Redefining overweight and obesity in rheumatoid arthritis patients


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Redefining overweight and obesity in rheumatoid arthritis patients
Extended Report

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Abstract

Objectives: To assess whether body mass index (BMI) and body fat (BF) differ between RA patients, patients with non-inflammatory arthritis (Osteoarthritis – OA) and healthy individuals, and whether disease-specific measures of adiposity are required to accurately reflect BF in these groups.

Methods: A total of 641 individuals were assessed for BMI (kg/m²) and BF (bioelectrical impedance). Of them, 299 [174 RA, 43 OA and 82 healthy controls (HC)] formed the observation group and 342 (all RA) the validation group. RA disease characteristics were collected.

Results: ANOVA revealed significant differences between disease groups for BMI (p<0.05) and BF (p<0.001). ANCOVA showed that age accounted for the differences in BMI (F1,294 = 5.10, p<0.05); age (F1,293 = 22.43, p<0.001), gender (F1,293 = 380.90, p<0.001) and disease (F2, 293 = 18.7, p<0.001) accounted for the differences in BF. For a given BF, patients with RA exhibited BMI levels reduced by 1.83kg/m² (p<0.001) compared to HC; there were no significant differences between OA and HC. A predictive model for BF was developed (R²=0.769, p<0.001) and validated using Limits of Agreement Analysis against measured BF in the validation group (95%LIMAG = 6.17; CV = 8.94).

Conclusions: In individuals with RA, BMI cut-off points should be reduced by 2 kg/m² (i.e. to 23 kg/m² for overweight and 28 kg/m² for obesity). The equation developed can be used to accurately predict body fat from BMI in RA patients. These findings may be important in the context of the cardiovascular comorbidity of RA.

Key Words: Rheumatoid arthritis, cardiovascular risk, body mass index, body composition, bioelectrical impedance.
**Introduction**

Excess body fat (BF) is a prominent health hazard [1] significantly contributing to the development of cardiovascular disease (CVD).[2] About two-thirds of patients who have had a myocardial infarction (MI) exhibit increased body weight.[3] Obesity increases the risk of coronary heart disease (CHD) through a number of different pathophysiological pathways, including insulin resistance, type 2 diabetes, hypertension and dyslipidaemia.[4, 5]

Assessments for overweight or obesity include the calculation of body mass index [6] (BMI - in kg/m2) or more accurate estimations of relative adiposity (BF percentage) through a number of techniques (e.g., skinfold thickness, hydrostatic weighing, and bioelectrical impedance).[7] BF estimations require sophisticated equipment and trained personnel, whereas BMI is easy to obtain and is widely used in the routine clinical setting.

In the general population, BMI of <25, 25-30, and >30kg/m2 indicate healthy, overweight, and obese individuals and associate with low, medium, and high CVD risk, respectively.[8, 9] However, BMI is only a proxy of body fat, [6] and over recent years its validity has been questioned.[3, 7, 10-13] Overweight as defined by BMI of >25, has poor specificity in detecting excess body fat in healthy men and women of all ages [6] as well as in patients with coronary heart disease.[3] In specific sub-populations, such as Indian-Asian,[10] women,[12-14] and large size athletes,[7] new BMI cut-off points have been suggested, that optimally reflect BF and may better predict CVD risk.

The weakness of BMI is that it does not distinguish between lean-body mass and fat mass. Consequently people of similar stature and weight, but different muscle content, will have the same BMI but different BF levels. This tends to be more evident in individuals with low BMI levels.[6] Such limitations of the BMI may explain the better cardiovascular outcomes observed in overweight and mildly obese patients with established CHD as compared to their normal-weight counterparts, who may have proportionately more BF.[3] Therefore, although it is well established that CHD risk increases with advancing BMI levels, [9] global cut-off points may be misleading for several populations.

Central obesity poses a great risk for cardiovascular disease.[15, 16] Regional fat distribution, as measured by waist-to-hip ratio, has been proposed as a more accurate predictor of CHD risk than BMI.[15, 16] Indeed, it has been suggested that obesity should be redefined based on waist-to-hip ratio instead of BMI, since waist-to-hip ratio is significantly associated with MI risk in most ethnic groups.[17] However, its predictive strength can be negatively affected by gender and overall body weight. [18] In a way that pear- shaped or obese individuals might have optimal waist-to-hip ratio but increased overall body weight. More research is necessary to identify the optimal definition of obesity as a predictor for CHD in the general population and specific sub-groups.[19]

Patients with rheumatoid arthritis (RA) have an increased risk for CHD events.[20] RA is a chronic inflammatory disease which affects predominantly synovial joints, causing pain, swelling, stiffness and eventually irreversible damage and deformity, all of which may lead to significant reduction in physical activity. RA associates with increased mortality particularly from CHD,[20] most probably due to accelerated atherogenesis secondary to the metabolic and vascular effects of systemic inflammation.[21] Nearly two thirds of all individuals with RA experience involuntary loss of fat-free mass and progressively increased fat mass in the presence of stable or even slightly decreased weight, a condition referred to as rheumatoid cachexia.[22] The exact mechanisms causing rheumatoid cachexia remain
undetermined, but muscle loss due to systemic inflammation and reduced physical activity may both contribute.[23]

We hypothesised that for a given BMI, RA patients exhibit significantly higher proportions of fat mass than healthy individuals, or even than patients with movement restriction due to a non-inflammatory arthritis, such as osteoarthritis (OA). The possible consequences of this, in the context of the increased CVD mortality in RA, are obvious. In the present study we aimed to investigate whether BMI and BF differ according to arthritic disease (OA vs RA) and within RA according to disease state (e.g. active vs inactive, early vs established disease). We also developed and validated RA-specific BMI cut-off levels and algorithms to calculate BF from BMI.

Methods

Participants

Consecutive patients attending routine rheumatology or orthopaedic outpatient clinics at the Dudley Group of Hospitals NHS Trust, UK, and healthy controls (Hospital and University staff) were invited to participate. The study had Local Research Ethics Committee approval by the Dudley Ethics Committee, and all volunteers provided informed consent. The observation group (n=299) included 174 volunteers with RA (1987 revised ACR criteria [24]), 43 with OA of the hip [25] or knee,[26] and 82 healthy controls (individuals who by self-report did not have any known clinical conditions and were taking no medication). The validation group (n=342) consisted of RA patients only. Demographic and disease characteristics from all subjects appear in Table 1.

Assessments

All volunteers were subjected to the same data collection procedures overseen by the same trained investigators. Specifically, standing height was measured to the nearest 0.5cm on a Seca 214 Road Rod portable stadiometer. Body composition was assessed by Bioelectrical Impedance, using a Tanita BC-418 MA Segmental Body Composition Analyzer, which incorporates 8 tactile electrodes (Tanita Corporation, Tokyo, Japan). This apparatus, measures total body mass and assesses body composition in terms of percentage body fat, fat mass, fat free mass and total body water, as well as fat distribution in different body segments (abdominal and peripheral fat) and has a standard error of <3.[27] After initial manual entry of their demographic details, participants stood bear-footed on the analyzer and held the handgrips provided until the apparatus printed the results. BMI (kg/m²) was calculated on the basis of measured height and weight. In RA patients, contemporary serological inflammation and clinical disease activity were assessed by the erythrocyte sedimentation rate (ESR), C-reactive protein (CRP) [using routine laboratory procedures] and the Disease Activity Score-28 (DAS28).[28] Disease duration was recorded from review of the patients’ hospital notes.
Table 1: Demographic and disease characteristics of all volunteers [mean (sd)]

<table>
<thead>
<tr>
<th></th>
<th>Observation group</th>
<th>Validation Group</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>MALE (n= 110)</td>
<td>FEMALE (n= 189)</td>
</tr>
<tr>
<td>N</td>
<td>RA</td>
<td>OA</td>
</tr>
<tr>
<td>56</td>
<td>15</td>
<td>39</td>
</tr>
<tr>
<td>Age</td>
<td>60.6 (11.8)**</td>
<td>56.7 (13.3)*</td>
</tr>
<tr>
<td>Height</td>
<td>173.6 (7)*</td>
<td>171.3 (6.7)*</td>
</tr>
<tr>
<td>Weight</td>
<td>83.6 (13.3)</td>
<td>78.4 (14.8)</td>
</tr>
<tr>
<td>BMI</td>
<td>27.7 (4.3)*</td>
<td>26.8 (4.7)</td>
</tr>
<tr>
<td>BF</td>
<td>28.7 (7.7)**</td>
<td>24.8 (7.9)*</td>
</tr>
<tr>
<td>Trunkal Fat</td>
<td>30.5 (8)**</td>
<td>26.6 (8.9)*</td>
</tr>
<tr>
<td>DAS28</td>
<td>4.2 (1.2)</td>
<td>4.3 (1.4)</td>
</tr>
<tr>
<td>ESR</td>
<td>23.2 (18.5)</td>
<td>26 (22.1)</td>
</tr>
<tr>
<td>CRP</td>
<td>15.6 (15)</td>
<td>15.8 (14.9)</td>
</tr>
<tr>
<td>Disease Duration</td>
<td>11.4 (10.2)</td>
<td>11.3 (9.9)</td>
</tr>
</tbody>
</table>

One-way ANOVA:  * Significant difference compared to HC (p<0.05)
** Significant difference compared to HC (p<0.001)
† Significant difference compared to OA (p<0.05)
§ Significant difference compared to experimental RA group (p<0.001)
Data management and analysis

Data were inserted in a purpose-designed spreadsheet (Microsoft Excel 2003) and audited for accuracy weekly. They were exported for analysis to The Statistical Package for Social Sciences version 11.0 (SPSS Inc. Chicago, IL, USA). Preliminary evaluation of the variables using a Kolmogorov-Smirnov test of normality revealed that none of them required logarithmic transformation to reach normality. Means (sd) were calculated for all variables.

The method of analysis was to define either BMI or BF as the dependent variable and then to incorporate all other known parameters thought to influence these measures of adiposity as either factors, in an ANOVA, or factors with covariates in an ANCOVA. Factors included gender and disease status (RA, OA and HC) while age, disease activity and duration, and serological inflammation were entered as continuous covariates. The initial ANCOVA analysis incorporated all these factors and covariates, but only those found to be significant were subsequently retained and reported in the prediction equation model below.

Within the RA population of the observation group, correlations of disease activity (DAS28, ESR, CRP) and disease duration with BMI and BF were obtained for each gender. RA patients were also sub-grouped according to their clinical disease activity (DAS remission < 2.6, mild 2.7 – 3.2, moderate 3.3 – 5.1, high > 5.1 [28]), serological inflammation (ESR [29] and CRP [30]), disease duration (early <3 years, established 3-10 years, longstanding >10 years), rheumatoid factor positivity (ever), or corticosteroid administration (yes/no ever): differences between these sub-groups in relation to BMI and BF were assessed using ANCOVA (Table 2). The level of significance was set at p<0.05.

The external validity of the predictive model was tested with the Limits of Agreement (LIMAG) method [31] against BF of the validation group. The Limits of Agreement were obtained as follows:

i) We calculated the mean (d) and the standard deviation (s) of the differences that indicate the level of bias and the random variation between the two measures of BF (i.e. the predicted BF and measured BF of the validation group, respectively).

ii) Provided the differences are normally distributed, the 95% ‘Limits of Agreement’ are given by: d ± (1.96×s).

Bland and Altman [31] argue that, provided that differences within these limits are not clinically important, the two measurement methods can be used interchangeably.

Results
Observation group:
Within the RA population of the observation group, no significant correlations were found between DAS28, ESR, CRP, disease duration and BMI or BF. Similarly, when RA patients were grouped according to these variables as well as rheumatoid factor positivity and corticosteroid use, no significant differences for BMI and BF were observed (p>0.05 in all cases, see Table 2).
Table 2: BMI and BF of RA patients (observation group) according to categorisation based on their disease characteristics.

<table>
<thead>
<tr>
<th>Disease Characteristics</th>
<th>Categories</th>
<th>BMI Male</th>
<th>BMI Female</th>
<th>BF Male</th>
<th>BF Female</th>
</tr>
</thead>
<tbody>
<tr>
<td>DAS28 (DAS28 score)</td>
<td>Remission (&lt;2.6)</td>
<td>27.2 (3.46)</td>
<td>27.2 (5.6)</td>
<td>26.5 (7.6)</td>
<td>39.5 (6.7)</td>
</tr>
<tr>
<td></td>
<td>Mild (2.7-3.2)</td>
<td>28 (4.3)</td>
<td>27.3 (4.6)</td>
<td>28 (6)</td>
<td>39.3 (6.6)</td>
</tr>
<tr>
<td></td>
<td>Moderate (3.3-5.1)</td>
<td>27.8 (4.5)</td>
<td>27 (5.3)</td>
<td>27.4 (6.8)</td>
<td>37.3 (7.7)</td>
</tr>
<tr>
<td></td>
<td>High (&gt;5.1)</td>
<td>25.3 (5.5)</td>
<td>27.3 (5.5)</td>
<td>26.1 (5.6)</td>
<td>37.7 (7.2)</td>
</tr>
<tr>
<td>ESR (mm/h)</td>
<td>Normal*</td>
<td>27.9 (4.4)</td>
<td>26.9 (4.8)</td>
<td>27.1 (7.2)</td>
<td>38.3 (6.3)</td>
</tr>
<tr>
<td></td>
<td>High</td>
<td>26.4 (4.6)</td>
<td>27.6 (6.1)</td>
<td>26.7 (5.9)</td>
<td>37.6 (8.9)</td>
</tr>
<tr>
<td>CRP (mg/L)</td>
<td>Low (&lt;3)</td>
<td>26.5 (2.4)</td>
<td>28.3 (6.2)</td>
<td>25.9 (5.4)</td>
<td>38.5 (8.7)</td>
</tr>
<tr>
<td></td>
<td>Normal (3-8)</td>
<td>27.8 (4.7)</td>
<td>26.5 (4.7)</td>
<td>26.7 (8)</td>
<td>37.6 (6.6)</td>
</tr>
<tr>
<td></td>
<td>High (&gt;8)</td>
<td>26.9 (4.6)</td>
<td>27.6 (5.7)</td>
<td>27.3 (5.8)</td>
<td>38.3 (7.9)</td>
</tr>
<tr>
<td>Disease Duration (years)</td>
<td>Early (&lt;3)</td>
<td>26.4 (5)</td>
<td>26.1 (5)</td>
<td>26.4 (7.9)</td>
<td>37.9 (8.3)</td>
</tr>
<tr>
<td></td>
<td>Established (3-10)</td>
<td>28.8 (4.1)</td>
<td>27.8 (5.7)</td>
<td>27.8 (6.3)</td>
<td>38.2 (7.5)</td>
</tr>
<tr>
<td></td>
<td>Longstanding (&gt;10)</td>
<td>26.8 (4.4)</td>
<td>27.1 (5.1)</td>
<td>27.7 (5.7)</td>
<td>38.8 (6.7)</td>
</tr>
<tr>
<td>Rheumatoid Factor</td>
<td>Positive</td>
<td>26.6 (3.6)</td>
<td>27.2 (5.7)</td>
<td>25.1 (6.7)</td>
<td>38.3 (7.3)</td>
</tr>
<tr>
<td></td>
<td>Negative</td>
<td>27.5 (5)</td>
<td>27.1 (5.1)</td>
<td>27.7 (6.4)</td>
<td>37.9 (7.3)</td>
</tr>
<tr>
<td>Corticosteroid Administration</td>
<td>Yes</td>
<td>27.1 (4.4)</td>
<td>27.3 (5.3)</td>
<td>26.2 (6)</td>
<td>38.1 (7.4)</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>24.5 (4.9)</td>
<td>26.7 (5.3)</td>
<td>27.8 (7.3)</td>
<td>37.7 (7)</td>
</tr>
</tbody>
</table>

For all differences between groups: p>0.05
* Normal ESR: < 50 years: male <15, female <20
> 50 years: male <20, female <30

Between the different disease groups, one-way ANOVA revealed significant differences in BMI (p<0.05) and BF (p<0.001; Table 1): RA males had higher BMI and BF (including trunkal fat) than HC males, and RA females had higher BF than HC females, even though their BMI did not differ significantly. ANCOVA revealed that BMI differences between the groups were mainly due to the significant effect of the covariate age (F1,294 = 5.10, p <0.05) and not due to disease (F2,294 = 1.00, p >0.05), gender (F1,294 = 0.59, p >0.05), or their interactions.

ANOVA also revealed that RA and OA patients exhibited lower BMI levels than their HC for a given BF. However, differences were only significant for the RA patients [RA:-1.826 kg/m² (p<0.001); OA: -0.352 kg/m² (p>0.05)]. BMI was significantly (p<0.001) predicted by age, disease, gender and BF (R² = 0.58).

When BF was adopted as the dependent variable, ANCOVA identified significant differences between disease groups (F2,293 = 18.70, p<0.001) and gender (F1,293 = 380.90, p<0.001) together with a significant covariate, age (F1,293 = 22.43, p<0.001). The contribution of BMI as a covariate in this analysis was also significant (F1,293 = 370.74, p<0.001). For a given BMI, RA patients exhibited significantly increased levels of BF (4.273, p<0.001) compared to healthy controls. The difference for OA patients was non-significant (1.648, p>0.05). The variation of BF was predicted by age, gender, BMI, and disease type (R² = 0.769, p<0.001). This was only very slightly improved (for RA) by the addition of RA disease duration (F1,293 = 0.340, p>0.05) in the equation (from 76.9% to 77.1%), so we did not include this variable in the final model. The predictive model obtained from this analysis is:
BF = Disease Status + Gender - 0.719 + 0.108 x Age + 1.059 x BMI

<table>
<thead>
<tr>
<th>Disease status</th>
<th>Gender</th>
</tr>
</thead>
<tbody>
<tr>
<td>RA = 4.273</td>
<td>Male = -11.294</td>
</tr>
<tr>
<td>OA = 1.648</td>
<td>Female = 0</td>
</tr>
<tr>
<td>HC = 0</td>
<td></td>
</tr>
</tbody>
</table>

Validation group:
To establish external validity of our predictive model, we assessed its agreement with the measured BF in 342 patients with RA. Preliminary analyses for LIM$_{AG}$ revealed no heteroscedasticity, thus the LIM$_{AG}$ can be reported as absolute measurements.[31] Our analyses suggested that the bias of our prediction is 0.4 (i.e., our model over-predicts BF by 0.4) with a standard error of 3.2 (95%LIM$_{AG}$ = 6.17, coefficient of variation = 8.9; Figure 1). The difference is statistically significant ($t = 2.3$, $p < 0.05$), but the coefficient variation (CV = 8.9) is within acceptable limits.

---Figure 1 about here---

RA-specific BMI cut-off levels:
The fact that patients with RA exhibited increased BF values for a given BMI compared to HC, suggested that BMI cut-off points in the RA population would be more appropriate if they were reduced by approximately 2 kg/m$^2$ (to 23 and 28 kg/m$^2$ for overweight and obesity respectively). We therefore compared the proportions of subjects in each group that would be correctly classified as overweight or obese using the widely accepted BMI cut-offs of 25 and 30 kg/m$^2$ vs. the proposed (for RA) 23 and 28 kg/m$^2$ vs. the age- and gender specific cut-off points of measured BF. This analysis showed that 9% of male and 15% of female RA patients would be misclassified as of normal weight based on traditional BMI cut-offs. Such misclassification was not a problem either for OA or HC, where if anything, BMI overestimated BF. Application of the proposed RA-specific BMI cut-offs of 23 and 28 kg/m$^2$ corrected this misclassification (Figure 2a). A modified, RA-specific BMI chart for the classification of patients with RA into underweight, normal, overweight and obese categories was developed and is provided in Figure 2b.

---Figure 2 about here---

Discussion
The validity of BMI as an acceptable measure of overweight or obesity, and as an accurate reflection of body fat (BF) content, has been repeatedly questioned and the need for population-specific BMI cut-off points has been highlighted.[7, 10-14] Ideally, individualized assessment of BF should be pursued in the clinical setting, as BF percentage is a more reliable measure of fatness than BMI, at least in the general population.[32] Indeed, our data indicate that only 58% of the variance in BMI can be predicted, as opposed to 77% in BF. BF in vivo can be determined via a number of methods such as underwater weighing, dual-energy x-ray absorptiometry, total body water, total body nitrogen, ⁴⁰K whole body counting, and urinary creatinine excretion.[33-35] BF can also be estimated from the thickness of partial subcutaneous fat, near-infrared rays, and ultrasound.[36] However, none of these methods can
be practically used in the routine clinical setting as they require sophisticated apparatus and specialised personnel.[34]

In recent years, a bioelectrical impedance method for the estimation of BF in different populations has become popular and widely recommended, as it is reliable, objective, practical, relatively inexpensive, and does not require highly trained personnel.[33, 34] The validity of this method has been confirmed in various studies.[33, 37-40] Devices with eight tactile electrodes using single frequency electrical current, similar to the one used in this study, generate highly reproducible measurements of total BF and segmental fat distribution. [41] Their correlation with the “gold standards” of dual-energy x-ray absorptiometry and hydrostatic weighing is 0.90 and 0.80 respectively, with a standard error of around 3.0, producing a co-efficient of variation of <10%. [33] This suggests that bioelectrical impedance measurements (especially when using eight electrodes) are valid and suitable for body composition studies.[33, 40, 41] Patients are usually happy to undergo such a measurement due to its simplicity and similarity to normal weighing.

In the absence of the necessary equipment or expertise, the predictive model presented here can be used to easily calculate BF of RA patients from BMI. The cross-validation of this predictive model in patients with RA is reassuring. Even though there was a statistically significant difference between the measured and the predicted BF, closer examination of the means indicates that this difference is at a level of less than 0.5% of BF with a co-efficient of variation of < 10%. The statistical significance of such a small difference can be attributed to the very large number of the validation group and is clinically not significant. However, the parts of the equation referring to OA patients and healthy individuals need further prospective validation in sufficiently large samples of the relevant populations.

BMI remains the most commonly used indicator of body fatness in the clinical setting, and the cut-off points of 25 and 30kg/m² (for overweight and obesity, respectively) used for the general population are also routinely applied in RA patients. This study shows that application of these BMI cut-off points misclassified 9% of male and 15% of female RA patients in terms of actual body fatness. For a given BMI, RA patients exhibited an average 4.3% increase in BF compared to healthy controls. In contrast, for the same level of BF, RA patients had BMI values almost 2 kg/m² lower than those of healthy controls. We propose that BMI cut-off points in the RA population should be lowered to 23kg/m² (from 25kg/m²) for overweight, and 28kg/m² (from 30kg/m²) for obesity. The lowest limit for normal BMI (i.e. 18.5kg/m²) should remain unaltered, as low BMI levels have been related to increased cardiovascular risk in patients with RA.[42, 43] We also provide a chart for the classification of RA patients in normal, overweight and obese categories according to these BMI cut-offs, for use in the routine clinical setting (Figure 2b).

The most likely explanation for the BMI and BF differences observed in RA is rheumatoid cachexia associated with the chronic inflammatory response, given that such differences were not as prominent in OA. RA patients experience accelerated involuntary loss of fat-free mass, predominantly in the skeletal muscle, in excess of what is normally expected due to the aging process.[44] Although the underlying mechanisms for rheumatoid cachexia remain unknown, possible contributing factors include the overproduction of inflammatory cytokines such as Tumour Necrosis Factor α and Interleukin-1 β.[44, 45] Our sub-analyses within the RA population revealed that neither BMI nor BF were associated with current clinical or serological disease activity, seropositivity for rheumatoid factor (which tends to associate with more severe disease) or corticosteroid administration. This is not totally
surprising as disease activity may vary within small periods of time, depending on medication and the disease itself, whereas changes in body composition are longer-term processes. On the other hand, disease duration appeared to be of some importance. It is possible that most alterations in body composition of RA patients occur in the first few years of the disease, as it has previously been reported,[22] irrespective of disease characteristics or medical treatment.

The results of the present study are reminiscent of the observations made for Asian populations, which have significantly higher CVD risk than Caucasians: BF in Asians has been found to be 3-5% higher than that of Caucasians with similar BMI, whereas BMI was 3-4 kg/m² lower than that of Caucasians with similar BF.[33] Differences in body build (trunk-to-leg-length ratio and slenderness) and in muscularity have been suggested as possible explanations for these discrepancies. As a result, new cut-off points for Asian populations have been set at 23 kg/m² and 27 kg/m² for overweight and obesity, respectively,[10] and have been shown to be more sensitive in identifying Asians at increased risk for CVD.[46]

In our participants, lowered BMI cut-off points would reflect an average reduction of 5-6 kg, or 8%, in the ideal weight (the weight one should have in order to be below the BMI cut-off for overweight). Such reductions in body weight are likely to lead to physiological benefits in the cardiovascular system: in the general population, even a 5% reduction of body weight is known to affect favourably most classical CVD risk factors.[47, 48]

The reduced BMI cut-off points for RA suggested here may be of significance both for the management of individual patients and for further research into the cardiovascular morbidity and mortality of RA. In the clinical arena, the reduction of these thresholds would identify an additional 10-15% of people with RA as overweight or obese, and may trigger closer scrutiny for other CVD risk factors and appropriate intervention, if necessary. Moreover, obesity, defined by the BMI, is one of the WHO criteria for the metabolic syndrome [47]. Aggressive identification and reduction of classical CVD risk factors in patients with RA is an obvious strategy for reducing the increased cardiovascular mortality of this disease.[20] From the research perspective, the new thresholds may trigger re-analysis of previously published cohorts or further analysis of prospective cohorts as to the importance of body fat as a predictor of CVD in RA and its association with other individual risk factors.

We conclude that, in the clinical setting, body fatness of RA patients should be evaluated based on the BMI cut-off points of 23 kg/m² for overweight and 28 kg/m² for obesity. In the absence of specialised equipment, if necessary, BF of patients with RA can be estimated from BMI using the equation provided.
List of Abbreviations
BF: Body Fat
BMI: Body Mass Index
CVD: Cardiovascular Disease
CHD: Coronary Heart Disease
MI: myocardial infarction
HC: Healthy Controls
OA: Osteoarthritis
RA: Rheumatoid Arthritis
ANOVA: Analysis of Variance
ANCOVA: Analysis of Co-variance
LIM_{AG}: Limits of Agreement

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Competing interests
The authors have no competing interest to declare

Authors' consent to publication
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Figure Legends

**Figure 1: Agreement between Predicted and Measured Fat in patients with RA.** Body fat was measured by bioelectrical impedance using a Tanita BC-418 MA Segmental Body Composition Analyzer. Predicted fat was assessed using the formula: \( BF = 4.273 + \text{Gender} - 0.719 \times \text{Age} + 1.059 \times \text{BMI} \). 95% Limits of Agreement were 6.17 with a coefficient of variation of 8.9.

**Figure 2a: Classification of male (top graph) and female (bottom graph) participants into obese, overweight, normal and underweight groups according to currently accepted BMI cut-off points (BMI), body fat content (BF) and RA-specific BMI cut-off points (RA-BMI).** Accepting BF as the most accurate assessment of body fatness, currently accepted BMI cut-off points misclassify a significant proportion of both males and females with RA (notice the difference in the respective bars). This misclassification is corrected when the proposed RA-specific BMI cut-off points are applied.

RA: patients with rheumatoid arthritis
OA: patients with osteoarthritis
HC: healthy controls
BMI: classification according to existing body mass index (BMI) cut-off points of 25kg/m\(^2\) for overweight and 30kg/m\(^2\) for obesity
BF: classification according to age and gender specific cut-off points for body fat percentage
RA-BMI: classification according to the proposed RA-specific BMI cut-off points of 23kg/m\(^2\) for overweight and 28kg/m\(^2\) for obesity

**Figure 2b: BMI chart developed specifically for patients with RA.** Values were calculated using the formula: \( BMI = \frac{\text{weight (in kg)}}{\text{height}^2 \ (\text{in m})} \) for the rheumatoid arthritis-specific BMI levels identified in the present study [23kg/m\(^2\) for overweight, 28kg/m\(^2\) for obesity]. The generally accepted lower threshold for normal BMI [18.5 kg/m\(^2\)] was not altered.
References


Male Participants

Female Participants
This graph illustrates the relationship between weight and height using BMI (Body Mass Index) categories. The BMI categories are:

- **Underweight**: BMI < 18.5
- **Normal**: BMI 18.5 - 22.9
- **Overweight**: BMI 23 - 28
- **Obese**: BMI > 28

The graph uses different shades to represent these categories: lighter shades for normal, medium shades for overweight, and darker shades for obese and underweight. The axes are labeled as follows:

- **Height**
- **Weight**

The grid lines help in visualizing the BMI categories for various weight and height combinations.