Insulin resistance in transgender individuals correlates with android fat mass

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Abstract

Background: Transgender individuals receiving gender-affirming hormone therapy (GAHT) are at increased risk of adverse cardiovascular outcomes. This may be related to effects on body composition and insulin resistance.

Aims: To examine relationships between body fat distribution and insulin resistance in transgender individuals on established GAHT.

Methods: Comparisons of body composition (dual energy X-ray absorptiometry) and insulin resistance [Homeostasis Model of Insulin Resistance (HOMA2-IR)] were made between transgender individuals (43 trans men and 41 trans women) on established GAHT (>12 months) and age-matched cisgender controls (30 males and 48 females). Multiple linear regressions were used to examine the relationship between HOMA2-IR and fat mass with gender, adjusting for age and total duration of GAHT and Pearson correlation coefficients are reported.

Results: Compared with control cisgender women, trans men had mean difference of +7.8 kg (4.0, 11.5), p < 0.001 in lean mass and higher android:gynoid fat ratio [0.2 [0.1, 0.3], p < 0.001], but no difference in overall fat mass or insulin resistance. Compared with control cisgender men, trans women had median difference in lean mass of −6.9 kg [−10.6, −3.1], p < 0.001, fat mass of +9.8 kg [3.9, 14.5], p = 0.001, lower android:gynoid fat ratio −0.1 [−0.2, −0.0], p < 0.05, and higher insulin resistance 1.6 (1.3–1.9), p < 0.001). Higher HOMA2-IR correlated with higher android (r² = 0.712, p < 0.001) and gynoid (r² = 0.572, p < 0.001) fat mass in both trans men and trans women.

Conclusion: Android fat more strongly correlates with insulin resistance than gynoid fat in transgender individuals. Higher fat mass and insulin resistance in trans women may predispose to increased cardiovascular risk. Despite adverse fat distribution, insulin resistance was not higher in trans men.

Keywords: body composition, gender dysphoria, gender identity, insulin resistance, transgender persons, transsexualism

Introduction

Background

It is estimated that 0.6–1.2% of the population identify as transgender (or trans)¹ ² and the number of individuals presenting to medical services for assistance with gender transition is rapidly rising.³ ⁴ Transgender individuals experience incongruence between the sex assigned to them at birth and their deeply held sense of gender identity. Gender-affirming hormone therapy (GAHT) is used by many transgender individuals to align physical characteristics with their gender identity. Masculinising hormone therapy with testosterone for trans men and feminising hormone therapy with oestradiol and anti-androgen agents for trans women are both associated with improvements in psychological outcomes and quality of life.⁵
GAHT is usually continued lifelong; however, little is known about the long-term effects. Two large cohort studies suggest higher rates of cardiovascular events in transgender individuals on hormone therapy compared with cisgender individuals. Trans men on testosterone had higher rates of myocardial infarction compared with cisgender women, and trans women on oestrogen had an increased risk of ischaemic stroke and venous thromboembolism when compared with both cisgender men and cisgender women.6,7

Regional fat distribution, in particular central adiposity, is an important contributor to cardiovascular risk and is heavily influenced by sex steroids.8 A recent systematic review confirmed that feminising hormone therapy is consistently associated with increases in fat mass and decreases in lean mass, while trans men experience decreases in fat mass and increases in lean mass with masculinising hormone therapy.9 In cisgender populations central abdominal fat, also referred to as android fat, is associated with high cardiovascular risk and one of its measures, the waist:hip ratio, is more strongly correlated with cardiovascular outcomes than is body mass index (BMI).10 Trans men, but not trans women, have an increase in waist:hip ratio 12 months after commencing hormone therapy,9 suggesting higher cardiovascular risk. Additionally, the route of administration of hormone therapy may influence fat mass. In postmenopausal women, oral conjugated equine oestrogen is associated with higher body fat and loss of lean tissue when compared with transdermal oestradiol, thought to be mediated via insulin-like growth factor 1 (IGF-1) production.11–13

Insulin resistance is also an important contributor to cardiovascular risk. Both oestrogen and testosterone are capable of altering insulin sensitivity via a direct effect on liver, muscle and endothelial tissues, as well indirect effects via changes in body fat distribution.14–18 Although central to the pathogenesis of type 2 diabetes mellitus, insulin resistance also independently predicts a variety of poor outcomes in otherwise well non-diabetic individuals including hypertension, obesity and dyslipidaemia, as well as cardiovascular and all-cause mortality.19–22

Regional fat distribution and insulin resistance in cisgender individuals are well correlated. Adipose tissue, particularly central adiposity, has been shown to induce insulin resistance through release of multiple mediators including free fatty acids, steroid hormones and proinflammatory cytokines.18 Given the significant body composition changes known to occur in transgender individuals with gain in fat mass and loss of lean mass with feminising hormone therapy and the reverse seen with masculinising hormone therapy,9 determining whether a correlation exists between regional fat mass and insulin sensitivity is important, yet not previously described. Understanding this link will provide insights into whether GAHT affects insulin resistance predominantly through direct versus indirect (via changes in body composition) mechanisms, and guide clinicians in providing more accurate preventative strategies.

We aimed to further investigate the effect of GAHT on insulin resistance and body composition as surrogate markers of cardiovascular risk. We hypothesised that, first, trans men on testosterone therapy would have higher lean mass, lower fat mass and greater android:gynoid body fat distribution compared with control cisgender women and that opposite effects would be seen in trans women on oestradiol therapy compared with control cisgender men. Second, we hypothesised that insulin resistance would correlate with higher android fat and, therefore, would be higher in trans men compared with control cisgender women, with opposite effects seen in trans women.

Materials and methods

Study design and participants

We conducted a cross-sectional study between 1 April 2017 and 30 April 2018 in transgender individuals aged 18 years and over who had been on continuous GAHT for 12 months or more. Trans men on standard dose testosterone therapy were compared with individuals of the same sex assigned at birth; cisgender female controls. Trans women receiving standard doses of oestriadiol-based therapy for feminisation were compared with cisgender male controls. Transgender participants were recruited from endocrinology outpatient clinics and from primary care general practice clinics specialising in transgender health in Melbourne, Australia. These participants were compared with age-matched cisgender control groups. Healthy control individuals were additionally recruited as control participants for a longitudinal study in bone health in transgender individuals and exclusion criteria included...
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diabetes, established osteoporosis, metabolic bone disease, glucocorticoid therapy, bisphosphonate therapy, antiepileptic medication, HIV pre-exposure prophylaxis, pregnancy, thromboembolic disease, liver disease, or any disease likely to lead to impairment in bone health. All participants provided written informed consent and the protocol was approved by the Austin Health Human Research Ethics Committee (approval no. HREC/17/Austin/74).

Data collection
All participants underwent fasting blood testing to measure oestradiol, testosterone, sex hormone binding globulin (SHBG), blood glucose, insulin, C-peptide and IGFB-1 levels. Where possible blood testing was undertaken as a trough level for those on depot medications (such as testosterone undecanoate). In cisgender female participants blood testing was not able to be timed to a particular point in the menstrual cycle. Oestradiol was measured using immunoassay (Cobas E801, Roche Diagnostics, inter-assay variation 25% at level of 100 pmol/L or less and 25% at a level of greater than 100 pmol/L). Those on unmeasurable forms of oestradiol (such as ethinyloestradiol) were not included in the calculation of median oestradiol levels. Testosterone was measured using immunoassay (Cobas E801, Roche Diagnostics, inter-assay variation 14.8% at level of 2.7 nmol/L or less and 15% at a level of greater than 2.7 nmol/L). SHBG was measured on immunoassay (Cobas E801, Roche Diagnostics, inter-assay variation 6% at a level of 21 nmol/L and 6% at a level of 40 nmol/L). Fasting plasma glucose was measured using hexokinase photometric assay (Cobas C8000, Roche Diagnostics, inter-assay variation 1.5 at levels of 4.8 and 15.5 mmol/L). Electrochemiluminescence immunoassay (Cobas C8000, Roche Diagnostics) was used to measure insulin (interassay variation 4% at 16.3 mIU/L and 5% at 154 mIU/L) and C-peptide (interassay variation 4.5% at a level of 2.5 nmol/L and 6.8% at 0.55 nmol/L). Fasting blood glucose and C-peptide were used to calculate insulin resistance using updated Homeostasis Model of Insulin Resistance (HOMA2-IR), which is available for download from The Oxford Centre for Diabetes, Endocrinology and Metabolism. This is a non-linear model, which accounts for variations in hepatic and peripheral glucose resistance. C-peptide can be used to model both beta-cell function and insulin resistance and compared with insulin is less likely to degrade if any haemolysis of the sample occurs. IGFB-1 was measured using chemiluminescence immunoassay (Liaison XL, DiaSorin); interassay at a level of 11.4 nmol/L is 10% and at 42.2 nmol/L is 8.5%. Body composition was measured using dual energy X-ray absorptiometry (DXA) (Prodigy Version 7.5 GE Lunar, Madison, WI, USA). Coefficient of variation was <2%.

Statistical analysis
Characteristics and body composition parameters of participants were summarised as median and interquartile ranges for each group. Multiple linear regressions were used to examine the relationship between HOMA2-IR and fat mass with sex, adjusting for age and total duration of GAHT. HOMA2-IR, android fat mass, gynoid fat mass and total fat mass were log-transformed to approximate normality, and results were back-transformed to estimate the ratio of geometric means with corresponding 95% confidence intervals (CIs). The mean difference with corresponding 95% CI (denoted in round brackets) were reported for fat mass measures that were not log-transformed. Separate analyses were done comparing females versus transgender men, and males versus transgender women. Further analysis of correlation between HOMA2-IR with fat mass was also performed using linear regression, and the Pearson correlation coefficients and t-tests for regression coefficient slope were reported. All statistical analyses were performed using R (version 3.6.0, R Foundation for Statistical Computing). A p-value of less than 0.05 was considered statistically significant. No adjustment for multiple comparisons was performed as the analysis is of an exploratory nature.

Results

Participant characteristics
This study recruited a total of 162 participants: 84 transgender individuals (41 trans women and 43 trans men) and 78 controls (30 cisgender females and 48 cisgender males). Participant characteristics are summarised in Table 1. Mean age in the cisgender male controls was younger than in trans women and as such analyses were adjusted for age.

All 43 trans men were receiving testosterone [intramuscular (IM) testosterone undecanoate
### Table 1. Results (participant characteristics and effect sizes).

| Parameter | Trans men \( n=43 \) | Control cisgender women \( n=48 \) | Effect (95% CI) |
|-----------|------------------------|-----------------------------------|----------------|
| **Age, years** | 28.8 [25.0–33.0] | 28.1 [24.0–38.7] | – |
| **BMI** | 25.2 [23.1–28.6] | 22.7 [20.9–26.1] | – |
| **Total duration of GAHT, months** | 44.0 [22.6–67.0] | – | – |
| **Oestradiol, pmol/L** | 115.0 [93.0–164.0] | 177.0 [32.5–359.2] | 1.12 (0.57, 2.22) |
| **Testosterone, nmol/L** | 15.6 [13.2–19.7] | 0.9 [0.4–1.2] | **22.62 [15.73, 32.53]** |
| **SHBG** | 31.5 [21.0–41.0] | 98.5 [73.8–132.0] | **0.30 [0.21, 0.44]** |
| **IGF-1** | 29.0 [22.0–33.5] | 36.2 [28.1–40.2] | ↓4.40 (–10.52, 1.73) |
| **HOMA2-IR** | 1.2 [1.0–1.6] | 1.1 [0.9–1.4] | 0.99 (0.75, 1.30) |
| **Total fat mass, kg** | 18.4 [14.3–28.0] | 20.1 [14.6–25.4] | ↑5.0 kg (–1.7, 11.8) |
| **Android fat mass** | 2.0 [1.3–2.7] | 1.4 [1.0–2.0] | 1.4 [1.0, 2.1] |
| **Gynoid fat mass** | 3.8 [2.9–5.4] | 4.7 [3.5–5.5] | 1.0 [0.8, 1.3] |
| **Android:gynoid fat ratio** | 1.0 [0.9–1.1] | 0.8 [0.7–0.9] | ↑0.2 (0.1, 0.3) |
| **Total lean mass, kg** | 48.1 [44.9–51.7] | 40.7 [37.0–43.7] | ↑7.8 kg (4.0, 11.5) |

| Parameter | Trans women \( n=41 \) | Control cisgender men \( n=30 \) | Effect (95% CI) |
|-----------|------------------------|-----------------------------------|----------------|
| **Age, years** | 41.1 [26.4–52.7] | 32.0 [26.3–40.9] | – |
| **BMI** | 23.6 [21.7–29.2] | 23.8 [23.1–25.8] | – |
| **Total duration of GAHT, months** | 39.0 [19.9–60.0] | – | – |
| **Oestradiol, pmol/L** | 327.0 [147.2–460.5] | 72.5 [49.5–93.8] | 5.12 [3.44, 7.61] |
| **Testosterone, nmol/L** | 0.6 [0.4–0.9] | 20.5 [16.0–24.1] | 0.04 [0.03, 0.06] |
| **SHBG** | 86.0 [59.5–116.8] | 51.5 [39.0–75.2] | 1.37 [1.06, 1.76] |
| **IGF-1** | 22.5 [16.5–27.5] | 28.3 [23.0–35.2] | ↓1.86 (–6.57, 2.86) |
| **HOMA2-IR** | 1.5 [1.3–2.2] | 1.1 [0.8–1.3] | 1.6 [1.3, 1.9] |
| **Total fat mass, kg** | 22.5 [17.3–34.2] | 15.7 [11.6–20.5] | ↑9.8 kg [3.9, 14.5] |
| **Android fat mass** | 2.1 [1.3–3.6] | 1.5 [1.1–2.0] | 1.40 [1.05, 1.87] |
| **Gynoid fat mass** | 4.5 [3.9–6.4] | 3.1 [2.5–4.1] | 1.53 [1.26, 1.85] |
| **Android:gynoid fat ratio** | 1.0 [0.8–1.0] | 1.0 [0.9–1.2] | ↓0.1 (–0.2, 0.0) |
| **Total lean mass, kg** | 51.5 [47.0–55.7] | 58.3 [54.2–64.0] | ↓6.9 kg [–10.6, –3.1] |

Results are presented as median (interquartile range).
Effect adjusted for age and total duration of GAHT is presented as a ratio of geometric means, or mean difference (where arrows are shown).
*\( p < 0.05 \).
**\( p < 0.001 \).
+Those on unmeasurable forms of oestradiol (such as ethinyloestradiol) were not included in the calculation of median oestradiol levels.
BMI, body mass index; CI, confidence interval; GAHT, gender-affirming hormone therapy; HOMA2-IR, Homeostasis Model of Insulin Resistance; IGF-1, insulin-like growth factor 1; SHBG, sex hormone binding globulin.
n = 30, IM testosterone enanthate n = 8, topical testosterone gel 1% n = 5] and all 41 trans women were receiving oestradiol (oral oestradiol valerate n = 34, oral ethinylestradiol n = 3, transdermal oestradiol n = 4). Seventy-eight per cent (n = 32) of the trans women were taking anti-androgen therapy in addition to oestradiol therapy (cyproterone acetate n = 21, spironolactone n = 5, progestogens n = 5) (levonorgestrel n = 3, medroxyprogesterone n = 1, micronised progesterone n = 1), gonadotropin releasing hormone analogue n = 1. Twenty-seven per cent (n = 11) of the trans women had undergone orchidectomy and 5% (n = 2) of the trans men had undergone oophorectomy. In both trans men and trans women, the median oestradiol and testosterone levels were within the target reference range for their affirmed gender. In trans men, median oestradiol was 115.0 (93.0, 164.0) pmol/L (laboratory male reference range for oestradiol was <160 pmol/L) and median testosterone was 15.6 nmol/L (13.2, 19.7) (laboratory male reference range for testosterone was 9.9–27.8 nmol/L). In trans women, median oestradiol concentration was 327.0 (147.2, 460.5) pmol/L (laboratory female reference range for oestradiol during follicular phase was 46–607 pmol/L) and mean testosterone concentration was 0.6 (0.4, 0.9) nmol/L (laboratory female reference range for testosterone was <1.8 nmol/L).

**Masculinising hormone therapy**

Trans men had significantly higher lean mass than cisgender women with mean difference +7.8 kg 95% CI (4.0, 11.5), p < 0.001) (Table 1). Other absolute body composition parameters (total fat mass, android and gynoid fat mass) were not significantly different from cisgender female controls; however, android:gynoid fat mass ratio was higher [mean difference +0.2 (0.1, 0.3), p < 0.001]. Total fat mass was lower and android: gynoid fat mass was higher in trans men compared with cisgender female controls.

There was no difference in HOMA2-IR in trans men compared with cisgender female controls. Insulin resistance as estimated by HOMA2-IR was significantly correlated with android fat mass ($r^2 = 0.712, \ p < 0.001$) and gynoid fat mass ($r^2 = 0.572, \ p < 0.001$); see Figures 1 and 2. HOMA2-IR was also weakly correlated with android lean mass ($r^2 = 0.449, \ p < 0.001$) and gynoid lean mass ($r^2 = 0.220, \ p = 0.01$) (data not shown).

Whilst not the primary aim of our analyses, when comparing trans men with cisgender male controls, there was also no difference in HOMA2-IR (Supplemental Material Appendix 1 online). Although trans men were younger [trans men median 28.8 years (25.0–33.0)] compared with cisgender...
male controls [32.0 years (26.3–40.9)], they had a higher BMI and lower lean mass. Testosterone levels between groups were similar, but trans men had higher median oestradiol than cisgender male controls [115.0 pmol/L (93.0–164.0) versus 72.5 (49.5–93.8), \( p < 0.01 \)] (Supplemental Appendix 1).

**Feminising hormone therapy**

Trans women had significantly lower lean mass [mean difference –6.8 kg (–10.6, –3.1), \( p < 0.001 \)] and all fat mass parameters were significantly higher than cisgender male controls (Table 1). Android fat mass and gynoid fat mass were 40% and 53% higher respectively in trans women compared with cisgender male controls. There was a significantly lower android:gynoid fat ratio [mean difference –0.1 (–0.2, –0.0), \( p < 0.05 \)].

Despite a lower android:gynoid fat mass, the total android fat mass was still high amongst trans women [median 2.1 kg (1.3–3.6)]. HOMA2-IR in trans women was 1.5; significantly higher than cisgender male controls (\( p < 0.001 \)).

As with trans males, insulin resistance as estimated by HOMA2-IR was significantly correlated with android fat mass and, to a lesser degree, gynoid fat mass. See Figures 1 and 2.

As an exploratory analysis, when trans women were compared with cisgender female controls, HOMA2-IR was significantly higher in trans women [mean difference 1.47 (1.22, 1.77), \( p < 0.01 \)] as was android fat mass [mean difference 1.39 kg (1.06, 1.83), \( p < 0.001 \)]. Trans women were, however, significantly older [trans women median 41.1 years (26.4, 52.7)] compared with cisgender female controls [28.1 years (24.0, 38.3)], had a significantly higher median oestradiol [327.0 pmol/L (147.2–460.5) versus 177.0 pmol/L (32.5–359.2), \( p < 0.01 \)], but median testosterone concentrations and BMI were not different (Supplemental Appendix 1).

Trans women who had undergone orchidectomy \( (n = 11) \) had a significantly lower HOMA2-IR than trans women who had not \( (n = 30) \) [1.3 (1.1–1.5) versus 1.8 (1.4–2.3) \( p < 0.03 \)]. This is despite trans women who had undergone orchidectomy being older in age [57.0 years (40.5, 68.4) compared with 33.5 (25.5, 48.9)] \( p < 0.03 \), having longer duration of GAHT [115.1 months (41.7, 180.9) versus 27.0 months (15.3, 47.1)] \( p < 0.002 \), yet similar body composition (no significant difference between BMI, fat mass, or lean mass). See Supplemental Appendix 2.
Discussion

This cross-sectional study showed a correlation between insulin resistance and fat mass in transgender individuals on established GAHT. Trans men had significantly higher lean mass as well as a higher android:gonad fat ratio with no significant differences in insulin resistance or overall fat mass compared with control cisgender women. Trans women were more insulin resistant than control cisgender men and had lower lean mass, higher fat mass and a lower android:gonad fat ratio. Insulin resistance correlated with android fat mass but, contrary to our original hypothesis, trans men did not have higher insulin resistance, most likely because higher lean mass may be protective in trans men.

Masculinising hormone therapy

Our findings of higher lean mass (median 7.8 kg) and a higher android:gonad fat ratio are consistent with previous studies investigating masculinising hormone therapy in trans men. Testosterone is known to increase the synthesis of muscle tissue by promoting differentiation of cells of the myogenic lineage and to inhibit the differentiation of adipocyte precursor cells. Moreover, testosterone also inhibits lipoprotein lipase activity in adipocytes, an enzyme that increases fat deposition by decreasing adipose tissue lipolysis.

We found no significant difference in insulin resistance between trans men and cisgender female controls, in keeping with all but one prior study in transgender men that showed either no change or a decrease in insulin resistance. All these studies were prospective longitudinal in design but only one had a control group. The lack of change in insulin resistance is consistent with data that found no change in incretin (glucagon-like peptide-1 and gastric inhibitory polypeptide) responses in trans men before and after 12 months of GAHT.

The importance of body composition – which takes many months to change with masculinising hormone therapy – is highlighted by a small study demonstrating an increase in insulin resistance measured by hyperinsulinaemic euglycaemic clamps in 13 transgender men over the first 4 months, but with follow-up over 12 months, no significant differences in insulin resistance over time emerged.

It is important to note that the roles of sex steroids in insulin sensitivity in cisgender populations are not fully understood. Men with hypogonadism have increased insulin resistance; however, exogenous testosterone replacement is associated only with a small, likely clinically insignificant, improvement in insulin sensitivity. Elevated testosterone levels in women, such as in polycystic ovary syndrome, are associated with increased, rather than decreased, insulin resistance, suggesting that the primary driver of insulin sensitivity may be due to the indirect, rather than direct, effects of testosterone.

The association observed between android fat mass and insulin resistance in trans men also supports the importance of body composition. Whilst there is also a significant correlation with gynoid fat mass and insulin resistance, it is stronger for android. This is in keeping with the predisposition to insulin resistance associated with abdominal adiposity in cisgender populations.

Feminising hormone therapy

Trans women in our study had a significantly higher fat mass (median 9.8 kg) and lower lean mass (median 6.9 kg) compared with cisgender male controls. These results are in line with previous studies that have evaluated body composition using DXA in trans women. Only four studies have also previously looked specifically at android and gynoid fat mass regions, either using DXA or magnetic resonance imaging, and, like this study, found an increase in fat mass in both regions. These findings support the theory that activation of oestrogen receptors can lead to stimulation of adipocyte proliferation as well as lipoprotein lipase activity. Oestrogen may also act indirectly via oestrogen receptors in the hypothalamus to regulate energy expenditure.

We found that trans women have significantly higher levels of insulin resistance estimated by HOMA2-IR compared with cisgender male controls. Nine studies have previously looked at insulin resistance in trans women on feminising hormone therapy and, of these, six similarly showed worsening insulin resistance. Three did not detect a significant change – one showed a trend towards increase insulin resistance but failed to reach statistical significance, another had a sample size of only six participants and the remaining study did not measure body composition changes so it is unclear whether changes to this occurred. All but one study was prospective longitudinal in design but none had a control group.
Only four studies have specifically looked at android and gynoid fat mass regions, and of these only one sought to correlate insulin resistance and regional fat mass. This 2003 case–control study found a small increase in visceral fat area in both trans males and trans females; however, it failed to find a correlation with insulin sensitivity or fasting insulin levels in either group, likely due to small sample size and lack of control group.34

It is important to note that our findings, and others’, contradict existing theories and animal models suggesting that oestradiol has direct beneficial effects on insulin sensitivity.14,16,56

In studies of cisgender women, oestrogen has generally been associated with a favourable effect on insulin sensitivity.57 Low oestrogen states such as menopause are associated with a decrease in insulin sensitivity, increased central adiposity and a higher risk of metabolic disease, and subsequent administration of oestrogen therapy in menopausal women is associated with an improvement in insulin sensitivity.58

One proposed explanation for the differences seen in trans women is the high amount of overall fat as well as retention of central fat, which may indirectly lead to increased insulin resistance. This may mitigate any potentially beneficial direct effect of oestrogen on the insulin receptor.16

The route of oestrogen administration may also be important, with the majority of trans women in this study taking oral oestradiol valerate. Oral, but not transdermal, oestrogen has been shown to impair the metabolic effect of growth hormone in the liver, resulting in lower IGF-1 production and fat oxidation with a subsequent gain of body fat and loss of lean tissue seen in postmenopausal women.11–13 Our study showed that trans women had a lower IGF-1 than both cisgender male and cisgender female controls. This is in contrast to the only other study investigating IGF-1 and body composition in trans women, which found serum IGF-1 levels at 24 months were similar to baseline, and that any changes were independent of the route of administration of oestrogen.59 All participants aged under 45 years (n = 34) were taking oral oestradiol and those aged 45 years and over (n = 15) were taking transdermal oestradiol, so participant age may have been a factor. Interestingly, another study showed that there was no difference in total regional fat mass between trans women on oral or transdermal oestrogen.45 Oestrogen doses in GAHT are generally higher than for menopausal hormone therapy, so further prospective studies in the transgender population are warranted.

Our findings of lower HOMA2-IR in the subset of trans women who had undergone orchidectomy are in keeping with a small prospective study from 2016,60 which hypothesised that orchidectomy in this context may be protective due to the ratio of circulating sex hormone levels; however, further research is needed to confirm this.

Limitations

Limitations of the study include its cross-sectional design. Characteristics were not assessed prior to GAHT and baseline differences may have existed. In fact, two previous studies suggest that trans women have lower muscle mass and higher fat mass than cisgender male controls at baseline and reported doing significantly less physical activity.47,50 The trans women were slightly older than the cisgender male controls. For simplicity the data were presented uniformly using the median and interquartile range; however, the mean age between the two groups was more closely matched – trans women were aged 40.8 ± 15.7 years versus cisgender male controls aged 36.0 ± 14.2 years. Data were adjusted for age. Trans men had a higher median BMI than cisgender female controls; however, if anything, this should lead to an overestimation of insulin resistance in the trans male group. Whilst our participants had undertaken GAHT for several years (median 44 months in trans men and 39 months in trans women) it is possible that changes to body composition are still ongoing. Participants were not on standardised GAHT regimens and we cannot discount that different hormone formulations, particularly oral versus transdermal oestradiol, may have differential effects on body composition and insulin resistance. Many participants were on a progestogen and this may affect the outcomes measured in this study. Testosterone and oestradiol assays used measured via immunoassay rather than liquid chromatography mass spectrometry. Although this study focuses on body composition and insulin resistance there are other contributors to cardiovascular risk. Additional research is needed and a prospective longitudinal study with a cisgender control group is needed to further investigate the impact of GAHT on body composition and insulin resistance.
Conclusion
We highlight the importance of lean and fat mass and the correlation with insulin resistance among transgender individuals and the relatively stronger correlation of insulin resistance with android over gynoid fat. Significantly higher levels of fat mass and lower lean mass in trans women is associated with insulin resistance, and whilst there is some degree of higher fat mass in trans men on established GAHT, the significantly higher lean mass relative to fat mass appears to be protective. These findings provide insights into sex hormone action and suggest a predominantly indirect mechanism of action (via changes in body composition) in mediating insulin resistance. Longitudinal studies are needed to further investigate this correlation and to better guide clinical practice. Until then, a proactive clinical approach to mitigate gain in fat as well maintain or increase lean mass, particularly in trans women, should be strongly encouraged.

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Author contributions
IB was involved in the conception, design, analysis, interpretation of data and writing the manuscript. CS and SYL were involved in data analysis and revising the manuscript. GP, MG, JDZ were involved in revising the manuscript. ASC was involved in the conception, design, funding acquisition, analysis, interpretation of data, as well as revising the manuscript.

Conflict of interest statement
AC has received speaker’s honoraria from Astra Zeneca and Merck Sharp & Dohme. MG has received research funding from Bayer, Weight Watchers, Lilly Otzuka, and speaker’s honoraria from Besins Health Care and Novartis. All other authors have no conflicts of interest to declare.

Data availability
The datasets generated during and/or analysed during the current study are not publicly available but are available from the corresponding author on reasonable request.

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Supplemental material
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