Improved adipose tissue function with initiation of protease inhibitor-only ART

Robert T. Maughan1*, Eoin R. Feeney1, Emilie Capel2, Jacqueline Capeau2, Pere Domingo3, Marta Giralt4,5, Joep M. A. Lange6,7†, Praphan Phanuphak7,8, David A. Cooper9, Peter Reiss6,7 and Patrick W. G. Mallon1 on behalf of the HIVNAT-019 Study Group‡

1HIV Molecular Research Group, School of Medicine, University College Dublin, Dublin, Ireland; 2Sorbonne Universities, UPMC Univ Paris 06, INSERM UMRS 938, Paris, France; 3Infectious Diseases Unit, Hospital de la Santa Creu/Sant Pau, Barcelona, Spain; 4Department of Biochemistry and Molecular Biology, University of Barcelona, Barcelona, Spain; 5CIBER Fisiopatología de la Obesidad y Nutrición, Barcelona, Spain; 6Academic Medical Center, Department of Global Health and Division of Infectious Diseases, and Amsterdam Institute for Global Health and Development, Amsterdam, The Netherlands; 7The HIV-Netherlands Australia Thailand Research Collaboration (HIV-NAT) - Thai Red Cross Aids Research Center (TRCARC), Bangkok, Thailand; 8Department of Medicine, Chulalongkorn University, Bangkok, Thailand; 9Kirby Institute, University of New South Wales, Sydney, Australia

*Corresponding author. Tel: +353-1-716-4584; Fax: +353-1-716-4539; E-mail: robert.maughan@ucdconnect.ie
†Deceased 17 July 2014.
‡Additional members of the HIVNAT-019 Study Group are listed in the Acknowledgements section.

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Objectives: Use of ART containing HIV PIs has previously been associated with toxicity in subcutaneous adipose tissue (SAT), potentially contributing to the development of lipodystrophy and insulin resistance. However, the effect of PIs on SAT function in ART-naive patients independent of other ART classes is unknown. This study aimed to elucidate the effect of initiating PI-only ART on SAT function in ART-naive subjects.

Methods: In the HIVNAT-019 study, 48 HIV-infected, ART-naive Thai adults commencing PI-only ART comprising lopinavir/ritonavir/saquinavir for 24 weeks underwent assessments of fasting metabolic parameters and body composition. In a molecular substudy, 20 subjects underwent SAT biopsies at weeks 0, 2 and 24 for transcriptional, protein, mitochondrial DNA (mtDNA) and histological analyses. ClinicalTrials.gov registration number: NCT00400738.

Results: Over 24 weeks, limb fat increased (+416.4 g, P = 0.023), coinciding with larger adipocytes as indicated by decreased adipocyte density in biopsies (-32.3 cells/mm², P = 0.047) and increased mRNA expression of adipogenesis regulator PPARG at week 2 (+58.1%, P = 0.003). Increases in mtDNA over 24 weeks (+600 copies/cell, P = 0.041), decreased NRF1 mRNA expression at week 2 (-33.7%, P < 0.001) and increased COX2/COX4 protein ratio at week 24 (+288%, P = 0.038) indicated improved mitochondrial function. Despite decreased AKT2 mRNA at week 2 (-28.6%, P = 0.002) and increased PTPN1 mRNA at week 24 (+50.3%, P = 0.016) suggesting insulin resistance, clinical insulin sensitivity [by homeostasis model assessment (HOMA-IR)] was unchanged.

Conclusions: Initiation of PI-only ART showed little evidence of SAT toxicity, the changes observed being consistent with a return to health rather than contributing to lipodystrophy.

Introduction

Metabolic comorbidities including lipodystrophy, insulin resistance and dyslipidaemia are prevalent amongst people living with HIV (PLWH), and not only affect adherence to therapy but also contribute to cardiovascular disease risk, a leading cause of death in treated PLWH. These conditions are in part caused by subcutaneous adipose tissue (SAT) toxicity due to HIV infection and exposure to certain classes of ART. Although toxicity in SAT has been demonstrated in untreated HIV infection, the development of clinical lipodystrophy (peripheral lipoatrophy, central lipohypertrophy, dyslipidaemia and insulin resistance) is associated with ART exposure, and once established is not fully reversible. Lipodystrophy was first attributed to HIV PIs; however, subsequent studies demonstrated that treatment with thymidine analogue NRTIs (NRTIs) is the main factor, primarily via initiation of SAT mitochondrial toxicity. Due to the routine use of PIs in combination with NRTIs,
their relative contribution to ART-mediated SAT toxicity remains unclear.

*In vitro*, PIs inhibit adipogenesis, alter lipid metabolism, impair glucose uptake insulin signalling, induce mitochondrial dysfunction and alter secretion of adipocyte-derived hormones and inflammatory cytokines. In clinical studies, PLWH receiving PI-containing ART had reduced expression of key genes required for the regulation of lipid metabolism, insulin sensitivity and adipogenesis in SAT. However, the concurrent use of tNRTIs, as well as the established lipodystrophy, in many of the subjects makes it difficult to determine the specific effects of PIs in these studies.

As no study to date has explored the effects of PI therapy in the absence of other ART classes on SAT function, we aimed to elucidate the effect of initiating PI-only ART through a comprehensive prospective analysis measuring clinical metabolic parameters in parallel with histological and molecular assessments of adipogenic, lipid metabolism, insulin signalling and mitochondrial function pathways in the SAT of ART-naive subjects.

**Methods**

**Ethics**

The study was approved by the Ethics Committee of the Faculty of Medicine, Chulalongkorn Hospital (approval number 235/2004), written informed consent was obtained for all patients prior to their inclusion and all study procedures were in accordance with standard ethical guidelines.

**Study design**

A molecular substudy was performed within the HIVNAT-019 trial, an open-label, randomized trial examining the virological efficacy and pharmacodynamics of four different dosing schedules of PI-only ART containing lopinavir boosted with ritonavir co-administered with saquinavir in ART-naive, HIV-1-infected subjects over 24 weeks. Dosing schedules comprised 400/100 mg of lopinavir/ritonavir with 1000 mg of saquinavir twice daily; 400/100 mg of lopinavir/ritonavir with 400 mg of saquinavir twice daily; 266/66 mg of lopinavir/ritonavir with 1000 mg of saquinavir twice daily; and 266/66 mg of lopinavir/ritonavir with 400 mg of saquinavir twice daily. Eligible adult (>18 years), HIV-1 antibody-positive and ART-naive subjects were recruited from the Thai Red Cross Society’s Anonymus Clinic and the HIV Outpatient Immune Clinic of King Chulalongkorn Memorial Hospital in Bangkok, Thailand. Relevant inclusion/ exclusion criteria and main clinical outcomes were described previously.

**Body composition and fasting blood parameters**

Limb and trunk fat were quantified by DXA (Discovery W, Hologic) at weeks 0, 12 and 24. Abdominal subcutaneous and visceral adipose tissue areas at the fourth lumbar vertebrae along with SAT at the right mid-thigh were quantified using single-slice CT (Aquilion ONE, Toshiba), described previously.

Fasting bloods (overnight) were drawn for total, LDL and HDL cholesterol, triglycerides, insulin and glucose at weeks 0, 12 and 24. Insulin resistance was estimated using the homeostasis model assessment (HOMA-IR).

**SAT biopsies**

Molecular substudy subjects underwent biopsies of abdominal flank SAT as described previously at weeks 0, 2 and 24. Biopsied tissue was immediately aliquoted and snap frozen in liquid nitrogen for DNA, RNA and protein extraction, and a tissue sample was formalin fixed for histology.

**DNA and RNA extraction from adipose tissue**

DNA was extracted from SAT using the QIAamp DNA Mini Kit (Qiagen) and RNA was extracted from homogenized SAT using TRIreagent (Ambion) according to the manufacturer’s instructions. The resulting RNA was treated with RNase-free DNase (Promega) and purified using the RNeasy Mini Kit (Qiagen).

**Gene expression**

Expression of 55 chosen mRNA targets was determined using a quantitative PCR (qPCR) array (RealTime Ready, Roche), with array details outlined in Table S1 and Table S2 (available as Supplementary data at JAC Online).

In brief, cDNA libraries were prepared using the Transcriptor First Strand cDNA Synthesis Kit (Roche) and sample quality was verified by qPCR measurement of reference gene actin β (ACTB). cDNA libraries underwent a linear 12-cycle pre-amplification using the RealTime Ready cDNA Pre-Amp System (Roche) and then assayed in duplicate with the appropriate controls on the LightCycler 480 (Roche). Gene expression was normalized to the average of reference genes ACTB, ribosomal protein L13a (RPL13A) and TATA box binding protein (TBP).

**Mitochondrial DNA (mtDNA) content**

Adipose tissue mtDNA copy number per cell was quantified as described previously. In brief, DNA samples were quantified against a standard curve of known copy number by qPCR (LightCycler 480, Roche) with primers targeting mitochondrially encoded cytochrome b (MT-CYB; region 1) and mitochondrially encoded cytochrome c oxidase I (MT-CO1; region 2). mtDNA levels were compared with nuclear genome DNA copy number with primers targeting peroxisome proliferator-activated receptor γ (PPARG) and mtDNA copies/cell was calculated as copy number of mtDNA/copy number of nuclear DNA.

**Protein expression**

SAT samples were homogenized in extraction buffer [10 mM HEPEs, pH 7.5/5 mM EDTA/5 mM dithiothreitol/5 mM MgCl2/PI (Complete Mini, Roche)]. Protein was analysed by immunoblot as previously described, using antibodies against cytochrome c oxidase subunit 2 and 4 (COX2 and COX4) (A-6404 and A-21347, Invitrogen); PPARG and sterol regulatory element binding transcription factor 1 (SREBP1) (Sc-1984X and Sc-367X, respectively, Santa Cruz Biotechnology); and β2-microglobulin (B2M) (P0163, Dako Cytomation). Chemiluminescence was developed using horseradish peroxidase-conjugated secondary antibodies (170-6510, Bio-Rad and 711-135-152, Jackson Immunoresearch) and Immobilon ECL Plus kit (Millipore). ODs were quantified using the Multigauge 3.0 software suite (Fujifilm) and normalized to total protein content.

**Adipose tissue adipocyte density**

Paraffin-embedded SAT was cut into 3 μm sections and stained with haematoxylin phloxine saffron by standard protocols. The number of adipocytes per field (>10 magnification) was quantified using Mercator software (Explora Nova) with adipocyte density expressed as the number of adipocytes/mm².

**Statistical analysis**

Subjects with more than one biopsy available were included in molecular analyses. Changes in gene and protein expression were compared using Wilcoxon signed rank tests. Parameters expected to have continuous longitudinal change (mtDNA, adipocyte density and clinical parameters) were analysed using longitudinal marginal models with appropriate covariance structures selected using Akaike’s information criterion (AIC). Data are...
Results

Between October 2004 and March 2006, 20 of 48 subjects randomized to the main study participated in the molecular substudy. Baseline characteristics of substudy subjects were broadly comparable to those of the main study (Table 1), with both genders represented and average baseline CD4+ T cell count suggesting advanced immunosuppression (14% of 20 subjects in the molecular substudy had a CD4+ T cell count <200 cells/mm³).

Of the substudy subjects, one was lost to follow-up (week 1), two withdrew for personal reasons (weeks 13 and 21) and one required ART intensification with the addition of NRTI. In the main study, one hepatitis B virus–co-infected subject stopped following a hepatic grade 4 transaminase elevation (week 5) and one required intensification with the addition of NRTI (week 6).

Changes in clinical metabolic parameters

Over 24 weeks, no changes were observed in BMI or trunk fat by DXA, but limb fat increased significantly [+416.4 (176.1) g, P = 0.023], as expected with ART initiation (Table 2). However, the trunk:limb fat ratio did not change, suggesting a generalized gain in adiposity with ART initiation. Consistent with this, both abdominal subcutaneous and visceral adipose tissue area by CT also increased [+9.5 (4.5) cm², P = 0.039 and +4.3 (2.2) cm², P = 0.058, respectively], with no significant change in their ratio (Table 2). CT evaluation of thigh SAT revealed a non-significant increase.

Fasting total, LDL and HDL cholesterol and triglycerides all increased, with no significant change in the total cholesterol:HDL cholesterol ratio. No changes were observed in fasting glucose, insulin or HOMA-IR (Table 2).

Table 1. Baseline characteristics of study participants

|               | HIVNAT study | Molecular substudy |
|---------------|--------------|--------------------|
| N             | 48           | 20                 |
| Male, n (%)   | 20 (42)      | 11 (55)            |
| Asian, n (%)  | 48 (100)     | 20 (100)           |
| Age (years), median (IQR) | 36 (31.7, 43.3) | 39 (30.7, 43.2) |
| BMI (kg/m²), median (IQR) | 21.5 (19.4, 24.3) | 21.8 (19.5, 23.2) |
| CD4+ T cell count (cells/mm³), median (IQR) | 113.5 (67.8, 193.5) | 119 (77.5, 217.3) |
| HIV RNA (log₁₀ copies/mL), median (IQR) | 4.9 (4.6, 5.1) | 4.8 (4.6, 5) |

HIV RNA, HIV viral load.

Gene expression analysis

The 55 genes examined reflected the following functions: (i) lipid metabolism and adipogenesis; (ii) insulin signalling; (iii) markers of inflammation and cell stress; (iv) adipocyte-secreted hormones; and (v) mitochondrial function. The full list of genes and their corresponding change in expression at weeks 2 and 24 are summarized in Table S1 and Table S3, respectively. Overall, biopsies from 15 subjects contributed to gene expression analyses, with 13 and 12 paired samples for week 2 and week 24 analyses, respectively; 1 sample in the week 24 analysis did not pass initial quality checks.

At week 2, expression of 13 genes changed significantly, 8 of which remained significant after correction for multiple comparisons. Of these 13 genes, 6 were related to lipid metabolism and adipogenesis, 3 to insulin signalling, 2 to mitochondrial function and 2 to cell stress markers. At week 24, only 3 target genes had significantly altered expression, none of which remained significant after correction. Two of these were related to insulin signalling and one to mitochondrial function. The smaller number of differentially expressed genes at week 24 was due to a higher level of variance in expression levels at this timepoint.

Adipogenesis and lipid metabolism genes

The initiation of PI-only ART resulted in significant increases in expression of PPARγ, the master transcriptional regulator of adipogenesis, at week 2 [+58.1% (+23.5, +93.4), P = 0.003, Figure 1a], with a persistent but non-significant increase at week 24 [+45.4% (-32.2, +88.6), P = 0.077, Figure 1a]. This
was accompanied by increases in week 2 expression (but not week 24) of the downstream PPARγ transcriptional targets involved in lipid metabolism: fatty acid binding protein 4 (FABP4) and lipoprotein lipase (LPL) [+58.8% (+27.3, +117.3), \(P<0.001\) and +36.3% (−2.9, +45.8), \(P=0.047\), respectively, Figure 1a]. Increases in PPARγ expression and activity indicate that initiation of PI-only ART facilitated increases in SAT lipid metabolism and adipogenesis from diminished baseline levels in ART-naïve subjects despite previous associations of PIs with impaired adipogenesis and lipid metabolism.\(^{11,12}\) In addition, expression of CCAAT/enhancer binding protein α (CEBPA) and sterol regulatory element binding transcription factor 1 (SREBF1), major transcriptional regulators of adipogenesis implicated in ART-mediated adipose tissue toxicity,\(^{11,16}\) did not change (Figure 1a).

The following were also detected at week 2: (i) decreases in hormone-sensitive lipase (LIPC) expression, a key enzyme involved in the triglyceride hydrolysis in adipocytes; (ii) decreases in nuclear receptor coactivator 1 (NCOA1), a transcriptional coactivator implicated in adipose tissue energy balance,\(^{28}\) and (iii) increases in α-2-glycoprotein 1, zinc-binding (AZGP1), a PPARγ transcriptional target\(^{25}\) (Table 3).

\[\text{Insulin signalling genes}\]

\textit{In vivo}, short-term exposure to certain PIs can reduce insulin sensitivity.\(^{30}\) Supporting this, changes in gene expression suggested an inhibition of insulin signalling in SAT with PI-only ART initiation. Expression levels of V-akt murine thymoma viral oncogene homologue 2 (AKT2), a key transducer of insulin signalling, and insulin-like growth factor 2 (IGF2), which activates insulin signalling via the insulin or IGF receptors, were both significantly reduced at week 2 [−28.6% (−46.8, −8.5), \(P=0.002\) and −39.2% (−66.9, −25.7), \(P=0.011\), respectively, Figure 1b], while expression of the insulin receptor (INSR) decreased non-significantly [−33% (−36.9, +16.7), \(P=0.094\), Figure 1b]. In addition, expression of two negative regulators of insulin signalling increased: protein tyrosine phosphatase, non-receptor type 1 (PTPN1) at both week 2 and 24 [+30.5% (+6.5%, +43.5%), \(P=0.052\) and +55.6% (+22.6, +72), \(P=0.022\), respectively, Figure 1b] as well as fororkhead box O1 (FOXO1) at week 24 [+29.1% (+5.7, +82.2), \(P=0.028\), Figure 1b].

In contrast, expression of ras homologue family member Q (RHOQ) involved in insulin-activated glucose uptake increased significantly at week 2 [+42% (+27.2, +74.2), \(P=0.022\), Table 3].
Genes encoding markers of cell stress and adipocyte-secreted hormones

Elevated levels of cell stress and inflammation markers among those in adipocyte-secreted hormones are features of ART-mediated SAT toxicity. However, in this study gene expression changes indicated a reduction in cellular stress inflammation with initiation of PI-only ART. Expression levels of tumour protein 53 (TP53), a major stress-responsive activator of apoptosis, and 1κB kinase β (IKKB), a key activator of inflammatory signalling, were reduced at week 2 (−15.9% (−36, −4.1), P=0.003 and −45.5% (−68.9, −25.8), P<0.001, respectively, Figure 1c). The expression of other inflammation genes, including TNF and IL6, did not change significantly, and no changes were detected in the five target genes encoding adipocyte-secreted hormones, including adiponectin (ADIPOQ) and leptin (LEP) (Figure 1c and Table S3).

Mitochondrial function genes and mtDNA

Both HIV infection itself and exposure to ART (principally nRTIs) and reduced mtDNA content in SAT. In contrast, the initiation of PI-only ART led to gene expression changes suggesting improvements in mitochondrial function; nuclear respiratory factor 1 (NRF1) and CREB regulated transcription coactivator 3 (CRTC3), both involved in mitochondrial response to cellular and oxidative stress, were significantly reduced at week 2 (−33.7% (−46.9, −18.4), P<0.001 and −32.3% (−38.1, −17.8), P<0.001, respectively, Figure 1d) and persisted to week 24, significantly so for CRTC3 (−22% (−32.4, −1.8), P=0.027, Figure 1d). No accompanying changes were observed in the expression of the key regulator of mitochondrial biogenesis PPARγ coactivator 1α (PPARGC1A) or in the mitochondrially associated uncoupling protein 2 (UCP2) (Figure 1d), genes dysregulated in nRTI-mediated SAT toxicity.

Analysis of the change in mtDNA content in a total of 19 sub-study subjects revealed a significant increase over 24 weeks [+600 (273.5) copies/cell, P=0.042 at region 1 and +1134.1 (528.4) copies/cell, P=0.046 at region 2, Figure 2] in keeping with the gene expression changes identified.

Adipose tissue protein

Overall, biopsies from 16 subjects contributed to protein analyses, although the numbers used in each protein analysis varied depending on sample abundance. No significant change was observed in PPARγ and SREBP1 protein (Figure 3a and b), although levels tended to increase. In keeping with gene expression and mtDNA results, improved mitochondrial function was evidenced by decreased levels of nuclear-encoded COX4 at week 24 (−56.3% (−65%, −29.6%), P=0.016, Figure 3c) and by an increasing trend in mitochondrially encoded COX2 at week 24 (−64.1% (−21.2%, +110.9%), P=0.203, Figure 3d), resulting in a significant increase in the COX2/COX4 ratio at week 24 (−288% (+42.2, +621.7), P=0.038), in contrast to the changes observed previously in nRTI-mediated SAT toxicity. Also observed were significantly reduced levels of the inflammatory marker B2M at week 2 (−68.7% (−60.2%, −22.6%), P=0.02) but not at week 24 (Figure 3e).

Adipocyte density

Biopsy sections from all 20 subjects contributed to adipocyte density analysis. Histological analysis revealed normal white adipose tissue structure at all timepoints. SAT adipocyte density decreased
significantly over 24 weeks (−32.3 (15.5) cells/mm², \(P = 0.047\)) (Figure 4a), implying larger adipocytes (Figure 4b), which is in keeping with the observed increases in body fat by DXA and CT.

**Discussion**

This is the first study (to our knowledge) to explore the in vivo effect of PI-only ART initiation on adipose tissue composition and function independently of other ART classes. Our findings suggest improved SAT function as reflected by increases in adipocyte size and molecular changes consistent with improvements in lipid metabolism, mitochondrial function and inflammation with PI-only ART initiation. Such changes indicate a return to health associated with the suppression of HIV replication and do not support the introduction of new toxicity with PI exposure. The use of focused transcriptional profiling, mtDNA, histological and protein...
analyses in parallel with the assessment of clinical metabolic parameters and body composition allowed a detailed evaluation of the metabolic effects of initiating PI-only ART. Using this approach, adipose tissue toxicity indicative of the development of lipodystrophy was not detected thus challenging the conventional view that PIs cause adipose tissue toxicity.

While the role of tNRTI in the development of SAT toxicity is well established,9,32 PIs have also historically been implicated, 7 with the primary mechanism proposed being the inhibition of PPARG, CEBPA and SREBF1 activity resulting in impaired adipogenesis.10,11 However, in the current study, initiation of PI-only ART did not lead to either the clinical changes suggestive of lipoatrophy or reductions in expression of these key regulators at either the mRNA or protein level in SAT (Figure 1a and Figure 3a and b). In fact, expression of PPARG and its downstream targets, FABP4 and LPL, increased (Figure 1a), suggesting an increase from diminished baseline levels in untreated HIV infection.4,34 Moreover, the observed increases in adipocyte size and decreases in the expression of the triglyceride lipase LIPE are indicative of increased adipocyte lipid uptake and storage, events that are consistent with increased PPARG activity.

Although our findings contrast with previous clinical studies,17,18 subjects in these studies had previous and concomitant exposure to tNRTI, thereby limiting the ability to determine the relative contribution of PI to the observed toxicities. Furthermore, our results are consistent with a recent study comparing two ART combinations (neither of which contained tNRTI) in which SAT adipogenic gene expression increased in subjects initiating PI-containing ART but not with non-PI ART initiation.15 These data, coupled with several clinical trials showing an absence of SAT loss with use of PI-containing ART without a tNRTI,24,36,37 reinforce our view that the use of PIs is not independently associated with SAT toxicity in vivo.

Exposure to certain PIs, including lopinavir/ritonavir, has also been associated with insulin resistance in vitro10,13 and in healthy volunteers.30,38 Despite the transcriptional changes observed suggesting insulin resistance in this study (Figure 1b), there was no effect on systemic insulin sensitivity (HOMA-IR), indicating that the transcriptional changes in SAT were insufficient to affect whole-body insulin sensitivity. Although use of the hyperinsulinaemic–euglycaemic clamp may have allowed more sensitive detection of changes in insulin sensitivity, the findings of previous studies using this method were consistent with ours.39,40

Cross-sectional studies have previously demonstrated increased expression of inflammation-related genes concurrent with reduced expression of PPARG and PPARG-target genes in the SAT of ART-naive HIV-infected subjects,6,34 presumably reflecting the pro-inflammatory effects of uncontrolled HIV infection. In this study, we found that initiation of PI-only ART was associated with the reverse—increased PPARG activity and decreased expression of inflammatory markers IKBKB (mRNA) and B2M (protein)—implying a recovery from HIV-induced effects
on SAT rather than PI-mediated toxicity. Furthermore, mitochondrial toxicity, another key feature of both untreated HIV and ART-mediated adipose tissue toxicity, was not observed in this study. In fact, initiation of PI-only ART led to improvements in markers of mitochondrial function in SAT; mtDNA and the COX2/COX4 ratio increased while mRNA expression of the stress-responsive regulators of mitochondrial biogenesis NRF1 and CRTC3 decreased. These outcomes contrast considerably with the mitochondrial dysfunction observed with tNRTI in previous studies and further support an overall improvement in SAT function with initiation of PI-only ART.

Taken together, our findings suggest that the initiation of PI-only ART, independent of the confounding effects of other ART classes, displays negligible SAT toxicity in clinical, transcriptional, protein and histological analyses. In the context of an ever-expanding number of PLWH and with the need to increase global access to ART, the safety and efficacy of novel and cheaper ART regimens including PI monotherapy are being increasingly explored. Within this clinical setting, our results are particularly relevant and reassuring considering the lack of significant PI-mediated SAT toxicity observed.

This study had limitations. The absence of a group initiating alternative ART regimens makes it difficult to differentiate between the effects of PIs on SAT function and the effect of suppressing HIV replication. In addition, the relatively advanced immunosuppression of substudy subjects at baseline may have contributed to an augmented ‘return to health’ phenomenon that could differ from cohorts beginning ART at higher CD4+ counts. However, our molecular and clinical data are consistent with the findings of recent randomized studies that also demonstrate generalized increases in adiposity in subjects of higher baseline CD4+ counts when initiating several types of tNRTI-sparing ART. With follow-up limited to 24 weeks, SAT dysfunction arising from longer periods of PI exposure may not have been detected. Despite this, PIs can induce adipocyte toxicity in vitro after as little as 5 days of exposure and in vivo transcriptional changes indicative of SAT toxicity have been demonstrated after even shorter periods of ART exposure. While it may be argued that these results may not apply to the PI class as a whole (only two types of PI were examined), initiation of other commonly used PIs, such as darunavir and atazanavir, has a similar effect on body composition. Finally, study subjects were exclusively Asian. Nevertheless, there is no reason to suggest that the effects of PI on SAT would be markedly different in other patient groups.

In summary, the initiation of PI-only ART in this study facilitated an improvement rather than a deterioration in SAT function, with the changes observed consistent with a return to health, including increased adiposity, reduced levels of inflammation and improved mitochondrial function in adipose tissue.

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Author contributions
P. P., D. A. C. and J. M. A. L. conceived and designed the main study, contributed to the substudy design and recruited patients to the study. J. C., P. R., P. D., M. G. and P. W. G. M. conceived and designed the substudy. R. T. M. conducted the gene expression experiments. E. R. F. collected the clinical data and performed the mtDNA experiments. E. C. and J. C. carried out the mtDNA experiments. J. C. and P. R. contributed to the substudy design and recruited patients to the study. J. M. A. L. conceived and designed the main study, contributed to the substudy design and recruited patients to the study. J. A. and J. C. performed the mtDNA experiments.

Supplementary data
Tables S1 to S3 are available as Supplementary data at JAC Online (http://jac.oxfordjournals.org/).

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