Fat-free weight prediction in morbidly obese females

Purpose: Precise estimation of creatinine clearance in obese individuals relies on the appropriate assessment of lean body weight (LBW). Anthropometric methods of predicting LBW have not been validated in morbidly obese populations.

Patients and methods: Using an existing dataset of anthropometric data for a female cohort with morbid obesity who had undergone measured FFW with dual energy absorptiometry, we evaluated the performance of five previously reported estimating equations for the prediction of LBW. Linear regression was used to derive a new LBW prediction formula and was then compared with the other formulae.

Results: Seventy females (mean [standard deviation] age, weight, and body mass index 43.0 [11.0] years, 128.1 [13.8] kg, and 48.3 [4.8] kg/m², respectively) were identified. LBW as estimated by the method of Garrow and Webster correlated well (r = 0.87) with measured mass while demonstrating the highest accuracy, best precision, and smallest bias (93%, 2.1 kg, and 2.9 kg, respectively; P < 0.0001 for all comparisons). The derived formula further improved bias, precision, and accuracy.

Conclusion: Among females with morbid obesity, most previously reported estimating equations for LBW predicted FFW poorly. These findings have important clinical implications for the assessment of kidney function and for safe and effective drug dosing.

Keywords: body composition, body weight, creatinine clearance, kidney function, lean body mass, obesity, pharmacokinetics

Introduction
At extremes of weight, misapplication of commonly used clinical tools developed in normal-weight populations may have serious adverse consequences. There is a need for clinically useful and accurate methods of estimating renal function among patients with morbid obesity (body mass index [BMI] ≥ 40 kg/m²) for the purpose of safe and effective drug dosing, monitoring of kidney function longitudinally, and effective risk counseling. In this population, estimation of renal function is problematic, as commonly used formulae perform poorly at extremes of weight. Estimations of creatinine clearance using adjusted body weight (ABW) may represent a better assessment of kidney function than estimations of glomerular filtration rate using the Modification of Diet in Renal Disease estimating formula among patients with morbid obesity. Recently, several investigators have reported improved renal function estimates in obese (BMI ≥ 30 kg/m²) individuals using the Cockcroft-Gault creatinine clearance formula with adjustment for lean body weight (LBW) either by direct fat-free weight (FFW) measurement or by way of an estimating formula.
Whether this applies to patients with morbid obesity is not clear, as the relationship between BMI and lean weight is not linear.9

Although direct measurement of FFW can be performed in several ways – to include dual energy absorptiometry (DEXA) – such methods are not practical for routine clinical use, due to complexity and cost.10–12 It is expected that LBW will overestimate fat-free mass (FFM) by 3%–5%, given that measurement of the latter does not include the normal adipose tissue surrounding muscle and nerves in a lean individual. However, use of LBW could serve as a clinically useful surrogate for FFM for calculation of clearances of endogenous creatinine, as this substance distributes mainly in muscle and interstitial and vascular spaces.5,13 Various formulae for estimating LBW have been reported,14–21 but none has been developed or validated for specific use in a population of patients with morbid obesity.

This study evaluates the ability of previously reported alternative body size descriptor (ABSD) estimating formulae to predict measured FFW by DEXA in a cohort of female, morbidly obese patients enrolled in a weight management program at a tertiary medical center in Central Pennsylvania. We did not limit the identification of previously reported formulae to those estimating LBW only, as the primary goal of this study was to identify a clinically useful estimating tool that could be employed in the Cockcroft and Gault creatinine clearance equation. As a secondary goal, we wished to improve upon the estimation of LBW in our own population through the development of a novel LBW estimating equation based on anthropometric and clinical data from our study population.

Material and methods

Subjects

The study was reviewed and approved under “exempt” status by the Geisinger Medical Center Institutional Review Board in May 2010. The study uses existing data from a cohort of 70 adult (aged ≥18 years) morbidly obese (BMI of ≥40.0 kg/m²) female patients who participated in a feasibility project evaluating the use of preoperative DEXA body composition scanning between May 2005 and January 2006 while undergoing evaluation in a comprehensive weight management clinic prior to bariatric surgery at Geisinger Medical Center.22 The database that served as the source for this study includes demographic, clinical, outcomes, and claims information, including detailed anthropometric measurements, and has been previously validated.23–27

Anthropometric measurements

Body composition measurements (fat mass, lean mass, and bone mass) were made with a Hologic Series 4500 W Fan Beam X-ray Bone densitometer (Hologic, Inc Bedford, MA). Measurements were performed on subjects following the removal of all metal accessories. Scanners were calibrated daily prior to patient use, using standard phantom measurements. Patients were weighed at each clinic visit in the weight management clinic wearing hospital gowns. All weights are expressed in kilograms (kg). Height (Ht) in centimeters (cm) was measured at the initial clinic visit and not repeated subsequently. From the above measurements, BMI = Wt/(Ht × 100)² and ideal body weight (IBW) (IBW = 45.4 + 0.89 × (Ht – 152.4))23 were calculated. LBW in the context of this study refers to the weight of lean tissue, including the normal amount of fat contained in a lean individual (approximately 3% in males and 5% in females),17 whereas FFW refers to the weight of lean tissue stripped of all fat, including the fat content of nervous system, cell membranes, and bone marrow.

Selection of LBW estimating equations and assessment of performance

In order to identify previously reported estimations for LBW, we performed a literature search using Ovid MEDLINE® (1950 to July Week 2, 2010) using the following search terms: body composition, morbid obesity, obesity, kidney function tests, glomerular filtration rate, lean body weight estimation, creatinine clearance, and pharmacokinetics. References of pertinent articles were also searched for studies describing LBW estimating equations for use in obese subjects. No method of estimating LBW exclusively in morbidly obese subjects was identified; five ABSD equations for the nonmorbidly obese populations were identified and assessed for the purposes of this analysis (Table 1). For the ABSD formula initially proposed by Karkeck,15 two versions were used in this study, one using the original reported equation with a correction factor of 0.25, and a second with a correction factor of 0.5, as the latter reportedly improves caloric expenditure prediction and drug clearance in obese individuals when compared with the original equation.16,29 In addition, a sixth equation developed in our patient population was used to estimate FFW (FFW<sub>GEISINGER</sub>). Thus, each of our subjects had one measured FFW (FFW<sub>DEXA</sub>) and six LBW estimations, each using a different ABSD.

To compare the ability of each individual ABSD to predict FFW<sub>DEXA</sub>, we calculated correlation, bias, precision, and accuracy.
Fat-free weight prediction

Table 1  Previously reported LBW estimating equations

<table>
<thead>
<tr>
<th>Author</th>
<th>Alternative body size descriptor</th>
<th>BMI, kg/m² (range)</th>
<th>Weight, kg mean ± SD (range)</th>
<th>Formulaa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Garrow et al14</td>
<td>FFM [garrow]</td>
<td>16.7–50.1</td>
<td>Variable, not reported in all studies</td>
<td>0.287 × TBW + 9.74 × Ht</td>
</tr>
<tr>
<td>Karkeck et al15</td>
<td>ABW0.25</td>
<td>NA</td>
<td>Variable, not reported in all studies</td>
<td>IBW + CF × (TBW-IBW)</td>
</tr>
<tr>
<td>Cutts et al16</td>
<td>ABW0.5</td>
<td>Variable, not reported in all studies</td>
<td>cF = 0.25 or 0.5</td>
<td>[9270 × TBW]/[8780 + 2.44 × BMI]</td>
</tr>
<tr>
<td>Glynn et al29</td>
<td>IBW + cF × (TBW-IBW)</td>
<td>17 ± 19.7 (44.7–123.9)</td>
<td>Variable, not reported in all studies</td>
<td>71 ± 18.7 (44.7–123.9)</td>
</tr>
<tr>
<td>Janmahasatian et al18</td>
<td>LBW [Janmahasatian]</td>
<td>17.1–69.6</td>
<td>DeXA NR</td>
<td>[9270 × TBW]/[8780 + 244 × BMI]</td>
</tr>
<tr>
<td>Hume and Weyers19</td>
<td>LBW [hume]</td>
<td>NR</td>
<td>Antipyrine space</td>
<td>1.07 × TBW - 0.0148 × BMI</td>
</tr>
</tbody>
</table>

Note: For each equation, only the formula used for females is shown. ABB, adjusted body weight (kg); BMI, body mass index (kg/m²); CF, correction factor; DeXA, dual energy absorptiometry; FFM, fat-free mass (kg); Ht, height (cm); IBW, ideal body weight (kg); LBW, lean body weight (kg); TBW, total body weight (kg).

Continuous data are reported as means and standard deviations (SDs), whereas categorical variables are expressed as frequencies. When appropriate, the distributions of each continuous variable were examined for a symmetrical, bell-shaped distribution (ie, the normal distribution) and homoscedasticity (ie, equality of variance). Bias, precision, accuracy, and the Pearson correlation coefficient (r) were calculated for each estimating equation using FFW [DEXA] as the gold standard. Bias was defined as the absolute value of the difference between the estimated LBW and FFW [DEXA] (also known as absolute bias). Precision was defined as the SD of the absolute bias. Accuracy was calculated as the percentage of patients who had an estimated FFM within 10% of the FFW [DEXA].

Differences in bias and accuracy were compared between ABSDs using a paired t-test and McNemar's test. This was done by comparing the ABS formula that performed the best (ie, with the least bias or with the highest accuracy) against all other ABS formulae in a pairwise fashion. Differences in precision among the various ABSDs were not tested.

To determine whether estimating formula performance was similar for different levels of obesity, the bias, precision, accuracy, and correlation coefficient were also calculated after stratifying by BMI subcategories (40.0–44.9 kg/m², 45.0–49.9 kg/m², ≥50.0 kg/m²).

Agreement between measured and estimated LBW values was assessed using the method of Bland and Altman,30 in which the difference between estimated LBW and FFW [DEXA] is plotted against the FFW [DEXA]. A positive difference suggests an overestimation of the formula, whereas a negative difference suggests an underestimation. The limits of agreement were calculated as the bias plus two times the precision.

In exploratory analysis, linear regression was used to determine whether a new ABS formula could be derived that outperformed existing ABSDs. BMI, total body weight (TBW), Ht, and IBW were considered for inclusion in the regression model and the minimal subset of measures that maximized the model r-squared was retained.

All statistical analyses were performed using SAS® software (version 9.2, Cary, NC).

Results

Seventy morbidly obese female patients underwent body composition DEXA. Patient characteristics and major comorbid conditions are presented in Table 2. All patients were Caucasian, with a mean age of 43 years. Thirty-six percent
Table 2 Characteristics of population of morbidly obese female patients undergoing DEXA (n = 70)

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (y)</td>
<td>43 (11)</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>128.1 (13.8)</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>162.8 (5.4)</td>
</tr>
<tr>
<td>Serum creatinine (mg/dL)</td>
<td>0.8 (0.2)</td>
</tr>
<tr>
<td>BMI (kg/m²)</td>
<td>48.3 (4.8)</td>
</tr>
<tr>
<td>Waist circumference (cm)</td>
<td>128.1 (10.2)</td>
</tr>
<tr>
<td>FFW_{DEXA} (kg)</td>
<td>63.4 (6.9)</td>
</tr>
</tbody>
</table>

Comorbid conditions

- Hypertension: 25 (36)
- Hyperlipidemia: 22 (31)
- Diabetes: 15 (21)
- Obstructive sleep apnea: 6 (9)
- Cerebrovascular disease: 2 (3)
- Atherosclerotic cardiovascular disease: 1 (1)
- Chronic renal disease: 1 (1)

Note: Data available for 53 patients.

Abbreviations: BMI, body mass index (kg/m²); DEXA, dual energy absorptiometry; FFW, fat-free weight (kg).

had hypertension, one-third had hyperlipidemia, and 21% were diabetics. The majority (94%) of patients had preserved renal function (estimated glomerular filtration rate) calculated using the four-variable Modification of Diet in Renal Disease equation (>$60 \text{ mL/min/1.73 m}^2$).

The remaining four patients had estimated glomerular filtration rates 42.9, 43.7, 51.2, and 55.4 \text{ mL/min/1.73 m}^2, respectively.

Results of individual formula performance are presented for the overall population in Table 3 and for subcategories of BMI in Table 4. FFM_{Garrow}^\text{a} the ABW formulae, and LBW_{Janmahasatian}^\text{b} correlated well with FFW_{DEXA} (r > 0.80). In contrast, LBW_{Hume} and LBW_{Moore}^\text{c} correlated poorly.

Table 3 Correlation, bias, and accuracy of reported estimating equations for LBW compared with DEXA-derived lean body weight for female patients with morbid obesity (BMI > 40 kg/m²)

<table>
<thead>
<tr>
<th>Formula</th>
<th>All patients (n = 70)</th>
<th>Bias $^a$</th>
<th>Precision $^b$</th>
<th>Accuracy $^c$</th>
<th>r</th>
</tr>
</thead>
<tbody>
<tr>
<td>FFM_{Garrow}</td>
<td>2.9</td>
<td>2.1</td>
<td>93%</td>
<td>0.87</td>
<td></td>
</tr>
<tr>
<td>ABW_{J325}</td>
<td>9.7</td>
<td>3.8</td>
<td>17%</td>
<td>0.84</td>
<td></td>
</tr>
<tr>
<td>ABW_{J51}</td>
<td>28.0</td>
<td>4.1</td>
<td>0%</td>
<td>0.87</td>
<td></td>
</tr>
<tr>
<td>LBW_{Janmahasatian}</td>
<td>6.0</td>
<td>3.6</td>
<td>54%</td>
<td>0.85</td>
<td></td>
</tr>
<tr>
<td>LBW_{Hume}</td>
<td>18.7</td>
<td>10.0</td>
<td>4%</td>
<td>-0.15</td>
<td></td>
</tr>
<tr>
<td>FFW_{DeBiase}</td>
<td>2.8</td>
<td>1.8</td>
<td>97%</td>
<td>0.87</td>
<td></td>
</tr>
</tbody>
</table>

Notes: $^a$Bias = mean absolute bias (mean absolute value of difference between formula- and DEXA-derived LBW for each individual); $^b$Precision = standard deviation of mean absolute bias; $^c$Accuracy = percentage of estimations within 10% of the FFW_{DEXA}.

Abbreviations: ABW, adjusted body weight (kg); BMI, body mass index (kg/m²); DEXA, dual energy absorptiometry; FFM, fat-free mass (kg); FFW, fat-free weight (kg); LBW, lean body weight (kg).

Beyond correlation, the performance of equations other than FFM_{Garrow}^a was quite poor. The overall mean bias was smallest for the FFM_{Garrow}^a equation ($P < 0.0001$ for pairwise comparisons). This equation also had the highest accuracy ($P < 0.0001$ for pairwise comparisons) and best precision. Bias, precision, and accuracy were lower for the BMI-based equations (LBW_{Janmahasatian}^b and LBW_{Hume}). LBW_{Janmahasatian}^b did maintain its level of accuracy in BMI categories between 40.0–44.9 kg/m² and 45.0–49.9 kg/m² but not in subjects with BMI ≥ 50.0 kg/m².

A Bland–Altman plot for the FFM_{Garrow}^a equation is shown in Figure 1. There is an inverse relationship between bias and incremental measured FFW ($r = -0.73$).

In regression analysis performed to identify an improved prediction formula, the combination of body composition measures that optimized the prediction of FFW was TBW and Ht. The resulting equation was FFW_{Geisinger} = -11.41 + (0.354 \times \text{TBW}) + (11.06 \times \text{Ht})$. Though not significantly different from FFM_{Garrow}^a, this equation had the least bias, the best precision, and the highest accuracy.

Discussion

LBW estimation in morbidly obese individuals presents unique challenges. In our exclusively Caucasian female population, not all previously reported ABSD estimating formulae performed well for the determination of FFW as measured by DEXA body composition testing.

One estimating formula in particular, FFM_{Garrow}^a outperformed others. This regression equation was developed in a predominantly (81%) female population with mean (SD) BMI 27.2 (7.8) kg/m² against a gold standard consisting of the average body fat measurement obtained by three different methods (tissue density, total body water, using a tritiated water dilution method, and total body potassium using the $^{40}$K isotope gamma spectrometry method). It is interesting that each of these methods measures a slightly different lean weight, making the combination of the three methods a potentially robust assessment of the true lean mass. In our cohort, this equation demonstrated the most optimal performance characteristics for the overall population with morbid obesity as well as within each subcategory of BMI.

Other ABSDs performed suboptimally, though several formulae incorporating BMI (LBW_{Janmahasatian}^b and LBW_{Hume}) performed modestly well across some of the study metrics. The formula developed by Janmahasatian et al\textsuperscript{18} was developed using FFW_{DEXA} as a gold standard in a gender-balanced Australian population that was predominantly (70%) overweight (BMI ≥ 25). This equation has subsequently been
shown to eliminate the bias of Cockcroft–Gault against both the creatinine and the inulin clearances when used instead of measured weight.7,8 Though this equation estimates inulin and creatinine clearance fairly well in obese populations,7,32 it did not estimate measured FFW as well as the FFM[garrow] in our population of morbidly obese females. Consistent with the observation of declining performance as BMI increases for this formula, our results indicate that at incrementally higher BMI, LBW[Janmahasatian] predicts FFW poorly. Thus, its applicability to individuals with morbid obesity may be limited.

Weight-based estimating equations had the poorest performance in our study. The ABW formulae with various correction factors – although commonly used in nutrition practice for various resting energy calculations – are based on the assumption that excess weight in obesity is linearly increased from the IBW, an assumption that is physiologically flawed. Amato et al33 evaluated the accuracy and precision of using ABW (with a correction factor of 0.5) in 113 morbidly obese (mean ± SD BMI 52 ± 9 kg/m²) patients in a surgical intensive care unit and observed significant bias and poor precision of this body size descriptor for the prediction of

![Figure 1](https://www.dovepress.com/)

**Figure 1** Bland–Altman plot representing absolute differences between lean weight estimations using FFM[garrow] equation vs measured FFW[DEXA] by BMI category.

**Abbreviations:** BMI, body mass index (kg/m²); DEXA, dual energy absorptiometry; FFM, fat-free mass (kg); FFW, fat-free weight (kg).
measured resting energy expenditure. Although a direct relationship between calorimetric testing and measured FFW cannot be assumed, our results are congruent with those of Amato et al.

The clinical significance of our study findings is illustrated by comparing the Cockcroft and Gault-estimated creatinine clearance using LBW obtained from each of the reported and derived formulae with the clearance measurement calculated using FFM from DEXA body composition scanning (Table 5). As is readily apparent, the range of calculated creatinine clearance using the various formulae is alarmingly broad. Finding and using the right lean weight estimation formula for the morbidly obese population is essential for safe health care delivery.

We derived a formula that improved the performance of FFM prediction and best approximated the estimated creatinine clearance calculation, at least among our own population of female patients with morbid obesity. This prediction formula should be tested and validated in other populations prior to clinical application.

We acknowledge that not all formulae used in our analysis were originally developed to measure the same body compartment, especially formulae such as ABW with various CF. Although this can explain the bias, it cannot account for the poor precision and accuracy in predicting FFW(DEXA). Our intention was to evaluate previously reported formulae developed to estimate alternative body weight descriptors for their potential to predict FFW in a population of morbidly obese females. The goal was not to assess the performance of these equations for their original intent. Also, our study population consists exclusively of Caucasian females, and the findings of this analysis should not be extrapolated to other patient populations without further analysis.

Table 5 Estimated creatinine clearance using the method of Cockcroft and Gault and LBW derived from DEXA and six alternative body size descriptor equations for a population of extremely obese females (n = 70)

<table>
<thead>
<tr>
<th></th>
<th>Mean (SD) estimated creatinine clearance, mL/min</th>
</tr>
</thead>
<tbody>
<tr>
<td>FFW(DEXA)</td>
<td>98.5 (27.9)</td>
</tr>
<tr>
<td>FFM(Garrow)</td>
<td>97.1 (26.5)</td>
</tr>
<tr>
<td>ABW(ABW25)</td>
<td>113.1 (30.7)</td>
</tr>
<tr>
<td>ABW(ABW50)</td>
<td>141.9 (39.4)</td>
</tr>
<tr>
<td>LBWmanaged</td>
<td>89.3 (23.7)</td>
</tr>
<tr>
<td>LBW(Heimburger)</td>
<td>68.1 (19.1)</td>
</tr>
<tr>
<td>FFW(USENAC)</td>
<td>98.3 (27.5)</td>
</tr>
</tbody>
</table>

Abbreviations: ABW, adjusted body weight (kg); DEXA, dual energy absorptiometry; FFM, fat-free mass (kg); FFW, fat-free weight (kg); LBW, lean body weight (kg).

Conclusion

In summary, our study describes the performance of several formulae for the estimation of FFW measured by DEXA body composition scanning in a cohort of morbidly obese females. The FFM(Garrow) equation provides the best estimation of the measured FFM. The findings have important potential clinical implications as they relate to effective and safe medication dosing and optimal nutritional care, both for the hospitalized patient and for the morbidly obese patient managed in the outpatient setting.

Acknowledgments

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Disclosure

The authors report no conflicts of interest in this work.

References