Estimation of body composition and water data depends on the bioelectrical impedance device

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Abstract

Overweight, obese and chronic kidney disease patients have an altered and negative body composition being its assessment important. Bioelectrical impedance analysis is an easy-to-operate and low-cost method for this purpose. This study aimed to compare and correlate data from single- and multi-frequency bioelectrical impedance spectroscopy applied in subjects with different body sizes, adiposity, and hydration status. It was a cross-sectional study with 386 non-chronic kidney disease volunteers (body mass index from 17 to 40 kg/m²), 30 patients in peritoneal dialysis, and 95 in hemodialysis. Bioelectrical impedance, body composition, and body water data were assessed with single- and multi-frequency bioelectrical impedance spectroscopy. Differences (95% confidence interval) and agreements (Bland-Atman analyze) between devices were evaluated. The intraclass correlation coefficient was used to measure the strength of agreement and Pearson’s correlation to measure the association. Regression analyze was performed to test the association between device difference with body mass index and overhydration. The limits of agreement between devices were very large. Fat mass showed the greatest difference and the lowest intraclass and Pearson’s correlation coefficients. Pearson’s correlation varied from moderate to strong and the intraclass correlation coefficient from weak to substantial. The difference between devices were greater as body mass index increased and was worse in the extremes of water imbalance. In conclusion, data obtained with single- and multi-frequency bioelectrical impedance spectroscopy were highly correlated with poor agreement; the devices cannot be used interchangeably and the agreement between the devices was worse as body mass index and fat mass increased and in the extremes of overhydration.

Keywords: Bioelectrical impedance; body composition; fat mass; hydration status; obesity; renal disease

Introduction

Overweight and obese individuals have a body composition similar to those with chronic kidney disease (CKD): increased body fat sometimes added to lean mass depletion [1,2]. These conditions have a negative effect on physical capacity and are related to a higher risk of mortality [1,3] and lower life expectancy [4]. Thus, body composition assessment is important for these subjects.

However, presence of edema and excess fat limit the application of classic methodologies, such as anthropometry [5]. Moreover, reference methods are expensive, time-consuming, and have low availability [5,6]. Therefore, bioelectrical impedance analyze is a promising method for body composition assessment as it is easy to operate, of low cost, and with good accuracy rates [5,7-12].

Several bioelectrical impedance methods exist, including single-frequency bioelectrical impedance analyze (SFBIA) and multi-frequency bioelectrical impedance spectroscopy (BIS) [8]. Because the devices are different in the range of frequencies and mathematical approaches applied for body composition and water estimation [7,9-11,13], the agreement between their measures is unclear.

Therefore, we investigated whether the use of different bioelectrical impedance devices influence the estimation of body composition and water data in a population with different body sizes, adiposity, and hydration status; factors that influence the difference between methods were also assessed.
Materials and methods
This study evaluated the data from 3 cross-sectional observational studies, including 386 non-CKD volunteers (professionals working at a university hospital, undergraduate students, and graduate students, 204 females and 182 males, aged from 20 to 40 years); 30 patients undergoing peritoneal dialysis (PD) treatment; and 93 patients in haemodialysis (HD) treatment. Both groups of patients were under treatment for at least 3 months, 53 were females and 70 males, aged from 15 to 81 years, recruited from a tertiary care hospital.

Convenience sampling was used to contact and screen potential candidates: for the non-CKD group, all subjects interested in participating and within the criteria of eligibility were evaluated. From the 36 initial PD outpatients, 3 were ineligible and 3 refused to participate. From the 310 HD patients, 162 were ineligible and 55 refused to participate.

Exclusion criteria for the non-CKD group were pregnancy, lactation, infectious diseases, inflammatory state, amputation, presence of prosthesis or pacemaker were known to influence body composition and for CKD groups, metabolic and/or endocrine diseases and use of medication that influence body composition other than CKD were evaluated. From the 36 initial PD outpatients, 3 were ineligible and 3 refused to participate. From the 310 HD patients, 162 were ineligible and 55 refused to participate.

Table 1. Descriptive data for non-CKD subjects stratified in BMI subgroups and for CKD patients stratified in PD and HD subgroups

| BMI<18.5 | 18.5<BM<25 | 25<BM<30 | BM>30 | PD | HD |
|----------|------------|----------|-------|----|----|
| n        |            |          |       |    |    |
| Women (%)|            |          |       |    |    |
| (80)     | (35)       | (37)     | (45)  |    |    |
| Age (years) |          |          |       |    |    |
| 26±4.2   | (90)       | (90)     | (90)  |    |    |
| Weight (kg) |          |          |       |    |    |
| 48±9.4   | (39)       | (39)     | (39)  |    |    |
| BMI (kg/m²) |          |          |       |    |    |
| 18±0.7   | (15)       | (15)     | (15)  |    |    |
| OH (L)   | -0.1±0.4   | -0.1±0.4 | -0.1±0.4 |    |    |
| OH<1.1 L (%) | (0)       | (0)      | (0)   |    |    |
| OH>1.1 L (%) | (12)      | (12)     | (12)  |    |    |
| OH+1.1 L (%) | (25)      | (25)     | (25)  |    |    |
| FFM (kg) | 34±16.3   | 34±16.3 | 34±16.3 |    |    |
| FFM (kg/m²) |          |          |       |    |    |
| 12±1.5   | (26)       | (26)     | (26)  |    |    |
| FM (kg)  | 14±4      | 14±4     | 14±4  |    |    |
| FMI (kg/m²) |          |          |       |    |    |
| 5±1.5   | (4 to 21)  | (4 to 21)| (4 to 21) |    |    |
| PA (%)   | 5±8±0.5   | 5±8±0.5 | 5±8±0.5 |    |    |
| Impairment Nutritional status (%) | 4 | 7 | 4 | 23 | 33 |
| (3)      | (6)        | (3)      | (2)   | (76) | (36) |
| DM (%)   | 0          | 0        | 0      | 10  | 61 |
| (0)      | (0)        | (0)      | (0)   | (33) | (66) |
| SAH (%)  | 0          | 0        | 0      | 19  | 61 |
| (0)      | (0)        | (0)      | (0)   | (63) | (66) |
| Residual Diuresis (ml) | 937±680*** | 170±330*** | 937±680*** | (50 to 2000) | (0 to 2000) |
| Kt/V1    | 2.6±0.9*** | 1.5±0.3*** | 2.6±0.9*** | (1.4 to 4.5) | (1 to 2.5) |
| Duration of dialytic treatment (months)2 | 33±32*** | 63±48*** | 33±32*** | (3 to 144) | (5 to 264) |
| Dialytic treatment >12 months (%) | 21 | 82 | 21 | 82 |
| (70) | (70) | (70) | (70) |

BMI, body mass index; CKD, chronic kidney disease; DM, Diabetes Mellitus; FFM, fat free mass; FMI, fat free mass index; FM, fat mass; FMI, fat mass index; HD, hemodialysis; OH, overhydration; PA, phase angle; PD, peritoneal dialysis; SAH, Systemic Arterial Hypertension. Data presented as mean ± SD (minimum value to maximum value) or in %. Values with different letters in the same line between BMI subgroups, PD and HD groups are significantly different, p<0.05 (ANOVA). 2According to the obtained PA and the cut-off points proposed by Kuchnia and collaborators [24]. 3Data analyzed with unpaired t test, PD vs HD, ***p<0.001. FFM, FMI, FM, OH and PA data from multifrequency bioelectrical impedance spectroscopy.
Distribution 

The unpaired t-test for comparison and frequency values. We applied Q-Q plot to analyze data to provide such information. For descriptive data of body respectively obtained with the BIS device, as SFBIA is unable overhydration (OH) values of > 1.1 L and ≤1.1 L [25], Hyperhydration and dehydration were determined by analyze according to Kuchnia and collaborators [24].

Status was assessed by phase angle (PA) obtained by BIS minus FFM and ICW as TBW minus ECW. The nutritional and TBW [23] for CKD groups. FM was calculated as weight for non-CKD group, FFM for PD [22] and for HD [23] groups applied in random order and both in hand-to-foot tetrapolar position [8]. Unless a fistula was present, the right side was used and measurements were done after being in supine position for 20 min. Using resistance (R) and reactance (Xc) from BIS, intracellular water (ICW) and extracellular water (ECW) [10], fat free mass (FFM), and fat mass (FM) [13] were estimated applying predictive equations previously developed. Total body water (TBW) was calculated by the sum of ICW and ECW. The appropriate predictive equations were used for SFBIA data to calculate ECW [20], FFM [21] and TBW [21] for non-CKD group, FFM for PD [22] and for HD [23] groups and TBW [23] for CKD groups. FFM was calculated as weight minus FFM and ICW as TBW minus ECW. The nutritional status was assessed by phase angle (PA) obtained by BIS analyse according to Kuchnia and collaborators [24]. Hyperhydration and dehydration were determined by overhydration (OH) values of > 1.1 L and ≤1.1 L [25], respectively obtained with the BIS device, as SFBIA is unable to provide such information. For descriptive data of body composition and nutritional status BIS data was used.

Data are presented as mean ± SD, minimum, maximum and frequency values. We applied Q-Q plot to analyze data distribution [26], the unpaired t-test for comparison between CKD groups, and ANOVA for comparison between non-CKD and CKD groups [27]. Differences between devices were evaluated as the difference between BIS and SFBIA (BIS – SFBIA). The 95% confidence interval (95%CI) for mean difference was calculated: if the interval included zero, the data measured with the 2 devices agreed on group level. Agreement on individual level was evaluated using Bland-Altman analyze with limits of agreement [28]. We applied intraclass correlation coefficient (ICC) to measure the strength of agreement and Pearson’s correlation to assess the association as previously proposed [29,30]. Regression analyze was performed to test the association between agreement with BMI and OH. Statistical significance was considered when p<0.05. Data analyze were performed using MINITAB, version 18.

Results

A total of 509 subjects were evaluated. Descriptive data are shown in Table 1. PD (43% ≥60 years old) and HD (23% ≥60 years old) groups were older than the non-CKD group. Weight, FM, and FMI significantly increased with increasing BMI subgroups. The PD group was classified according to BMI as 3% underweight, 40% overweight, and 3% obese and the HD group as 3, 31 and 19%, respectively for the same categories. The FFM and FFMI were similar between CKD groups and the underweight subgroup. PD and HD groups were similar to overweight subgroup for FM and FMI.

As groups have differences in age and sex distribution, we evaluated the association of age and sex with body composition by Pearson’s correlation: in the non-CKD group, age did not correlated with BMI, FFMI or FMI (p>0.05), but sex had a correlation coefficient of 0.60 with FFMI and of 0.48 with FMI; for the PD group, sex was not correlated (p>0.05), but age had a correlation of 0.15 with BMI, -0.22 with FFMI, and 0.19 with FMI; for HD, sex had a correlation of 0.24 with FFMI and 0.15 with FMI; age was correlated with FFMI (-0.11). Therefore, the greater FM and FMI and lower FFMI and FFMI observed in CKD when compared with the non-CKD group are partially explained by sex and age differences given the observed correlation coefficients in each group.

Almost 80% of the PD group and 40% of HD group had nutritional impairment. Hyperhydration was more common in PD and dehydrated in HD group. SAH affected more than half of CKD patients and DM was more common among HD patients. Residual diuresis was higher in the PD group with 7% anuric, 7% oliguric, and 86% with residual diuresis. For the HD group, the same classification was 62, 15, and 23%, respectively. Kt/V was greater in PD and the duration of dialytic treatment was longer in the HD group.
Table 2. Statistics of BIS vs SFBIA data in non-CKD BMI stratified subgroup

| Data analysed | R (ohm) | Xc (ohm) | PA (°) | TBW (L) | ECW (L) | ICW (L) | FFM (kg) | FM (kg) |
|---------------|---------|----------|--------|---------|---------|---------|---------|---------|
| **Underweight subjects (n=40)** | | | | | | | | |
| | 848±74  | 596±103  | -0.02  | -0.12  | 252±134 | (35.5±18.9) | 209 to 294 | (29 to 42) | -10 | 513 |
| | 869±39  | 63±6.5   | -0.03  | -0.12  | 23±14   | (31.1±18.1) | 19 to 28   | (25 to 37) | -4.6 | 51 |
| Bland-Altman | 5.8±0.5  | 6.1±0.6  | 0.12   | 0.09   | -0.3±0.8 | (4.6±12.6) | -0.5 to -0.05 | (-8.9 to -0.7) | -1.8 | 1.2 |
| 95% limits of agreement | 27±13.9 | 30±4.1   | 0.38** | 0.47** | -3.4±4.1 | (11.7±14.3) | -4.7 to -2 | (-16 to -7.1) | -11 | 4.7 |
| 95% limits of agreement (%): Biasa (%)b 95%CI Biasc (%)d Lower | 11±1.4  | 14±2.2   | 0.20** | 0.47** | -2.9±2 | (22.8±14.8) | -3.6 to -2.3 | (-28 to -18) | -6.8 | 0.9 |
| 95% limits of agreement (%): Biasa (%)b 95%CI Biasc (%)d Upper | 16±2.6  | 16±4.9   | 0.45** | 0.44** | -0.4±2.5 | (-3.1±14.8) | -1.2 to 0.4 | (-7.9 to 1.7) | -5.2 | 4.4 |
| **Normal weight subjects (n=120)** | | | | | | | | |
| | 718±99   | 621±97   | -0.02  | -0.05  | 97±141  | (14.6±20.6) | 71 to 122 | (11 to 18) | -180 | 374 |
| | 82±9.4   | 63±8.9   | 0.04   | 0.13   | 19±12   | (25.8±16.9) | 16 to 21 | (23 to 29) | -5.3 | 43 |
| Bland-Altman | 6.6±0.8  | 5.9±0.5  | 0.17   | 0.17   | 0.7±0.9 | (11.1±14.3) | 0.5 to 0.9 | (8.6 to 14) | -1 | 2.5 |
| 95% limits of agreement | 35±7.3  | 32±4.4   | 0.62** | 0.62** | 2.1±5.7 | (4.9±16.7) | 1.0 to 3.1 | (1.8 to 7.9) | -9.1 | 13 |
| 95% limits of agreement (%): Biasa (%)b 95%CI Biasc (%)d Lower | 14±2.6  | 16±2.4   | 0.58** | 0.58** | -1.3±2.1 | (-9.4±14.3) | -1.7 to 0.9 | (-12 to -6.8) | -5.4 | 2.7 |
| 95% limits of agreement (%): Biasa (%)b 95%CI Biasc (%)d Upper | 20±4.8  | 17±2   | 0.27** | 0.56** | 3.4±16.4 | (19.2±37.6) | 2.7 to 4.1 | (13 to 20) | -4.4 | 11 |
| **Overweight subjects (n=118)** | | | | | | | | |
| | 650±78   | 54±710   | 0.01   | -0.02  | 102±128 | (18±22) | 79 to 126 | (14 to 22) | -148 | 352 |
| | 38±7.7   | 37±6.9   | 0.52** | 0.52** | 0.8±8.6 | (1.4±18) | -0.4 to 2.1 | (-1.8 to 4.7) | -13 | 14 |
| Bland-Altman | 6.8±0.8  | 6.6±0.9  | 0.01   | 0.01   | 0.2±1.2 | (3±18) | -0.03 to 0.4 | (-0.4 to 6.3) | -2.2 | 2.6 |
| 95% limits of agreement | 16±2.8  | 17±2.8   | 0.65** | 0.65** | -0.9±2.2 | (-6.1±13) | -1.4 to 0.6 | (-8.4 to -3.7) | -5.2 | 3.3 |
| 95% limits of agreement (%): Biasa (%)b 95%CI Biasc (%)d Lower | 22±5.1  | 20±3.5   | 0.30** | 0.30** | 1.8±5.1 | (7.4±23) | 0.9 to 2.8 | (3.1 to 12) | -8.1 | 12 |
| 95% limits of agreement (%): Biasa (%)b 95%CI Biasc (%)d Upper | 45±13   | 47±5.6   | 0.52** | 0.52** | -0.9±2.2 | (-6.1±13) | -1.4 to 0.6 | (-8.4 to -3.7) | -5.2 | 3.3 |
| **Obese subjects (n=108)** | | | | | | | | |
| | 596±77   | 48±5±3   | 0.07   | 0.19   | 114±85  | (20.8±14.8) | 98 to 130 | (17.9 to 23.6) | -52 | 280 |
| | 71±9.3   | 61±6.9   | -0.01  | -0.01  | 9.8±12  | (14.5±17.4) | 8 to 12 | (11.2 to 17.9) | -13 | 33 |
| Bland-Altman | 6.8±0.8  | 7.2±6.0  | 0.01   | 0.01   | 0.4±5.1 | (-5.7±15.3) | -0.6 to -0.2 | (-8.7 to -2.8) | -2.5 | 1.7 |
| 95% limits of agreement | 42±5.8  | 42±5.4   | 0.70*** | 0.70*** | -0.6±5.4 | (-2.4±12.5) | -1.6 to 0.4 | (-4.8 to 0.01) | -11 | 9.9 |
| 95% limits of agreement (%): Biasa (%)b 95%CI Biasc (%)d Lower | 18±3.1  | 19±2.9   | 0.79*** | 0.79*** | -0.8±1.9 | (-4.5±9.7) | -1.2 to 0.4 | (-6.4 to -2.7) | -4.4 | 2.8 |
| 95% limits of agreement (%): Biasa (%)b 95%CI Biasc (%)d Upper | 24±5.3  | 23±2.4   | 0.53** | 0.53** | 0.2±4.9 | (0.9±16.6) | -0.5 to 1.4 | (-4.1 to 2.3) | -7.6 | 8.1 |
| **BMI, body mass index; ECW, extracellular water; FFM, fat free mass; FM, fat mass; ICC, Intraclass Correlation Coefficient; ICW, intracellular water; PA, phase angle; r, Pearson correlation coefficient; R, resistance; SFBIA, single-frequency bioelectrical impedance; TBW, total body water; Xc, reactance. Data presented as mean ± SD or minimum to maximum value. ICC, *p<0.05, **p<0.01 BIS vs SFBIA. r, *p<0.05, **p<0.01 BIS vs SFBIA. Mean error between BIS and SFBIA: BIS – SFBIA. Mean percentage error between BIS and SFBIA: Bias/[(BIS + SFBIA)/2] X 100. 95%CI of difference between BIS and SFBIA: BIS – SFBIA. 95%CI of difference between BIS and SFBIA (%): Bias/[(BIS + SFBIA)/2] X 100. 95%CI that include zero are unbiased.** |
Concerning agreement between BIS and SFBIA, in the non-CKD group (see table 2), SFBIA underestimated resistance (R), reactance (Xc), and FM, and overestimated ECW and FFM. For the underweight subgroup, the greatest difference occurred for ECW, FFM, and FM, and the best agreement was for TBW and ICW, which did not show difference between devices. For normal weight, FFM, ICW, and FM had the greatest differences and ECW had the best agreement. For overweight, the greatest differences were for FFM and FM, and the best agreement for ECW; PA and TBW did not show difference between devices. For obese, FM and FFM had the greatest differences, ECW had the best agreement, and TBW and ICW did not show a difference between devices. For PD group (see table 3), SFBIA underestimated FM and overestimated TBW, ECW, ICW, and FFM. The greatest differences were for FFM and FM, and the best agreement for TBW; R, Xc, and PA did not show a difference between devices. For the HD group (see table 3), SFBIA underestimated R, Xc, and FM and overestimated TBW, ECW, and FFM. The greatest differences were for FFM and FM, the best agreement for FFM and TBW, and PA and ICW did not show a difference between devices.

For all variables in all groups, the limits of agreement were very large; data generated by SFBIA and BIS are not interchangeable. In addition, FM had the highest difference and limits of agreement, and the lowest correlation and agreement coefficients. In addition, a proportional agreement was observed as the difference between devices were greater in extreme values of BMI or as BMI increased (see Fig. 1) and agreement decreased in extremes values of water imbalance or as OH increased (see Fig. 2).

Discussion

Body composition is important given its role in survival, clinical outcomes, quality of life, and risk of mortality [5,31]. Although the existence of reference methods for body composition analyze, the low accessibility and high costs direct the efforts for bedside procedures. However, it is still unclear which bedside tool is most useful to estimate body composition and hydration status in epidemiological studies or in-patient groups, specifically obese and CKD subjects. Thus, bioelectrical impedance is a promising tool for body composition analyze, but whether the different technologies and mathematical procedures in bioelectrical devices generate similar results needs clarification.

The great differences and wide limits of agreement found in this study indicate that the results obtained with both tested devices are not interchangeable, as concluded by others [32,33].

The difference between bioimpedance devices increased with increasing BMI or were higher in BMI extremes, showing an influence of body size on measurements. As shown in the present study, the increase in BMI was due to an increase in FM, standing out as an interfering factor. Some studies evaluated the ability of BIS and SFBIA to measure body fluid or body composition compared with reference methods and observed systematic errors positively correlated with BMI [34].

Due to these errors, new equations for estimating water content and body composition by BIS were developed [10,13]; these mathematical models promised a better fit with body size as a correction for BMI is applied [9]. In the present study, such mathematical innovation was applied and partly explained the broad limits of agreement between BIS and SFBIA.

In addition, the differences between devices were higher in water imbalance status. SFBIA has as a principle that ICW-to-ECW ratio is constant with no variation of specific resistivity across different tissues [35]. However, specific resistivity is related to electrolyte concentration [35] as well as ICW and ECW distribution, factors altered by nutritional status and in disease state, as in CKD and obesity [8,14]. SFBIA has a single frequency of 50 kHz and it is unable to penetrate the cell membrane and properly compute ICW. This is the major limitation for adequately measuring TBW [11,36], ICW, and ECW as well as differentiate one parameter from another, interfering in FFM predictive capacity and overestimating fat free tissue [32].

On the other hand, the Cole model and Hanai mixture theory are mathematical models shown to best describe the physiological alteration in tissues bioelectric properties [10]. Thus, the BIS approach, with high and low frequencies, can directly measure ICW, ECW, and TBW [8,9,35]. However, BIS is based on some principles not always respected across the range of body composition, especially in states of hyperhydration and excess adiposity [37]; many constants are employed, such as fixed values for specific resistivity of ECW and ICW compartments, body density, and shape [9,10].

Thus, these limiting factors present in each equipment but with different natures can justify the wide limits of agreement between the devices, as well as the greater differences in OH, body size, and FM extremes.

PD had older individuals, as it was shown by the Brazilian National Base in Renal Substitutive Therapies [38]. SAH and DM, main risk factors for CKD [14], were the most prevalent diseases among individuals with CKD, corroborating findings from the literature [39,40].

In non-CKD subgroups, the high BMI was largely due to the participation of FM, as observed by others [41]; a worrying information considering the young age of the group and the cardiometabolic risks that the excess body fat can exert. Regarding body composition of individuals in PD and HD, an excess FM and low FFM suggest the presence of sarcopenia, obesity, and sarcopenic obesity. The prevalence of sarcopenia in renal population in dialysis therapy varies from 20 to 44% in CKD final stages [42,43], and around 10% in the CKD under conservative treatment [44].
This prevalence is much higher than that observed in the general population [45] and in patients with early CKD stages, suggesting that the loss of muscle mass increases as renal function decreases [1].

This deleterious body composition of CKD groups can explain the high percentage of individuals with low PA values, predicting a worse clinical prognosis; PA is considered a marker of cellular integrity and associated with nutritional status. It is also an independent risk factor for long-term mortality [46]. Advanced age is a risk factor for sarcopenia and nutritional impairment [47,48] and hyperhydration is associated with inflammation and increased risk of mortality [49].

The guidelines published by the National Kidney Foundation [50] suggest that the adequacy of dialysis treatment should be interpreted considering not only the clearance of small solutes, but also a careful analyze that encompasses several aspects within each nutritional status and fluid volume. Thus, patients in PD and HD here evaluated presented nutritional risk either due to compromised body composition or to presence of dehydration and hyperhydration status [49].

This study has several strengths and limitations. All measurements were standardized and the adherence to the protocol was verified prior to measurements. The choice of predictive equations applied to raw data from BIS and SFBIA was based on the analyze of the greatest similarity with the original sample characteristics, as suggested by Mulasi and collaborators [12], being a way to improve accuracy. The evaluation of the agreement of body size, FM, and water imbalance measures between the devices with a deep statistical analyze allowing collective and individual assessment, best detailed the main interfering factors for agreement between methods. Also, individuals from PD and HD groups achieve clinical stability after 3 months on dialysis therapy and the majority remain for more than one year in renal replacement therapy [14]. However, the limitation of this study is the lack of a reference method for body composition and water content data. Thus, it is not possible to indicate which device is the most reliable.

### Conclusion

SFIBIA and BIS generated data that are not interchangeable. This study highlights the limitations of both technologies showing that body size, fat mass, and hydration status are interfering factors in the results and influenced the differences between methods. The limitations in BIS and SFIBIA should be considered when assessing body composition and hydration status especially in obese individuals and in those with water imbalance status, such as renal patients. Future studies are needed to improve these limiting factors.
Figure 1: Regression analysis between BIS and SFBIA bias (BIS-SFBIA) with BMI. Data analyzed only for BMI subgroups. BIS, multifrequency bioelectrical impedance spectroscopy; BMI, body mass index; ECW, extracellular water; FFM, fat free mass; FM, fat mass; ICW, intracellular water; PA, phase angle; R, resistance; SFBIA, single-frequency bioelectrical impedance; TBW, total body water; Xc, reactance. (a) R; (b) Xc; (c) PA; (d) TBW; (e) ICW; (f) ECW; (g) FFM; (h) FM. Circle: underweight; Square: normal weight; Trapezium: overweight; Triangle: obese.
Figure 2: Regression analysis between BIS and SFBIA bias (BIS-SFBIA) with OH. Data analyzed only for CKD groups. BIS, multifrequency bioelectrical impedance spectroscopy; BMI, body mass index; CKD, chronic kidney disease; ECW, extracellular water; FFM, fat free mass; FM, fat mass; ICW, intracellular water; PA, phase angle; R, resistance; SFBIA, single-frequency bioelectrical impedance; TBW, total body water; OH, overhydration state; Xc, reactance. (a) R; (b) Xc; (c) PA; (d) TBW; (e) ICW; (f) ECW; (g) FFM; (h) FM.
Conflict of interest

Authors state no conflict of interest

References

1. Sharma D, Hawkins M, Abramowitz MK. Association of sarcopenia with eGFR and miscategorization of obesity in adults with CKD in the United States. Clin J Am Soc Nephrol. 2014;9:2079-2788. https://doi.org/10.2215/CJN.02140214

2. Johansen KL, Lee C. Body composition in chronic kidney disease. Curr Opin Nephrol Hypertens. 2015;24:268-275. https://doi.org/10.1097/MNH.0000000000000120

3. Johansen KL, Dalrymple LS, Delgado C, Kaysen GA, Kornak J, Grimes B, et al. Association between body composition and frailty among prevalent hemodialysis patients: a US Renal Data System special study. J Am Soc Nephrol. 2014;25:381-389. https://doi.org/10.1681/ASN.2013040431

4. Olshansky SJ, Passaro DJ, Hershov RC, Layden J, Carnes BA, Brody J, et al. A potential decline in life expectancy in the United States in the 21st century. N Engl J Med. 2005;352:1138-1145. https://doi.org/10.1056/NEJMsr043743

5. Earthman CP. Body composition tools for assessment of adult malnutrition at the bedside: A tutorial on research considerations and clinical applications. JPEN J Parenter Enteral Nutr. 2015;39:787-822. https://doi.org/10.1177/0148607115595227

6. Sergi G, Trevisan C, Veronese N, Lucato P, Manzato E. Imaging of sarcopenia. Eur J Radiol. 2016;85:1519-1524. https://doi.org/10.1016/j.ejrad.2016.04.009

7. Lukaski HC. Evolution of bioimpedance: a circuitous journey from estimation of physiological function to assessment of body composition and a return to clinical research. Eur J Clin Nutr. 2003;59:258-66. https://doi.org/10.1016/S0955-2863(03)00300-4

8. Kyle Ug, Bosaeus I, De Lorenzo AD, Deurenberg P, Elia M, Gómez JM, et al. Composition of the ESPEN Working Group Bioelectrical impedance analysis—part 1: review of principles and methods. Clin Nutr. 2004;23:195-214. https://doi.org/10.1016/j.clnu.2004.06.004

9. Matthie JR. Bioimpedance measurements of human body composition: critical analysis and outlook. Expert Rev Med Devices. 2008;5:239-261. https://doi.org/10.1586/17434440.5.2.239

10. Moissl U, Wabel P, Chamney PW, Bosaeus I, Levin NW, Bosy-Westphal A, et al. Body fluid volume determination via body composition spectroscopy in health and disease. Physiol Meas. 2006;27:921-933. https://doi.org/10.1088/0967-3334/27/9/012

11. Jaffrin MY, Morel H. Body fluid volumes measurements by impedance: a review of bioimpedance spectroscopy (BIS) and bioimpedance analysis (BIA) methods. Med Eng Phys. 2008;30:1257-1269. https://doi.org/10.1016/j.medengphy.2008.06.009

12. Mulasi U, Kuchnia AJ, Cole AJ, Earthman CP. Bioimpedance at the bedside: current applications, limitations and opportunities. Nutr Clin Pract. 2015;30:180-193. https://doi.org/10.1177/0884536314568155

13. Chamney PW, Wabel P, Moissl UM, Müller MJ, Bosy-Westphal A, Korth O, et al. A whole-body model to distinguish excess fluid from the hydration of major body tissues. Am J Clin Nutr. 2007;85:80-89. https://doi.org/10.1093/ajcn/85.1.80

14. Improving Global Outcomes (KDIGO) CKD Work Group. KDIGO 2012 clinical practice guideline for the evaluation and management of chronic kidney disease. Kidney Inter Suppl 2013;3:1-150.

15. American Diabetes Association. Diagnosis and classification of diabetes mellitus. Diabetes Care. 2014;1:81-90. https://doi.org/10.2337/dc14-0801

16. McCormack T, Krause T, O’Flynn N. Management of hypertension in adults in primary care: NICE guideline. Br J Gen Pract. 2012;62:163–164. https://doi.org/10.3399/bjgp12X630232

17. Kellum JA, Levin N, Bouman C, Lameire N. Developing a consensus classification system for acute renal failure. Curr Opin Crit Care 2002; 8:509–514. https://doi.org/10.1007/s0075198-200212000-00005

18. Heymsfield SB. Anthropometric measurements: application in hospitalized patients. Infusionstherapie. 1990;17:48-51.

19. World Health Organization. Physical status: the use and interpretation of anthropometry, report of a WHO Expert Committee. Geneva, Switzerland: WHO Technical Report Series 854. 1995.

20. Sergi G, Bussolotto M, Perini P, Calliari I, Gantin V, Ceccan A, et al. Accuracy of bioelectrical impedance analysis in estimation of extracellular spaces in healthy subjects and in fluid retention. Ann Nutr Metab. 1994;38:158-165. https://doi.org/10.1159/000177806

21. Sun SS, Chumlea WC, Heymsfield SB, Lukaski HC, Schoeller D, Fried K, et al. Development of bioelectrical impedance analysis prediction equations for body composition with the use of a multicomponent model for use in epidemiologic surveys. Am J Clin Nutr. 2003;77:331-340. https://doi.org/10.1093/ajcn/77.2.331

22. Deurenberg P, Weststrate JA, Hautvast JG. Changes in fat free mass during weight loss measured by bioelectrical impedance and by densitometry. Am J Clin Nutr. 1989;49:33-36. https://doi.org/10.1093/ajcn/49.1.33

23. Kushner RF, Schoeller DA. Estimation of total body water by bioelectrical impedance analysis. Am J Clin Nutr. 1986;44:417-424. https://doi.org/10.1093/ajcn/44.3.417

24. Kuchnia AJ, Teigen LM, Cole AJ, Mulasi U, Gonzalez MC, Heymsfield SB, et al. Phase Angle and Impedance Ratio: reference cut-points from the United States National Health and Nutrition Examination Survey 1999–2004 from bioimpedance spectroscopy data. JPEN J Parenter Enteral Nutr. 2016;41:1310-1315. https://doi.org/10.1177/0148607116670378

25. Ronco C, Verger C, Crepaldi C, Pham J, De Los Rios T, Gauly A, et al. Baseline hydration status in incident peritoneal dialysis patients: the initiative of patient outcomes in dialysis (iPOD-PD study). Nephrol Dial Transplant. 2015;30:849-858. https://doi.org/10.1093/ndt/gfv013

26. Wilk MB, Gnanadesikan R. Probability plotting methods for the analysis of data. Biometrika, Biometrika Trust. 1968;55:1-17.

27. Pagano M, Gauvreau K. Princípios de Bioestatística. 1st ed. Sao Paulo, SP: Thomson; 2004.
28. Bland JM, Altman DG. Statistical methods for assessing agreement between two methods of clinical measurement. Lancet. 1986;1:307-310. https://doi.org/10.1016/S0140-6736(86)90837-8

29. Landis JR, Koch GG. The Measurement of observer agreement for categorical data. Biometrics. 1977;33:159-174. https://doi.org/10.2307/2529310

30. Zou KH, Tuncali K, Silverman SG. Correlation and simple linear regression. Radiology. 2003;227:617-622. https://doi.org/10.1148/radiol.2273011499

31. Baracos V, Caserotti P, Earthman CP, Fields D, Gallagher D, Hall KD, et al. Advances in the science and application of body composition measurement. JPN J Parenter Enteral Nutr. 2012;36:96-107. https://doi.org/10.1177/0148607111417448

32. Jochen GR, Samer RA, Li L, Zhu F, Larive B, Kotanko P, et al. Agreement of single- and multi-frequency bioimpedance measurements in hemodialysis patients: an ancillary study of the frequent hemodialysis network (FHN) daily trial. Nephron Clin Pract. 2014;128:115-126. https://doi.org/10.1159/000366447

33. Yalın SF, Gulcicek S, Avci S, Erkalma Senates B, Altiparmak MR, Trabulus S, et al. Single-frequency and multi-frequency bioimpedance analysis: What is the difference? Nephrology. 2018;23:438-445. https://doi.org/10.1111/nap.13042

34. Sun G, French CR, Martin GR, Younghusband B, Green RC, Xie YG, et al. Comparison of multifrequency bioelectrical impedance analysis with dual energy X-ray absorptiometry for assessment of percentage body fat in a large, healthy population. Am J Clin Nutr. 2005;81:74-81. https://doi.org/10.1093/ajcn/81.1.74

35. Ellis RJ, Bell SJ, Chertow GM, Chamlea WC, Knox TA, Kotler DP, et al. Bioelectrical impedance methods in clinical research: a follow-up to the NIH Technology Assessment Conference. Nutrition. 1999;15:874-880. https://doi.org/10.1016/S0899-9007(99)00147-1

36. Seoane F, Abtahi S, Abtahi F, Elleégård L, Johannsson G, Bosaeus I, et al. Mean expected error in prediction of total body water: a true accuracy comparison between bioimpedance spectroscopy and single frequency regression equations. BioMed Res Int. 2015. https://doi.org/10.1159/000356323

37. Popovic V, Zerahn B, Heaf JG. Comparison of dual energy X-ray absorptiometry and bioimpedance in assessing body composition and nutrition in peritoneal dialysis patients. J Ren Nutr. 2017;27:355-363. https://doi.org/10.1053/j.jrn.2017.03.003

38. Cherchiglia ML, Machado EL, Szuster DA, Andrade EL, Assis Acúrcio Fd, Cailaffa WT, et al. Epidemiological profile of patients on renal replacement therapy in Brazil, 2000-2004. Rev Saúde Pública. 2010;44:639-649. https://doi.org/10.1590/S0034-89102010000400007

39. Cabrera C, Brunelli SM, Rosenbaum D, Anum E, Ramakrishnan K, Jensen DE, et al. A retrospective, longitudinal study estimating the association between interdialytic weight gain and cardiovascular events and death in hemodialysis patients. BMC Nephrol. 2015;16:113. https://doi.org/10.1186/s12882-015-0110-9

40. Sesso RC, Lopes AA, Thomé FS, Lugon JR, Martins CT. Brazilian Chronic Dialysis Census 2014. J Bras Nefrol. 2016;38:54-61. https://doi.org/10.5935/0101-2800.20160009

41. Anjos LA, Wahrlich V, Vasconcellos MT. BMR in a Brazilian adult probability sample: the Nutrition, Physical Activity and Health Survey. Public Health Nutr. 2013;17:853-860. https://doi.org/10.1017/S1368950012005381

42. Kim JK, Choi SR, Choi MJ, Kim SG, Lee YK, Noh JW, et al. Prevalence of and factors associated with sarcopenia in elderly patients with end-stage renal disease. Clin Nutr. 2014;33:64-68. https://doi.org/10.1016/j.clnu.2013.04.002

43. Lamarca F, Carrero JJ, Rodrigues JC, Bigogno FG, Petter RL, Avesani CM. Prevalence of sarcopenia in elderly maintenance hemodialysis patients: the impact of different diagnostic criteria. J Nutr Health Aging. 2014;18:710-717. https://doi.org/10.1007/s12603-014-0505-5

44. Pereira RA, Cordeiro AC, Avesani CM, Carrero JJ, Lindholm B, Amparo FC, et al. Sarcopenia in chronic kidney disease on conservative therapy: prevalence and association with mortality. Nephrol Dial Transplant. 2015;30:1718-1725. https://doi.org/10.1093/ndt/gfv133

45. Beaudart C, Reginster JY, Slomian J, Buckinx F, Dardenne N, Quabron A, et al. Estimation of sarcopenia prevalence using various assessment tools. Exp Gerontol. 2015;61:31-37. https://doi.org/10.1016/j.exger.2014.11.014

46. Oliveira CM, Kubrusly M, Mota RS, Silva CA, Choukroun G, Oliveira VN. The phase angle and mass body cell as markers of nutritional status in hemodialysis patients. J Ren Nutr. 2010;20:314-320. https://doi.org/10.1053/j.jrn.2010.01.008

47. Fielding RA, Vellas B, Evans WJ, Bhasin S, Morley JE, Newman AB, et al. Sarcopenia: an undiagnosed condition in older adults. Current consensus definition: prevalence, etiology and consequences. International working group on sarcopenia. J Am Med Dir Assoc. 2011;12:249-256. https://doi.org/10.1016/j.jamda.2011.01.003

48. Dodds RM, Roberts HC, Cooper C, Sayer AA. The epidemiology of sarcopenia. J Clin Densitom. 2015;18:314-320. https://doi.org/10.1016/S1368-9800(15)00255-X

49. Noordzij M, Jager KJ. Survival comparisons between hemodialysis and peritoneal dialysis. Nephrol Dial Transplant. 2012;27:3385-3387. https://doi.org/10.1093/ndt/gfs031

50. National Kidney Foundation. KDOQI Clinical Practice Guidelines and Clinical Practice Recommendations for 2006 Updates: Hemodialysis Adequacy, Peritoneal Dialysis Adequacy and Vascular Access. Am J Kidney Dis. 2006;48:1-322.