

Advanced body composition assessment: from body mass index to body composition profiling

Magnus Borga,^{1,2,3} Janne West,^{2,3,4} Jimmy D Bell,⁵ Nicholas C Harvey,^{6,7} Thobias Romu,^{1,2,3} Steven B Heymsfield,⁸ Olof Dahlqvist Leinhard^{2,3,4}

¹Department of Biomedical Engineering, Linköping University, Linköping, Sweden

²Center for Medical Image Science and Visualization (CMIV), Linköping University, Linköping, Sweden

³Advanced MR Analytics AB, Linköping, Sweden

⁴Department of Medical and Health Sciences, Linköping University, Linköping, Sweden

⁵Research Centre for Optimal Health, University of Westminster, London, UK

⁶MRC Lifecourse Epidemiology Unit, University of Southampton, Southampton, UK

⁷NIHR Southampton Biomedical Research Centre, University of Southampton, University Hospital Southampton NHS Foundation Trust, Southampton, UK

⁸Pennington Biomedical Research Center, Baton Rouge, Louisiana, USA

Correspondence to

Professor Magnus Borga, Department of Biomedical Engineering, Linköping University, Linköping 581 83, Sweden; magnus.borga@liu.se

Accepted 8 March 2018

ABSTRACT

This paper gives a brief overview of common non-invasive techniques for body composition analysis and a more in-depth review of a body composition assessment method based on fat-referenced quantitative MRI. Earlier published studies of this method are summarized, and a previously unpublished validation study, based on 4753 subjects from the UK Biobank imaging cohort, comparing the quantitative MRI method with dual-energy X-ray absorptiometry (DXA) is presented. For whole-body measurements of adipose tissue (AT) or fat and lean tissue (LT), DXA and quantitative MRIs show excellent agreement with linear correlation of 0.99 and 0.97, and coefficient of variation (CV) of 4.5 and 4.6 per cent for fat (computed from AT) and LT, respectively, but the agreement was found significantly lower for visceral adipose tissue, with a CV of >20 per cent. The additional ability of MRI to also measure muscle volumes, muscle AT infiltration and ectopic fat, in combination with rapid scanning protocols and efficient image analysis tools, makes quantitative MRI a powerful tool for advanced body composition assessment.

INTRODUCTION

The human body—as well as the body of every other animal—is mainly composed of four molecular-level components: water, fat, proteins and minerals, usually in that order of decreasing amounts.¹ The substance that has attracted the most attention, from laypeople to medical professionals, is fat. This is, of course, motivated by the well-established fact that an excessive amount of body fat is related to increased morbidity and mortality. But also because adipose tissue (AT) is, by far, the most varying compartment—between individuals, but also within an individual over time. The most widely used way to estimate body fat is the body mass index (BMI)—body weight normalized by height squared (kg/m^2). Being a very simple and inexpensive method, it is the basis for WHO's definition of overweight ($25 \leq \text{BMI} < 30$) and obesity ($\text{BMI} \geq 30$). However, for a given BMI, the body fat percentage changes with age, and the rate of this change is different depending on sex, ethnicity and individual differences.² And while BMI correlates with fat accumulation and metabolic health in large

populations, it is insensitive to the actual distribution of body fat.³

When comparing methods for body composition analysis, it is important to distinguish fat (triglyceride) from AT,⁴ which contains approximately 80 per cent fat, the rest being water, protein and minerals.⁵ While most of the body fat is stored in AT, fat is also present in organs such as liver and skeletal muscle. Today, it is well known that the metabolic risk related to fat accumulation is strongly dependent on its distribution. Central obesity and, in particular, ectopic fat accumulation are important metabolic risk factors.^{6–8} Large amounts of visceral AT (VAT) are related to increased cardiac risk,^{8–9} type 2 diabetes,^{10–11} liver disease¹² and cancer.^{13–14} High levels of liver fat increase the risk for liver disease and type 2 diabetes,¹⁵ and increased muscle fat has been associated with increased risk for insulin resistance and type 2 diabetes¹⁶ and reduced mobility.¹⁷ While there are other anthropometric measures, such as waist circumference and waist-to-hip ratio, which more strongly correlate with metabolic risk,^{18–19} it is now well recognized that BMI and other anthropometric surrogate measures are poor predictors for individual fat distribution and metabolic risk.^{3–20–21}

Besides fat, acting as the body's long-term energy storage, skeletal muscles are of great interest to study, and the balance between the energy-consuming muscles and the energy-storing fat compartments is, of course, highly relevant in order to understand the metabolic balance of the body. Cachexia, involuntary loss of body weight, usually with disproportionate muscle wasting, is a life-threatening condition, often related to the progression of an underlying serious disease (eg, cancer²²). In cancer, cachexia is defined as weight loss of >5 per cent over 6 months, BMI <20 kg/m^2 or appendicular muscle mass normalized by body height squared of <7.26 kg/m^2 or 5.45 kg/m^2 for males and females, respectively.²³ Sarcopenia, which can be related to cachexia, but is also associated with aging, is often defined as reduced physical performance following loss of muscle mass, usually accompanied by increased fat infiltration of the muscles.²⁴ When diagnosing sarcopenia, muscle strength tests combined with muscle volume measurements are needed.²⁵



To cite: Borga M, West J, Bell JD, et al. *J Investig Med* Epub ahead of print: [please include Day Month Year]. doi:10.1136/jim-2018-000722

Furthermore, Willis *et al* showed that muscle pathology progression over 1 year could be detected by quantitative MRI but not by assessing muscle strength or function.²⁶ These examples illustrate the need for more sophisticated body composition analysis tools that go beyond simple anthropometric measures.

Since the early part of the last century, scientists have tried to determine the body composition in different ways, with a wide range of different physical principles and devices, and using different models and assumptions. Today, local *in vivo* measurements of different fat depots and fat infiltration in organs can be made using tomographic imaging techniques such as CT and MRI that were not even invented when the first scientific studies on body composition were published. These techniques are now recognized as golden standard for body composition analysis.^{25 27}

The purpose of this paper is to give a brief introduction to the most commonly used methods for body composition analysis and a review of an MRI-based body composition analysis technique, comparing its performance to other methods. This includes a previously unpublished validation study of the agreement between this method and dual-energy X-ray absorptiometry (DXA).

TECHNOLOGY OVERVIEW

A number of different techniques for body composition assessment have been developed, from very simple indirect measures such as waist-to-hip ratio and calipers to sophisticated direct volumetric measurements based on three-dimensional imaging techniques. There are also a range of invasive or *in vitro* methods for body composition analysis such as inhalation or injection of water-accumulating or fat-accumulating agents, or dissection and chemical analysis of cadavers. This overview will, however, focus solely on non-invasive *in vivo* measurement techniques.

Hydrostatic weighing (densitometry)

Hydrostatic weighing (underwater weighing), or densitometry, is based on Archimedes' principle. The difference of the body weight in air and water is used to compute the body's density. Assuming a two-component model with different densities for fat mass and fat-free mass and correcting for the air volume in the lungs, the total body fat percentage can be estimated. Obviously, this technique cannot give any measurements of the distribution of AT or lean tissue (LT).

Air displacement plethysmography (ADP)

ADP is perhaps better known under its commercial brand name BOD POD (Life Measurement, Concord, California, USA). Similar to hydrostatic weighing, ADP measures the overall body density and hence total body fat and LT but not their distributions. By putting the body in an enclosed chamber and changing the chamber's volume, the volume of the displaced air (ie, the volume of the body) can be determined from the changes in air pressure. Since ADP is based on the same two-component model as hydrostatic weighing, it is also affected by the same confounders, mainly variations in bone mineral content (BMC) and hydration. Due to the limitations of the two-component model used in densitometry and ADP, a four-component (4C) model is often recommended.^{28 29} In addition to fat and LT, the 4C

model also takes BMC and total body water (TBW) into account. However, these two additional components have to be measured by other techniques (eg, DXA for the BMC and deuterium oxide dilution for TBW³⁰). The repeatability (coefficient of variation (CV)) of ADP for body fat has been reported to be between 1.7 and 4.5 per cent when measured within 1 day.³¹ Obviously, ADP, as well as hydrostatic weighing, is limited to gross body composition analysis, not making any estimates of regional fat or muscles.

Bioelectrical impedance analysis (BIA)

BIA uses the electrical properties of the body to estimate the TBW and from that the body fat mass.^{32 33} The body is modeled as five cylindrical LT compartments; the trunk and the four limbs, while fat is considered to be an insulator. The impedance is assumed to be proportional to the height and inversely proportional to the cross-sectional area of each compartment, and the electrical equivalent is a resistor (extracellular water) in parallel with a capacitor and a resistor in series (intracellular water). The model of uniform distribution of fat and water fits better to the extremities than the trunk,³⁴ and while there are BIA measurements that correlate well with total abdominal AT, BIA cannot be used for measuring VAT.³⁵ Potential error sources are variations in limb length (usually estimated from body height), recent physical activity, nutrition status, tissue temperature and hydration, blood chemistry, ovulation and electrode placement.³² BIA requires different model parameters to be used depending on age, gender, level of physical activity, amount of body fat and ethnicity in order to be reliable.^{36 37}

Dual-energy X-ray absorptiometry

DXA is a two-dimensional imaging technique that uses X-rays with two different energies. The attenuation of an X-ray is dependent on the thickness of the tissue and the tissue's attenuation coefficient, which is dependent on the X-ray energy. By using two different energy levels, the images can be separated into two components (eg, bone and soft tissue). DXA is mainly used for bone mineral density measurements, where it is considered as the gold standard,³⁸ but it can also be used to estimate total and regional body fat and LT mass. Pixels, where the ratio between attenuations of the two energies falls below a certain threshold, are classified as soft tissue (ie, without bone), and in those pixels, the attenuation is linearly dependent on the fat fraction of the soft tissue. Pixels above the threshold contain a mixture of bone and soft tissue, and there the soft tissue properties need to be interpolated from surrounding soft tissue pixels.³⁹ Approximately one-third of the pixels of the projected body contains bone.⁴⁰

DXA has been found to be more accurate than density-based methods for estimating total body fat.⁴¹ A possible confounder is that the DXA analysis assumes a constant hydration of lean soft tissue, which is not always true as hydration varies with age, gender and disease.⁴² Excellent repeatability (CV) in the range 1–2 per cent for body fat and 0.5–2 per cent for LT has been reported for DXA.

Since DXA only gives a two-dimensional (coronal) projection, it is not possible to obtain direct compartmental volumetric measurements, so regional volume estimates are obtained indirectly using anatomical models.

For example, VAT and parts of the subcutaneous adipose tissue (SAT) are mixed and cannot be separated in the DXA image. The distribution between VAT and SAT then needs to be estimated from an anatomical model predicting the SAT thickness. Furthermore, the physical properties of the technology do not allow for measurements of ectopic fat in organs such as liver fat or muscle fat infiltration. However, due to its ability to estimate regional fat and measure LT, in combination with relatively high availability, DXA has been used for body composition analysis in a wide range of clinical applications.⁴³

CT

CT gives a three-dimensional high-resolution image volume of the complete or selected parts of the body, computed from a large number of X-ray projections of the body from different angles. The known differences in attenuations of X-rays between lean soft tissue and AT can then be used to separate these tissues, as well as to determine mixtures between them. As opposed to the previously described techniques, CT can accurately determine fat in skeletal muscle tissue¹⁶ and in the liver.⁴⁴ It is, however, significantly less accurate for liver fat <5 per cent which limits its use to diagnose low-grade steatosis.⁴⁴ Being a three-dimensional imaging technique, CT has the potential of giving direct volumetric measurements of organs and different AT depots. In practice, however, CT-based body composition analysis is in most cases limited to two-dimensional analysis of one or a limited number of axial slices of the body, leading to the utilization of the area measured as a proxy for the volume. There are two reasons for this limitation: first, it is important to keep the part of the body being scanned to a minimum in order to minimize the ionizing radiation dose.⁴⁵ This is particularly important in the ethical considerations of research studies on healthy subjects. Second, manual segmentation of different compartments in the images is a very labor intensive task, which can be reduced by limiting the analysis to a few slices rather than a complete three-dimensional volume. This approach, however, limits its precision since the exact locations of slices, in relation to internal organs, cannot be determined a priori and will therefore vary between scans. Nevertheless, CT, together with MRI, is today considered the gold standard for body composition analysis, in particular regional.

MRI

MRI uses the different magnetic properties of the nuclei of certain chemical elements (normally hydrogen in water and fat) in the cells to produce images of soft tissue in the body. A number of MRI-based methods for quantification of AT (eg, see the review by Hu *et al.*⁴⁶) and muscles^{47–52} have been developed and implemented in the past.

By using so-called 'quantitative fat water imaging', precise measurements of regional AT and LT, as well as diffuse fat infiltration in other organs, can be obtained. The basis for quantitative fat water imaging is fat water separated, or Dixon, imaging,⁵³ where the different magnetic resonance frequencies of protons in fat and water are used for separating the two signals into a fat image and a water image. Due to a number of undeterminable factors affecting the MR signal, an MR image is not calibrated on an absolute

scale and therefore not quantitative in itself. But by using different postprocessing techniques, the image can be calibrated to quantitatively measure fat or AT. Examples of such methods are proton density fat fraction (PDFF)⁵⁴ measuring the fraction of fat in MR-visible soft tissue and fat-referenced MRI^{55–57} measuring the amount of AT in each voxel.

As opposed to CT and DXA, MRI does not use ionizing radiation, which enables true volumetric three-dimensional imaging even in healthy volunteers and infants. Still, many studies using MRI for body composition analysis have used one or a limited set of two-dimensional slices, mostly due to the lack of efficient image analysis tools for handling three-dimensional image segmentation. However, since there is no ionizing radiation limiting the image acquisition, the slices can be selected from a complete image volume, thereby reducing the uncertainty in their locations. Still, using a sparse set of slices as a proxy for the complete volume will inevitably negatively affect accuracy and precision as only a fraction of the data is used. It has, for example, been shown that single-slice MRI is poor at predicting VAT and SAT changes during weight loss.^{58 59}

BODY COMPOSITION PROFILING USING FAT-REFERENCED MRI

Body composition profiling implies the simultaneous collection and analysis of a number of body composition parameters, including subcutaneous and visceral AT, ectopic fat such as liver and skeletal muscle fat and muscle volumes. Fat-referenced MRI is a methodology that enables all such measurements in one single rapid examination. This section gives a brief introduction to body composition profiling using fat-referenced MRI, together with a review of published validation results of the method. Finally, a previously unpublished validation study of the agreement between this method and DXA for measurements of body fat/AT, body LT and VAT is presented.

The body composition profiling methodology combines fat-referenced MRI with automated image segmentation of different compartments and was first described by Dahlqvist Leinhard *et al.*⁵⁵ Different aspects of the method have been further described in other publications.^{47 60–62} The two key features of this method are that it produces quantitative fat-referenced images and that it uses a supervised automated segmentation tool.

In a quantitative fat-referenced image, the value in each image volume element (voxel) represents the amount of fat in that voxel in relation to the amount of fat in pure AT. Hence, a voxel in pure AT has a value of one and a voxel without any fat has the value zero. This means that the following can be measured: the total amount of AT in any given region by summation of the voxel values in that region, AT-free volume by removal of amount of AT from volume measurements of regional LT (eg, muscles) and fractions of fat in specific internal organs, such as the liver.

The supervised automated segmentation tool enables an efficient way of segmenting different AT compartments, as well as different muscle groups, reducing the manual work to a few minutes, rather than hours, for analyzing a whole-body data set. Anatomical compartments, such as the visceral compartment and different muscle groups, are automatically segmented using predefined anatomical



Figure 1 Example of segmentation of abdominal subcutaneous AT (ASAT), visceral AT (VAT) and 10 muscle groups from fat water separated MRI using fat-referenced MRI and multi-atlas image segmentation. To the left is the fat image with ASAT (blue) and VAT (red), and to the right is the water image with the different muscle groups colored. Reproduced with permission from AMRA Medical AB.

atlases and the operator can then adjust the segmentations if needed. An example of such segmentations is illustrated in [figure 1](#).

See online supplementary appendix 1 for a summary of how fat-referenced MRI is implemented in AMRA Profiler (AMRA Medical AB, Linköping, Sweden), which is the tool for body composition profiling that was used in the validation studies of fat-referenced MRI.

Precision and accuracy

In a previous study,⁶¹ the accuracy of body composition profiling using fat-referenced MRI, in terms of agreement with manual quantification of T1-weighted MR images, was evaluated on 23 (11 females, 12 males) subjects with an average BMI of $31.7 \pm 5.1 \text{ kg/m}^2$ (range 22–46 kg/m^2); age 36–66 years. There was no significant difference in the measured amount of VAT (4.73 ± 1.99 vs $4.73 \pm 1.75 \text{ L}$, $P=0.97$). Furthermore, the agreement between the methods was excellent for both VAT (95 per cent limits of agreement (LoA) -1.06 to 1.07 L) and abdominal subcutaneous

AT (ASAT) (-0.36 to 1.60 L). However, a very small yet statistically significant difference in ASAT was observed (10.39 ± 5.38 vs $9.78 \pm 5.36 \text{ L}$, $P<0.001$). Clearly this small difference has no clinical significance.

Test-retest repeatability and agreement with manual quantification for VAT was evaluated by Newman *et al*.⁶³ The study included 30 subjects with five subjects from each gender for each of the following categories of BMI: 18–25 kg/m^2 , 25–30 kg/m^2 and $>30 \text{ kg/m}^2$. Each subject was scanned twice with at least 20 min interval, during which the subject left the scanner room. There was no significant difference between the evaluated method and manual quantification of VAT ($P=0.73$). Bland-Altman analysis of the test-retest repeatability showed a bias of -0.04 L (95 per cent LoA -0.12 to 0.13 L) for VAT and 0.05 L (95 per cent LoA -0.55 to 0.64 L) for ASAT. The CV was 1.80 per cent for VAT and 2.98 per cent for ASAT using the method above. The CV for manual quantification of VAT was 6.33 per cent as a comparison.

Middleton *et al* evaluated the accuracy and repeatability of VAT, ASAT and thigh muscle quantification by comparing with manual segmentation on 20 subjects.⁶⁴ Due to the laborious work with manual segmentation, 15 two-dimensional axial slices were manually segmented in the abdominal region for VAT and ASAT and 5 slices over the thigh muscles. For repeatability assessment, the subjects were scanned three times, with the subject remaining in the same position on the scan table between scans 1 and 2 and with the subject removed from the table between scans 2 and 3. The intraexamination (scans 1–2) repeatability test obtained a CV of 3.3 per cent for VAT, 2.2 per cent for ASAT and 1.5 per cent for total thigh muscle volume. For the inter-examination test (scans 2–3), the CVs were 3.6, 2.6 and 1.5 per cent for VAT, ASAT and thigh muscle volume, respectively. Good agreement with the manual measurements in the 20 slices was observed for all measurements. Neither the slopes nor the intercepts of the regression lines were significantly different from those of the identity lines.

Test-retest repeatability of muscle quantification of left and right abdominal muscles, left and right, anterior and posterior thigh muscles and left and right lower limb muscles, as well as accuracy of lower leg muscle quantification, were evaluated by Thomas *et al*.⁶⁵ comparing the method above with manual segmentation. The study included 15 subjects of each gender, ranging from normal weight to obese. Each subject was scanned twice with at least 20 min interval, during which the subject left the scanner room. The intra-class correlation (ICC) between the first and second scan was almost perfect (between 0.99 and 1.0) for all muscle groups. The 95 per cent LoA ranged from -0.04 to 0.02 L for the posterior thigh muscles to -0.15 to 0.08 L for the left lower limb. The lowest accuracy for the lower limbs was a bias of -0.08 L with 95 per cent LoA of -0.25 to 0.09 L .

Test-retest repeatability of measurements of VAT and ASAT volumes and volumes and fat infiltration of left and right posterior and anterior thigh muscles, lower leg muscles and abdominal muscles were evaluated by West *et al* on 36 sedentary postmenopausal women.⁶⁶ Each subject was scanned twice, and the subjects were removed from the scanner room between the acquisitions. The intraexamination CV was 1.54 per cent for VAT, 1.06 per cent for ASAT, 0.8–1.9 per cent for volumes of muscle groups (thigh,

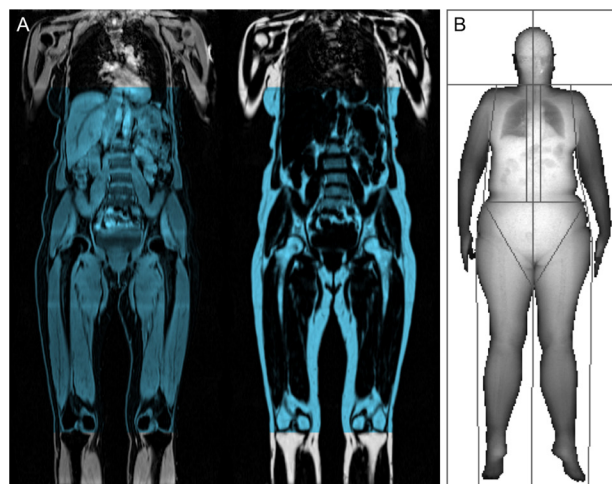


Figure 2 (A) The definition of lean and adipose tissue measured by MRI from the bottom of the thigh muscles to top of vertebrae T9 marked in blue color in the water (left) and fat (right) image. (B) An example of a dual-energy X-ray absorptiometry (DXA) image from the study cohort. DXA image copyright UK Biobank. Reprinted with permission.

lower leg and abdomen) and 2.3–7.0 per cent for individual muscle volumes. The 95 per cent LoA was -0.13 to 0.10 L for VAT, -0.38 to 0.29 L for ASAT. The LoA for liver PDFF were within ± 1.9 per cent, and for muscle fat infiltration, they were within ± 2.06 per cent for muscle groups and within ± 5.13 per cent for individual muscles.

The method's reproducibility of fat-free muscle volume quantification between 1.5 T and 3 T MR scanners, as well as the agreement with manual segmentation, was investigated on 11 different muscle groups.⁴⁷ The ICC between the automated method and manual measurements was at least 0.97 for all muscle groups except in the arms. Except for the arms, the ICC between 1.5 T and 3 T data ranged from 0.97 (left lower leg) to 1.00 (left posterior thigh) with a mean difference volume ranging from 0.39 L (95 per cent LoA 0.01 to 0.77 L) (left abdomen) to 0.0 L (95 per cent LoA -0.10 to 0.09 L) (right lower leg). The muscles of the arms had worse accuracy and reproducibility due to difficulties to include the arms in the field of view.

Agreement with ADP

A previous study⁶⁷ compared AT measured using fat-referenced MRI with total body fat measured by ADP. The ICC was 0.984. After converting the ADP body fat measures to AT volume (assuming that most of the fat resided in AT and a density of 0.9 kg/L for AT), a Bland-Altman analysis showed that ADP underestimated AT by 0.78 L on average, but the bias was strongly dependent on the level of adiposity with significant underestimation for lean subjects and significant overestimation for subjects with higher amounts of AT. Similar bias dependence has been observed when ADP has been compared with DXA³¹ and MRI.⁶⁸

Agreement with BIA

Ulbrich *et al*⁶⁹ investigated the agreement between fat-referenced MRI and BIA on 80 subjects between 20 and 62 years with a BMI range from 17.5 to 26.2 kg/m². The linear

correlation between body fat mass measured by BIA and AT volume measured by MRI was 0.75 and 0.81 for females and males, respectively. The total AT measured by MRI was converted to total fat mass (again assuming that most of the fat resided in AT and using a constant density of 0.94 kg/L). Compared with MRI, the BIA underestimated the total fat with approximately 5 kg (± 7 kg LoA) on average, this despite the fact that the MRI-based measurements of total body fat excluded the arms and lower legs. The highest linear correlation found between BIA and MRI-derived measures was 0.75 and 0.81 for females and males, respectively. These correlations were found between BIA-derived body mass percentage and the MRI-derived 'total AT index' (total AT divided by body height squared).

Agreement with DXA

Methods and materials

The agreement between DXA and the fat-referenced MRI technique was assessed using data from the UK Biobank study,⁷⁰ approved by the North West Multicenter Research Ethics Committee, UK, and with written informed consent obtained from all subjects prior to study entry. The age range for inclusion was 40–69 years of age. For the present analysis, participants were selected, out of the first 6214 scanned, who had both DXA and MRI scans. One subject with obviously erroneous DXA values (2.7 kg total fat and 6.8 kg LT) was excluded, yielding a total 4753 subjects (2502 females and 2251 males). All included MRI images were analyzable for VAT, ASAT and both thigh muscles according to the predefined quality criteria.⁶² The BMI range was 16.4 – 54.3 with a mean of 26.2 kg/m².

The MR images were acquired using a Siemens Aera 1.5 T scanner (Syngo MR D13) (Siemens, Erlangen, Germany) with the dual-echo Dixon Vibe protocol, covering neck to knees as previously described.⁶² The MR images were analyzed using AMRA Profiler. The body AT and LT were measured from the bottom of the thigh muscles to level of the top of vertebrae T9 (figure 2). The LT was defined as the volume of soft tissue subtracted by the volume of AT.⁴⁷

Whole-body DXA data were acquired using a GE-Lunar iDXA (GE Healthcare, Madison, Wisconsin, USA) with the subjects in supine position.⁷¹ The images were analyzed using the GE enCORE software by the radiographer at, or soon after, the scan. The GE iDXA estimates VAT within an automatically segmented region with the lower border at the top of the iliac crest and its height is set to 20 per cent of the distance from the top of the iliac crest to the base of the skull.⁷²

Since the DXA and MRI analyses measure different entities (fat and LT mass vs AT and LT volume, respectively) and they do not cover the same part of the body, a linear model was estimated by linear regression between the MRI and DXA measurements using a training data set of 2376 randomly selected subjects. The remaining 2377 subjects were then used for estimating the agreement between the techniques after linear transformation using the linear model (ie, validating the linear model). The MRI-based measurements (L) were transformed to predict the DXA measurements (kg) using the linear regression coefficients from the training data, and a Bland-Altman analysis was performed to investigate the agreement between MRI-derived and

DXA-derived measurements in the validation data. To investigate the agreement between DXA and MRI-derived VAT measurements, a linear model was estimated between the DXA and MRI measurements. Of the 4669 subjects with available DXA VAT measurements, 2334 cases were used to estimate the model and the remaining 2335 subjects were used to validate the agreement between VAT measured by MRI and the transformed DXA measurements using Bland-Altman analysis.

Results

The linear regression between MRI and DXA was $1.23x - 0.12$ (kg/L) for body fat/AT and $1.88x + 1.82$ (kg/L) for body LT. The linear correlation coefficient, r , between DXA and the transformed MRI measurements was 0.99 for body fat and 0.97 for LT. The 95 per cent LoA from the Bland-Altman analysis were -2.25 to 2.31 kg for fat and -4.33 to 4.31 kg for LT (figure 3). The prediction error SD relative to the mean (CV) was 4.5 per cent for body fat and 4.6 per cent for LT. The correlation between VAT measured by MRI and VAT as predicted by DXA was 0.97 and the LoA were -1.02 to 1.05 L, with CV=21 per cent (figure 4).

DISCUSSION

Densitometry, including ADP, shows relatively good precision and high correlation with MRI-based measurements of whole-body AT, but with a significant volume-dependent bias. Since these methods only measure the volume

or density of the body, they cannot be used for regional measurements and body composition profiling.

BIA is highly available and its relatively low cost is an advantage, which also makes it useful for consumer products. Furthermore, it can differentiate intracellular water from extracellular water, which is a unique capability of BIA. BIA can also, in principle, be used for regional measurements, but it is severely limited when it comes to measuring VAT or ectopic fat in internal organs.

DXA techniques have shown good accuracy when evaluated against MRI for whole-body measurements and very good repeatability. The prediction of whole-body fat and LT from MRI agrees well with DXA after a linear transformation, but less so for VAT. While the correlation between DXA and MRI-derived VAT was high ($r=0.97$), the agreement after a linear transformation was, however, much lower than for total body fat and body LT, with a CV >20 per cent. The high linear correlation, despite a modest agreement, can be explained by the very wide range of measured VAT volumes, ranging from almost 0 to >14L. The CV for VAT is in line with the results by Kaul *et al* with a CV of 15.6 per cent for females and 25.9 per cent for males when comparing the same DXA model with CT.⁷² Park *et al* found a linear correlation of 0.85 between VAT measured by DXA and MRI in a study including 90 non-obese men.⁷³ However, Kamel *et al* found that the correlation was much lower ($r=0.46$) for obese men.⁷⁴ The fact that the agreement is lower for obese subjects can also be observed in figure 4 where the prediction

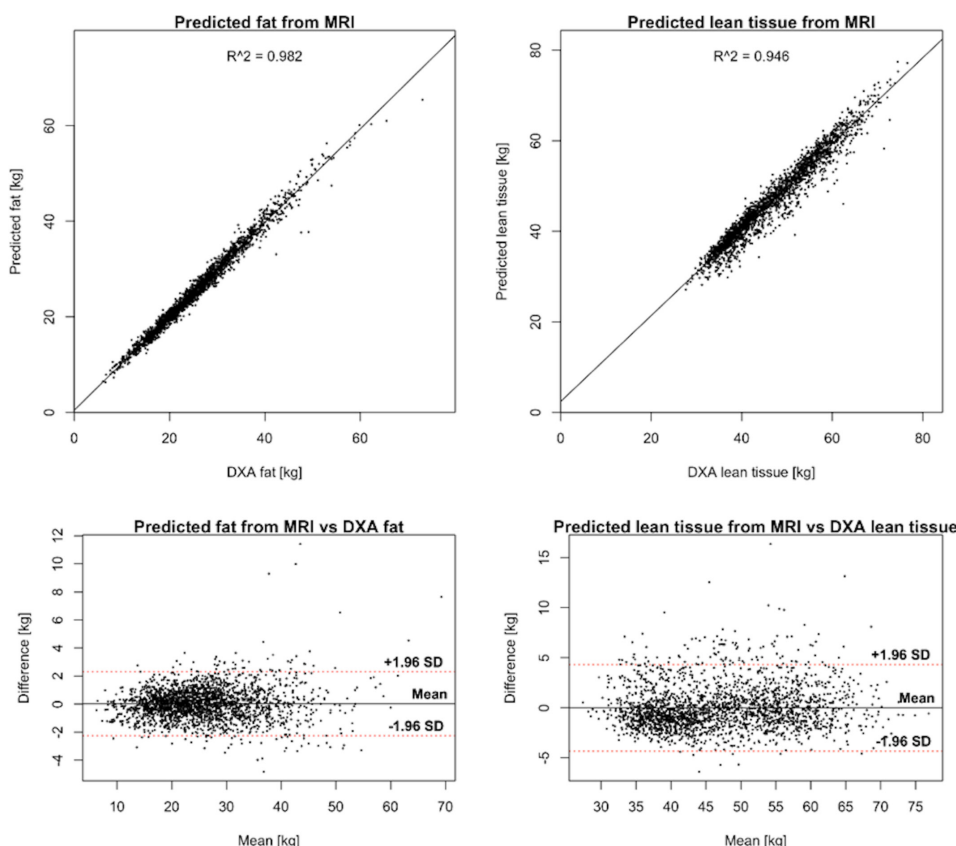


Figure 3 Correlation plots (upper row) between dual-energy X-ray absorptiometry (DXA) and corresponding measurement predicted from MRI using a linear transformation for body fat (left) and body lean tissue (right). The bottom row shows Bland-Altman plots of the agreement between DXA and corresponding measures predicted from MRI.

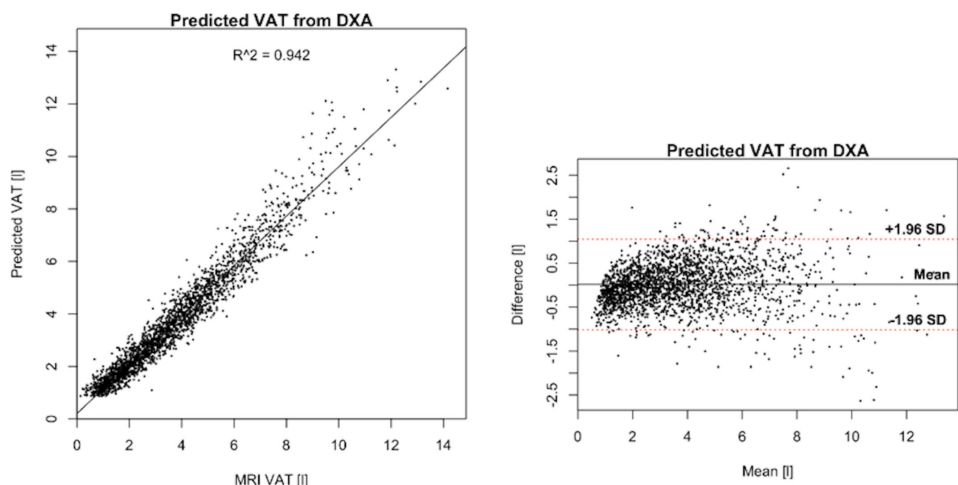


Figure 4 Correlation between visceral adipose tissue (VAT) predicted by dual-energy X-ray absorptiometry (DXA) and VAT measured by MRI (left) and Bland-Altman plot showing the agreement (liters) between the methods (right).

error increases with increased VAT volume. Silver *et al* found an excellent correlation without significant bias between fat water MRI and DXA for ‘gross body adipose tissue’ but with a significant negative bias (MRI – DXA) for ‘total trunk adipose tissue’ as well as total and trunk LT.⁷⁵ Interestingly, for DXA, the lowest precision is for fat in the arms, with reported CV up to 11 per cent.⁷⁶ This is the same compartment that is difficult to measure with MRI due to signal loss in the outer parts of the field of view. A strength with DXA, compared with MRI, is the simultaneous assessment of bone mineral density and mass.

When comparing different technologies, both accuracy and precision are important. Accuracy, however, can be rather difficult to compare between technologies for several reasons. First, there is no ground truth available. Even though there is a growing consensus that tomographic methods are the gold standard that can be used to assess accuracy for other methods, they differ between themselves and are difficult to compare in terms of accuracy. Using physical phantoms is one way to assess accuracy, but they miss the difficulties caused by anatomical variations that we know can lead to different measurement errors. Automated tomographic imaging methods can be evaluated against manual methods, but this addresses only one of several important components in the measurement system—the segmentation of different compartments. Second, not all methods measure the same thing, so even if two technologies correlate strongly, there may be a significant bias if they measure different physical entities. For example, AT is not equivalent to fat—besides fat AT also contains water, protein and minerals. When comparing a method that measures AT in volume units, such as MRI, to a method that measures fat in weight units (eg, DXA), we have to convert one unit to the other using a density that is assumed to be constant, which again may not be always accurate.

Although this review has not focused on measurements of ectopic fat, this is an important component in body composition profiling, especially for understanding metabolic status and assessing risk. Among the techniques discussed here, CT and quantitative MRI are the only methods that can quantify local diffuse infiltration of AT and ectopic fat.

(Non-invasive measurements of ectopic fat, in particular liver fat, are commonly done by MR spectroscopy (MRS), but since MRS only measures local substance concentrations and not absolute amounts of fat, AT or LT, this technology was not included in this study.) While it is possible—and sometimes necessary—to use different equipment for different measurements in a study, it is often desirable to keep the number of different examinations and modalities to a minimum in order to optimize the work flow. By using quantitative MRI, or CT if the radiation dose is not a concern, a large number of metabolically relevant body composition parameters can be measured with high accuracy and precision in a single examination.

A comparison of the capabilities of different measurements of the techniques discussed above is summarized in [table 1](#).

CONCLUSION

There are several methods available that can measure whole-body AT or fat and LT. In terms of precision and accuracy, DXA and MRI are comparable as they show excellent agreement after a linear transformation. However, the agreement is much lower for compartmental measurements such as VAT. Moreover, MRI gives access to accurate and direct measurements of diffuse infiltration of AT in muscles and ectopic fat (eg, liver fat). Rapid MRI scanning protocols, in combination with efficient image analysis methods, have promoted

| Table 1 Comparison of the capabilities of different techniques for body composition analysis | | | | | |
|--|-----|-----|-------------|-----|-----|
| | ADP | BIA | DXA | CT | MRI |
| Total fat | Yes | Yes | Yes | Yes | Yes |
| Total lean tissue | Yes | Yes | Yes | Yes | Yes |
| VAT | No | No | Approximate | Yes | Yes |
| Volume of individual muscles | No | No | No | Yes | Yes |
| Diffuse fat infiltration | No | No | No | Yes | Yes |
| Without ionizing radiation | Yes | Yes | No (low) | No | Yes |
| ADP, air displacement plethysmography; BIA, bioelectrical impedance analysis; DXA, dual-energy X-ray absorptiometry; VAT, visceral adipose tissue. | | | | | |

MRI to a competitive option for advanced body composition assessment, thus enabling a more complete description of a person's body composition profile from a single examination.

Acknowledgements This research has been conducted using the UK Biobank Resource under Data Access Application 6569. For full acknowledgements, see online supplementary appendix 2.

Contributors MB, JW and ODL planned the work. MB, TR and ODL developed and applied the MR analysis methods used. JDB was responsible for the UK Biobank body MRI and NCH for the UK Biobank DXA scans. MB and JW performed the statistical analyses. SBH contributed with expertise on body composition. MB drafted the manuscript. All authors contributed to editing the text.

Funding Funding support for analysis of UK Biobank data was provided by Pfizer.

Competing interests MB, JW, TR and ODL are employees and stockholders of AMRA Medical AB.

Patient consent Obtained.

Ethics approval North West Multicenter Research Ethics Committee (MREC), UK.

Provenance and peer review Commissioned; externally peer reviewed.

Data sharing statement The underlying data used in this study are available through the UK Biobank resource. All bonafide researchers can apply to use the UK Biobank resource for health-related research that is of public interest by applying in the Access Management System (AMS). For details, see: www.ukbiobank.ac.uk/

© American Federation for Medical Research (unless otherwise stated in the text of the article) 2018. All rights reserved. No commercial use is permitted unless otherwise expressly granted.

REFERENCES

- Wang ZM, Pierson RN, Heymsfield SB. The five-level model: a new approach to organizing body-composition research. *Am J Clin Nutr* 1992;56:19–28.
- Gallagher D, Heymsfield SB, Heo M, et al. Healthy percentage body fat ranges: an approach for developing guidelines based on body mass index. *Am J Clin Nutr* 2000;72:694–701.
- Thomas EL, Frost G, Taylor-Robinson SD, et al. Excess body fat in obese and normal-weight subjects. *Nutr Res Rev* 2012;25:150–61.
- Shen W, Wang Z, Punyanita M, et al. Adipose tissue quantification by imaging methods: a proposed classification. *Obes Res* 2003;11:5–16.
- Snyder WS. *Report of the Task group on reference man: a report*. Oxford: Pergamon 1975;1975.
- Britton KA, Fox CS, Depots EF, et al. *Circulation* 2011;124:e837–e841.
- Demerath EW, Reed D, Rogers N, et al. Visceral adiposity and its anatomical distribution as predictors of the metabolic syndrome and cardiometabolic risk factor levels. *Am J Clin Nutr* 2008;88:1263–71.
- Liu J, Fox CS, Hickson DA, et al. Impact of abdominal visceral and subcutaneous adipose tissue on cardiometabolic risk factors: the Jackson Heart Study. *J Clin Endocrinol Metab* 2010;95:5419–26.
- Neeland IJ, Ayers CR, Rohatgi AK, et al. Associations of visceral and abdominal subcutaneous adipose tissue with markers of cardiac and metabolic risk in obese adults. *Obesity* 2013;21:n/a–E447.
- Iwasa M, Mifuji-Moroka R, Hara N, et al. Visceral fat volume predicts new-onset type 2 diabetes in patients with chronic hepatitis C. *Diabetes Res Clin Pract* 2011;94:468–70.
- Kurioka S, Murakami Y, Nishiki M, et al. Relationship between visceral fat accumulation and anti-lipolytic action of insulin in patients with type 2 diabetes mellitus. *Endocr J* 2002;49:459–64.
- van der Poorten D, Milner KL, Hui J, et al. Visceral fat: a key mediator of steatohepatitis in metabolic liver disease. *Hepatology* 2008;48:449–57.
- Britton KA, Massaro JM, Murabito JM, et al. Body fat distribution, incident cardiovascular disease, cancer, and all-cause mortality. *J Am Coll Cardiol* 2013;62:921–5.
- Doyle SL, Donohoe CL, Lysaght J, et al. Visceral obesity, metabolic syndrome, insulin resistance and cancer. *Proc Nutr Soc* 2012;71:181–9.
- Ekstedt M, Franzén LE, Mathiesen UL, et al. Long-term follow-up of patients with NAFLD and elevated liver enzymes. *Hepatology* 2006;44:865–73.
- Goodpaster BH, Kelley DE, Thaete FL, et al. Skeletal muscle attenuation determined by computed tomography is associated with skeletal muscle lipid content. *J Appl Physiol* 2000;89:104–10.
- Marcus RL, Addison O, Dibble LE, et al. Intramuscular adipose tissue, sarcopenia, and mobility function in older individuals. *J Aging Res* 2012;2012:1–6.
- Chan DC, Watts GF, Barrett PH, et al. Waist circumference, waist-to-hip ratio and body mass index as predictors of adipose tissue compartments in men. *QJM* 2003;96:441–7.
- Hsu CH, Lin JD, Hsieh CH, et al. Adiposity measurements in association with metabolic syndrome in older men have different clinical implications. *Nutr Res* 2014;34:219–25.
- Prentice AM, Jebb SA. Beyond body mass index. *Obes Rev* 2001;2:141–7.
- Tomiya AJ, Hunger JM, Nguyen-Cuu J, et al. Misclassification of cardiometabolic health when using body mass index categories in NHANES 2005–2012. *Int J Obes* 2016;40:883–6.
- Morley JE, Thomas DR, Wilson MM. Cachexia: pathophysiology and clinical relevance. *Am J Clin Nutr* 2006;83:735–43.
- Fearon K, Strasser F, Anker SD, et al. Definition and classification of cancer cachexia: an international consensus. *Lancet Oncol* 2011;12:489–95.
- Cruz-Jentoft AJ, Morley JE. *Sarcopenia*. 2012. Chichester, West Sussex: Hoboken NJ: Wiley-Blackwell.
- Cruz-Jentoft AJ, Baeyens JP, Bauer JM, et al. *Sarcopenia: European consensus on definition and diagnosis: report of the European Working Group on Sarcopenia in older people*. United States, North America: Oxford University Press, 2010.
- Willis TA, Hollingsworth KG, Coombs A, Eagle M, Mayhew A, et al. Quantitative muscle MRI as an assessment tool for monitoring disease progression in LGMD21: a multicentre longitudinal study. *PLoS One* 2013;8:e70993.
- Thomas EL, Fitzpatrick JA, Malik SJ, et al. Whole body fat: content and distribution. *Prog Nucl Magn Reson Spectrosc* 2013;73:56–80.
- Baumgartner RN, Heymsfield SB, Lichtman S, et al. Body composition in elderly people: effect of criterion estimates on predictive equations. *Am J Clin Nutr* 1991;53:1345–53.
- Bergsma-Kadijk JA, Baumeister B, Deurenberg P. Measurement of body fat in young and elderly women: comparison between a four-compartment model and widely used reference methods. *Br J Nutr* 1996;75:649–57.
- Forbes GB. *Human body composition: growth, aging, nutrition, and activity*. Verlag: Springer, 1987.
- Fields DA, Goran MI, McCrory MA. Body-composition assessment via air-displacement plethysmography in adults and children: a review. *Am J Clin Nutr* 2002;75:453–67.
- Stahn A, Terblanche E, Gunga H-C. Use of bioelectrical impedance: general principles and overview. In: Preedy VR, ed. *Handbook of anthropometry: physical measures of human form in health and disease*. New York, NY: Springer New York, 2012:49–90.
- Khalil SF, Mohktar MS, Ibrahim F. The theory and fundamentals of bioimpedance analysis in clinical status monitoring and diagnosis of diseases. *Sensors* 2014;14:10895–928.
- Kyle UG, Bosaeus I, De Lorenzo AD, et al. Bioelectrical impedance analysis--part I: review of principles and methods. *Clin Nutr* 2004;23:1226–43.
- Browning LM, Mugridge O, Chatfield MD, et al. Validity of a new abdominal bioelectrical impedance device to measure abdominal and visceral fat: comparison with MRI. *Obesity* 2010;18:2385–91.
- Dehghan M, Merchant AT. Is bioelectrical impedance accurate for use in large epidemiological studies? *Nutr J* 2008;7:26.
- Haroun D, Taylor SJ, Viner RM, et al. Validation of bioelectrical impedance analysis in adolescents across different ethnic groups. *Obesity* 2010;18:1252–9.
- Garg MK, Kharb S. Dual energy X-ray absorptiometry: pitfalls in measurement and interpretation of bone mineral density. *Indian J Endocrinol Metab* 2013;17:203–10.
- Webber CE. Reproducibility of DXA measurements of bone mineral and body composition: application to routine clinical measurements. In: Preedy VR, ed. *Handbook of anthropometry: physical measures of human form in health and disease*. New York, NY: Springer New York, 2012:151–65.
- Laskey MA. Dual-energy X-ray absorptiometry and body composition. *Nutrition* 1996;12:45–51.
- Prior BM, Cureton KJ, Modlesky CM, et al. In vivo validation of whole body composition estimates from dual-energy X-ray absorptiometry. *J Appl Physiol* 1997;83:623–30.
- Siri WE. The gross composition of the body. *Adv Biol Med Phys* 1956;4:239–80.
- Albanese CV, Diessel E, Genant HK. Clinical applications of body composition measurements using DXA. *J Clin Densitom* 2003;6:75–85.
- Kramer H, Pickhardt PJ, Kliever MA, et al. Accuracy of liver fat quantification with advanced CT, MRI, and ultrasound techniques: prospective comparison with MR spectroscopy. *AJR Am J Roentgenol* 2017;208:92–100.

- 45 Yu L, Liu X, Leng S, *et al.* Radiation dose reduction in computed tomography: techniques and future perspective. *Imaging Med* 2009;1:65–84.
- 46 Hu HH, Chen J, Shen W. Segmentation and quantification of adipose tissue by magnetic resonance imaging. *Magnetic Resonance Materials in Physics, Biology and Medicine* 2016;29:259–76.
- 47 Karlsson A, Rosander J, Romu T, Borga M, *et al.* Automatic and quantitative assessment of regional muscle volume by multi-atlas segmentation using whole-body water-fat MRI. *J Magn Reson Imaging* 2015;41:1558–69.
- 48 Andrews S, Hamarneh G. The generalized log-ratio transformation: learning shape and adjacency priors for simultaneous thigh muscle segmentation. *IEEE Trans Med Imaging* 2015;34:1773–87.
- 49 Lareau-Trudel E, Le Troter A, Ghattas B, *et al.* Muscle quantitative MR imaging and clustering analysis in patients with facioscapulohumeral muscular dystrophy type 1. *PLoS One* 2015;10:e0132717–16.
- 50 Orgiu S, Lafortuna CL, Rastelli F, *et al.* Automatic muscle and fat segmentation in the thigh from T1-Weighted MRI. *J Magn Reson Imaging* 2016;43:601–10.
- 51 Ugarte V, Sinha U, Malis V, *et al.* 3D multimodal spatial fuzzy segmentation of intramuscular connective and adipose tissue from ultrashort TE MR images of calf muscle. *Magn Reson Med* 2017;77:870–83.
- 52 Yang YX, Chong MS, Tay L, *et al.* Automated assessment of thigh composition using machine learning for Dixon magnetic resonance images. *MAGMA* 2016;29:723–31.
- 53 Dixon WT. Simple proton spectroscopic imaging. *Radiology* 1984;153:189–94.
- 54 Reeder SB, Hu HH, Sirlin CB. Proton density fat-fraction: a standardized MR-based biomarker of tissue fat concentration. *J Magn Reson Imaging* 2012;36:1011–4.
- 55 Dahlqvist Leinhard O, Johansson A, Rydell J, *et al.* Quantitative abdominal fat estimation using MRI. *International conference on pattern recognition*, 2008:2137–40. Conf: 19.
- 56 Hu HH, Nayak KS. Quantification of absolute fat mass using an adipose tissue reference signal model. *J Magn Reson Imaging* 2008;28:1483–91.
- 57 Romu T, Borga M, Dahlqvist Leinhard O. MANA - Multi scale adaptive normalized averaging. *Proceedings of the 2011 IEEE International Symposium on Biomedical Imaging: From Nano to Macro*, 2011:361–4.
- 58 Thomas EL, Bell JD. Influence of undersampling on magnetic resonance imaging measurements of intra-abdominal adipose tissue. *Int J Obes Relat Metab Disord* 2003;27:211–8.
- 59 Shen W, Chen J, Gantz M, *et al.* A single MRI slice does not accurately predict visceral and subcutaneous adipose tissue changes during weight loss. *Obesity* 2012;20:2458–63.
- 60 Romu T, Dahlqvist Leinhard O, Forsgren M, *et al.* Fat water classification of symmetrically sampled two-point dixon images using biased partial volume effects. *Proceedings of the annual meeting of the International Society for Magnetic Resonance in Medicine (ISMRM 2011)*, 2011.
- 61 Borga M, Thomas EL, Romu T, Rosander J, *et al.* Validation of a fast method for quantification of intra-abdominal and subcutaneous adipose tissue for large-scale human studies. *NMR Biomed* 2015;28:1747–53.
- 62 West J, Dahlqvist Leinhard O, Romu T, *et al.* Feasibility of MR-based body composition analysis in large scale population studies. *PLoS One* 2016;11:e0163332.
- 63 Newman D, Kelly-Morland C, Leinhard OD, *et al.* Test-retest reliability of rapid whole body and compartmental fat volume quantification on a widebore 3T MR system in normal-weight, overweight, and obese subjects. *J Magn Reson Imaging* 2016;44:1464–73.
- 64 Middleton MS, Haufe W, Hooker J, *et al.* Quantifying abdominal adipose tissue and thigh muscle volume and hepatic proton density fat fraction: repeatability and accuracy of an mr imaging-based, semiautomated analysis method. *Radiology* 2017;283:438–49.
- 65 Thomas MS, Newman D, Leinhard OD, *et al.* Test-retest reliability of automated whole body and compartmental muscle volume measurements on a wide bore 3T MR system. *Eur Radiol* 2014;24:2279–91.
- 66 West J, Romu T, Thorell S, *et al.* Precision of MRI-based body composition measurements of postmenopausal women. *PLoS One* 2018;13:e0192495.
- 67 Thunón P, Romu T, Zanjani S, *et al.* Body composition volumetry by whole-body water-fat separated MRI. *ISMRM 22nd Annual Meeting & Exhibition*, 10 - 16 May 2014, Milan, Italy, 2014.
- 68 Ludwig UA, Klausmann F, Baumann S, *et al.* Whole-body MRI-based fat quantification: a comparison to air displacement plethysmography. *J Magn Reson Imaging* 2014;40:1437–44.
- 69 Ulbrich EJ, Nanz D, Leinhard OD, *et al.* Whole-body adipose tissue and lean muscle volumes and their distribution across gender and age: MR-derived normative values in a normal-weight Swiss population. *Magn Reson Med* 2018;79.
- 70 Sudlow C, Gallacher J, Allen N, *et al.* UK biobank: an open access resource for identifying the causes of a wide range of complex diseases of middle and old age. *PLoS Med* 2015;12:e1001779.
- 71 Harvey NC, Matthews P, Collins R, *et al.* Osteoporosis epidemiology in UK Biobank: a unique opportunity for international researchers. *Osteoporos Int* 2013;24:2903–5.
- 72 Kaul S, Rothney MP, Peters DM, *et al.* Dual-energy X-ray absorptiometry for quantification of visceral fat. *Obesity* 2012;20:1313–8.
- 73 Park YW, Heymsfield SB, Gallagher D. Are dual-energy X-ray absorptiometry regional estimates associated with visceral adipose tissue mass? *Int J Obes Relat Metab Disord* 2002;26:978–83.
- 74 Kamel EG, McNeill G, Van Wijk MC. Usefulness of anthropometry and DXA in predicting intra-abdominal fat in obese men and women. *Obes Res* 2000;8:36–42.
- 75 Silver HJ, Niswender KD, Kullberg J, *et al.* Comparison of gross body fat-water magnetic resonance imaging at 3 Tesla to dual-energy X-ray absorptiometry in obese women. *Obesity* 2013;21:765–74.
- 76 Cordero-MacIntyre ZR, Peters W, Libanati CR, *et al.* Reproducibility of DXA in obese women. *J Clin Densitom* 2002;5:35–44.