Phase angle from bioelectrical impedance analysis is a useful indicator of muscle quality

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Abstract

Background Methods that facilitate muscle quality measurement may improve the diagnosis of sarcopenia. Current research has focused on the phase angle (PhA) obtained through bioelectrical impedance analysis (BIA) as an indicator of cellular health, particularly cell membrane integrity and cell function. The current study therefore aimed to evaluate the relationship between the PhA and muscle quality and muscle-related parameters and to determine factors associated with the PhA. Moreover, we attempted to determine the cut-off value of PhA for predicting sarcopenia.

Methods First-year university students (830 male students, 18.5 ± 0.6 years old; 422 female students, 18.3 ± 0.5 years old) and community-dwelling elderly individuals (70 male individuals, 74.4 ± 5.5 years old; 97 female individuals, 73.1 ± 6.4 years old) were included. PhA and other body composition data were measured using BIA, while muscle quality was calculated by dividing handgrip strength by upper limbs muscle mass. The relationship between PhA and the aforementioned parameters were then analysed, after which the cut-off value of PhA for predicting sarcopenia was examined.

Results Multiple linear regression analysis revealed that age, skeletal muscle mass index (SMI), and muscle quality were independently associated with PhA in both sexes [male (age: standardized regression coefficient (β) = −0.43, P < 0.001, SMI: β = 0.61, P < 0.001, muscle quality: β = 0.13, P < 0.001) and female (age: β = −0.56, P < 0.001, SMI: β = 0.52, P < 0.001, muscle quality: β = 0.09, P = 0.007)]. Participants with sarcopenia had a significantly lower PhA compared with those without it (sarcopenia vs. non-sarcopenia: young male participants, 5.51 ± 0.41° vs. 6.25 ± 0.50°, P < 0.001; young female participants, 4.88 ± 0.16° vs. 5.37 ± 0.44°, P = 0.005; elderly male participants: 4.14 ± 0.29° vs. 4.63 ± 0.42°, P = 0.009). Although no significant findings were observed in elderly male participants, the same tendency was noted. Receiver operating characteristic (ROC) curve analysis indicated that PhA had good predictive ability for sarcopenia in young male, elderly male, young female, and elderly female participants (area under the ROC curve of 0.882, 0.838, 0.865, and 0.850, with cut-off PhA values of 5.95°, 5.04°, 5.02°, and 4.20° for predicting sarcopenia, respectively).

Conclusions The PhA reflected muscle quality and exhibited good accuracy in detecting sarcopenia, suggesting its utility as an index for easily measuring muscle quality, which could improve the diagnosis of sarcopenia.

Keywords Phase angle; Sarcopenia; Muscle quality; Body composition; Bioelectrical impedance

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Introduction

Reports have shown that the prevalence of sarcopenia can range from 5% to 13% among individuals between the age of 60 and 70 and from 11% to 50% among elderly individuals over 80 years.\(^1\) According to a recent systematic review and meta-analysis, the prevalence of sarcopenia in the world is 10% \(\pm\) 95% confidence interval (CI) = 8–12%\(^2\) in male participants and 10% \(\pm\) 95% CI = 8–13%\(^2\) in female participants, respectively.\(^3\) The prevention and improvement of sarcopenia, which remains a serious problem worldwide, has become an important issue. Sarcopenia is mainly diagnosed based on guidelines proposed by the European Working Group on Sarcopenia in Older People (EWGSOP)\(^4\) in 2010\(^5\) and subsequently revised in 2018 (EWGSOP2).\(^6\) Essentially, sarcopenia has been diagnosed based on whether ‘muscle strength’, ‘muscle quantity’, and ‘physical performance’ satisfy certain criteria.

Although muscle mass remains the primary factor of muscle strength, studies have shown that muscle strength is only moderately correlated with muscle cross-sectional area and muscle thickness among living bodies.\(^7\)\(^8\) Furthermore, both muscle mass and strength decline along with age; however, the decline in muscle strength is substantially greater than the decline in muscle mass.\(^7\)\(^8\) Thus, a decrease in muscle mass alone cannot fully explain the loss of muscle strength.\(^9\)

The skeletal myocyte gap contains not only muscle fibres (muscle cells) but also fibrous tissue, intramuscular extracellular fat, and extracellular fluid. During muscle atrophy, qualitative changes, such as an increase in intramuscular fat and fibrous tissue, occur in addition to quantitative changes (e.g. a decrease in muscle cross-sectional area),\(^10\) thereby suggesting the importance of assessing muscle quality. Since the effects of muscle quality deterioration have also been considered when diagnosing sarcopenia, guidelines have indicated the need for examining muscle quality.\(^4\) Accordingly, the EWGSOP2 guideline has defined ‘muscle quality’ as that ‘referring both to micro- and macroscopic changes in muscle architecture and composition, and to muscle function delivered per unit of muscle mass’.\(^4\) Nonetheless, no clear criteria have been established for diagnosing sarcopenia.

Highly sensitive imaging modalities, such as computed tomography\(^11\) and magnetic resonance imaging,\(^12\) have been used to assess muscle quality by measuring fat infiltration into the muscle and evaluating muscle attenuation. Moreover, muscle biopsy has been used to directly examine the muscle architecture and composition.\(^13\) However, although these methods can provide precise measurements, obtaining such measurements has remained challenging given the need for invasive procedures, large-scale equipment, prolonged restraint, and skilled technicians. Although echo intensity obtained from ultrasonography images has also been used to assess muscle quality,\(^14\) reproducibility of results has remained a concern given the differences in measurement techniques between each examiner.

Alternatively, the ratios of muscle strength to appendicular skeletal muscle mass (ASM)\(^15\) or muscle volume\(^16\) have also been used as indicators of muscle quality. Despite being simple ratios, they have attracted much attention given studies associating with cardiovascular diseases.\(^17\) Nonetheless, given the lack of a universal consensus thus far, methods that can easily assess muscle quality in clinical practice have been desired.

Interestingly, bioelectrical impedance analysis (BIA) has become quite a popular method for estimating body composition, including muscle mass, considering that it is non-invasive, inexpensive, portable, and easy and quick to use.\(^18\) To determine body composition, BIA measures the human body’s impedance (Z), which is the electrical opposition to the alternating current (AC) of the body composed of resistance (R) and reactance (Xc) represented by the following formula: \(Z^2 = R^2 + Xc^2\).\(^19\) After substituting the obtained impedance values and participant height into the regression equations for each studied population, BIA can estimate lean body mass and body water content, among others.

The phase angle (PhA), which is calculated using the arc-tangent value of the ratio of Xc to R, is independent of conventional regression equations for estimating body composition.\(^20\) When an AC flows through the human body, healthy cell membranes function as capacitors that store electrical energy, consequently causing a delay in its flow. This lag in the current that penetrates cell membranes and tissue interfaces creates the phase difference between the current and voltage, which is expressed as the PhA.\(^21\)

Recent studies have focused on the PhA as an indicator of cellular health, particularly reflecting cell membrane integrity and cell function.\(^22\) Accordingly, the PhA has been reported to be associated with age,\(^23\)\(^–\)\(^25\) sex,\(^23\)\(^–\)\(^25\) and nutrition.\(^26\) Currently, studies have reported low PhA values among patients with HIV\(^27\)\(^,\)\(^28\) and cancer,\(^29\)\(^–\)\(^31\) as well as those on haemodialysis,\(^32\) with PhA being used in their treatment. Moreover, one study reported that lower PhA was correlated with poor muscle function, particularly low muscle volume or strength.\(^33\)

However, no study has determined whether muscle quality and quantity are associated with PhA. As such, the present study aimed to evaluate the association between PhA and muscle parameters and determine factors associated with PhA. Moreover, we attempted to determine cut-off values of PhA for detecting sarcopenia. Granting that muscle quality is associated with PhA, we believe that PhA could be a useful and convenient index of muscle quality, which can help improve the diagnosis of sarcopenia.
Methods

Participants

Participants were recruited during health check-ups held by the town or universities. Young participants included first-year university students of Kyoto Institute of Technology or Shiga University aged 18 to 20, while the elderly participants comprised community-dwelling elderly individuals residing in Seika town, Kyoto.

The exclusion criteria were as follows: (i) unable to maintain a standing position, (ii) presence of cardiac pacemakers, (iii) presence of artificial joints, (iv) apparent oedematous; (v) pregnant or suspected of being pregnant, and (vi) considered unsuitable for study participation.

Consent to participate was obtained from all participants before study inclusion. This study was reviewed and approved by the Ethics Committee of National Hospital Organization Kyoto Medical Center (approval number: 18-106 in young participants and 18-107 in elderly participants) and was conducted in compliance with the ethical principles stated in the Declaration of Helsinki.

Data collection

Bioelectrical impedance and phase angle
A multi-frequency segmental body composition analyser (MC-780A-N, TANITA Co., Ltd., Tokyo, Japan) was used to measure bioelectrical impedance and obtain whole and segmental body composition data. This system uses three different frequency currents (5, 50, and 250 kHz) for high accuracy. An estimation formula for ASM in this model has been published, and a previous validation study demonstrated that body composition data measured using this device was highly correlated with that measured using dual-energy X-ray absorptiometry measurements.34 Data were collected while participants stood on foot-electrodes and held hand-electrodes.

Thereafter, the skeletal muscle mass index (SMI, kg/m²) was calculated by dividing the ASM (kg) by the square of the height (m²). Body mass index (BMI, kg/m²) was calculated by dividing the body weight (kg) by dividing the square of the height (m²).

Moreover, the PhA was calculated using the following formula: PhA (°) = arctangent (Xc/R) * (180/π). Reactance and resistance values measured at 50 kHz current were used to calculate the PhA.19

Muscle quality
The EWGSOP2 sarcopenia guidelines define muscle quality as ‘referring both to micro- and macroscopic changes in muscle architecture and composition, and to muscle function delivered per unit of muscle mass’.4 In this study, muscle quality (kg/kg) was calculated by dividing handgrip strength (HGS) (kg) by upper limbs muscle mass (kg) according to previous reports,17,35,36 focusing on ‘muscle function delivered per unit of muscle mass’. HGS was measured using a Smedley spring-type dynamometer (Grip-D, Takei Scientific Instruments Co., Ltd., Niigata, Japan), while upper limb muscle mass was determined using BIA.

Sarcopenia definition
Sarcopenia was determined using HGS and the SMI based on algorithms proposed by EWGSOP2.4 Cut-off values for HGS were set to <28 and <18 kg, while those for SMI were set to <7.0 and <5.7 kg/m² for Asian male and female participants according to the guideline of Asian Working Group for Sarcopenia, respectively.37 Participants having a HGS and SMI below both criteria were determined to have sarcopenia.

Statistical analysis

Data are expressed as mean ± standard deviation. All statistical analyses were performed using BellCurve for Excel Version 3.20 (Social Survey Research Information Co., Ltd., Tokyo, Japan), with a two-sided P value of <0.05 indicating statistical significance.

The Mann–Whitney U test was used for assessing differences between young and elderly groups according to sex.

Single linear regression analysis was conducted to evaluate the association between PhA and each measured item. For multiple linear regression analysis of PhA, age, SMI, HGS, and muscle quality were selected as independent variables: Model 1 (independent variables: age and SMI), Model 2 (independent variables: age, SMI, and HGS), and Model 3 (independent variables: age, SMI, and muscle quality).

The Mann–Whitney U test was used to examine differences in PhA between the sarcopenia and non-sarcopenia groups (young male, elderly male, young female, and elderly female participants). Moreover, receiver operating characteristic (ROC) curve and the area under the ROC curve (AUC) were used to determine the cut-off values of PhA that indicated the presence of sarcopenia in each group. Cut-off values were determined using the point on the ROC curve located closest to the (0, 1) point.

Results

Initially, 1456 individuals, comprising 1287 young (850 male and 437 female participants) and 169 elderly participants (72 male and 97 female participants) were registered in the present study. After conducting measurements, 23 young participants (12 males and 11 female participants) and 2 elderly male participants with incomplete data and 12 young participants (8 male and 4 female participants) with
impedance data errors were excluded. Ultimately, 1419 participants composed of 830 young male, 70 elderly male, 422 young female, and 97 elderly female participants were included in the study population (Figure 1).

Participant characteristics are summarized in Table 1. Accordingly, young and elderly male participants had a mean PhA of 6.25 ± 0.50° and 5.26 ± 0.50° respectively, with the difference being significant (P < 0.001). Similarly, young female participants had a significantly higher the mean PhA compared with elderly ones (young: 5.37 ± 0.44°, elderly: 4.61 ± 0.43°; P < 0.001). Table 2 details the results of single linear regression analysis on the relationship between PhA and measured items. In both male and female participants, a negative correlation was observed between PhA and age (male: r = −0.48, P < 0.001; female: r = −0.57, P < 0.001). PhA was positively correlated with both the SMI and HGS, which were diagnostic indicators for sarcopenia [SMI (male: r = 0.59, P < 0.001; female: r = 0.50, P < 0.001) and HGS (male: r = 0.40, P < 0.001; female: r = 0.47, P < 0.001)]. Furthermore, PhA was positively correlated with muscle quality (male: r = 0.09, P = 0.006; female: r = 0.36, P < 0.001).

The results of multiple linear regression analysis are provided in Table 3. First, age and SMI, which were strongly correlated with PhA, were selected as independent variables. Thereafter, HGS (a diagnostic indicator for sarcopenia) and muscle quality were also selected as independent variables. In this study, we particularly focused on the relationship between muscle quality and PhA. Model 1 showed that both of age and SMI were significantly associated with PhA. Model 2 showed that only HGS was not significantly associated with PhA, although age and SMI were significantly associated. Model 3 showed that all of age, SMI, and muscle quality significantly associated with PhA in both male and female participants. Age was negatively associated, and SMI and muscle quality were positively associated, respectively [male (age: standardized regression coefficient (β) = −0.43, P < 0.001, SMI: β = 0.61, P < 0.001, muscle quality: β = 0.13, P < 0.001); female (age: β = −0.56, P < 0.001, SMI: β = 0.52, P < 0.001, muscle quality: β = 0.09, P = 0.007)].

Table 4 shows comparisons of the PhA values between the sarcopenia and non-sarcopenia groups. Sarcopenia was diagnosed in eight young male (1.0%), two elderly male (2.9%), five young female (1.2%), and five elderly female (5.2%) participants. Young male, young female, and elderly female participants in the sarcopenia group had significantly lower mean PhA than did those in the non-sarcopenia group. Although no significant difference was observed among elderly male participants, the same tendency was noted.

Finally, results of the ROC analyses are presented in Figure 2 and Table 5. The results showed moderate predictive accuracy of PhA for sarcopenia in young male (AUC = 0.882, 95% CI = 0.796–0.967), elderly male (AUC = 0.838, 95% CI = 0.516–1.160), young female (AUC = 0.865, 95% CI = 0.804–0.926), and elderly female (AUC = 0.850, 95% CI = 0.674–1.026) participants, respectively. The cut-off values of PhA to discriminate sarcopenia from non-sarcopenia were 5.95° in young male participants, 5.04° in elderly male participants, 5.02° in young female participants, and 4.20° in elderly female participants, respectively.

**Figure 1** Study flow for participant registration.
Table 1  Characteristics of participants

| Variable                  | Male All (n = 900) | Young (n = 830) | Elderly (n = 70) | P value |
|---------------------------|-------------------|----------------|-----------------|---------|
| Age (year)                | 22.8 ± 15.1       | 18.5 ± 0.6     | 74.4 ± 5.5      | <0.001  |
| Height (m)                | 1.71 ± 0.06       | 1.71 ± 0.06    | 1.65 ± 0.06     | <0.001  |
| Body weight (kg)          | 62.4 ± 9.7        | 62.2 ± 9.8     | 63.8 ± 9.4      | 0.069   |
| BMI (kg/m²)               | 21.4 ± 3.1        | 21.2 ± 3.0     | 23.4 ± 3.6      | <0.001  |
| Total fat mass (kg)       | 11.5 ± 5.4        | 11.2 ± 5.3     | 14.2 ± 5.2      | <0.001  |
| Body fat percentage (%)   | 17.7 ± 6.7        | 17.4 ± 6.6     | 21.7 ± 5.8      | <0.001  |
| Total muscle mass (kg)    | 48.3 ± 5.1        | 48.4 ± 5.2     | 47.0 ± 4.4      | 0.037   |
| ASM (kg)                  | 23.1 ± 2.9        | 23.3 ± 2.8     | 21.6 ± 3.0      | <0.001  |
| SMI (kg/m²)               | 7.9 ± 0.7         | 7.9 ± 0.7      | 7.9 ± 0.8       | 0.893   |
| Total bone mass (kg)      | 2.7 ± 0.3         | 2.7 ± 0.3      | 2.6 ± 0.2       | 0.039   |
| Body water content (kg)   | 34.6 ± 4.1        | 34.5 ± 4.1     | 35.9 ± 4.1      | 0.006   |
| HGS (kg)                  | 38.4 ± 6.1        | 38.6 ± 6.0     | 35.7 ± 6.7      | <0.001  |
| Muscle quality (kg/kg)    | 8.3 ± 1.1         | 8.4 ± 1.1      | 7.5 ± 1.1       | <0.001  |
| Phase angle (°)           | 6.17 ± 0.57       | 6.25 ± 0.50    | 5.26 ± 0.50     | <0.001  |

Table 2  Single linear regression analysis on the relationship between phase angle and measured items

| Variable                  | Male              | Female             |
|---------------------------|-------------------|--------------------|
| Age (year)                | −0.48 <0.001      | −0.57 <0.001       |
| Height (m)                | −0.04 0.250       | 0.15 <0.001        |
| Body weight (kg)          | 0.28 <0.001       | 0.13 0.002         |
| BMI (kg/m²)               | 0.32 <0.001       | 0.05 0.229         |
| Total fat mass (kg)       | 0.08 0.014        | −0.03 0.564        |
| Body fat percentage (%)   | −0.02 0.540       | −0.12 0.004        |
| Total muscle mass (kg)    | 0.35 <0.001       | 0.33 <0.001        |
| ASM (kg)                  | 0.41 <0.001       | 0.40 <0.001        |
| SMI (kg/m²)               | 0.59 <0.001       | 0.50 <0.001        |
| Total bone mass (kg)      | 0.42 <0.001       | 0.23 <0.001        |
| Body water content (kg)   | 0.43 <0.001       | 0.25 <0.001        |
| HGS (kg)                  | 0.40 <0.001       | 0.47 <0.001        |
| Muscle quality (kg/kg)    | 0.09 0.006        | 0.36 <0.001        |

ASM, appendicular skeletal muscle; BMI, body mass index; HGS, handgrip strength; SMI, skeletal mass index.
Results are expressed as correlation coefficients (r).

Discussion

To the best of our knowledge, this is the first study demonstrating that PhA can be an independent useful indicator of muscle quality defined as HGS divided by upper limbs muscle mass. This study showed that PhA was independently associated with age, SMI, and muscle quality. Our results also showed that the sarcopenia group had a significantly lower PhA than the non-sarcopenia group. Moreover, ROC analysis revealed that PhA exhibited good accuracy in predicting sarcopenia while indicating the best cut-off values of PhA.

Indeed, studies have shown that PhA decreased with age\(^{23–25}\) was lower in women than in men\(^{23–25}\) and was proportional to BMI.\(^{23–25}\) Moreover, other reports have shown that the PhA was positively correlated with muscle mass, HGS,\(^{38}\) lower limb muscle strength,\(^{33}\) and physical function assessed using the speed at which individuals walked and rose from the sitting position.\(^{39}\) The present study found that PhA was lower in women than in men, was negatively correlated with age, and was positively correlated with BMI, muscle mass, and HGS. While these results were consistent with those presented in previous reports, our finding showed that SMI had the strongest correlation with PhA among the body composition and physical function parameters (Table 2).

One study showed that PhA was moderately positively correlated with muscle quality defined as HGS divided by upper limbs muscle mass. The results showed that PhA was independently associated with age, SMI, and muscle quality. Our results also showed that the sarcopenia group had a significantly lower PhA than the non-sarcopenia group. Moreover, ROC analysis revealed that PhA exhibited good accuracy in predicting sarcopenia while indicating the best cut-off values of PhA.
upper limbs muscle mass among elderly women.39 Similarly, the present study found a moderately positive correlation between PhA and muscle quality among female participants \( (r = 0.36, P < 0.001) \), although a very weak positive correlation had been noted among male participants \( (r = 0.09, P = 0.006) \). However, multiple linear regression analysis conducted herein revealed that PhA was significantly associated with age, SMI, and muscle quality in both male \( (\beta = 0.13, P < 0.001) \) and female \( (\beta = 0.09, P = 0.007) \) participants.

One interesting finding obtained herein was that muscle quantity and quality were each independently positively correlated with PhA, suggesting that higher PhA indicated higher muscle quantity and quality.

Evidence has shown that the PhA reflects cell membrane structure, cell mass, cellular integrity, and cell function, with higher PhA levels indicating better parameters.20,22,40 Although the relationship between the PhA and muscle parameters has yet to be fully elucidated, the PhA values were independently associated with muscle quality defined as HGS divided by upper limb muscle mass. The decrease in PhA has been attributed to a reduction in reactance due to muscle mass loss and/or increased resistance caused by increased fat mass.22 Increased intramuscular fat and fibrous tissue has been considered to decrease muscle quality.10 As such, changes in muscle mass and quality have been considered to be independently associated with the PhA. In line with this, sarcopenia guidelines have indicated the need for examining not only muscle quantity but also muscle quality.4 Considering that PhA measurements can simultaneously determine both muscle quality and quantity, they can perhaps be utilized to satisfy requirements established by the guidelines.

Our findings showed that elderly female participants in the sarcopenia group had lower PhA values than those in the non-sarcopenia group. Although no significant difference had been observed in elderly male participants, the same tendency had been noted. This could have been attributed to the fact that only two participants were diagnosed with sarcopenia. These results were consistent with those presented in a previous report, wherein elderly individuals with sarcopenia had lower PhA values than those without sarcopenia.42 The present study also showed that young

### Table 3

| Sex   | Age   | PhA (°) | 95% CI Low  | 95% CI High |
|-------|-------|---------|-------------|-------------|
| Male  | Young | 0.019   | -0.028      | 0.066       |
|       | Elderly | 0.026   | -0.035     | 0.086       |
| Female| Young | 0.026   | -0.039     | 0.091       |
|       | Elderly | 0.026   | -0.039    | 0.091       |

$\beta$, standardized regression coefficient; CI, confidence interval; HGS, handgrip strength; SMI, skeletal mass index.

### Table 4

Comparisons of PhA values between the non-sarcopenia and sarcopenia groups

| Sex | Age   | Non-sarcopenia N | Sarcopenia N | P value |
|-----|-------|------------------|--------------|---------|
| Male| Young | 6.25 ± 0.50      | 5.51 ± 0.41  | <0.001 |
|     | Elderly | 5.28 ± 0.48     | 4.48 ± 0.78  | 0.603 |
| Female| Young | 5.37 ± 0.44     | 4.88 ± 0.16  | 0.005 |
|      | Elderly | 5.37 ± 0.42   | 4.14 ± 0.29  | 0.009 |

N.A., not analysed; PhA, phase angle.

Data are expressed as mean ± standard deviation. Significant differences between the non-sarcopenia and sarcopenia groups were determined using the Mann–Whitney U test.
Participants in the sarcopenia group had lower PhA values than those in the non-sarcopenia group. Moreover, ROC analyses revealed that all young male, young female, elderly male, and elderly female participants had an AUC exceeding 0.8, indicating that PhA had good accuracy in predicting sarcopenia, with cut-off values of 5.95°, 5.04°, 5.02°, and 4.20°, respectively.

Interestingly, elderly individuals included herein had higher cut-off PhA values compared with those included in previous studies, perhaps because our elderly participants were younger. Given that the current study and previous studies have shown that PhA decreases with age, cut-off values may be considered to differ depending on the age of the target population. Therefore, using PhA for...
sarcopenia diagnosis requires establishing cut-off values for not only sex but also 5 or 10 year age groups.

Although differences in PhA had been observed and factors associated with PhA had been identified, the current study could not determine how the decrease in PhA over time adversely affects health given the cross-sectional design of the current study. Studies have suggested that PhA may increase following short-term resistance training. Therefore, to evaluate the effects of changes in PhA over time on physical health, further longitudinal studies are necessary.

The EWGSOP2 guideline has defined ‘muscle quality’ as that ‘referred to both micro- and macroscopic changes in muscle architecture and composition, and to muscle function delivered per unit of muscle mass’. However, these definitions are not identical. Although increased levels of intramuscular fat have been reported to lead muscle dysfunction, such as decreased muscle strength, their minute association remains to be elucidated in the future. Investigating fat infiltration into muscles using computed tomography and magnetic resonance imaging, as well as muscle echo intensity using ultrasonography images, improve the relationship between PhA and changes in muscle architecture and composition, would certainly improve the utility of PhA as a marker of muscle quality.

In conclusion, the current study revealed that PhA was associated with age, SMI, and muscle quality and that higher PhA reflected higher muscle quality. Moreover, this study demonstrated that PhA had good accuracy for detecting sarcopenia. Cut-off PhA values for predicting sarcopenia were 5.95°, 5.04°, 5.02°, and 4.20° in young male, elderly male, young female, and elderly female participants, respectively. Taken together, these novel findings indicate that PhA can be a useful index for easily measuring muscle quality, which has been desired when diagnosing sarcopenia.

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Conflict of interest

All authors of this manuscript declare no conflicts of interest.

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