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Comparison of body fat calculations by sex and puberty status in obese schoolchildren using two and four compartment body composition models

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Abstract

Introduction: Determine body composition changes in adiposity can assess an individual.

Objective: The objective of this study was to evaluate total body fat percentage based on two and four compartment models in obese Chilean school children, adjusting for differences in sex and puberty status.

Methods: Sixty-one obese school children (33 boys and 28 girls) between 8 and 13 years of age were evaluated. Two compartment measurements of body fat percentage considered isotopic dilution, plethysmography, radiographic absorptiometry and bioelectrical impedance; using the four compartment model as a benchmark.

Results: Each method explained between 43-87% of the variance in body fat percentage in Tanner stage I and II children and between 78-96% in Tanner stage III and V children. In both groups of children differences were significant for stage I, with the exception of plethysmography. High R² values were observed for girls in all Tanner stages. Each method explained between 34-92% of the variance in body fat percentage for girls in stages I and II and between 63-93% for stages III and V. In obese boys, R² values were high for stages I and V. In girls and boys in Tanner stage III and V, the smallest differences were observed for isotopic dilution, and DXA (dual-energy X-ray absorptiometry) scan for stages I and II.

Conclusions: For obese boys and girls, the two compartment model with isotopic dilution and DXA had the best precision and smallest differences in determining body fat percentage compared to the benchmark.

Palabras clave:

Resumen

Introducción: determinar la composición corporal permite valorar cambios en la adiposidad de un individuo.

Objetivo: el objetivo de este estudio fue evaluar la grasa corporal total basada en modelos de dos compartimentos (2C) y compararlos con el modelo de cuatro compartimentos (4C) en escolares chilenos obesos, considerando potenciales diferencias por sexo y desarrollo puberal.

Métodos: 61 escolares obesos (33 niños y 28 niñas), de entre 8 y 13 años. La medición de la grasa corporal por 2C consideró dilución isotópica, pletismografía, absorciometría radiográfica y bioimpedanciometría; utilizando como patrón de referencia el modelo de 4 compartimentos.

Resultados: cada método dio cuenta de 43-87% de la varianza para determinar el porcentaje de grasa corporal en niños en etapa I y II y 78-96% en etapa III y IV. En ambos grupos de niños, las pendientes difirieron significativamente en la etapa I, con la excepción de plethysmography. En niñas los valores de R² altos se observaron en todas las etapas del desarrollo puberal. Cada método dio cuenta de 34-92% de la varianza para determinar el porcentaje de grasa corporal en niñas en etapa I y II y 63-93% en etapa III y IV. En niños obesos, los valores de R² fueron altos, principalmente en el grupo de etapa III y IV. En niños y niñas de Tanner I y II, las menores diferencias con el patrón de referencia fueron con dilución isotópica; en cuanto a Tanner III y IV, las menores diferencias se obtuvieron con DXA.

Conclusiones: en ambos sexos, el modelo de dos compartimentos con dilución isotópica y DXA tuvo la mayor precisión y las menores diferencias para determinar la grasa corporal en los niños y adolescentes obesos, en comparación con el patrón de referencia.
INTRODUCTION

The prevalence of obesity in Chilean school children is 25.3%, with slightly higher levels among boys compared to girls (28.3% vs. 22.3%) (1). Increases in adiposity associated with childhood obesity is a risk factor for glucose intolerance, insulin resistance, dyslipidemia, non-alcoholic fatty liver disease, hypertension, heart attack, stroke and premature death (2-6). Although body mass index (BMI) is widely used as an index of body fat (BF), it is not a direct measurement of adiposity. BMI cannot distinguish between different types of body mass (e.g., fat mass, fat free mass, bone mass), thus the use of BMI can lead to errors in the estimation of BF, especially in the context of obesity. Changes in weight and height associated with normal growth are responsible for a 50% increase in BMI, which further complicates the interpretation of this index for children and adolescents. The increase in BMI during adolescence is primarily a result of an increase in fat free mass (7,8). BMI is a global indicator of nutritional status and does not distinguish between lean and fat mass (9). Thus, measuring fat mass would allow for the quantification of metabolic risk associated with an increase in obesity.

In the classic two compartment model of body composition, body weight is divided into fat mass and fat free mass. This model is used widely for clinical practice and nutritional follow-up. BODPOD uses the relationship between pressure and volume to calculate body volume and density (10) Isotopic dilution quantifies total body water, which can be used to predict fat free mass, as a proportion of water known in fat free mass by age and sex (11) DXA differentiates between fat and fat free mass based on the differential attenuation of x-rays (12). Bioelectrical impedance, BIA, is an indirect method to measure the total body water and fat free mass (13).

The most precise method, considered the gold standard, for determining body composition is the four compartment model. For this method, fat free mass is divided into water, minerals and proteins (10). Although multi-compartment models of body composition have better precision, few studies have used them to validate simpler methodology in obese children and adolescents (14). The current study aimed to determine the predictive capacity of the two compartment model of BF% (body fat percentage) compared to the four compartment model in a sample of obese Chilean school children, adjusting for possible differences by sex and pubertal stage.

METHODS

SUBJECTS

We worked with a sample of 61 obese children years (males = 33 and females = 27) between 8 and 13, from a school in the Macul neighborhood of Santiago, Chile. The school was chosen for convenience, given the proximity of the school to the place of measurements. Inclusion criteria included: BMI ≥ 95th percentile according to CDC-NCHS references (15), full-time attendance at an educational institution, parental consent and child assent. The exclusion criteria included: medical diagnosis of psychomotor dis-order, use of drugs that can alter body composition, performing physical activity, and/or biochemical parameters. This research was approved by the Ethics Committee of the University of Chile.

BIOLOGICAL AGE

Pubertal development was classified using Tanner staging, considering breast development in females and genital in males (16). Developmental stages were determined by visual inspection during a physical examination by a pediatrician.

ANTHROPOMETRY

Weight and height were assessed in the morning after an overnight fast. Children wore minimal clothing, standing in front of the scale, with feet together at the center of it, arms attached to the body, the head forming a straight line parallel to the floor to join the corner of the eye and the birth of the ear. An electronic balance (SECA® Model 767) was used with sensitivity of 10 grams for weight and Holtain stadiometer (SECA) with sensitivity 0.1 cm for height, both imported by Precision Hispana. Four skin folds (biceps, triceps, subscapular and suprailiac), with a Lange caliper millimeter (1 mm), were assessed in triplicate using the technique described by Lohman et al. (17).

ISOTOPIC DILUTION

Total body water was determined with deuterium dilution. The isotope (4 grams of deuterium oxide 99.8%) was administered orally according to body weight of the subject. The amount of body water was measured by determining the concentration of deuterium oxide, according to the Plateau method. This required that the subjects were in total fasting for a period of three hours, which corresponds to the period of equilibrium and minimizes changes in total body water content (11). After the fast, a saliva sample (2 mL, baseline) was taken. Subsequently, the deuterium dose and an additional 20 ml of tap water were given to ensure dose ingestion. After three hours, during which participants were not allowed to urinate, eat or drink anything additional, the second saliva sample (post dose) was taken and frozen at -20 °C. For analyzing the concentration of deuterium in saliva, the sample was thawed, equilibrated in hydrogen gas, adding 5% platinum on aluminum with time of three days to reach equilibrium. The deuterium/hydrogen ratio in the gas released was analyzed by mass spectrometry (Hydra, Europe Scientific, Crewe, Cheshire, United Kingdom).

PLETHYSMOGRAPHY

Volume and body density were measured with an air displacement plethysmograph (BODPOD, mod 2000, Life Measurement,
Inc., Concord, USA). Children were tested with underwear, without metal objects and a swimming cap to compress the hair. Later, children were weighed on a calibrated scale with an accuracy of 5 g. The system performs a pressure measurement with the chamber empty, then the equipment is calibrated using a 50 liter calibration cylinder, after which the subject is measured 2-3 times. Body size obtained by this method was used for the 4C (four compartment) equation.

**DUAL-ENERGY X-RAY ABSORPTIOMETRY**

Bone mineral density was estimated using dual energy x-ray absorptiometry using Lunar Prodigy Gsc DPX-NT (Lunar Radiology, WI, USA) technology, which assesses the entire body in a five-minute sweep. Children were placed supine wearing a light robe.

**BIOELECTRICAL IMPEDANCE**

Bioelectrical impedance was measured using Tanita BC-418MA, eight-electrode, hand-to-foot system, manufactured by the Tanita Corporation (Tokyo, Japan). Measurements were collected according to manufacturer’s guidelines using a 50 kHz frequency. Height, sex and age were entered manually, whereas weight was recorded automatically. Measurements were taken in the morning after limited physical exercise and empty bladders.

**4C MODEL**

The 4C model divides the body in fat, water, protein, and minerals (18-20). The ability of the model to adjust core body mineral mass can result in a more accurate estimate of hydration and lean mass density compared with the 3C model. The 4C model is considered the “gold standard” because it takes into account the variability of its components. The equation has been previously validated in children of the same group (21).

The 4C equation used was a follows:

\[ BF (kg) = [(2,747*BV) – (0,710*TBW)] + [(1,460*BMC) – (2,050*W)] \]

\[ BV = \text{body volume in liters (plethysmography)}, \ TBW = \text{total body water in liters (isotope dilution)}, \ BMC = \text{bone mineral content in kg}, \ (DXA) \text{and W = body weight (kg)}. \]

**STATISTICAL ANALYSIS**

Descriptive statistics were used: minimum, maximum and frequency tables. Continuous variables were analyzed with the goodness of fit test of Shapiro Wilk test of homogeneity of variance. For variables that met normality assumptions we reported the average and standard deviation, otherwise the median and interquartile range were shown. Differences by gender and pubertal development were analyzed using Student’s t test.

Each of the methods (isotopic dilution, DXA and plethysmography BIA) were compared with the results of the 4C model. This comparison was made using the Lin (22) concordance coefficient and Bland-Altman method (23). The Bland-Altman analysis was calculated as the mean difference value between the reference (4C model) and each of the methods and the 95% distribution (confidence intervals).

A regression analysis was done to compare the 4C model and the simplest methods (isotope dilution, DXA, plethysmography and BIA) for determining BF%. The slopes and intercepts were assessed and the standard error of the estimate (SEE) was calculated. p < 0.05 was established as the cutoff for statistical significance. The study data were analyzed using STATA program version 10.1 (24).

**RESULTS**

Physical and body composition of the sample by gender and pubertal development characteristics are shown in table I. There was no interaction between sex and pubertal development. However, several significant sex differences were found. Boys had significantly higher values in the variables: age, weight, height, total body water and bone mineral density. As well, in body composition for both BF (kg) and (fat free mass) FFM in kg and percentage for the 4C model, isotopic dilution, DXA, BIA and plethysmography. Also, boys had higher values in the determination of BF% by BIA. Similarly, there were significant differences associated with pubertal development. Both males and females with advanced puberty, showed significantly higher results in age, weight, height, total body water and bone mineral density. BF (kg) and FFM (kg) in the 4C model, isotopic dilution, DXA, BIA and plethysmography. Lean mass (%) for boys only in the BIA and 4C model. Girls with pubertal development I and II had significantly higher values of FFM (%).

Lin coefficients for the different methods estimating BF% compared to the “gold standard” 4C model, by sex and pubertal development are shown in table II. Males in stage I and II, had concordance coefficients ranging between 0.352 and 0.866 and between 0.721 and 0.959 for stage III and V. In males, the greatest agreement was obtained with DXA (stage I and II) and isotopic dilution in stage III and V. The lowest concordance was observed for the BIA (0.352 and 0.721 for boys and girls, respectively). In females of all stages, the greatest agreement was with plethysmography.

Table III shows the R² value, intercepts and slopes for the regressions for BF% according to the 4C model and each of the different methods, along with SEE stratified by gender and pubertal development. R² values were high for males, mainly in stage III and V. Each method explained between 43-87% of the variance in BF% for males in stage I and II and 78-96% for males in stage III and V.

In both groups of children, the slopes differed significantly from 1 except for plethysmography. Lower values of SEE were observed in stage III and V males.
## Table I. Body composition and physical characteristics of the sample, by sex and pubertal development

<table>
<thead>
<tr>
<th>Variable</th>
<th>Boys</th>
<th>Girls</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Genital I &amp; II (n = 19)</td>
<td>Genital III &amp; V (n = 14)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>11.5 ± 1.1</td>
<td>13.6 ± 1.0</td>
</tr>
<tr>
<td>Weight (kg)</td>
<td>60.9 ± 13.5</td>
<td>76.3 ± 11.7</td>
</tr>
<tr>
<td>Height (cm)</td>
<td>149.4 ± 12.0</td>
<td>161.4 ± 4.8</td>
</tr>
<tr>
<td>Total body water (L)</td>
<td>26.9 ± 5.5</td>
<td>35.6 ± 5.0</td>
</tr>
<tr>
<td>Bone mineral density (kg)</td>
<td>1.8 ± 0.5</td>
<td>2.4 ± 0.3</td>
</tr>
</tbody>
</table>

### 4-Component model

<table>
<thead>
<tr>
<th>Variable</th>
<th>Boys</th>
<th>Girls</th>
</tr>
</thead>
<tbody>
<tr>
<td>BF (%)</td>
<td>41.2 ± 6.0</td>
<td>36.9 ± 7.9</td>
</tr>
<tr>
<td>BF (kg)</td>
<td>25.3 ± 7.9</td>
<td>28.6 ± 9.3</td>
</tr>
<tr>
<td>FFM (%)</td>
<td>58.8 ± 6.0</td>
<td>63.1 ± 7.9</td>
</tr>
<tr>
<td>FFM (kg)</td>
<td>35.6 ± 7.7</td>
<td>47.7 ± 6.4</td>
</tr>
<tr>
<td>Isotopic dilution BF (%)</td>
<td>40.3 ± 5.7</td>
<td>36.8 ± 6.7</td>
</tr>
<tr>
<td>Isotopic dilution BF (kg)</td>
<td>24.8 ± 7.8</td>
<td>28.4 ± 8.2</td>
</tr>
<tr>
<td>Isotopic dilution FFM (%)</td>
<td>59.7 ± 5.7</td>
<td>63.2 ± 6.7</td>
</tr>
<tr>
<td>Isotopic dilution FFM (kg)</td>
<td>36.1 ± 7.3</td>
<td>47.9 ± 6.8</td>
</tr>
<tr>
<td>DXA BF (%)</td>
<td>41.5 ± 5.6</td>
<td>39.0 ± 7.0</td>
</tr>
<tr>
<td>DXA BF (kg)</td>
<td>25.6 ± 8.2</td>
<td>30.1 ± 8.8</td>
</tr>
<tr>
<td>DXA FFM (%)</td>
<td>58.5 ± 5.6</td>
<td>61.0 ± 7.0</td>
</tr>
<tr>
<td>DXA FFM (kg)</td>
<td>35.3 ± 6.7</td>
<td>46.2 ± 6.5</td>
</tr>
<tr>
<td>Plethysmography BF (%)</td>
<td>44.4 ± 6.7</td>
<td>39.2 ± 9.0</td>
</tr>
<tr>
<td>Plethysmography BF (kg)</td>
<td>27.2 ± 8.3</td>
<td>30.6 ± 10.5</td>
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<tr>
<td>Plethysmography FFM (%)</td>
<td>55.6 ± 6.7</td>
<td>60.8 ± 9.0</td>
</tr>
<tr>
<td>Plethysmography FFM (kg)</td>
<td>33.7 ± 8.0</td>
<td>45.7 ± 5.9</td>
</tr>
<tr>
<td>BIA BF (%)</td>
<td>34.8 ± 4.4</td>
<td>36.3 ± 4.1</td>
</tr>
<tr>
<td>BIA BF (kg)</td>
<td>21.3 ± 6.6</td>
<td>28.0 ± 7.0</td>
</tr>
<tr>
<td>BIA FFM (%)</td>
<td>61.6 ± 4.4</td>
<td>63.7 ± 4.1</td>
</tr>
<tr>
<td>BIA FFM (kg)</td>
<td>39.6 ± 8.1</td>
<td>48.3 ± 5.6</td>
</tr>
</tbody>
</table>

x: mean; SD: standard deviation.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Boys</th>
<th>Girls</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>r*</td>
<td>Diff [95% CI]</td>
</tr>
<tr>
<td>Isotopic dilution</td>
<td>0.851</td>
<td>-0.04-6.92</td>
</tr>
<tr>
<td>DXA</td>
<td>0.866</td>
<td>-0.10-6.12</td>
</tr>
<tr>
<td>Plethysmography</td>
<td>0.819</td>
<td>-0.17-7.98</td>
</tr>
<tr>
<td>BIA</td>
<td>0.352</td>
<td>2.24-15.31</td>
</tr>
</tbody>
</table>

* r: correlation; 95% CI: confidence interval.

In girls, the highest R² values were observed with the plethysmography method for all stages of pubertal development. Each method explained 34-92% of the variance in female BF% for stage I and II and 63-93% for girls in stage III and V, respectively.

Figure 1 presents the analysis of specific agreement by Bland and Altman test in estimating BF% between the 4C model and the other methods (isotopic dilution, DXA and plethysmography BIA). In males, an underestimation of BF% with DXA and plethysmography BIA was observed. In girls, the highest R² values were observed with the plethysmography method for all stages of pubertal development. Each method explained 34-92% of the variance in female BF% for stage I and II and 63-93% for girls in stage III and V, respectively.
mography was observed. Isotope dilution overestimated by 0.941 for stage I and II and 0.155 for stages III and V, implying a lack of agreement. Also in males, we observed an overestimation of FM (fat mass) by 6.449 for stages I and II and 0.596 for stages III & V,

**Table III.** Regression analysis modeling different methods to estimate total body fat mass, adjusted for sex and pubertal development

<table>
<thead>
<tr>
<th>Variable</th>
<th>Boys Genital I &amp; II (n = 19)</th>
<th>Genital III &amp; V (n = 14)</th>
<th>Girls Breast I &amp; II (n = 5)</th>
<th>Breast III &amp; V (n = 23)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$R^2$ Intercept Slope SEE</td>
<td>$R^2$ Intercept Slope SEE</td>
<td>$R^2$ Intercept Slope SEE</td>
<td>$R^2$ Intercept Slope SEE</td>
</tr>
<tr>
<td>Isotopic dilution</td>
<td>0.75 -1.29 1.06 0.07 0.94</td>
<td>-1.19 1.17 0.02 0.81</td>
<td>-11.33 1.22 0.22 0.77</td>
<td>-19.08 1.44 0.06</td>
</tr>
<tr>
<td>DXA</td>
<td>0.75 -2.79 1.06 0.06 0.87</td>
<td>-6.93 1.12 0.07 0.66</td>
<td>-18.71 1.46 0.23 0.69</td>
<td>-14.39 1.32 0.07</td>
</tr>
<tr>
<td>Plethysmography</td>
<td>0.87 1.63 0.89 0.07 0.96</td>
<td>2.61 0.87 0.03 0.92</td>
<td>9.09 0.72 0.23 0.93</td>
<td>7.13 0.78 0.03</td>
</tr>
<tr>
<td>BIA</td>
<td>0.43 -5.51 1.34 0.12 0.78</td>
<td>-32.34 1.91 0.08 0.34</td>
<td>6.23 0.88 0.34 0.63</td>
<td>-13.91 1.31 0.09</td>
</tr>
</tbody>
</table>

All slopes are significantly different from 1 ($p < 0.05$).

**Figure 1.**

A: Difference in BF% from 4 compartment (4C) model - Isotopic dilution (D2O dilution) model vs. Average BF% (4C + D2O dilution)/2. B: Difference in BF% from 4C model - Dual-energy X-ray absorptiometry (DXA) model vs. Average BF% (4C + DXA)/2. C: Difference in BF% from 4C model - Plethysmography (BOD POD) vs. Average BF% (4C + Plethysmography)/2. D: Difference in BF% from 4C model - BIA vs. Average BF% (4C + BIA)/2.

○, Breast I & II female subjects; ●, Breast III & V female subjects; ∆, Genital I & II male subjects; ▲, Genital III & V male subjects.
DISCUSSION

Most methods for assessing body composition are 2C and involve measuring one compartment and body weight difference with another is estimated (e.g., total BF is evaluated and fat free mass is obtained). These methods incorporate theoretical assumptions about the composition of lean tissue, which are not valid in all cases, especially not in the context of obesity and during growth. Therefore, to minimize error, a combination of measurements is suggested to determine body composition. The 4C model allows the quantification of BF and lean mass, with a degree of precision not achievable with a single method (21). Methods have been demonstrated that there are significant differences in the properties of fat-free tissue in obese, compared to normal weight, children and adolescents have been demonstrated (25-27). Hydration increases and density is lower in lean tissue of obese, compared to normal weight, individuals (25). These differences have been attributed in part to the expansion of extracellular water space (28,29); but also to lower bone mineralization (30). In this study, the 4C model was used as the gold standard to verify the validity of methods for 2C models in estimating total BF% in obese boys and girls with different levels of development of secondary sex characteristics. It is important to have methods that estimate BF% in school age and adolescent children because, even at the same weight, the risk of developing cardiovascular problems is increased with higher proportion of BF and lower lean mass ratio (31).

In our study, the best matching methods of the 2C and 4C models were obtained with DXA and isotopic dilution. Males in stage I & II with DXA (r = 0.87) and stage III & V males with isotope dilution (r = 0.95). In contrast, girls in both categories of pubertal development, the best agreement was obtained with plethysmography (I & II females r = 0.84 and r = 0.89 for the group III & V). These results are comparable to those obtained in a British study, conducted with 30 obese adolescents aged 14.10 ± 1.83 years. In boys, the correlation with plethysmography was r = 0.97 and r = 0.96 for DXA and isotope dilution, respectively. For girls, highest agreement was with plethysmography (r = 0.94) (32). An American study, which evaluated 25 children 11.4 ± 1.4 years, the model showed greater concordance with isotope dilution was 4C (r = 0.98), plethysmography (r = 0.97) and DXA (r = 0.95) (33). A study conducted with Mexican adolescents in school-age children (12.2 ± 2.0 years), showed greatest concordance with DXA, regardless of gender and pubertal stage (r = 0.95) (34).

In turn, in both sexes and stages of development, less agreement and correlation was obtained between the BIA and the 4C model. Our findings are similar to those of Aguirre et al. (35) who studied 424 Chilean students between 7-10 years and described the inaccuracy of the BIA for body composition estimation in pre-pubertal children. Bray et al. (36) also concluded that the BIA method was less acceptable.

The differences in BF% by pubertal stage, between the 2C and 4C models ranged from -3.17 to 6.50 in boys and girls and between -2.68 for 2.060 for both sexes.

Goran et al. (37) found underestimates similar to those reported in our study, from -6.45 (-7.37, -5.53) and in the case of Deurenberg et al. differences ranged from +1.91 (1.18 to 2.63) (38).

One of the possible explanations for the limited functionality of the BIA for obese subjects may relate to the characteristics of the population in which the equations were created from. Differences in ethnicity, nutritional status, bone geometry, body composition and pubertal maturation may influence the relevance of the equations for other populations (21,39-41).

Our data are of particular interest because they allow for the comparison of validity of the models versus the gold standard (4C) to determine which methods are more reliable for assessing BF in obese adolescents of different pubertal development. In this cross-sectional study, comparing DXA and isotope dilution to the 4C model, we found that Tanner III and V stage children had minor differences for isotope dilution compared with the gold standard and that DXA had the best comparison to the gold standard for Tanner I & II children. Thus, isotopic dilution and DXA seem to have the highest accuracy and reliability for measuring BF in obese children and adolescents.

One of the strengths of this research was utilizing four compartments, which yields high precision and accuracy and is considered the gold standard in determining BF in vivo. Another advantage of the study was in the evaluation of differences in BF estimation using different methods, while considering gender and pubertal stage of development. However, a potential limitation is the narrow age range of the study group, which prevents extrapolation to a larger population.

In conclusion, it has been shown that isotopic dilution and DXA are methods sufficiently reliable compared to the gold standard method (4C) for determining BF% in obese children and adolescents, considering sex and stage of pubertal development. In the case of isotope dilution, it is a simple method that can be performed on site (e.g., schools, health centers) and only requires that samples be sent to a laboratory for validation, if sites do not have equipment. DXA requires equipment that is available in clinical centers or hospitals, which facilitates the measurement of fat and bone mineral density in obese patients.

Being able to reliably assess BF, which is associated with other metabolic variables, in obese adolescents could encourage better monitoring and thus help in the treatment of obese subjects. Moreover, we believe that having the proper methods is a contribution to the nutritional surveillance of pediatric obesity, providing reliable information to estimate the change in adiposity, in the global assessment of nutritional status for obese adolescents.

ACKNOWLEDGEMENTS

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