Bioelectric impedance analysis for body composition measurement and other potential clinical applications in critical illness

Hanneke Pierre Franciscus Xaverius Moonen\textsuperscript{a} and Arthur Raymond Hubert Van Zanten\textsuperscript{a,b}

**Purpose of review**
Insight into body composition is of great value in the ICU. Bioelectric impedance analysis (BIA) is the most applicable bedside technique. However, bioimpedance has not been validated in the critically ill, and the interpretation of the measurements poses challenges. This review discusses the potential clinical applications of BIA and explores caveats and solutions to its use in the intensive care setting.

**Recent findings**
A correlation is repeatedly found between raw impedance parameters, fluid ratios, overhydration, and adverse outcome of critical illness. However, cut-off and reference values remain elusive. Experience with BIA-guided fluid management in the ICU is limited. BIA-derived muscle mass appears a promising biomarker for sarcopenia, correlating well with CT-analysis. Body cell mass and fat-free mass provide potential use in estimation of metabolic rate, protein requirements and pharmacokinetics. Several methods of reducing bias in BIA parameters in critical illness require validation.

**Summary**
There are currently too many uncertainties and discrepancies regarding interpretation of bioimpedance in critical illness, to justify therapeutic consequences. However, there are several promising areas of research, concerning some of the most urgent clinical problems in intensive care, emphasizing the need to evaluate further the use and interpretation of bioimpedance in the intensive care setting.

**Keywords**
bioelectric impedance, bioimpedance, body composition, critical illness, phase angle

**INTRODUCTION**
Body composition describes the relative contribution of fat, muscle, bone and water to an individual’s body volume. In the ICU, real-time knowledge of body composition is advantageous to the individualization and optimization of fluid balances, nutrition regimes and medication dosing. Several body composition techniques are available, based on assumptions of weight (hydrostatic weighing), water content (isotope dilution), volume (air displacement plethysmography), energy attenuation (Dual-Energy X-Ray Absorptiometry; DXA), and imaging techniques like computer tomography (CT) and MRI. Although extensively validated, all techniques have limitations when applied during critical illness, because of costs, impracticality or radiation exposure.

Bioelectric impedance analysis (BIA) is quick, noninvasive and relatively inexpensive, making it ideal for bedside use. However, BIA assumes static ratios, most notably a fixed hydration of tissues, which often do not apply to critically ill patients, making interpretation less straightforward. Nevertheless, it is worth exploring potential applications, as BIA currently seems the most feasible body composition measurement technique in the ICU.
Angles for future research will be indicated throughout this manuscript with an asterix (*) and are summarized in Table 1.

**PRINCIPLES OF BIOIMPEDANCE ANALYSIS**

Impedance is the vector analysis of resistance, the opposition to flow of a current, and reactance, the opposition to a current change because of a material's capacitance. When an electrical current is sent through the body, tissues present varying resistance levels. Electrolyte-rich body water is highly conductive; therefore, muscles, having a higher water content, will encounter less resistance than relatively anhydrous tissues, such as fat. Conversely, reactance increases proportionally to cell numbers and their integrity, because of membrane capacitance.

Single-frequency BIA devices (SF-BIA) use a single frequency (usually 50 kHz) to measure impedance. However, low-frequency currents will not penetrate cell membranes, and thus will only measure extracellular water (ECW) impedance. Total body water (TBW) is then estimated through proportional equations. High-frequency currents will go through cells. This impedance reflects combined ICW and ECW : TBW (Fig. 1).

Multifrequency BIA devices (MF-BIA), therefore, provide a more direct portrayal of water compartments, making them more reliable in case of altered hydration status or electrolyte imbalances. Bioimpedance spectroscopy (BIS) applies a more extensive frequency range than MF-BIA. The increase in information obtained from BIS potentially improves predictive power. However, it still requires extrapolation based on population references. Superiority of BIS to the SF-BIA and MF-BIA techniques has not been proven in nonhealthy populations [1–3,4*].

**Table 1.** Suggestions for future research angles concerning bioelectric impedance analysis in critical care

| Subject           | Research angle                                                                 |
|-------------------|--------------------------------------------------------------------------------|
| Internal validity | Influence of overhydration and rapid hydration shifts on BIA measurements      |
|                   | Influence of overhydration and rapid hydration shifts on predictive value of BIA parameters |
|                   | Influence of body temperature on BIA measurements                               |
|                   | Influence of osmotic shifts on BIA measurements                                 |
| External validity | Reference values for BIA measurements in (subgroups of) critically ill patients |
|                   | Cutoff values for outcome predictive qualities of BIA measurements in (subgroups of) critically ill patients |
|                   | Validation of overhydration adjustment of derived parameters in (subgroups of) critically ill patients |
| Safety            | Possible interference of BIA electrical current with electrical implants other than internal ICDs |
| Clinical use      | Development and validation of predictive scoring systems including raw BIA parameters for (subgroups of) critically ill patients |
|                   | Assessment of predictive qualities of BIA measurements for malnutrition         |
|                   | Development and validation of BIA-derived metabolic rate equations with gold-standard methods |
|                   | External validation of method to predict glomerular filtration rate based on BIA-derived body cell mass (BIA-eGFR) |
|                   | Pharmacokinetic models using BIA-eGFR and effect on outcome parameters          |
|                   | Pharmacokinetic models using BIA-derived body composition and effect on outcome parameters |
|                   | Development and validation of equation for protein dosing to BIA-FFM and effect on outcome parameters |
|                   | Exploring options to calculate derived BIA parameters omitting body weight and possibly height |
|                   | Effect of BIVA/BIA-guided fluid management on ICU patient-centered outcomes     |
The phase angle (PhA) shows the relationship between reactance and resistance (Fig. 2).

The greater the number of cell membranes the signal has to pass through, the greater the reactance, and therefore, the PhA. Thus, a large PhA is consistent with a large body cell mass (BCM) relative to ECW, as seen in healthy individuals, whereas ICU patients tend to have a lower PhA. A PhA greater than 6 is assumed normal in health, although PhA varies with sex (men) and age (because of loss of fat-free mass; FFM), and should ideally be related to a reference population, or converted to standardized PhA (SPhA) before comparing across populations [5,6]. PhA measured at 50 kHz is most frequently used, and most reference data are available for this frequency, as this is the frequency at which both resistance and maximum reactance are best measured [7,8].

Bio-electrical impedance vector analysis (BIVA) represents impedance as a vector of reactance and resistance in an x–y plot referring to reference population’s tolerance ellipses (Fig. 3).

BIVA allows simultaneous interpretation of direction (phase) and length of the impedance vector; through which changes in tissue hydration and BCM can be appreciated, independent of regression equations, or body water.

**FIGURE 1.** Low-frequency currents will not penetrate cell membranes, and as such will measure extracellular water impedance. High-frequency currents will go through cells, at which point the impedance reflects total body water (TBW).

**FIGURE 2.** When an electric current passes a cell membrane, reactance causes a time delay, creating a phase shift between voltage and current. The phase angle describes this difference between the voltage and the current. A high-phase angle is, therefore, consistent with large quantities of intact cell membranes and body cell mass.
**DERIVED PARAMETERS**

Reactance, resistance, impedance and PhA, are often referred to as ‘raw’ BIA parameters, that is, not reliant upon empirical modeling. BIA defines the water volumes using impedance and body height, upon which other body composition parameters are based. Earlier BIA devices regarded the body as one cylinder and extrapolated impedance measured on one side of the body. However, this simplification overlooks possible asymmetry and the proportional difference between the trunk and the limbs. Segmental BIA (SM-BIA) devices consider the body as five separate cylinders and use electrodes on all limbs, improving accuracy (Fig. 4).

Various body composition parameters are derived from thereon, using regression analyses with multiple variables obtained through reference measurements. Figure 5 provides an overview of the relationship between several frequently used parameters. SM-BIA can provide additional values, such as

**FIGURE 3.** Bioelectric impedance vector analysis relates the length and direction of the phase angle to that of a reference population, enabling a visual interpretation of the clinical relevance of the raw bioelectric impedance analysis values.

**FIGURE 4.** Earlier bioelectric impedance analysis devices regarded the body as one cylinder, calculating body water volumes based on whole-body impedance and body height. Segmental BIA devices consider the body as five separate cylinders and use electrodes on all limbs, improving accuracy. BIA, bioelectric impedance analysis.
the appendicular skeletal muscle mass (ASMM), the sum of the four limbs’ muscle masses.

**OUTCOME PREDICTION WITH BIOELECTRIC IMPEDANCE ANALYSIS**

Several raw and BIA-derived body composition parameters have been validated as mortality and morbidity predictors in various patient groups and are now being researched as predictors of critical illness outcome [9,10].

**Raw parameters**

Diminished cell count, membrane integrity and altered hydration status in critical illness can lead to changes in reactance and resistance, thereby decreasing PhA compared with healthy individuals [11,12]. Decreased PhA at ICU admission has been associated with hospital, 28-day, 90-day and 12-month mortality [6,13–16]. Concordantly, PhA improved over the first 5 days of ICU stay in ICU survivors, while decreasing significantly in non-survivors [17]. Furthermore, negative correlations have been observed between admission PhA and the length of ward stay, ICU stay and hospital stay, mechanical ventilation duration the APACHE-II score, and recently with the severity of disease of coronavirus disease 2019 (COVID-19) [15,16,18–20].

However, the cut-off values for the predictive value of PhA vary across these studies. The heterogeneity of the ICU populations studied might in part explain these discrepancies. A study comparing sepsis patients with other critically ill patients found that PhA was negatively correlated with the APACHE-II score only in the nonsepsis group [21]. Additionally, in addition to the acute changes because of the current illness, PhA inherently also reflects poor underlying health, muscle wasting and frailty, which are independently associated with outcome.

One study using Segmental Multifrequency BIA (SMMF-BIA) found that impedance, reactance and PhA showed more predictive power for mortality than the SAPS, APACHE-II and SOFA severity scoring systems. Similarly, the landmark Phase Angle Project showed that a combined multivariable score improved the discriminative power in predicting mortality, compared with PhA alone [22].

**Hydration parameters**

Overhydration in ICU patients is positively correlated with adverse outcomes but current methods to assess volume status [in-bed weighing, cumulative fluid balance (CFB), central venous pressure] have their limitations. Marked BIVA-OH on the first 5 days after ICU admission was shown in ICU and 60-day nonsurvivors [19,23,24]. Notably, BIVA predicted mortality better than CFB [23].

Fluid distribution can also be assessed by BIA-derived ECW/TBW ratio. A healthy ECW/TBW ratio varies slightly between sources and device manufacturers but ranges from 0.36 to 0.40. An ECW/TBW ratio of more than 0.40 is considered indicative of overhydration of the extracellular compartment. ECW/TBW-ratio is higher among ICU nonsurvivors and correlates with a longer mechanical ventilation duration [25]. Slobod et al. found that a SF-BIA ECW/TBW-ratio greater than 0.39 on ICU-day 1, associated with an increased number of ventilation days, independent of the APACHE-II score [26]. In CRRT ICU patients, a cut-off for SMMF-BIA ECW/TBW-ratio of 0.413 predicted 28-day mortality, with 71.4% sensitivity and 70.6% specificity [27].

On the basis of the assumption that excess volume accumulates primarily as ECW, the quantity of overhydration can be calculated as the difference between expected ECW, based on the euvolemic ECW/TBW ratio, and the measured ECW [28,29]. On ICU days 1 and 3, BIS-OH (>1l) associated significantly with hospital mortality in 140 ICU patients with 23 nonsurvivors. Day 3 volume status correlated with the duration of ventilation and ICU stay. More ICU-free and ventilator-free days were observed among patients with normal hydration status on day 3 (OH <1 to 1l) [30]. We showed...
increased SMMF-BIA-OH, and ECW/TBW ratio were associated with mortality in COVID-19 [20*].

**Muscle mass**

Determining muscle mass is essential in distinguishing the sarcopenic, from the nonsarcopenic obese, as the former are at higher risk of adverse outcome in the ICU. Furthermore, rapid wasting of muscle mass is a major clinical conundrum, as it is a strong independent predictor for morbidity, mortality, physical functioning and quality of life. PhA is often considered a proxy for LBM. Indeed, two studies found that low BIA/BIS-PhA corresponded to low CT-muscle mass (CT-MM) and muscle density in the critically ill [31**,32**]. Additionally, BIA provides several derived muscle parameters, including FFM, soft lean mass (SLM), LBM, SMM, SMM index (SMI) and segmental values. Two groups studied agreement between CT-MM and BIA-SMM in the ICU. One used the SMM automatically generated by the SMMF-BIA software, and found a high correlation, regardless of patients’ sex, or edema status [33*]. Another group calculated SMM, ASMM and total muscle mass based on raw SF-BIA measurements, using three different equations, and found that although the BIA and CT measurements correlated significantly, the agreement was low, with increasing overestimation of muscle mass by BIA at higher CT-MM. However, BIA did correctly identify patients with low CT-MM [31**]. Therefore, BIA might be clinically useful to identify sarcopenic patients at risk for adverse outcome. However, there was a time difference between the BIA and CT evaluation in these studies, potentially inducing bias. Furthermore, increased muscle mass in ICU patients should not be interpreted as muscle mass of good quality, as intramuscular edema will be classified as muscle mass by both BIA and CT analysis. However, a recent pilot-study comparing CT-MM at ICU admission and BIS-FFM adjusted for overhydration, using an algorithm developed for dialysis patients, found significant correlations and good agreements between the two techniques [32**]. The unadjusted BIS-FFM correlated with CT-MM but performed poorly in classifying muscularity status [32**].

**NUTRITION MANAGEMENT**

Critically ill patients are at increased risk of malnutrition. Several BIA parameters can potentially provide information on nutrition status and requirements.

**Body cell mass**

BCM is the metabolically active part of FFM, in contrast to bone and ECW. As such, a decrease of BCM resulting from critical illness is a marker for malnutrition. Logically, increased ECW is associated with a lower BCM/FFM-ratio. A study comparing BIA measurements before and after hemodialysis in AKI patients (mean weight loss 3.8 kg), suggested hydration shifts have little effect on the BCM measurement, theoretically making it more reliable in critically ill patients [34,35]*.

**Raw parameters**

PhA inherently reflects BCM. In 89 ICU patients, a PhA less than less than 5.5° showed an accuracy of 79% in identifying patients at high nutrition risk (NUTRIC score ≥5) [19*]. In renal replacement therapy patients, a PhA cut-off of 4.6° has been shown to predict malnutrition, defined by protein–energy wasting [36,37]. A study comparing the accuracy of BIVA, versus the definition according to ESPEN in hospitalized patients, in predicting malnutrition, found that BIVA might be the superior method [38].

**Fat-free mass**

Assessment of muscularity by BIA is recommended by the Global Leadership Initiative on Malnutrition (GLIM) [39]. A prospective study among 60 ventilated ICU patients found that a cumulative energy deficit during ICU stay was independently associated with loss of BIS-FFM between inclusion and ICU discharge, as well as with ICU-acquired weakness [40**]. In a retrospective post hoc analysis, including this study, these associations disappeared [41*]. However, raw parameters remained related to muscle weakness [41*].

FFM is closely related to energy expenditure, and some BIA devices offer options to estimate basal metabolic rate (BMR), using FFM-based equations (e.g. Cunningham, or Katch–McArdle). However, based upon derived FFM, these calculations are subject to caveats and have proven to be inferior to indirect calorimetry in several populations, albeit still more accurate than weight-based equations [42,43].

Potentially, BIA-FFM could facilitate targeted protein dosing. Protein targets are usually set to measured actual weight or calculated FFM*. However, these methods do not incorporate changes in body composition and weight gain because of overhydration, as such masking the decrease of FFM during ICU stay.

**FLUID MANAGEMENT**

BIA is commonly used in dialysis patients to guide fluid management by calculating dry weight goals [11,44]. Likewise, in critical illness PhA, ECW/TBW
ratio and overhydration could be used to monitor the effect of fluid management strategies. A prospective, clinician-blinded study was conducted to assess the feasibility and validity of BIVA as a measure of hydration in critically ill patients. The study showed that clinicians blinded to the BIVA results, achieved a mean CFB that was concordant with the prior BIVA classification (i.e. positive for patients’ BIVA classified as dehydrated, negative for overhydrated patients and neutral for normally hydrated patients), proving feasibility [45]. Moreover, directional BIVA changes correlated with directional changes in fluid balance. However, the study showed that vector length increased in parallel with 2.4-l fluid loss, suggesting BIVA might be insensitive to smaller changes [45]. The effect of BIVA/BIA-guided fluid management on patient-centered outcomes has not yet been researched.

GLOMERULAR FILTRATION RATE AND PHARMACOKINETICS

Adequate dosing of renally excreted drugs is challenging in critically ill patients because of changes in kidney function. Most equations to estimate glomerular filtration rate are based on serum creatinine measurement. However, significant limitations arise when these formulas are applied to patients with altered body composition, like low muscle mass [9,10–13]. A Dutch group recently developed and
validated a formula to predict creatinine/urea clearance based on 24 h urine collection (currently the gold-standard in ICU) using serum creatinine and MF-BIA-BCM and ECW/TBW ratio, with good results [46–**]

BIA also provides interesting theoretical ways for pharmacokinetic characterization and medication dosing through real-time appreciation of the changing body composition and volumes of distribution [47]. However, no recent attempts for predictive pharmacokinetic models using BIA in the ICU have been published.

CAVEATS

One main drawback of BIA is the incorporation of reference population values, for all but the raw parameters, which might not apply to the individual patient or population. Although they are validated against standard methods (usually MRI and DXA), the exact equations used by BIA software are rarely released by manufacturers, impairing judgment of applicability [48]**. Several other caveats impair routine use of BIA in the ICU, such as use of inexact input parameters, the lack of ICU reference and cutoff values, and the possible bias introduced by a rapidly changing clinical status. Evidence regarding other considerations to use and interpretation of BIA in the ICU setting are summarized in Table 2.

CONCLUSION

There are several promising areas of BIA research concerning some of the most urgent clinical problems in intensive care. A correlation is repeatedly found between raw impedance parameter, fluid ratios, hydration and adverse outcomes in critical illness. BIA-derived muscle mass appears a promising biomarker for sarcopenia, as it correlates well with CT-analysis. BCM and fat-free mass provide potential parameters, which might not apply to the individual reference population values, for all but the raw parameters, which might not apply to the individual patient or population. Although they are validated against standard methods (usually MRI and DXA), the exact equations used by BIA software are rarely released by manufacturers, impairing judgment of applicability [48]**. Several other caveats impair routine use of BIA in the ICU, such as use of inexact input parameters, the lack of ICU reference and cutoff values, and the possible bias introduced by a rapidly changing clinical status. Evidence regarding other considerations to use and interpretation of BIA in the ICU setting are summarized in Table 2.

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