JBMR MS# M1611079 revised. Hamill et al.

Associate Editor:

In its present form the manuscript is too long - as noted by Reviewer 1, the Methods and Results in particular could be shortened.

We have responded to all the reviewers’ comments, shortened the paper and reduced the Methods and Results by 2 typed pages. We have highlighted the changes and used track-changes to indicate inserts and significant deletions.

Reviewer: 1

Comments to the Author

The authors present a longitudinal study over 12 months of follow-up amongst pre-menopausal women in South Africa comparing women with/without HIV infection and on/off ART. The authors are to be congratulated for their thorough investigation among young women in South Africa - these are indeed rare data.

We thank the reviewer for these comments

A few thoughts:

1. As currently presented, this manuscript is rather lengthy and contains a great deal of detail in Methods/results. There seems to be an imbalance between Methods /Results and Discussion with too much of the first two and too little Discussion. I wonder if Methods /Results could be simplified perhaps with some additional material on line?

We have reduced the Methods by deleting details available in published papers and moving information on lab procedures to supplementary material. Likewise we have reduced the Results section (see next point).

Figure 2 is much easier to take in than the rather turgid text. I would particularly suggest attention is paid to 'Results' which are rather confusing and dense to read. Changes in anthropometry, age variance, CD4 count variance and ART duration are listed in a way that is difficult to make sense of.

To simplify the Results section, we have removed the data on anthropometry, age, CD4 count and ART exposure from the text and tabulated them in new Table 1; we have shortened the text and divided it into sub-sections, and illustrated the results for all women as well as those in the ART restricted dataset in a revised Figure 2.

2. Discussion smooths over a number of weaknesses, including (a) any mention of reliability of the serum assays used

We have added a comment in the Discussion on the limitations of TmP/GFR as a marker of proximal tubule damage. We have also included more detail on the reproducibility of the serum assays used in the supplemental material.

(b) that some DEXAs were performed by a different observer(s)

Our original comment overstated the point because, in fact, >90% of scans were performed by only one operator and only one individual scrutinised, graded and analysed the scans for definition of region of interest. We have deleted the comment.

(c) that women crossed over during the study due to change in HIV status, pregnancy and lactation - could there be any systematic differences causing bias?

There are two different sets of women who ‘changed-groups’ or were pregnant/lactating during the study. Firstly, as depicted in Figure 1, there are those women measured at baseline who had seroconverted or been pregnant/lactating. The first paragraph of the Results details the general similarity of the CD4 count, anthropometry, bone and biochemical measures at baseline in women not included at 12 months for these and other reasons versus those included, giving reassurance of no obvious systematic bias in the follow-up at 12 months.

Secondly, there are those HIV-positive women that ‘changed group’ because of initiation or non-initiation of ART counter to original expectations. Relatively few Ppres women initiated ART (n=11) and those that did were exposed for less time than for ART-exposed Plow women, whereas there were only few Plow women who did not start ART (n=9). We therefore considered there were insufficient numbers to re-classify these women and conduct an analysis comparing 4 groups of HIV-positive women (Ppres±ART, Plow±ART). Instead we investigated the possibility of systematic bias and potential dilution of an effect within the full dataset, by comparing the results obtained with those after restricting the data to women whose HIV/ART status had not changed by 12 months (i.e. PresN and PlowY). We have sought to clarify the approach taken by expanding the explanation in the statistical methods.

Why does total body BMD not change in the ART group?

Figure 2 presents the change in total body BMD for ART-exposed group (ie PlowY) and new Table 4 gives the differences in change over time relative to the other two groups. Both demonstrate that there was a significant decrease in total body BMD in the ART-exposed group, with and without adjustment for changes in bone and body size. In Tables 2 and 3 that present the changes in Plow by original designation (ie includes the 9 women who did not start ART) there is an indication of significant total body bone loss (with a lower effect size than for just the ART women in Fig 2) that is attenuated slightly and becomes not significant after adjustment. Taken together these results imply that the change in total body BMD in the 9 unexposed women was less or in the opposite direction to those exposed to ART, in line with the lack of bone loss in the unexposed Ppres group, thus diluting the observed effect and significance when tested in the full dataset. However with only n=9 in the non-exposed Plow group, this cannot be formally tested, and we have not drawn out this point in the paper. It is of note that the decreases in total body BMD in the ART-exposed group is numerically less than those seen at the spine and femoral neck, likely indicating greater loss over 12 months at sites richer in trabecular bone than in the whole body, as would be anticipated.

3. The Discussion could merit clarifying the probable importance of TDF in the ART group with mention of other ART agents that might be of lesser impact on bone health and perhaps that the new pro-drug TAF may have differential effects?

Thank you, we have added a comment to this effect to the last paragraph of the Discussion.

4. How reliable are BALP and TALP and how good are the tubular function markers employed in this study?

(a) We are unsure whether the reviewer is referring to reproducibility for BALP/TALP assays or the reliability of these indices as markers of effects on bone. To address the first possibility, we have retained information about assay reproducibility in the section now moved to the supplemental material and indicated the low cross-reactivity in the BALP assay for the liver enzyme. As to the second, while recognising that BALP/TALP alone has limitations as a marker of effects on bone, taken together with the results for the bone turnover markers P1NP and CTX, the results add to the overall impression of increased bone turnover in the ART-exposed women in association with the bone loss. We have sought not to over-emphasise the bone marker results in the Discussion beyond support of the DXA findings (paragraph 2) and, to avoid doing so, we have not expanded the text to discuss the reliability of BALP/TALP as an index of effects on bone.

(b) As alluded to by the reviewer, serum phosphate, TmP/GFR and eGFR are indicators of renal function and phosphate handling, but are not specific markers of proximal tubular function such as retinol-binding protein of beta2 microglobulin. We have rephrased paragraph 2 in the Discussion and added this as a limitation in paragraph 4. We have also taken the opportunity to recalculate the eGFR using the CKD-EPI formula rather than MDRD because the majority had eGFR in the normal range.

**Reviewer: 2**

Comments to the Author

In this manuscript the authors reported the changes in BMD, vit D and body composition in premenopausal HIV south African women. The topic is interesting, and relevant. However the description of the methodology and statistical analysis are difficult to understand.

Main comments

There are some problems with the methodology and statistical analysis. I don’t understand how the BMD is analyzed : use of T score, definition of BMD loss and how many patients had a significant bone loss (> 0,03 g/cm²) and the statistical models description in not understandable  …

We have responded to all these points below.

Introduction

P 3 l 39-54 Can the authors add references for the association between poor vitamin D status and use of NNRTI ?

In addressing the need to shorten the text, we have deleted lines P3, 39-54, because of limited relevance to the paper. However, for the reviewer’s interest, references supporting this comment include Boura et al. J Int AIDS Soc 2014; 17: 19826 doi: 10.7448/IAS.17.4.19826ecollection2014.

P4 can the authors give their hypothesis before the aims

We have added an overarching hypothesis to the last paragraph of the introduction before stating the aims.

P4 what is mineral metabolism?

We have clarified in the text that this refers to calcium and phosphate metabolism.

Methods

P7 Body composition parameters lines 39-40: what are the measured parameters?

The body composition measures were lean mass (g) and fat mass (g). We have added extra clarification in this section.

Do the authors have some data on risk factors for osteoporosis , or bone loss? Smoking, alcohol, amenorrhea, other comorbidies (coinfection hepatitis B, or C).

Is there any data on fractures?

We have included information on lifetime fracture and pertinent lifestyle characteristics in new Table 1. Diagnosed hepatitis B and C was an exclusion criterion for the baseline study. We have not described this or other design aspects, because of the requirement to shorten the Methods, but have referred the reader to published papers where these details are given.

Statistical methods

P10: statistical methods : how is analysed BMD? What is SD-scores: T or Z scores? How is defined BMD loss? how many patients had a significant bone loss (> 0,03 g/cm²) …

1. Change over time in BMD was analysed on a group basis by using repeat measures ANOVA and ANCOVA utilising the Linear Model Software in DataDesk and by regressing final BMD as dependent variable against baseline value. We have shortened and refashioned the Statistics section in an endeavour to simplify the explanation. Full details of the models are also given in the footnotes to the tables.
2. The SD-scores quoted in the original manuscript were the equivalent of Z-scores, calculated with mean and SD data from the local reference group, as described in the statistics section of the original version. However, as these added little to the paper and were a cause of confusion, all mention of SD-scores has been deleted from the methods and results and a brief referenced comment about the z-score of these young adult women added to the discussion. We have commented further on this in response to a comment about the results section below.
3. We defined BMD loss over time on a group basis as per the original design of the study, using conventional statistical tests (pairwise Scheffe post-hoc tests, by group, testing the significance of mean within-individual differences between 12 month and baseline BMD values using Linear Models). By using a contemporaneous local reference group measured on the same instrument we minimised the possibility of misinterpreting BMD changes over time in HIV-positive women due to instrument imprecision and drift. Nevertheless, to address the reviewer’s question about how many individuals had bone loss that exceeded the least significant change of >0.03 g/cm2 (a conventional figure based on a notional instrument precision of 1%, as per new reference 20), we have added these data for the lumbar spine in the Results section. The proportion of individuals exceeding this threshold in our dataset is a conservative estimate of measurable bone loss because the precision of our instrument at the lumbar spine was 0.7%. Nevertheless, these data support the conclusion of bone loss in the Plow group, especially among women exposed to ART, compared to little or no change in the other two groups, and serve as an illustration.

What is the main criteria; what are the outcomes and the covariates used in the different models. It is not really clear. Authors should give their list of outcomes and covariates

We had detailed the covariates used in the models in both in the statistical section of the methods and the footnotes to the table (the footnotes were complied at the back of the original script but have been moved below each table in the revised version to avoid further confusion). The outcomes are also described in the last paragraph of the introduction and itemised in the Methods and Tables. We did not adjust for any other factor, as this was not necessary to determine whether there was evidence of differences between the groups of women in changes in our variables of interest over time. We have added this clarification to the statistical section.

I don’t understand the modalities of adjustment

We have expanded and simplified the description of our statistical approaches and adjustments in the revised Methods section (see the two responses above)

What is the number of missing data, how the authors manage this problem?

The design of the follow-up, as described in the Method section, was such that there were no missing data for the DXA and anthropometric measurements, as only the baseline data for women measured at 12 months were included. The number of missing biochemical data points is provided per analyte in the footnote to Table 4 (now new Table 5). The nature of the models used in this study is such that group comparisons at each time-point are similar whether individuals with a missing value are excluded or not, and comparisons of change over time are identical. For the sake of brevity, we have not added a comment on this point.

Finally I’m not sure that so many analyses should be performed with small sample size

We recognise that a large number of tests have been presented, but the data and p-values have been given in each case to allow the reader to form a judgement on the statistical probability of a biological effect.In addition, we have used Scheffe post hoc tests to minimise the problem of multiple-testing within each model. For the separate models examining different outcome variables, the significance level is less than 1:100 and 1:1000 in many instances, indicative of a low probability of a chance result. Supportive evidence for the changes over time in bone mineral density and anthropometry being genuine and not by chance come from both the biochemistry and the similar patterns of change in the different, independently measured, variables.

Results

This paragraph is difficult to follow and very long: authors may present clearly their data on body weight, BMD loss, body composition vit D and bone turnover markers changes.

We have shortened and simplified the results, as we have detailed in our response to reviewer 1 above.

Additive results should be added : Prevalence of patients with low bone mass (Z or T < -2) , % of patients with significant bone loss (> 0.03 g/cm²).

We have responded to this point above and revised the manuscript by adding a comment about individual bone loss. We wish to remind the reviewer that the study did not involve osteoporosis patients but young adult women from the community, for whom T-scores are not appropriate, and that both groups of HIV-positive women at baseline had mean±SD BMDs very similar to those of the local reference group, i.e. SD-scores (locally derived Z-scores) close to zero. This was described in Hamill et al 2013 and we have now reiterated this in the discussion.

Interest of the paragraph about albumin?

Albumin data provide valuable information in three ways (1) as an indicator of frailty, including advanced disease in HIV-positive individuals and protein malnutrition (2) as an acute phase marker that is reduced during infection and (3) as the main carrier protein for serum calcium such that albumin adjustment is necessary for interpreting calcium status. We consider it is important to retain the albumin data for these reasons and also because we have shown that the HIV-positive women with low CD4 counts had low albumin which was improved with ART.

Discussion

The authors should discuss more about their objectives: results of vit D, body composition?

The discussion has been expanded to include brief mention of possible mechanisms for the rapid fat accumulation in the ART exposed women. We have not expanded the discussion of the vitamin D result because in our study neither HIV infection nor ART exposure affected vitamin D status. Given the need to shorten the text, we consider that giving a critique of the design of other studies that have suggested that vitamin D status is compromised (which we mention briefly in the introduction), is beyond the scope of this paper.

What are the explanations and hypothesis for the authors for the body composition results?

We have expanded the paragraph in the discussion on body composition to cover this point (as discussed above)

Authors should add in the limitations, the small sample size of each group, the absence of extrapolation to other population

We have added these points to the paragraph on limitations and strengths in the Discussion

Table 3 BMD results are adjusted on size ;any other adjustments?

The models used in all the tables are included in their footnotes. As we have responded above, we made no adjustments other than size and have added this comment to the statistical section

Table 4 : The content of the table is not clear ; authors should clarify the bone remodeling markers results

A description of the table contents, results and abbreviations (now Table 5) is in the footnote which is now located beneath the table rather than at the end of the script.

**JBMR MS# M16110779 revised**

**Changes in bone mineral density, body composition, vitamin D status and mineral metabolism in urban HIV-positive South African women over 12 months**

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**ABSTRACT**

HIV infection and antiretroviral therapy (ART) are associated with bone loss and poor vitamin D status in Caucasian populations, though their relative roles are not known. No previous studies have examined longitudinal changes in areal bone mineral density (aBMD), measured by DXA, or in vitamin D status in HIV-positive African women. Of 247 premenopausal, urban, black African women from Soweto, South Africa, initially recruited, 187 underwent anthropometry, DXA scanning and blood and urine collections at both baseline and 12 months. Of these, 67 were HIV-negative throughout (Nref), 60 were HIV-positive with preserved CD4 counts at baseline (Ppres) and 60 were HIV-positive with low CD4 counts at baseline, eligible for ART by South African standards of care at the time (Plow). No participant had been exposed to ART at baseline. By 12 months, 51 Plow women had initiated ART, >85% of whom took combined tenofovir disoproxil fumarate (TDF), lamivudine and efavirenz. By 12 months, Plow and Nref, but not Ppres, increased in body weight and fat mass (group-by-timepoint *p* ≤0.001, *p*=0.002 respectively). Plow had significant decreases in aBMD of 2-3%, before and after size adjustment, at the femoral neck (*p* ≤0.002) and lumbar spine (*p* ≤0.001), despite significant weight gain. These decreases were associated with increased bone turnover but there were no significant differences or changes over time in vitamin D status, serum phosphate concentrations or renal phosphate handling. Excluding data from 9 Plow women unexposed to ART and 11 Ppres women who had initiated ART accentuated these findings, suggesting the bone loss in Plow was related to ART exposure. This is the first study describing DXA-defined bone loss in HIV-positive Sub-Saharan African women in association with ART. Further work is required to establish if bone loss continues with on-going ART and, if so, whether this results in increased fracture rates.

Key words: AFRICA; ANTIRETROVIRAL THERAPY; BONE HEALTH; HIV; PREMENOPAUSAL WOMEN; VITAMIN D.

**Introduction**

HIV-infection is associated with weight loss and with increased morbidity and mortality related to infectious complications of immune suppression. Since the advent of effective combination antiretroviral therapy (ART) survival has increased dramatically,(1) and in more recent years there has been a shift in focus from HIV-associated communicable diseases to non-communicable diseases including osteoporosis and associated fragility fractures.(2)

HIV and ART have both been associated with bone loss and increased risk of fracture.(3-6) Evidence is mixed but the general consensus is that ART-exposure, particularly to tenofovir disoproxil fumarate (TDF), is associated with bone loss.(7,8) Even though the mechanism(s) is not entirely understood, it is hypothesised that the bone loss is related to TDF-induced renal phosphate wasting with subsequent skeletal demineralisation.(9) In a meta-analysis(3) HIV-positive individuals were three times more likely to have lower areal bone mineral density (aBMD) than controls and the prevalence of dual-energy X-ray absorptiometry (DXA) defined osteoporosis was as high as 15% .

A potential contributor to bone loss in HIV-infected individuals is poor vitamin D status, which has been described in several HIV-positive populations, both in ART-naïve patients(10) and in association with ART-exposure.(11,12) The precise mechanism underlying these observations is unknown but it has been suggested that it may be a consequence of decreased dermal synthesis and/or as a result of ART-induced up-regulation of the metabolism of 25-hydroxyvitamin D (25(OH)D). Adequate vitamin D status, as measured by serum 25(OH)D, has a well-established role in skeletal health.

To date, available data about the possible effects of HIV-infection on bone health and vitamin D status are heavily biased towards Caucasian males in resource-rich societies who have a much lower lifetime risk of osteoporotic fracture than women. Some of these studies are also limited by retrospective design and lack of an HIV-negative control group. The effects of HIV and ART on bone health and vitamin D status in Sub-Saharan Africa, where the burden of HIV-infection lies, and where osteoporotic fracture rates are predicted to rise,(13) are largely unstudied(14) and, although data are emerging,(15-17) there are important gaps and an urgent need for targeted research.(14)

We have shown, in a cross-sectional study of urban South African premenopausal women which compared those with HIV-infection but ART-naïve with those who were HIV-negative, that there were no significant differences in aBMD or vitamin D status related to HIV-status, despite HIV-positive women with low CD4 counts having less body fat than both HIV-positive women with preserved CD4 counts and HIV-negative women.(18) HIV-positive women in the low CD4 group were eligible to start ART under the South African medical guidelines current at the time of the study (2010). We hypothesised, based on studies from Western countries, that HIV-positive African women would lose bone mineral over time, accompanied by changes in weight and body composition, in vitamin D status and in calcium and phosphate metabolism, and that the bone loss would be accentuated in those exposed to ART. The aim of the study presented in this paper was to follow these three groups of women for 12 months, in order to investigate the effects of HIV-infection and ART over time on bone mass, body composition, vitamin D status and markers of calcium and phosphorus metabolism.

**Participants and Methods**

Study design

The study was designed as a 12-month longitudinal investigation of urban South African premenopausal women with and without HIV-infection. Participants attended for study visits at the SAMRC/Wits Developmental Pathways for Health Research Unit (DPHRU) in Soweto, Johannesburg, SA, at baseline and 12 months. At each visit anthropometry and DXA scans were performed, and blood and urine samples were collected for laboratory analysis, as described below. The participants with HIV-infection continued to attend their usual primary health care facilities for monitoring and management of their disease.

Study details, inclusion and exclusion criteria at enrollment, baseline characteristics and dietary intakes have been described in full elsewhere.(18,19) In brief, 247 urban, black South African women were recruited into three groups of approximately equal size from clinics in Soweto, Johannesburg, between February and July 2010. All women were premenopausal and not pregnant or lactating at the time of enrollment. The three groups were: HIV-negative women to act as the reference group (Negative-reference: Nref, *n* = 98); HIV-positive women with preserved CD4 counts (≥ 350 x 106 cells/l) anticipated not to require ART-initiation for at least 12 months (Positive-preserved: Ppres, *n* = 74) and HIV-positive women with low CD4 counts (≤200 x 106 cells/l) who were eligible to commence ART soon after the baseline visit (Positive-low: Plow *n* = 75). At the 12-month visit, Nref participants were offered repeat HIV-antibody testing using the Alere Determine™ rapid HIV-antibody test (Alere San Diego, Inc. San Diego, CA, USA). Those who had a reactive HIV test were referred to a local clinic for confirmatory testing and CD4 count.

The University of the Witwatersrand Human Research Ethics Committee (HREC number: M101525) and the Gauteng Department of Health, South Africa, approved the study. All participants provided informed written consent prior to enrollment.

Anthropometry

Height was measured to the nearest 0.1 cm using a permanent wall-mounted stadiometer (Holtain, Crosswell, UK). Weight was measured to the nearest 0.1kg using an electronic digital scale (Tanita, TBF-410 MA Body Composition Analyzer, Tanita Corporation of America, Inc., Illinois, USA) with participants wearing light clothing. Body Mass Index (BMI) was calculated as weight in kilograms divided by the square of height in metres (kg/m2). Waist and hip circumferences were measured to the nearest 0.1cm using a non-stretchable plasticised tape measure.

Bone mineral density and body composition by DXA

DXA was performed using an Hologic QDR 4500A DXA (Model: Discovery W (S/N 71201) software version 12.5:7 Hologic, Inc., Waltham, MA, USA). Scans of the whole body, lumbar spine L1-L4, total hip and femoral neck were conducted with participants wearing light clothing, and performed using the automatic scan mode. Whole body was analysed as ‘whole body less head’ (WBLH). For each individual, their follow-up scans were compared to baseline to ensure consistent placement of regions of interest. DXA bone measures were bone mineral content (BMC, g), bone area (BA, cm2) and aBMD (g/cm2) at each site and body composition measures were whole body lean mass (g) and fat mass (g). Daily calibration and long-term DXA scanner stability monitoring were conducted using manufacturer phantoms. Short-term repeat scan precision on 30 participants with repositioning was 0.65% for lumbar spine and 0.97% for proximal femur; the coefficient of variation on daily quality control scans during the period was <0.5%. The extent to which individuals experienced aBMD loss in excess of the least significant change was determined using the conventional DXA 0.03g/cm2 threshold, which is based on a notional instrument precision of 1%.(20)

Laboratory measures

Full details of the blood and urine collection, processing and analytical procedures are given in the supplemental information. In brief, blood was collected in the morning after an overnight fast by venepuncture and processed as EDTA plasma for parathyroid hormone (PTH) analysis and as serum for other analytes relating to calcium, phosphorus and vitamin D metabolism (calcium, phosphate, magnesium, albumin, 25(OH)D) and bone turnover (total alkaline phosphatase (TALP); bone alkaline phosphatase (BALP); serum type 1 procollagen N-terminal (P1NP); and serum collagen type 1 cross-linked β-C-telopeptide (β-CTX). All plasma and serum samples were stored frozen, initially at -20oC and subsequently at -80oC. Urine was collected into a sterile container at the second void of the day after an overnight fast acidified with concentrated hydrochloric acid and stored at -20oC. Serum was analysed for 25(OH)D in the laboratory at DPHRU in duplicate using a chemiluminescent immunoassay (Liaison, DiaSorin Inc., Stillwater, MN, USA). The DPHRU laboratory participates in the international Vitamin D External Quality Assessment Scheme (DEQAS, www.deqas.org) and holds the certificate of proficiency. All other analyses were conducted at MRC Human Nutrition Research (HNR), Cambridge, UK; the methods and assay performance are given in the supplementary material. The number of samples successfully collected, transported and analysed varied depending on the analyte and timepoint. The footnote to Table 5 details the numbers of biochemical datapoints for each analyte by group and timepoint.

Serum calcium was corrected for albumin (calciumcorr) by normalising to an albumin concentration of 40 g/l:(21) Calciumcorr (mmol/l) = SCa +[0.02 x (40 – Salbumin)], where SCa and Salbumin are the serum concentrations of calcium (mmol/l) and albumin (g/l) respectively.

The ratio of tubular maximum reabsorption rate of phosphate to glomerular filtration rate (TmP/GFR) was derived using the following equations after calculation of the tubular resorption of phosphate (TRP):(22)

(a) if TRP ≤0.86 then TmP/GFR mmol/l = SP x TRP or

(b) if TRP >0.86 then TmP/GFR mmol/l = 0.3 x SP x [TRP/(1-(0.8 x TRP))]

where TRP = [(UP/SP) x (SCr/UCr)] and SP and SCr are the serum phosphate (mmol/l) and creatinine (mmol/l) concentrations and UP (mmol/l) and UCr (mmol/l) are their respective fasting urine concentrations.

Estimated glomerular filtration rate was calculated using the Chronic Kidney DiseaseEpidemiology Collaboration (CKD-EPI) formula for females(23) but without the factor for African-American ethnicity,(24) as per the South African guidelines: eGFR (ml/min/1.73m2) **= 141 × min (SCr/0.7, 1)-0.329 × max(SCr/0.7, 1)-1.209 × 0.993age × 1.018**, where SCr is serum creatinine concentration in mg/dl (i.e **SCr** in µmol/l x 0.0113) and age is in years.

Statistical methods

Data were analysed using DataDesk 6.3.1 (Data Description Inc, Ithaca, NY). Summary statistics are presented as mean ± standard deviation (SD) for normally distributed data or median [25th percentile, 75th percentile (IQR)] for skewed distributions. Based on findings from the baseline study, fat mass-to-lean mass2 (fat:lean2) was used to compare body composition between the groups.(18) All continuous variables were transformed to natural logarithms prior to data manipulation and analysis. This enabled the differences between groups and between timepoints to be expressed as a sympercent ([difference/mean] x 100) (25) and, for positively skewed data, normalised the distribution. Summary sympercent data are presented as percentage mean difference ± SE.

Two approaches were used to evaluate and compare the changes in each variable over time within the 3 groups, utilising Linear Model software in DataDesk, with Scheffé post hoc tests.

1. Firstly, using repeat-measures ANOVA and ANCOVA in hierarchical models constructed for each variable of interest, with individual identifier (nested by group), timepoint, group and a group-by-timepoint interaction term. Weight and bone area were included to adjust bone mineral data for the possible effects of bone and body size(26); height was not included because it was not anticipated to change in these women over 12 months. These models evaluated the size and significance of between-group differences at baseline and 12 months and of within-group differences between values at baseline and 12 months. They also tested whether the change from baseline differed significantly between groups.
2. Secondly, ANCOVA models were constructed with the value at 12 months of each variable of interest as the dependent variable and baseline value and group as independent variables. Adjustment for the possible influence of differences and change in bone and body size was achieved by including mean height, mean and change in weight, and mean and change in bone area between baseline and 12 months. These models quantified the size effect of the difference between each pair of groups in the change from baseline to 12 months.

The summary data and models presented are only for those women included in the dataset at 12 months, as detailed in the results section. Including all women who had participated at baseline, but not at 12 months, made no material difference to the results described. We also present models with no additional adjustment for covariates such as lifestyle factors, because they did not differ significantly between the groups. A number of women in Ppres had initiated ART during the 12 months since baseline (*n* = 11 of 60), while some Plow women who had been expected to initiate ART had not done so (*n* = 9 of 60). To more closely consider the possible effects of ART on the measured outcomes the models were repeated excluding these 18 women. The two groups of HIV-positive women in the dataset restricted by ART status are designated as PpresN (Ppres not ART exposed) and PlowY (Plow exposed to ART).

**Results**

The flow of participants through the study and the reasons for loss-to-follow up are detailed in **Figure 1**. Of the 247 women measured at baseline, 39 were not available at 12 months for, generally because they could not be contacted. In addition, data from 21 women were excluded because of pregnancy/lactation in the interim period or, in the Nref group, because they had become HIV-positive. There were no significant differences in baseline CD4 count, bone and biochemical variables and most of the anthropometry between women in the same group who were included at 12 months and those who were not. The exceptions were that women included at 12 months in the Nref and Ppres groups were significantly older, and in both the Nref and Plow groups were heavier with greater BMI, fat mass, waist circumference and hip circumference at baseline than those in the same group who were not included at 12 months (data not shown).

**Table 1** gives the ages and other characteristics at baseline and 12 months of the participants included in the follow-up study by their original group at baseline. On average, Ppres women were older and had more pregnancies that Nref but there were no significant differences in other characteristics. Table 1 also gives, for HIV-positive women, data on CD4 counts and ART initiation and duration. Over 85% of participants requiring ART were treated with a combination of TDF, lamivudine and efavirenz.

Changes in anthropometry and bone measures by HIV status at baseline

**Table 2** presents the anthropometric, body composition and aBMD data at baseline and 12 months for participants in the follow-up study. Table 2 also details the statistical significance in the hierarchical linear models (Method 1) of within-individual change over time in each variable by group and of group-by timepoint interactions which indicate whether the time effect differed significantly between groups. **Table 3** gives the percentage changes over time within each group from the models presented in Table 2. **Figure 2** illustrates these changes over time by group for all women and for women defined by their ART status at 12 months. **Table 4** gives the percentage differences in change over time between groups from the ANCOVA models (Method 2).

Plow women were significantly lighter at baseline with lower fat mass, lean mass and fat:lean2 ratio than Ppres or Nref. Nref and Plow gained significant amounts of weight and body fat over the 12 months, but Ppres had no significant anthropometric changes. Despite their larger increases in weight and fat mass, Plow remained significantly lighter with less fat mass than Nref.

The mean baseline aBMD values of Ppres and Plow women at the different skeletal sites were generally slightly lower than Nref. These differences were not significant in cross-sectional models, but were statistically significant at the hip and WBLH in the hierarchical longitudinal models. The aBMD of Nref and Ppres women increased significantly at the total hip and aBMD at the lumbar spine was also increased in Nref. These changes were largely associated with the increase in body weight, and were diminished and not significant after size adjustment. Conversely, despite their increase in body weight, by 12 months the aBMD of Plow women had significantly decreased by 2-3% at the femoral neck and lumbar spine, before and after size adjustment, with smaller decreases at the total hip and WBLH.

With the dataset restricted by ART-exposure status at 12 months that included only Ppres women who had not initiated ART (PpresN) and Plow women who had (PlowY), the size effects and statistical significance of the changes within each group (Method 1, Figure 2 A and B) and the differences in change over time between groups (Method 2, Table 4) were generally similar or slightly greater than in the full dataset, despite the smaller numbers of women. These analyses indicate that both PlowY and Nref had gained weight and fat mass relative to PpresN, whereas PlowY experienced significant decreases in aBMD relative to the other two groups, both before and after size adjustment, the greatest differences occurring at the lumbar spine (difference ± SE: -3.9 ± 0.8% relative to Nref, -3.0 ± 0.9% relative to PpresN). A greater number of women in PlowY lost more than 0.03 g/cm2 of aBMD than the other two groups, for example at the lumbar spine (Nref = 8%, PpresN = 4%, PlowY = 50%, *p* ≤0.0001).

Changes in markers of calcium and phosphorus metabolism

**Table 5** presents the data and results of the hierarchical models for 25(OH)D and markers of calcium and phosphorus metabolism. There was no significant group-by-timepoint interaction for 25(OH)D. There were also no significant differences in 25(OH)D over time or between groups other than that Ppres had significantly higher concentrations at 12 months than at baseline and than Plow. There were also no significant group-by-timepoint interactions for serum phosphate, TmP/GFR or eGFR-MDRD. Similar findings were obtained using the dataset restricted by ART-exposure status.

In contrast, a significant group-by-timepoint interaction was observed for TALP (*p* ≤0.001, Table 5). TALP increased over time in all 3 groups (change over time ± SE: Nref = +10.8 ± 4.4%, *p* = 0.053; Ppres = +15.6 ± 4.5%, *p* = 0.003; Plow = +35.2 ± 3.9%, *p* ≤0.001) but significantly more so in Plow (Plow - Nref = +28.7 ± 5.5%, *p* ≤0.001; Plow - Ppres = +26.5 ± 5.4%, *p* ≤0.001). The magnitude of the differences in change over time between Plow and the other groups was accentuated in the dataset restricted by ART-exposure status, (PlowY - Nref = +34.4 ± 5.3%, *p* ≤0.001; PlowY - PpresN = +39.5 ± 5.6%, *p* ≤0.001). BALP measured only at 12 months showed similar group differences to TALP (Table 4 and, in the restricted dataset, PlowY – Nref: BALP = +29.0 ± 5.3%, *p* ≤0.001). The two measures were closely correlated (R2 = 58.9%; *p* ≤0.001), indicating that the increase in TALP in Plow at 12 months was predominantly due to an increase in the bone isoenzyme.

P1NP and β-CTX were also significantly higher in Plow compared to Nref at 12 months (Table 5). There was no significant difference in PTH at 12 months between Plow or PlowY and Nref. However, PTH at 12 months in Ppres was significantly lower than Plow (Table 4). For all three of these analytes, restricting the dataset by ART-exposure status increased the magnitude and significance of these differences (PlowY-Nref; P1NP = +47.6 ± 8.2%, *p* ≤0.001 and CTX = +48.4 ± 11.9%, *p* ≤0.001; Ppres-Plow: PTH = -32.4 ± 10.5%, *p* = 0.009).

Significant group-by-timepoint interactions were seen for serum albumin and fasting UMg/Cr (Table 4). Serum albumin was lower in Plow at baseline than the other 2 groups but had increased significantly by 12 months towards Nref values. Serum albumin in Ppres was also lower than Nref at baseline, but higher than Plow, and was little changed at 12 months. There was a marked increase in UMg/Cr in Plow from baseline to 12 months compared to the other two groups (difference over time ± SE: +40.2 ± 10.4%, *p* ≤0.001).

**Discussion**

This is the first prospective study comparing DXA-defined changes in aBMD, vitamin D status and biochemistry in HIV-positive women not on ART at baseline and HIV-negative women in Africa. Women who had a CD4 count low enough to warrant ART initiation, based on South African health department guidelines at the time, lost significant amounts of bone mineral over 12 months when assessed using DXA. Bone loss averaged 2-3% over one year, a rate which exceeds the 1-2% annual decreases in aBMD seen in older women in early menopause. The women in this study were premenopausal and, on average, in their early thirties, when no bone loss would be expected and their average aBMD values at baseline were within -0.5 SD (i.e. z-score close to zero) of the HIV-negative reference group.(18). Indeed, increases or no change in aBMD over 12 months were observed in both the reference women and HIV-positive women with preserved CD4 counts.

The magnitude and the statistical significance of the aBMD decreases in Plow were unchanged or increased after excluding Plow women who remained unexposed to ART at 12 months. In addition, the bone loss observed in Plow women, especially those exposed to ART, was in spite of increased body weight and improved CD4 count and serum albumin concentration. These findings, plus the lack of difference in bone measures at baseline between the two HIV-positive groups when they were not exposed to ART, strongly suggests that the observed bone loss in Plow was a result of ART exposure rather than severity of HIV-infection. Decreases in aBMD due to TDF have been reported in longitudinal studies of HIV-negative African women and men receiving pre-exposure prophylaxis to prevent HIV-infection, although generally of a smaller magnitude to those observed in the HIV-positive women in this study, possibly because of lower treatment adherence.(27,28)

Unlike previous reports e.g.(10-12) there was no indication that either HIV-infection or ART-exposure was associated with compromised vitamin D status in these South African women. The mean 25(OH)D concentration in all 3 groups exceeded 50 nmol/l at both baseline and 12 months with no decrease over time in Ppres or Plow. Similarly, there was no evidence from the serum phosphate, TmP/GFR and eGFR data that either HIV-infection or ART-exposure was associated with renal damage, as has been reported by others in connection with TDF exposure.e.g.(29,30) In contrast, the decreases in aBMD in Plow were associated with higher bone turnover at 12 months as shown by serum concentrations of TALP, BALP, P1NP and β-CTX, effects that were accentuated when Plow women not exposed to ART were excluded. This suggests a direct or indirect effect of ART on bone, resulting in an increased rate of bone remodelling and loss of skeletal mineral despite no difference in PTH between the groups. The observed increase in fasting urinary magnesium excretion (UMg/Cr) in Plow also fits with this possibility, although this may also reflect the magnesium stearate present in many TDF preparations.

There were very high rates of overweight and obesity in this cohort of urban, black South African women.(18) Women who initiated ART gained weight rapidly, largely due to an increase in fat mass rather than in lean mass. Greater fat accumulation in relation to ART-associated weight gain has been reported in other settings and it has been suggested that this is due to mitochondrial dysfunction.(31) In the South African women, the deposition of fat rather than muscle with ART may also be a consequence of their poor diet and low physical activity.(19) Whatever the mechanism, such rapid increases in adiposity are likely to increase the risk of poor cardiometabolic outcomes.

This study is limited by its observational design, the absence of data on HIV-viral load, HIV-clade and duration of HIV-infection, and the relatively small numbers. It was conducted in women from Soweto, a poor township of South Africa at latitude 26oS, and the results may not extrapolate to other populations. In addition, we had no data on markers specific for proximal tubule damage as reported in connection with TDF-exposure,(29,30) such as retinol-binding protein or beta-2 microglobulin. The strengths are the longitudinal design, that no participant was exposed to ART at baseline, that an HIV-negative reference group of similar age and background was included, and that >85% of women exposed to ART received standard South African first-line treatment removing the confounding effects of various ART combinations.

In summary, this study suggests that, in urban, black South African women, HIV-infection *per se* has no discernible effects on bone mineral status over a 12-month period but that exposure to TDF-based ART is associated with loss of bone mineral and an increase in bone turnover. Newer prodrugs of TDF such as Tenofovir Alafemamide Fumarate (TAF) have been demonstrated in controlled trials to be more bone-sparing than TDF.(32) TAF has been licensed in the US and Europe but is not commonly available in Africa. It is unlikely to replace TDF for some time because of its higher cost, a difference in affordability that will increase greatly when generic TDF becomes available after its patent expires in 2017. Given that TDF is currently part of first-line ART in many African nations and other low- and middle-income countries, there is a need to develop better awareness among clinicians of possible decreases in bone mineral density, and to ascertain whether such bone loss is progressive with longer duration of ART exposure, ultimately resulting in osteoporosis and increased fracture risk.

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Authors responsibilities: MMH, AP and JMP designed the study; MMH conducted the study; JMP and SAN provided senior oversight of the study in South Africa; KW provided senior oversight and interpretation of the DXA data; AP provided senior oversight and interpretation of the biochemical data; MMH and AP jointly conducted the statistical analysis and drafted the paper; all authors had full access to the data and contributed to interpretation of the findings. MMH and AP have responsibility for data integrity. All authors approved the manuscript for publication.

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Table 1. Subject characteristics at baseline and 12 months by initial HIV status

Timepoint Nref Ppres Plow

months *n* = 67 *n* = 60 *n* = 60

Age (y) 0 31.4±8.5 34.5±6.0a 33.2±6.2

12 32.2±8.5 35.5±6.0a 34.1±6.2

Gravidity 0 1 [0,2] 2 [2,3]b 2 [1,3]

Regular menses (%) 0,12 69, 81 83, 82 75, 77

Hormonal contraception (%) 0,12 39, 36 33, 27 35, 32

Any fracture to date (%) 0,12 28, 28 23, 23 13, 13

Current smoker (%) 0,12 7, 3 15, 15 8, 8

Alcohol consumer (%) 0 15 30 22

Calcium intake (mg/d) 0 670±372 656±306 727±576

ART initiated (*n*,%) 12 - 11 (18) 51 (85)c

ART duration (d)¶ 12 - 198 [118,279] 330 [298,347]c

CD4 counts (x106/l)

All HIV-positive women 0 - 416 [343,466] 176 [102,224]c

12 - 402 [338,474] 233 [179,333]c

Restricted set# 0 - 428 [352, 478] 151 [93,212]c,d

12 - 419 [350,482] 230 [178,348]c,d

Footnote to Table 1

Data are mean ± SD, median [IQR] or percentage of subjects reporting ‘Yes’.

Nref = HIV-negative throughout; Ppres = HIV-positive with preserved CD4 counts at baseline; Plow = HIV-positive with low CD4 counts at baseline.

¶ In women who had initiated ART by 12 months (PPres n=11, Plow, n=51)

# Restricted set by ART-exposure status at 12 months: PpresN = Ppres not exposed to ART (n=59), PlowY = Plow exposed to ART (n=51).

Significance of group differences at same timepoint, or within-individual timepoint differences in the same group using ANOVA (continuous variables) or Chi-square test (frequencies):

a *p* ≤0.05 different to Nref

b *p* ≤0.01 different to Nref

c *p* ≤0.001 different to Ppres

d *p* ≤0.001 different to baseline

There were no other significant differences.

Table 2. Anthropometry and bone mineral densities at baseline and 12 months by initial HIV status

Nref Ppres Plow Group-by-

timepoint

Baseline 12 months Baseline 12 months Baseline 12 months *p*

*Anthropometry*

Height (m) 1.58±0.06 1.58±0.06 1.59±0.06 1.60±0.06 1.59±0.06 1.59±0.06 0.66

Weight (kg) 72.2±17.4 74.1±17.5h 72.1±17.4 71.6±17.3a 64.5±15.7a,d 66.8±15.4a,d,g **≤0.001**

BMI (kg/m2) 29.1±7.1 29.9±7.2h 28.3±6.5c 28.1±6.6a 25.5±6.0a,d 26.4±5.9a,d,g **≤0.001**

Fat mass, WBLH (kg) 27.8±12.1 29.4±12.2h 26.0±9.7c 26.0±10.3a 21.3±9.3a,d 23.7±10.7a,d,g  **0.002**

Lean mass, WBLH (kg) 39.0±5.8 39.2±5.8 39.5±5.9 39.0±5.8 36.8±5.0a,d 37.4±5.6a,d **0.05**

Fat:lean2 (1000\*kg/kg2) 17.8±4.9 18.7±4.9i 16.4±4.6a 16.8±5.2a 15.4±5.2a,d 16.7±5.3a,g 0.07

Waist (cm) 88.5±14.7 91.4±14.7h 89.8±15.2 90.9±16.6 85.1±14.3a,d 87.5±13.7a,e,h 0.20

Hip (cm) 109.7±14.1 111.7±14.7i 107.5±14.2c 108.4±13.5a 100.7±13.7a,d 104.7±12.2a,d,g **0.01**

Waist:hip (cm/cm) 0.81±0.07 0.82±0.07 0.84±0.09a 0.84±0.08 0.85±0.06a 0.83±0.06 0.11

*Bone mineral density (aBMD g/cm2)*

Total hip 1.018±0.151 1.042±0.153g 0.989±0.134a 1.008±0.135a,h 0.985±0.128a 0.993±0.128a **0.04**

Femoral neck 0.941±0.120 0.939±0.130 0.918±0.133a 0.916±0.134a 0.923±0.135b 0.901±0.134a,f,h **0.02**

Lumbar spine 1.019±0.132 1.032±0.127i 1.023±0.116 1.020±0.112 1.011±0.135f 0.990±0.140a,d,g **≤0.001**

WBLH 0.967±0.088 0.966±0.087 0.948±0.077a 0.952±0.075a 0.949±0.083a 0.942±0.082a,e,i **0.02**

**Footnote to Table 2**

Nref, *n* = 67, HIV-negative women; Ppres, *n* = 60, HIV-positive with preserved CD4 counts at baseline; Plow, *n* = 60, HIV-positive with low CD4 counts at baseline, WBLH = whole body less head. Data are means ± SDs, aBMD data are unadjusted. Significance of differences from Scheffé *post hoc* tests from hierarchical linear models of the variable in natural logarithms with timepoint (0/12 months), group (Nref/Ppres/Plow), ID (nested within group) and a group-by-timepoint interaction, as follows: between Ppres or Plow and Nref at each timepoint a ≤0.001, b≤0.01, c ≤0.05; between Ppres and Plow at each timepoint: d ≤0.001, e ≤0.01, f ≤0.05; between baseline and 12 months in each group: g ≤0.001, h≤0.01, i ≤0.05.

Table 3. Percentage change over 12 months within each group by initial HIV status

Nref Ppres Plow

%∆±SE *P* %∆±SE *P* %∆±SE *p*

*Anthropometry*

Weight **+2.7±0.8 0.004** -0.7±0.9 0.70 **+3.8±0.9 ≤0.001**

BMI **+2.6±0.80.007**-1.0±0.9 0.50 **+3.8±0.9≤0.001**

Fat mass **+6.2±2.0 0.008** -0.8±2.1 0.93 **+9.4±2.1 ≤0.001**

Lean mass +0.5±0.6 0.63 -1.2±0.6 0.17 +0.7±0.6 0.50

Fat:lean2 **+5.1±1.9 0.03** +1.5±2.0 0.75 **+8.0±2.0 ≤0.001**

Waist **+3.3±0.9 0.002** +1.0±1.0 0.56 **+2.8±1.0 0.01**

Hip **+1.9±0.7 0.04** +1.0±0.8 0.47 **+4.1±0.8 ≤0.001**

Waist:hip +1.4±0.9 0.28 +0.1±1.0 0.99 -1.3±1.0 0.37

*Bone mineral density, unadjusted*

Total hip **+2.5±0.6 ≤0.001** **+1.9±0.6 0.007** +0.4±0.6 0.78

Femoral neck -0.1±0.6 0.99 -0.3±0.7 0.92 **-2.4±0.7 0.002**

Lumbar spine **+1.4±0.5 0.03** -0.2±0.5 0.94 **-2.0±0.6 ≤0.001**

WBLH-0.2±0.3 0.86 +0.4±0.3 0.51 **-0.9±0.3 0.03**

*Bone mineral density, size-adjusted*

Total hip +0.7±0.6 0.49 -0.3±0.6 0.84 **-1.7±0.6 0.02**

Femoral neck -0.4±0.7 0.84 -0.2±0.7 0.94 **-2.7±0.7 ≤0.001**

Lumbar spine +1.2±0.5 0.07 -0.2±0.6 0.92 **-2.3±0.6 ≤0.001**

WBLH -0.0±0.3 0.99 +0.3±0.3 0.62 -0.7±0.4 0.14

**Footnote to Table 3**

Nref, *n* = 67, HIV-negative women; Ppres, *n* = 60, HIV-positive with preserved CD4 counts at baseline; Plow, *n* = 60, HIV-positive with low CD4 counts at baseline; WBLH, whole body less head. Data are mean percentage changes (%∆) ± SEs expressed as sympercents from hierarchical linear models. All continuous variables were transformed to natural logarithms. Predictor variables were time (0/12 months), group (Nref/Ppres/Plow) and ID (nested within group) and a group-by-timepoint interaction. Adjustment of aBMD for bone and body size was achieved by including bone area and weight in the models. The *p*-values for the interaction are in Table 1.

Table 4. Percentage difference between groups in change over 12 months by initial HIV status and by ART status at 12 months

HIV status at baseline ART status at 12 months

Ppres-Nref Plow-Nref Plow–Ppres Group PpresN-Nref PlowY-Nref PlowY-PpresN Group

%∆±SE %∆±SE %∆±SE *P* %∆±SE %∆±SE %∆±SE *p*

*Anthropometry*

Weight (kg) -3.5±1.2b +0.4±1.2 +3.9±1.2b **0.002** -3.9±1.3b +1.4±1.3 +5.3±1.4a **≤0.001**

Fat mass (kg) -7.4±2.8c +1.3±3.0 +8.7±3.0c **0.006** -8.7±3.0c +3.3±3.1 +12.1±3.4b **≤0.001**

Lean mass (kg) -1.6±0.8 -0.2±0.8 +1.4±0.9 0.12 -1.6±0.9 +0.1±0.9 +1.7±1.0 0.13

Fat:lean2 (1000\*kg/kg2) -4.3±2.7 +1.4±2.8 +5.7±2.8 0.11 -5.7±2.9 +2.2±3.0 +7.9±3.1c **0.03**

Waist (cm) -2.1±1.3 -0.9±1.3 +1.2±1.3 0.25 -2.0±1.4 -0.5±1.3 +1.5±1.4 0.32

Hip (cm) -1.2±1.0 +0.9±1.0 +2.1±1.0 0.12 -1.7±1.0 +1.6±1.1 +3.3±1.1c **0.02**

Waist:hip (cm/cm) +0.0±1.2 -0.9±1.2 -0.9±1.2 0.70 +0.2±1.2 -1.4±1.2 -1.6±1.3 0.38

*Bone mineral density, unadjusted*

Total hip (g/cm2) -0.8±0.8 -2.3±0.8c -1.5±0.8 **0.02** -0.5±0.9 -2.5±0.9c -1.9±1.0 **0.02**

Femoral neck (g/cm2) -0.3±0.9 -2.4±0.9c -2.1±0.9 **0.02** -0.1±1.0 -2.5±1.0c -2.5±1.0 **0.02**

Lumbar spine (g/cm2) -1.5±0.8 -3.5±0.8a -1.9±0.8c **≤0.001** -0.9±0.8 -3.9±0.8a -3.0±0.9b **≤0.001**

WBLH (g/cm2) +0.4±0.4 -0.8±0.4 -1.3±0.5c **0.02** +0.6±0.5 -1.2±0.5c -1.8±0.5a **≤0.001**

*Bone mineral density, size-adjusted*

Total hip (g/cm2) -1.1±0.7 -2.1±0.7c -1.0±0.7 **0.02** -1.0±0.8 -2.5±0.8b -1.4±0.9 **0.01**

Femoral neck (g/cm2) +0.2±0.9 -1.9±0.9 -2.1±1.0 0.06+0.5±1.0 -2.1±1.0 -2.7±1.1 **0.04**

Lumbar spine (g/cm2) -1.4±0.8 -3.2±0.8a -1.8±0.8 **≤0.001** -0.9±0.8 -4.1±0.8a -3.2±0.9b **≤0.001**

WBLH (g/cm2) +0.1±0.5 -1.0±0.5 -1.1±0.5 **0.04** +0.2±0.5 -1.3±0.5c -1.6±0.5b **0.005**

**Footnote to Table 4**

Nref, *n* = 67, HIV-negative women; Ppres, n = 60, HIV-positive with preserved CD4 counts at baseline; Plow, n = 60, HIV-positive with low CD4 counts at baseline; PpresN, *n* = 49, HIV-positive with preserved CD4 counts at baseline not started on ART by 12 months; PlowY, *n* = 51, HIV-positive with low CD4 counts at baseline and started on ART during the 12 months; WBLH, whole body less head. Data are mean percentage differences (%∆) ± SEs expressed as sympercents from Scheffé *post-hoc* tests from ANCOVA models (y = aBMD at 12 months, x = aBMD at baseline; group (Nref/Ppres/Plow)). Adjustment for bone and body size was achieved by including height, change and mean bone area, change and mean weight in the models. All continuous variables were transformed to natural logarithms prior to data analysis. Significance of difference between the groups in each pair: a ≤0.001, b≤0.01, c ≤0.05.

Table 5. Biochemistry at baseline and 12 months by initial HIV status

Nref Ppres Plow Group-by-

timepoint

Baseline 12 months Baseline 12 months Baseline 12 months *p*

*Serum*

25(OH)D nmol/l 58.0±15.7 63.3±16.6 60.5±17.0 67.1±18.6h 60.6±22.6 61.3±20.3e 0.24

Phosphate mmol/l 1.10±0.19 1.16±0.20h 1.10±0.20 1.20±0.19h 1.15±0.16b 1.22±0.21 0.26

Calciumcorr mmol/l 2.49±0.10 2.48±0.10 2.51±0.10 2.48±0.10 2.52±0.13 2.49±0.11 0.44

Magnesium mmol/l 0.80±0.06 0.81±0.06 0.80±0.06 0.80±0.07 0.78±0.07 0.80±0.06 0.60

Albumin g/l 41.8±3.9 40.7±3.3 39.6±3.0c 38.7±4.5b 36.0±6.0a,d 38.7±4.8b,g **≤0.001**

Creatinine µmol/l 84.0±7.7 84.5±8.3 80.7±7.8 81.9±8.9 80.5±8.5 80.8±9.0b 0.70

eGFR ml/min/1.73m2 80.4±11.9 79.7±11.4 82.8±10.4 81.1±11.9 84.2±12.3 83.0±11.2 0.40

TALP¶ U/l43.4[37.7,55.6] 51.0[40.7,61.1]i 45.5[36.5,53.8] 50.0[40.7,59.9]h 48.7[36.4,60.9]e 79.1[56.7,92.8]a,d,g **≤0.001**

BALP¶ U/l - 18.3[15.4,21.1] - 16.9[14.4,19.5] - 23.3[17.5,30.8]a,d **≤0.001#**

P1NP¶ µg/l - 47.5[37.0,65.5] - 51.8[39.5,73.9] - 72.8[55.4,100.3]a,e **≤0.001#**

β-CTX¶ ng/l - 117[73,179] - 153[98,231]c - 171[117,254]a **≤0.001#**

PTH¶ ng/l - 23.6[19.1,31.6] - 20.2[12.7,27.9]c - 22.9[17.3,37.5] **0.04#**

*Urine*

Phosphate:Cre¶ 0.97[0.76,1.34] 0.92[0.68,1.34] 1.23[0.94,1.49] 1.32[0.89,1.84]b 0.76[0.61,1.14] 1.29[0.87,1.63] 0.31

Calcium:Cre¶ 0.08[0.03,0.17] 0.07[0.03,0.15] 0.08[0.04,0.15] 0.08[0.03,0.12] 0.07[0.03,0.14] 0.07[0.03,0.18] 0.63

Magnesium:Cre 0.16±0.07 0.17±0.08 0.15±0.09 0.15±0.07 0.16±0.07 0.24±0.11d **0.002**

TmP/GFR mmol/l 1.19±0.29 1.27±0.29 1.14±0.30 1.22±0.26 1.33±0.32 1.30±0.30 0.30

**Footnote to Table 5**

Data are means ± SDs for normal distributions, for those with positive skew (¶) are median [25,75percentile]. # indicates *p-*value is for group effect at 12 months. Abbreviations are 25(OH)D, 25-hydroxyvitamin D; eGFR-MDRD, estimated glomerular filtration rate using the MDRD formula; TALP, total alkaline phosphatase; BALP, bone alkaline phosphatase; P1NP, serum type 1 procollagen N-terminal; β-CTX, serum collagen type 1 cross-linked β-C-telopeptide; and serum type 1 procollagen N-terminal; PTH, parathyroid hormone; Cre, urine creatinine used to develop urine mineral ratios in mmol/mmol; TmP, tubular maximum reabsorption rate of phosphate. Values for BALP, P1NP, CTX and PTH are available at 12 months only. Significance of differences from Scheffé *post-hoc* tests from hierarchical linear models of the variable in natural logarithms with timepoint (0/12 months), group (Nref/Ppres/Plow), ID (nested within group) and a group-by-timepoint interaction, as follows: between Ppres or Plow and Nref at each timepoint

a ≤0.001, b≤0.01, c ≤0.05; between Ppres and Plow at each timepoint: d ≤0.001, e ≤0.01, f ≤0.05; between baseline and 12 months in each group:  g ≤0.001, h≤0.01, i ≤0.05. Numbers of biochemical datapoints are:25(OH)D - baseline: Nref = 67; Ppres = 60; Plow = 60; 12 months: Nref = 64; Ppres = 60; Plow = 59; Other blood analytes - baseline: Nref = 41; Ppres = 37; Plow = 51; 12 months: Nref = 65; Ppres = 60; Plow = 60; Urine mineral ratios - baseline: Nref = 50; Ppres = 42; Plow = 38; 12 months: Nref = 60; Ppres = 55; Plow = 42; TmP/GFR - baseline: Nref = 28; Ppres = 27; Plow = 33; 12 months: Nref = 58; Ppres = 55; Plow = 42.

**Figure Legends**

**Figure 1.**

Flow through of study participants in the three groups from recruitment to 12 months.

Nref = HIV-negative women; Ppres = HIV-positive women with preserved CD4 counts at baseline; Plow = HIV negative women with low CD4 counts at baseline; ART = antiretroviral therapy.

**Figure 2.**

Percentage changes in weight and unadjusted aBMD from baseline to 12 months in HIV-negative and HIV-positive women (A) according to group designation at baseline and (B) ART status at 12 months.

ART = antiretroviral therapy; Nref = HIV-negative at baseline and 12 months, *n* = 67; Ppres = HIV-positive women with preserved CD4 counts at baseline, *n* = 60; Plow = HIV negative women with low CD4 counts at baseline *n* = 60; PpresN = Ppres women not exposed to ART by 12 months, *n* = 49; PlowY = Plow women exposed to ART by12 months, *n* = 51;. Data are mean percentage changes from baseline to 12 months obtained from Scheffé *post-hoc* tests from hierarchical linear models of the variable in natural logarithms with timepoint (0/12 months), group (Nref/Ppres/Plow), ID (nested within group) and a group-by-timepoint interaction; error bars are SEs. Significance of change at 12 months from baseline: \**p* ≤0.05; \*\**p* ≤0.01; \*\*\**p* ≤0.001.

Figure 1 (separate file provided)

Figure 2 (separate file provided)