Independent associations of insulin resistance with high whole-body intermuscular and low leg subcutaneous adipose tissue distribution in obese HIV-infected women1–3

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ABSTRACT

Background: Obesity and insulin resistance are growing problems in HIV-positive (HIV+) women receiving highly active antiretroviral therapy (HAART).

Objective: The objective was to determine the contribution of adipose tissue (AT) enlargement and distribution to the presence of insulin resistance in obese HIV+ women.

Design: Whole-body intermuscular AT (IMAT), visceral AT (VAT), subcutaneous AT (SAT), and SAT distribution (leg versus upper body) were measured by whole-body magnetic resonance imaging. Insulin sensitivity (SI) was measured with an intravenous glucose tolerance test in obese HIV+ women recruited because of their desire to lose weight (n = 17) and in obese healthy controls (n = 32).

Results: The HIV+ women had relatively less whole-body SAT and more VAT and IMAT than did the controls (P < 0.05 for all). A significant interaction by HIV status was observed for the relation of total SAT with SI (r = 0.0001 for the regression’s slope interactions after adjustment for age, height, and weight). However, relations of IMAT, VAT, and SAT distribution (leg SAT as a percentage of total SAT; leg SAT%) with SI did not differ significantly between groups. For both groups combined, the best model predicting a low SI included significant contributions by both high IMAT and low leg SAT% independent of age, height, and weight, and no interaction between groups was observed (overall r² = 0.44, P = 0.0003).

Conclusion: In obese HIV+ women, high whole-body IMAT and low leg SAT% distribution are independently associated with insulin resistance.

KEY WORDS Subcutaneous adipose tissue, intermuscular adipose tissue, adipose tissue distribution, insulin resistance, HIV infection

INTRODUCTION

Insulin resistance is a central feature of the metabolic syndrome (1, 2). Generalized and regional lack of body fat in congenital lipodystrophies (3) and upper-body fat accumulation in obesity (4) have been shown to be associated with insulin resistance in HIV-positive (HIV+) individuals. A syndrome of fat redistribution consisting of peripheral fat loss (face, limbs, and buttocks) and central fat accumulation (abdomen, breast, and dorsocervical area) has been described in HIV+ men and women (5, 6). HIV+ men and women with fat redistribution were more insulin resistant than were those without such redistribution (7–19). Obesity has emerged as a growing problem in HIV+ individuals receiving highly active antiretroviral therapy (HAART) (20), particularly in women (21, 22). The influence of the enlargement and distribution of various adipose tissue (AT) compartments on the presence of insulin resistance has not been clearly established in obese HIV+ women. However, its influence could be important for assessing the response to interventions that alter body fat, which may be useful in HIV-negative (HIV−) but not in obese HIV+ women (5, 23–25).

Most studies reporting on the relation between fat distribution and insulin resistance have compared groups of HIV+ individuals with or without lipodystrophy as determined a priori by observational criteria (7, 8, 11–13, 17–19). Studies using continuous variables (9–11, 14–16) found correlations between lipodystrophy measured by dual-energy X-ray absorptiometry (DXA), ie, more trunk fat or less leg fat as a percentage of total body fat and insulin resistance in both HIV+ men and women (9, 11, 15–19). Whether these associations were similar in HIV+ and HIV− cohorts has not been noted (15). DXA did not separate fat in subcutaneous AT (SAT) from fat in the visceral AT (VAT) compartment or inside muscle and organs (26–28). Separation of AT depots at the waist level by computed tomography (CT) or magnetic resonance imaging (MRI) provided additional information (26, 27); however, the findings of both positive and negative associations of SAT and VAT areas at the waist level with insulin resistance (14, 16, 26, 27, 29, 30) may have been due to differences in sex or degree of overweight (29, 31). In previous studies in patients with HIV lipodystrophy, in which whole-body AT depots were measured by whole-body MRI, relations to corresponding insulin resistance indexes were not reported (32–35). In particular, a relation of insulin resistance to SAT distribution (ie, SAT of legs versus upper body) and to IMAT (ie, subfascial, subcutaneous, and intermuscular) has not been noted (15).

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intermuscular AT ([IMAT]) measured by whole-body MRI has not been reported previously in HIV + women (32–35). Higher IMAT was found to be independently associated with insulin resistance in healthy premenopausal women (28), and decreases in leg fat or thigh SAT were shown to be independently associated with unfavorable glucose and lipid concentrations in larger cohorts not characterized by HIV status (36, 37).

Therefore, the overall aim of this study was to determine the relations of total SAT, VAT, IMAT, and SAT distribution (legs versus upper body) measured by whole-body MRI in obese HIV + women with insulin resistance. Specifically, we asked whether IMAT and SAT distribution were independently associated with insulin resistance in obese HIV + women. We also determined whether such relations were similar to those found in healthy controls.

SUBJECTS AND METHODS

HIV + subjects

The HIV + women were recruited for a study of weight loss and exercise. The data presented here are the baseline data from a subset of the recruited subjects who were not diabetic and who had adequate venous access for testing (n = 17). The body mass index (BMI; in kg/m²) of the women was 30–38. Three of the women were non-Hispanic white, 3 were Hispanic, and 11 were African American. The HIV + women were obese (BMI > 30) and clinically stable. Most of the subjects were on an antiretroviral drug regimen for ≥4 wk before enrollment and had no plans to change the regimen during the study period. The women were allowed to take their HIV medications as prescribed. Of the 17 HIV + women whose data are presented here, 4 women were not receiving HAART, 13 were taking nucleoside reverse transcriptase inhibitors, 9 were taking protease inhibitors (PIs), and 9 were taking non-nucleoside reverse transcriptase inhibitors. Exclusion criteria were as follows: 1) any active opportunistic infection or malignancy, 2) pregnancy or breastfeeding, 3) uncontrolled hypertension, 4) history of MRI or any condition that would prevent exercise, 5) diabetes mellitus, and 6) a self-reported medical history of an eating disorder (eg, anorexia nervosa or bulimia nervosa), gallbladder disease, renal disease, active substance abuse, or methadone treatment. The women chosen for the present analysis were either pre- or perimenopausal and none of them were receiving estrogen replacement therapy. All subjects had an intravenous glucose tolerance test (IVGTT) performed within 10 d of the beginning of a regular menstrual period. The women were also weight stable for ≥6 wk before the measurements.

Healthy controls

The women in the control group 1 (n = 12) and in control group 2 (n = 20) were recruited for 2 other studies. Data from measurements that were identical to those performed for the HIV + women (except for the insulin sensitivity measurements in control group 2; see below) were used in the analyses. For both control groups, everyone with a BMI ≥ 30 (obese, as in the HIV + cohort) and for whom data were available was included in the study. In group 1, 5 of the women were non-Hispanic white, 1 was Hispanic, and 6 were African American. In group 2, 7 of the women were non-Hispanic white and 13 were African American. The women in both control groups were weight stable for 6 mo before the studies, were healthy, were premenopausal, were non-diabetic, had regular menstrual cycles, reported no medical problems, reported no symptoms, were taking no medications, and had normal results from blood count and chemistry panels. No HIV test was performed in these groups. All subjects signed an informed consent form; the protocol and consent form were approved by the St Luke’s–Roosevelt Hospital Institutional Review Board and Radiation Safety Committee.

Anthropometric measures

Body weight was measured to the nearest 0.1 kg (Avery Weigh-Tronix, Fairmont, MN) and height to the nearest 0.5 cm with a stadiometer (Holtain, Crosswell, United Kingdom).

Whole-body magnetic resonance imaging

AT compartments and skeletal muscle (SM) volumes were measured on a 1.5T MRI scanner (6X Horizon; General Electric, Milwaukee, WI) as described previously (38–40). The entire body was visualized on a scout coronal image (6X Horizon), and the axial level of L₄–L₅ was identified. The scans were acquired using contiguous axial slices of 10-mm thickness at 40-mm intervals below L₄–L₅ to the toes, and above this level to the fingertips (40–50 images for women of average height). Images were then analyzed on a PC platform as described (28, 32–35, 38–40). Briefly, the procedure for calculating AT volume is to first measure the relevant tissue area in each slice with the use of threshold methods and manual delineation to draw boundaries among different tissues. The volume between slices is extrapolated from the area measurements. The following volumes were calculated: VAT, total SAT, and IMAT. We defined IMAT as the AT visible between the muscle groups and beneath the muscle fascia (38). The gray level intensity (threshold value) of the AT in the SAT region was first determined and used as reference. This threshold value was reduced by 20% to identify IMAT threshold.

In addition to whole-body results, regional values for subcutaneous AT were determined for the arms, legs, and upper and lower trunk. The regional subcutaneous AT volumes used in this article delineated SAT volume in the legs (leg SAT, ie, all SAT inferior to the greater trochanter, including the greater trochanter area) and SAT volume in the upper body (upper SAT, ie, all SAT superior to the level of the trochanter, including the arms). The MRI scans were read at the New York Obesity Research Center Image Reading Center at St Luke’s Roosevelt Hospital Center. The CV on repeated readings of the same 2 scans by observers analyzing the images was 3.8% for TAT, 3.4% for SAT, 9.7% for VAT, 2.2% for SM, and 7.3% (estimate) for IMAT.

Insulin sensitivity index determined by intravenous glucose tolerance test

The Bergman minimal model was used to quantify S_i (MINMOD 2.0; copyright R N Bergman, 1986; 41). This measurement was made in all subjects during the follicular phase of the menstrual cycle. Glucose (0.3 g/kg, 50% dextrose injection; Abbott, North Chicago, IL) was administered intravenously at time 0 min. This was followed by an injection of tolbutamide (Orinase Diagnostic, Upjohn, Kalamazoo, MI) at time 20 min in control group 2. In the HIV + group and in control group 1, 0.03 units insulin/kg (Humulin R; Lilly Inc, Indianapolis, IN) was used due to the lack of availability of tolbutamide. Blood samples, collected through a catheter placed in the contralateral arm,
occurred at fasting and at 26 time points over the 3 h after glucose administration. Plasma glucose and insulin were measured in all samples, and the \( S_t \) was calculated from these values with the nonlinear mathematical model of glucose disappearance. Studies in the literature have shown that the \( S_t \) measured at 20 min in the same subject with the use of IVGTT with intravenous insulin is lower than the \( S_t \) measured with the use of IVGTT with intravenous tolbutamide; however, the difference between methods (14%) appears to be constant throughout the range of insulin sensitivity in nondiabetic subjects (42). Therefore, we used a 14% lower value than that determined by the computer program for the subjects who had an IVGTT with tolbutamide at 20 min (control group 2), and the data were pooled. Furthermore, we computed possible interactions with the IVGTT method using a categorical factor denoting the IVGTT method for all relevant analyses (see Data analysis below).

Data analysis

Data were expressed as means ± SDs for \( t \) test comparisons in the 3 experimental groups and were expressed as means ± SEMs for the analysis of covariance (ANCOVA) results. Log transformations were used for variables for which deviation from normality of distribution was found. Independent \( t \) tests were used to compare measurements in the HIV+ group with those of control groups 1 and 2. Because there were 3 groups being compared by \( t \) test, a Bonferroni adjustment was used. Regression lines between dependent variables and covariates were tested for interactions before ANCOVA was performed. A general linear model was used to test interactions between slopes of regression for continuous variables in the HIV+ group compared with the control groups. Although use of a mathematical correction to compute \( S_t \) in control group 2 does not ensure that values would be comparable between groups, the literature suggests that rank-order correlation comparisons are preserved independently of the IVGTT method (42). In addition, as described above, a categorical factor denoting the IVGTT method (HIV+, HIV− group 1, and HIV− group 2) was entered in the ANCOVA and general linear model analyses of \( S_t \) and possible interactions were computed and reported if present. Analyses were done by using STATISTICA 6.0 (Statsoft Inc, Tulsa, OK). \( P < 0.05 \) was considered to be statistically significant.

RESULTS

Subject characteristics and whole-body MRI and metabolic measurements are shown in Table 1. The HIV+ women were less heavy and had less SAT than did the women in control group 1 but had significantly more VAT than did the women in both control groups (Table 1). Fasting glucose and insulin and insulin sensitivity values in the HIV+ group did not differ significantly from those of either control group.

The relative AT distribution (relative accumulation of AT in SAT, VAT, and IMAT compartments) in HIV+ women compared with that in the women in the 2 control groups combined is shown in Table 2. In contrast with Table 1, the data in the control groups were combined because measurement methods and group variances were homogenous. Data for individual AT compartment sizes (SAT, VAT, and IMAT) were adjusted for degree of overweight and for total AT. SAT was lower and both VAT and IMAT were higher in the HIV+ group than in the controls after adjustment for age, height, weight, and total AT (\( P < 0.05 \)).

### TABLE 1

| Subject characteristics and whole-body magnetic resonance imaging (MRI) and metabolic measurements in HIV-positive (HIV+) and control women\(^1\) |
|----------------------------------|----------------|----------------|----------------|
| **HIV+ group** (\( n = 17 \)) | **Control group 1** (\( n = 12 \)) | **Control group 2** (\( n = 20 \)) |
| Age (y) | 39.5 ± 7.5 | 36.7 ± 6.3 | 37.2 ± 6.4 |
| Weight (kg) | 91.5 ± 10.1 | 106.8 ± 9.9 | 91.7 ± 10.6 |
| BMI (kg/m²) | 34.8 ± 3.1 | 38.6 ± 2.8 | 34.1 ± 3.4 |
| SM (L)\(^1\) | 23.2 ± 3.6 | 25.0 ± 3.7 | 24.4 ± 3.3 |
| SAT (L)\(^1\) | 41.2 ± 9.8 | 52.9 ± 8.7 | 40.1 ± 6.8 |
| Upper SAT (L)\(^1\) | 24.3 ± 4.7 | 32.1 ± 5.0 | 23.4 ± 4.9 |
| Leg SAT (L)\(^2\) | 16.9 ± 5.6 | 20.8 ± 4.8 | 16.7 ± 3.0 |
| VAT (L)\(^2\) | 3.7 ± 1.2 | 2.4 ± 0.9 | 2.7 ± 1.0 |
| IMAT (L)\(^2\) | 2.3 ± 0.7 | 2.2 ± 0.5 | 1.9 ± 0.7 |
| Fasting glucose (pmol/L) | 5.4 ± 0.6 | 5.3 ± 0.4 | 5.2 ± 0.4 |
| Fasting insulin (pmol/L) | 132 ± 107 | 84 ± 21 | 86 ± 32 |
| \( S_t (\mu U \cdot mL^{-1} \cdot 10^{-4} \cdot min^{-1})^6 \) | 1.6 ± 1.2 | 1.3 ± 0.9 | 2.2 ± 1.2 |

\(^1\) All values are \( x \) ± SD. SM, skeletal muscle; SAT, subcutaneous adipose tissue; VAT, visceral adipose tissue; IMAT, intermuscular adipose tissue; \( S_t \), insulin sensitivity index.

\(^2\) Significantly different from the HIV+ group, \( P < 0.0166 \).

\(^3\) Measured by whole-body MRI.

\(^4\) Includes SAT of the arms and trunk above the level of the greater trochanter.

\(^5\) Includes all SAT inferior to the level of the greater trochanter.

\(^6\) Calculated by the minimal model program (copyright RN Bergman, 1986; 41) from data obtained during intravenous glucose tolerance tests (insulin modified for the HIV+ group and control group 1 and tolbutamide modified for control group 2). Because a mathematical correction was used to calculate \( S_t \) in control group 2 (42) and because of a lack of homogeneity of variances for fasting insulin, \( t \) tests (Bonferroni adjusted) were used for comparisons between the 3 groups.

Differences persisted after additional adjustment for SM volume. Upper- versus lower-body SAT distribution was calculated as leg SAT as a percentage of total SAT (leg SAT%). In unadjusted analyses, leg SAT% was not significantly different between the HIV+ group and the control group. However, the relation between leg SAT% and body weight or total AT differed significantly between the HIV+ and control groups (\( P < 0.05 \) for both). In the controls, correlations were \( r = -0.2 (P = 0.27) \) for body

### TABLE 2

Adipose tissue distribution in HIV-positive (HIV+) and control women\(^1\)

<table>
<thead>
<tr>
<th><strong>SM (L)</strong> (^1)</th>
<th><strong>SAT (L)</strong> (^3)</th>
<th><strong>Leg SAT% (of total AT)</strong> (^2)</th>
<th><strong>Leg SAT% (of total AT)</strong> (^3)</th>
<th><strong>VAT (L)</strong> (^2)</th>
<th><strong>IMAT (L)</strong> (^3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(n = 17)</td>
<td>(n = 32)</td>
<td>(n = 17)</td>
<td>(n = 32)</td>
<td>(n = 17)</td>
<td>(n = 32)</td>
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<tr>
<td>23.9 ± 0.7</td>
<td>24.4 ± 0.5</td>
<td>0.566</td>
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<tr>
<td>42.8 ± 0.3</td>
<td>44.1 ± 0.2</td>
<td>0.003</td>
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<tr>
<td>35.0 ± 1.3</td>
<td>37.0 ± 1.0</td>
<td>0.21</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>40.1 ± 1.3</td>
<td>40.8 ± 1.0</td>
<td>0.65</td>
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<tr>
<td>3.5 ± 0.2</td>
<td>2.6 ± 0.2</td>
<td>0.007</td>
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<tr>
<td>2.4 ± 0.1</td>
<td>2.0 ± 0.1</td>
<td>0.043</td>
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</tr>
</tbody>
</table>

\(^1\) All values are \( x \) ± SEM. SM, skeletal muscle; SAT, subcutaneous adipose tissue; VAT, visceral adipose tissue; IMAT, intermuscular adipose tissue.

\(^2\) Values adjusted for age, height, and weight.

\(^3\) Measured by whole-body magnetic resonance imaging.

\(^4\) Values adjusted for age, height, weight, total AT, and SM.

\(^5\) Leg SAT as a percentage of total adipose tissue (SAT + VAT + IMAT) or as a percentage of total SAT. Values were unadjusted.
of slopes by HIV status.

sured by whole-body magnetic resonance imaging. VAT was measured by whole-body magnetic resonance imaging. P < 0.001 for interaction of slopes by HIV status.

and r = −0.24 (P = 0.18) for total AT. In the HIV+ group, correlations were r = 0.47 (P = 0.054) for body weight and r = 0.64 (P = 0.006) for total AT, ie, lower values for leg SAT% were seen at lower degrees of adiposity in the HIV+ group, whereas this was not the case for the controls. Similar results were obtained for upper SAT as a % of total SAT (data not shown).

The relations of AT compartment sizes and AT distribution with SI are shown in the figures. Specifically, the relations of SI—and adjusted for age, weight, and height—with total SAT are shown in Figure 1, with VAT are shown in Figure 2, with IMAT are shown in Figure 3, and with VAT:SAT are shown in Figure 5. Analyses were also performed for upper SAT as a percentage of total SAT and for the absolute values of leg SAT and upper SAT, but no additional information was obtained. We computed the residual values of SI in a general linear model after adjustment for age, height, and weight. Because SI was measured with 2 different methods (using insulin for the HIV+ women and the women in control group 1 and using tolbutamide for those in control group 2), a group factor (HIV+ group, control group 1, and control group 2) was initially entered in the analyses. Because no interactions between the control groups were observed, the results are shown for the control groups combined, with HIV status (positive or negative) entered as a factor in the model. We determined whether any of the AT measures significantly contributed to the additional variance in SI after these adjustments and whether significant interactions were observed in these relations between the HIV+ and healthy controls.

A significant interaction by HIV status was observed for the relations of total SAT with residual SI (after adjustment for age, height, and weight; Figure 1; P < 0.001 for slope interaction). For total SAT, a positive relation was observed in the HIV+ women (Figure 1; r = 0.64, P = 0.006), whereas the relation in

FIGURE 1. Relation between residual insulin sensitivity (SI) and total subcutaneous adipose tissue (SAT) in HIV-positive (HIV+; △) and control (●) women. SI values were log transformed for normality. Residual SI is the difference between the observed and the expected values of SI, calculated through multiple regression as a function of age, height, and weight. VAT was measured by whole-body magnetic resonance imaging. P < 0.001 for interaction of slopes by HIV status.

FIGURE 2. Relation between residual insulin sensitivity (SI) and visceral adipose tissue (VAT) in HIV-positive (HIV+; △) and control (●) women. SI values were log transformed for normality. Residual SI is the difference between the observed and the expected values of SI, calculated through multiple regression as a function of age, height, and weight. VAT was measured by whole-body magnetic resonance imaging. P = 0.54 for interaction of slopes by HIV status.

FIGURE 3. Relation between residual insulin sensitivity (SI) and intermuscular adipose tissue (IMAT) in HIV-positive (HIV+; △) and control (●) women. SI values were log transformed for normality. Residual SI is the difference between the observed and the expected values of SI, calculated through multiple regression as a function of age, height, and weight. IMAT was measured by whole-body magnetic resonance imaging. P = 0.25 for interaction of slopes by HIV status.

FIGURE 4. Relation between residual insulin sensitivity (SI) and subcutaneous adipose tissue (SAT) as a percentage of total SAT (leg SAT%) in HIV-positive (HIV+; △) and control (●) women. SI values were log transformed for normality. Residual SI is the difference between the observed and the expected values of SI, calculated through multiple regression as a function of age, height, and weight. Leg SAT (inferior to the greater trochanter) was measured by whole-body magnetic resonance imaging. P = 0.22 for interaction of slopes by HIV status.
FIGURE 5. Relation between residual insulin sensitivity ($S_I$) and the ratio of visceral to subcutaneous adipose tissue (VAT:SAT) in HIV-positive (HIV+; △) and control (○) women. $S_I$ values were log transformed for normality. Residual $S_I$ is the difference between the observed and the expected values of $S_I$, calculated through multiple regression as a function of age, height, and weight. VAT:SAT was measured by whole-body magnetic resonance imaging. $P = 0.08$ for interaction of slopes by HIV status.

the controls was negative albeit not significant (Figure 1; $r = -0.22$, $P = 0.2$). No significant interactions by HIV status were found for the relations of VAT, IMAT, leg SAT%, and VAT:SAT with residual $S_I$; $P = 0.54$ (Figure 2), $P = 0.25$ (Figure 3), $P = 0.22$ (Figure 4), and $P = 0.08$ (Figure 5) for slope interactions, respectively. For both groups combined, the best model predicting a low $S_I$ included significant contributions by both high IMAT and low leg SAT% (independent of age, weight, and height) and with no interaction between groups (overall $r^2 = 0.44$, $P = 0.0003$). Neither VAT nor VAT:SAT was independently associated with $S_I$ in any of the models.

**DISCUSSION**

Associations between insulin resistance and both fat reduction and fat accumulation (7–9, 11–19) have been reported in previous cross-sectional studies in HIV+ individuals with lipodystrophy. Determining which aspect of fat distribution best represents increased insulin resistance in this HIV+ population is more difficult to assess in obese individuals, and a separate evaluation by sex is needed (34). Therefore, in this study we reported the relation between AT distribution and insulin resistance in obese HIV+ women and in obese healthy controls. A unique aspect of our study was the measurement of whole-body IMAT and of SAT distribution, which allowed the separation of superficial SAT in the trunks and legs (upper-body SAT above the level of the greater trochanter and leg SAT below this level), independent of accumulation of IMAT, VAT, or other fat depots, such as intramyocellular lipid (28).

Consistent with previous reports, we found no decrease in the absolute amounts of AT but significant AT redistribution with relatively less SAT and more VAT and IMAT in the obese HIV+ women than in the controls (35). In previous studies that used DXA, higher absolute and relative amounts of trunk fat accumulation were found in HIV+ women with lipodystrophy (9–11, 32, 43), in contrast with lower absolute amounts of total body and lower relative amounts of limb fat in HIV+ men with lipodystrophy (10, 13, 14, 32, 44). By CT or MRI, the increase in trunk fat in HIV+ women was attributed to an increase in VAT (32–35). Although lower absolute amounts of SAT have not been reported in HIV+ women as they have been in men (13, 14, 33), a lower ratio of abdominal SAT to VAT area (13, 14, 32), which suggests fat redistribution, was found in both sexes. This is the first report to address IMAT measurements in obese HIV+ women. The limb fat characterized in previous studies that used DXA included both leg IMAT and intramyocellular lipid in addition to SAT (32–35). We found a relative increase in IMAT and a relative decrease in leg SAT in our obese HIV+ women, which suggests that IMAT may have biologic characteristics different from those of SAT.

We also found that the relation between whole-body IMAT and insulin resistance did not differ significantly between the obese HIV+ group and the control group and that a high IMAT was independently associated with insulin resistance in both groups. These findings confirm our previous reports in HIV− women, which suggest that IMAT plays an important role in influencing insulin sensitivity independent of HIV status (28).

Although reports regarding differences between VAT and SAT in HIV− individuals have been published (45), reports regarding the biologic characteristics of IMAT in HIV− or HIV+ are lacking (28, 38–39). The influence of IMAT on insulin action may be due to its proximity to the muscle cells, similar to the effects of intramyocellular lipids (27). However, relatively greater amounts of IMAT may also reflect other characteristics of whole-body AT, such as abnormalities in the largest storing depot, SAT. Further studies are clearly needed to characterize whole-body IMAT in both HIV+ and HIV− individuals.

We found that the association between absolute SAT accumulation and $S_I$ was significantly different between the HIV+ and control groups. In contrast with results in the control group and with previous reports in HIV− obese women (4), the relation of absolute SAT accumulation with $S_I$ was positive in the HIV+ obese women in the present study. These findings were replicated even if absolute values of regional SAT (upper body or leg) were used (not shown). However, SAT distribution, ie, a relatively low amount of leg SAT (expressed as a percentage of total SAT), was similarly related to a low $S_I$ (insulin resistance) in both the obese HIV+ and the control groups. These relations were independent of VAT or IMAT. Differential characteristics of SAT in the upper versus the lower body could explain this finding. Gluteal and leg AT display increased glucose transport capacity, decreased sensitivity to lipolytic agents, and increased sensitivity to antilipolytic agents (46, 47)—in short, a higher capacity to store fat than the abdominal SAT. Less leg SAT could therefore reflect qualitative changes in overall SAT (more insulin resistance with decreased capacity to store), which could underlie the relation between SAT distribution and insulin resistance in both upper-body obese HIV− and HIV+ lipodystrophy. Indeed, recent data from 2 large studies in individuals not characterized by HIV status show associations between lower relative amounts of leg fat or thigh SAT and unfavorable glucose and lipid concentrations, independent of higher abdominal fat (36, 37).

We found VAT accumulation was not independently associated with insulin resistance in the groups of women studied. The relation between VAT and $S_I$ was negative in the controls and positive in the HIV+ group but was not statistically different in the 2 groups; neither of the calculated slopes was statistically different from zero. Although HIV+ women with lipodystrophy were reported to have increased VAT and to be more insulin resistant than HIV+ women without lipodystrophy (9, 11, 32),
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except for earlier studies in lean men and women (16, 30), most studies have not shown an independent relation between increased VAT and insulin resistance in HIV+ women (15, 26, 31). Therefore, our results in obese women could not be generalized to HIV+ men with lipodystrophy or to leaner HIV+ women. Still, it is of note that none of the previous studies made measurements of IMAT or SAT distribution.

Because HAART may directly alter both insulin sensitivity (48, 49) and AT metabolism (35, 50), it is not surprising that we found differences between the HIV + and control groups in the relations of certain AT depots (ie, whole-body SAT) with insulin resistance; one could argue that such relations in HIV – lipodystrophy should not be compared with those in healthy obese controls. However, the similarities we found in the relations of IMAT and SAT distribution with insulin resistance in both obese HIV+ and obese controls suggest that a partly common pathophysiologic mechanism may underlie these relations for both groups. For example, the relation between low leg SAT% and insulin resistance in both the HIV lipodystrophy and control subjects may be a consequence of a relative decreased amount of protective SAT (leg SAT) in both groups. Loss of limb fat, specifically of SAT [whereas intramyocellular fat (27) and probably IMAT are spared], has been attributed to HAART in longitudinal studies (51, 52). The heterogeneity of HAART in our HIV+ subjects and the small sample size precludes us from determining any specific effect of an antiretroviral drug class on insulin resistance and fat redistribution. Longitudinal studies are needed to further clarify these relations.

Finally, we acknowledge that the population of HIV+ women that we studied was small and self selected for the desire to lose weight, making generalizations to the larger HIV+ population was difficult. Nevertheless, similarities of findings in these obese HIV+ women as in reasonably representative healthy controls decrease the risk of such findings being biased by the selection of the study’s HIV+ cohort. In addition, the attributes of our population highlight the importance of our results because the women we studied were mostly African American and Hispanic; these women tend to be overweight or obese and tend to remain obese when infected with HIV (21, 22). Our study needs to be extended to other HIV-infected individuals, specifically men and HIV+ women with a larger range of body weight.

In conclusion, aspects of AT distribution associated with insulin resistance in obese HIV+ women seeking weight loss are a high whole-body IMAT and a low leg SAT distribution. Clinical recognition of these aspects of AT distribution may be difficult but important for studies of interventions aimed at improving insulin resistance in obese HIV+ women.

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