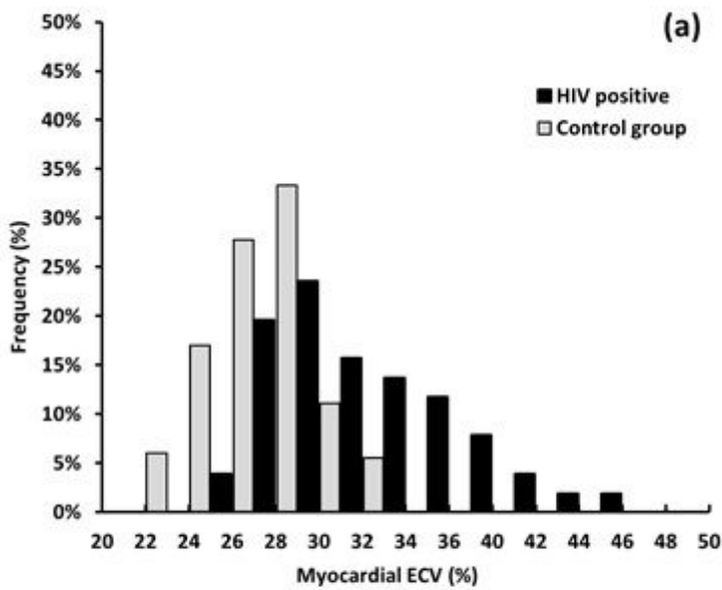


Figure 1

(a) Frequency histograms of ECV fraction. (b) Boxplot comparing ECV in controls and HIV+ patients: median, interquartile range (25%-75%) and p-value of the Mann-Whitney test for significance are also reported.

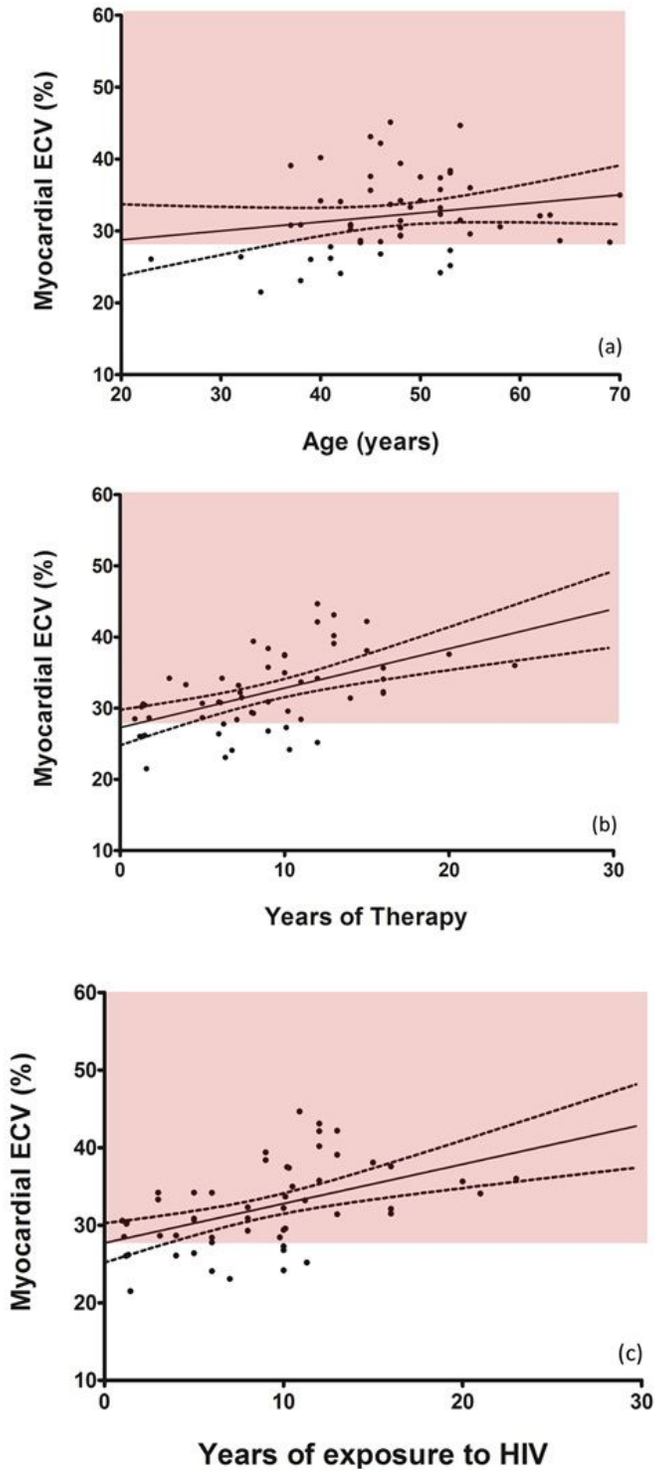


Figure 2

Scatter plots and best-fit regression curves (Solid line) with 95% CI bands (shown as dashed lines) of ECV versus age (a), years of therapy (b), and years of exposure to HIV (c) are shown for HIV+ patients. r and p -value are also reported. Red shading denotes the ECV range of the disease zone. Reference intervals are defined from the work of Scully et al. (3).

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- [Equations.pdf](#)