## Review Article

# Methods, Diagnostic Criteria, Cutoff Points, and Prevalence of Sarcopenia among Older People 

Valéria Pagotto ${ }^{1}$ and Erika Aparecida Silveira ${ }^{2}$<br>${ }^{1}$ Nursing School, Federal University of Goiás, 227 Street, Block 68, s/n, Setor Leste Universitário, 74605-080 Goiânia, GO, Brazil<br>${ }^{2}$ School of Medicine, Federal University of Goiás, Postgraduate Program in Health Sciences, 235 Street s/n, Setor Leste Universitário, 74605-020 Goiânia, GO, Brazil<br>Correspondence should be addressed to Valéria Pagotto; valeriapagotto@gmail.com

Received 10 September 2014; Accepted 11 November 2014; Published 17 December 2014
Academic Editor: Cristiano Capurso
Copyright © 2014 V. Pagotto and E. A. Silveira. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.


#### Abstract

Aim. To identify methods, index, diagnostic criteria, and corresponding cutoff points used to estimate the prevalence of sarcopenia in older people in different countries. Methods. A systematic review was carried out in accordance with PRISMA Statement. The search encompassed the MEDLINE and LILACS databases and was executed during March 2012 using the keyword sarcopenia. Results. A total of 671 studies were identified by the search strategy, and 30 meet all inclusion criteria. Specifically for dual-X-ray absorptiometry, prevalence ranged from $2.2 \%$ to $95 \%$ in men and from $0.1 \%$ to $33.9 \%$ in women. For bioelectrical impedance analysis, the range was from $6.2 \%$ to $85.4 \%$ in men and $2.8 \%$ to $23.6 \%$ in women. Regarding anthropometric and computed tomography, prevalence rates were, respectively, $14.1 \%$ and $55.9 \%$. Conclusions. Heterogeneity in prevalence of sarcopenia was identified, due to diagnostic method choice, cutoff points, and, characteristics of the population as well as reference population. These factors should be considered in research designs to enable comparison and validation of results. Despite the limitations of most studies that indicated high prevalence rates, the results indicate the need for early detection of this syndrome.


## 1. Introduction

Sarcopenia was conceptualized in the last two decades [1] and since then, several studies have been carried out as attempts to clarify definitions for estimation of the issue in the elderly, resulting in a wide diversity of methods and diagnostic criteria [2-4]. As a consequence of such diversity, international research groups have proposed definitions to enable the recommendation of parameters used in the evaluation of sarcopenia [5-8]. The Working Group on Sarcopenia in Older People (EWGSOP) defined it as a syndrome characterized by the progressive and generalized loss of muscle mass, strength, and performance [6]. In 2011, the International Working Group on Sarcopenia (IWGS) defined it as the loss of skeletal muscle mass and strength, associated with the aging process [7].

The study of sarcopenia is important in the areas of public health, geriatrics, and gerontology due to its contribution to adverse outcomes in the elderly $[4,9,10]$, hospitalizations
[10], and early death [11]. However, knowledge on the magnitude of sarcopenia in the elderly population is limited or at least controversial due to the variety of definitions and diagnostic parameters utilized $[3-8,12]$. We did not find systematic reviews focusing on the analysis of sarcopenia magnitude throughout different continents, nor on the different definitions and diagnostic methods for muscle mass evaluation. Analysis of existing studies, including a comparison of the aforementioned aspects, can contribute to the knowledge base on the use of methods and diagnostic criteria and even help direct towards a more operational and less theoretical definition of sarcopenia not only in clinical-epidemiological research but also for health services. Such research can contribute to the efforts to standardize diagnostic criteria applied in different continents and establish the magnitude of sarcopenia in the elderly.

Therefore, the objective of this study was to identify the methods, index, diagnostic criteria, and corresponding cutoff points used to estimate the prevalence of sarcopenia in older
people in different countries, defining global panorama of the issue.

## 2. Methods

A systematic review was carried out following the recommendations for reporting systematic reviews and meta-analyses of the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (The PRISMA Statement) [13].

Searches were carried out in the MEDLINE and LILACS databases until December, 2012, with no restriction on year of publication. Sarcopenia was the only keyword used due to the variety of methods for diagnosing muscle mass, strength, and performance. The following search limits were established: research on humans, in English, French, German, Spanish, or Portuguese, and age group over 45 years, with the keyword in any field. Within the search results obtained, manual searches were then carried out on the bibliographic references cited within the articles.

Subsequently for the identification of studies in the databases, duplicates were excluded and the titles and abstracts of the remaining results were screened, following the eligibility criteria: inclusion of prevalence rates of sarcopenia as well as the methods for measuring muscle mass and for diagnosing sarcopenia. All titles and abstracts were independently screened by two authors. The eligible articles were read in full and those that met all criteria were included.

A tool was developed to analyze data, which contained the following information: authors, year of publication, geographical location of the population, study type, sample size, age range, measurements and techniques for diagnosis of sarcopenia, population used as reference to compare muscle mass, type of evaluation method for muscle mass, indices, diagnostic criteria, cutoff points, cutoff values, and prevalence according to each technique by sex and number of men and women in each study. The confidence intervals (95\% CI) were calculated in STATA 12.0 and were presented in graphtype high-low. If the article did not have the information for calculating the CI, we contacted the corresponding author by e-mail requesting the data. Due to heterogeneity between studies, meta-analysis was not performed.

To assess the quality of articles, we adopted the methodology proposed by Downs and Black [14], whose purpose is to guide auditors in identifying the methodological features of most relevant observational studies. The proposed score is composed of 27 questions that assess clarity of writing (nine items), external validity (two items), internal validity (seven items), confounders (four items), and power of the study (one item). This tool was adapted as described by Monteiro and Victora [15]; because those criteria were originally designed for the evaluation of clinical trials being excluded four questions apply only to this type of study. Thus, the maximum possible score for each item was 24 .

## 3. Results

Through the utilization of search strategies, 854 unduplicated articles were identified: 808 in MEDLINE and 46 in LILACS. After reading of titles and abstracts, 794 articles
were excluded, of which 430 were off-topic and 384 were excluded due to the type of study (review, meta-analysis, clinical trials, and case control). Sixty articles were eligible for a full reading, and after a second round of exclusions, the final sample size was of 28 articles (Figure 1). Critical appraisal of the studies included in our analysis revealed that they were of high quality and credibility.

The mean score of methodological quality was 17.5. Regarding the general characteristics of the articles, crosssectional studies were predominant ( $60 \%$ ), with publications dating from 1998 to 2012. Ten studies were conducted in Asia, nine in North America, and eight in Europe and one was conducted in South America (Brazil) (Table 1).

Twenty-six studies (92.8\%) used exclusively muscle mass for the definition of sarcopenia, while two studies (7.2\%) included mass, strength, and performance, as recommended by the European Sarcopenia Consensus. For this reason we decided to evaluate only muscle mass for comparison purposes (Tables 1, 2, and 3).

To estimate muscle mass, eighteen studies (64.3\%) used dual-energy X-ray absorptiometry (DXA) (Table 1), eight used bioelectrical impedance analysis (BIA) (Table 2), and two used anthropometric measurements (calf circumference) (Table 3).

Within the DXA studies, all 18 used the appendicular muscle mass index (AMMI), defined as the sum of fat-free arm and leg mass in kg (appendicular muscle mass, AMM) divided by the square of the height in meters (AMMI = AMM/height ${ }^{2}$ ). Nine of the studies compared AMMI with other indices: three with total muscle mass (TMM), defined as AMM $\times 1.33 /$ height $^{2}$, and five with mass from a regression analysis adjusting fat mass and height (Table 1).

Three different BIA indices were found. The skeletal muscle index (SMI) adjusted for squared height was used in six studies and adjusted for weight multiplied by 100 was used in three studies (37.5\%). For the calculation of SMI, muscle mass was estimated by the equation: Skeletal Muscle Mass $(\mathrm{SMI})=\left[\left(\right.\right.$ Height ${ }^{2} /$ Resistance $\left.\times 0.401\right)+($ sex $\times 3.825)+($ age $\times-0.071)]+5$, where height is given in cm and resistance in ohms, female $=0$, male $=1$, and age is expressed in years (Bahat et al. 2010). Only one index estimated muscle mass using the DuBois formula: Body Surface Area (BSA) = $\left(\mathrm{kg}^{0.425} \times \mathrm{m}^{0.725}\right) \times 0.007184$ (Table 2).

The two anthropometric studies measured muscle mass using calf circumference, with a cutoff point of 31 cm (Table 3).

Four different criteria for sarcopenia diagnosis were identified: sarcopenia was defined when AMM was two standard deviations (SD) below the mean of a young reference population, by sex ( 20 studies), or when 20th percentile was below the elderly sample distribution (3 studies); sarcopenia was defined by the residual method ( 5 studies); and finally using a cutoff point by Roc curve analysis. Seven studies used the reference population of the USA. Rosetta study has a cutoff point of $7.26 \mathrm{~kg} / \mathrm{m}^{2}$ for men and $5.45 \mathrm{~kg} / \mathrm{m}^{2}$ for women. The other studies used their own young population, with ages ranging from 18 to 40 years. Three studies classified sarcopenia into Class 1 for muscle mass between -1 and -2 standard
Table 1: Diagnostic criteria and prevalence of sarcopenia according to gender estimated by Dual-X-ray-absorptiometry (DXA) method ( $n=18$ ).

|  |  | Country | Study design ( $n$ ) | Age (y) | Index* | Diagnostic criteriaDescription and reference <br> population | Cutoff points |  | Prevalence (\%) |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Authors, y |  |  |  |  |  |  | points <br> Female <br> $\left(\mathrm{kg} / \mathrm{m}^{2}\right)$ | Male <br> (\%) | Female <br> (\%) |
| 1 | Baumgartner et al. $1998 \text { [4] }$ | USA | Cross-sectional <br> Population-based study <br> (808) | >60 | AMMI | -2 SD below gender-specific mean of young adults from Rosseta study (18-40 y) | 7.26 | 5.45 | 28.5 | 33.9 |
| 2 | Melton III et al. 2000 [16] | USA | Cross-sectional <br> Population-based study (300) <br> (community-dwelling) | $\geq 60$ | $\begin{aligned} & \text { TSMI } \\ & \text { AMMI } \end{aligned}$ | -2 SD below gender-specific mean of young adults from Rochester (20-50 y) <br> -2 SD below gender-specific mean of young adults from Rosseta study ( $18-40 \mathrm{y}$ ) | 6.77 7.26 | 4.51 5.45 | 11.3 5.3 | 7.3 4.6 |
| 3 | Tankó et al. 2002 [17] | Denmark | Cross-sectional (754 healthy women) | >60 | AMMI | -2 SD below the normal sex-specific means for young persons of Danish population (19-39 y) | - | 5.40 | - | 12.3 |
| 4 | Iannuzzi-Sucich et al. 2002 [18] | USA | Cross-sectional <br> Community-dwelling <br> (337) | $\geq 65$ | AMMI TSMI | -2 SD below gender-specific mean of young adults from Rosseta study ( $18-40 \mathrm{y}$ ) <br> -2 SD below gender-specific mean of young adults from Rochester (18-40 y) | 7.26 6.77 | 5.45 4.51 | 26.8 11.3 | 22.6 1.5 |
| 5 | Kenny et al. 2003 [19] | USA | Cross-sectional Women users ERT (189) | >59 | AMMI | -2 SD below gender-specific mean of young adults from Rosseta study (18-40 y) | - | 5.45 | - | 23.8 |
| 6 | Gillette-Guyonnet et <br> al. 2003 [20] | France | Cohort <br> EPIDOS <br> (1.321 women) | 75-80 | AMMI | -2 SD below gender-specific mean of young adults from Rosseta study (18-40 y) | - | 5.45 | - | 8.9 |
| 7 | Visser et al. 2003 [21] | Amsterdam | $\begin{aligned} & \hline \text { Cohort } \\ & \text { LASA } \\ & (1.008) \\ & \hline \end{aligned}$ | 55-85 | MMA | Lowest sex-specific 15th percentile of the cohort (loss muscle mass greater than $3 \%$ ) | - | - | 15.7 |  |
| 8 | Newman et al. 2003 [12] | USA | Cohort <br> (2.984) <br> Health ABC Study | 70-79 | AMMI <br> $A M M I^{R}$ | $<20$ th percentile sex-specific distribution <br> $<20$ th percentile sex-specific residuals distribution adjusted for fat mass and height | $\begin{gathered} \hline 7.23 \\ -2.29^{\dagger} \end{gathered}$ | $\begin{gathered} 5.67 \\ -1.73^{\dagger} \end{gathered}$ | 14.1 20.0 | 12.4 24.0 |
| 9 | Lau et al. 2005 [22] | China | Cross-sectional <br> (527) <br> Community-dwelling | >70 | $\begin{gathered} \text { AMMI } \\ \text { TSMI } \end{gathered}$ | -2 SD below gender-specific mean of Chinese young adults (20-40 y) <br> -2 SD below gender-specific mean of Chinese young adults (20-40 y) | $<5.72$ 9.9 | $<4.82$ 8.5 | 12.3 23.4 | 7.2 36.7 |
| 10 | Delmonico et al. 2007 [23] | USA | $\begin{aligned} & \text { Cohort } \\ & \text { (2.976) } \\ & \text { Health ABC Study } \\ & \hline \end{aligned}$ | 70-79 | AMMI <br> $A M M I^{R}$ | $<20$ th percentile sex-specific distribution $<20$ th percentile sex-specific residuals distribution adjusted for fat mass and height | 7.25 | 5.67 | 20.3 20.2 | 20.2 20.3 |
| 11 | Lee et al. 2008 [24] | China | Cross-sectional <br> (4.000) <br> Community-dwelling | $\geq 65$ | AMMI | -2 SD below gender-specific mean of Chinese young adults (20-40 y) | 7.19 | 6.05 | 7.1 | 6.1 |
| 12 | Kim et al. 2010 [25] | Korea | KSOS <br> (526) <br> Community-dwelling | $\geq 60$ | AMMI | -2 SD below gender-specific mean of Korean young adults (20-39 y) | 6.58 | 4.59 | 6.3 | 4.1 |
| 13 | Woo et al. 2009 [26] | China | $\begin{aligned} & \hline \text { Cohort } \\ & (3.153) \\ & \text { Community-dwelling } \\ & \hline \end{aligned}$ | $\geq 65$ | AMMI | -2 SD below gender-specific mean of Chinese young adults (20-40 y) | <7.4 | $<6.4$ | 2.2 | 2.6 |

Table 1: Continued.

*AMMI: appendicular skeletal muscle mass index (appendicular skeletal muscle mass/height ${ }^{2}$ ).

* $A M M I^{\mathrm{R}}=$ ap .
${ }^{\dagger}$ Residuals method.
EPIDOS: Epidemiologie de l'Osteoporose Study.
LASA: longitudinal aging study Amsterdam.
Health ABC Study: The Health, Aging, and Body Composition Study.
KSOS: Korean Sarcopenic Obesity Study.
KNHANES IV: Fourth Korean National Health and Nutritional Examination Surveys.
TABLE 2: Diagnostic criteria and prevalence of sarcopenia according to gender estimated by bioelectrical impedance analysis (BIA) method ( $n=8$ ).

|  | Authors, y | Country | Study design <br> ( $N$ ) | Age (y) | Index* | Diagnostic criteria |  |  | Prevalence (\%) |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  | Description and reference population | Cutoff points |  |  |  |
|  |  |  |  |  |  |  | $\begin{gathered} \text { Male } \\ \left(\mathrm{kg} / \mathrm{m}^{2}\right) \end{gathered}$ | Female $\left(\mathrm{kg} / \mathrm{m}^{2}\right)$ | Male (\%) | Female (\%) |
|  | Janssen et al. 2002 | USA | Cross-sectional NHANES III | $\geq 60$ | SMI ${ }^{\text {\% }}$ | Class 1: -1 to 2 SD below gender-specific mean of young adults (18-39 y) | 37-31\% | 28-22\% | 44.0 | 60.3 |
|  |  |  | (4.504) |  |  | Class 2: -2 SD below gender-specific mean of young adults (18-39 y) | <31\% | <22\% | 7.5 | 12.2 |
| 2 | $\begin{aligned} & \text { Castillo et al. } 2003 \\ & \text { [33] } \end{aligned}$ | USA | Cross-sectional (1.700) community-dwelling | 55-98 | SMI ${ }^{\text {\% }}$ | -2 SD below gender-specific mean of young adults from Pichard study | 47.9\% | 34.7\% | 6.2 | 5.9 |
| 3 | Janssen et al. 2004 [9] | USA | Cohort <br> NHANES III (4.499) | $\geq 60$ | SMI ${ }^{\text {kg/m} 2}$ | Based on ROC curve analysis for moderate dysfunction <br> Based on ROC curve analysis for elevated dysfunction | $\begin{gathered} 8.51-10.75 \\ \leq 8.50 \end{gathered}$ | $\begin{gathered} 5.76-6.75 \\ \leq 5.75 \end{gathered}$ | 53.1 11.2 | 21.9 9.4 |
| 4 | Chien et al. 2008 [34] | Taiwan | Cross-sectional (302) community-dwelling | $\geq 65$ | SMI ${ }^{\mathrm{kg} / \mathrm{m} 2}$ | -2 SD below gender-specific mean of Taiwanese young adults (18 a 40 y) | <8.87 | <6.42 | 23.6 | 18.6 |
| 5 | Tichet et al. 2008 [35] | France | Cross-sectional (218) volunteers of healthcare centers | 60-78 |  | -2 SD below gender-specific mean of French young adults (18 a 39 y) <br> -2 SD below gender-specific mean of young adults (18 a 39 y) | $\begin{gathered} 34.4 \% \\ 8.6 \end{gathered}$ | $\begin{gathered} 26.6 \% \\ 6.2 \end{gathered}$ | 12.5 3.6 | 23.6 2.8 |
| 6 | Bahat et al. 2010 [36] | Turkey | Cohort (217) male nursing home residents | >60 | ASC | -2 SD below gender-specific mean of young adults from control group (24-45 y). | $29.6 \mathrm{~kg} / \mathrm{ASC}$ | - | 85.4 | - |
| 7 | $\begin{aligned} & \text { Landi et al. } 2012 \\ & {[10,37]} \end{aligned}$ | Italy | Cross-sectional (122) <br> Nursing home residents | >70 | SMI ( $\mathrm{kg} / \mathrm{m}^{2}$ ) | -2 SD below gender-specific mean of Italian young adults (18 a 40 y) | <8.87 | <6.42 | 68.0 | 21.0 |
| 8 | $\begin{aligned} & \text { Masanes et al. } 2012 \\ & \text { [38] } \end{aligned}$ | Spain | Cohort <br> (200) | 70-80 |  | -2 SD below gender-specific mean of Spanish young adults (20 a 40 y ) | <8.25 | 6.68 | 10.0 | 33.0 |

* $\mathrm{SMI}^{\mathrm{kg} / \mathrm{m} 2}$ : skeletal muscle index: muscle mass calculated using the bioelectrical impedance analysis equation of Janssen et al. (2000) [3]/height ${ }^{2}$. *SMI ${ }^{*}$ : skeletal muscle index: muscle mass calculated using the bioelectrical impedance analysis equation of Janssen et al. (2000) [3]/weight $\times 100$. NHANES III: National Health and Nutritional Examination Surveys.
TAbLe 3: Diagnostic criteria and prevalence of sarcopenia according to gender estimated by anthropometry $(n=2)$.

| Authors, $y$ | Country | Study design <br> $(N)$ | Age (y) |  | Diagnostic criteria <br> Index | Description and reference <br> population |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |

[^0] EPIDOS: European Patient Information and Documentation Systems (EPIDOS) Study.


Figure 1: Flow diagram for identification, screening, eligibility, and inclusion of articles in systematic review.
deviations from the mean and Class 2 for muscle mass below -2 standard deviations from the mean, both for their reference population. The variations in cutoff points for estimation of muscle mass are shown in Tables 1, 2, and 3.

Considering all methods and diagnostic criteria, prevalence of sarcopenia in the elderly ranged from $0.0 \%$ to $85.4 \%$ in men and $0.1 \%$ to $33.6 \%$ in women. For DXA, prevalence ranged from $0.0 \%$ to $56.7 \%$ in men and $0.1 \%$ to $33.9 \%$ in women (Table 1). For BIA, the range was from $6.2 \%$ to $85.4 \%$ and from $2.8 \%$ to $23.6 \%$, in men and women, respectively (Table 2). Figures 2 and 3 summarize all prevalence and confidence intervals ( $95 \%$ ) of 21 studies for men and 25 studies for women.

## 4. Discussion

This systematic review provides a broad panorama of sarcopenia prevalence in elderly people from five continents, allowing for comparisons of different diagnostic methods and cutoff points, thus contributes to defining the magnitude of the problem in different parts of the world, highlighting lacunae in some geographic areas and the lack of uniformity in diagnostic criteria, and so encourages reflections and propositions on the study of sarcopenia.

The first sarcopenia prevalence studies were only published 10 years after the term was coined in 1989. The first
index proposed for diagnosing sarcopenia by muscle mass was the appendicular muscle mass index (AMMI), which is currently widely used in studies from different countries [12, 22, 23, 25, 26, 31, 41, 42]. When using AMMI, muscle mass is measured by DXA (kg) and the result is compared to a young reference population [4].

The use of the AMMI classification criterion for other populations ( $n=18$ ) has provided a wide range of prevalence, varying from $0.0 \%$ to $56.7 \%$ in men and from $0.0 \%$ to $33.9 \%$ in women. These results can be attributed to racial characteristics, highlighting physical constitution and cultural aspects that imply physical activity levels, dietary regimes, and life quality of the elderly in different countries. This can be exemplified by the low prevalence encountered among the Chinese population, $[25,26,41]$ leading to the authors conclusion that AMMI is not an appropriate method to diagnose sarcopenia in this specific population. The cutoff points for the Chinese [41] population are lower than for Americans [4] ( $<5.72$ versus 7.26 in men and $<4.82$ versus $<5.45$ in women, resp., for Chinese and North-Americans), with young people of the same ethnic group as reference. The mean AMMI of young Asians was approximately 15\% lower than that of Caucasians even after height adjustments [22, 41]. Therefore, low muscle mass in young Asians will result in lower prevalence rates in the elderly. Moreover, sarcopenia may be less prevalent in Asians due to differences


FIGURE 2: Sarcopenia prevalence and CI 95\% in women. (A: DEXA; A1: appendicular skeletal muscle mass index (AMMI); A2: total skeletal muscle mass index (AMMT); and A3: residuals method; B: BIA; B1: skeletal muscle index/height; B2: skeletal muscle index/weight; C: calf circumference).
in risk factors such as a better dietary profile and higher levels of physical activity than Western populations, which act as protective factors against sarcopenia [26].

Although several studies apply AMMI $(n=15)$ and the recommendations to use it [6], other criteria and indices have been proposed to diagnose sarcopenia. Newman et al. [12] proposed a criterion based on the appendicular muscle mass adjusted for body weight (fat mass) and height, where the cutoff point was the 20th percentile of the distribution of linear regression residuals. This method was used in five studies $[12,23,26,41,42]$ and presented better sensitivity in the identification of sarcopenic individuals, particularly among elderly patients with a high prevalence of overweight and obesity [31, 37]. This method is recommended for sarcopenia studies, in overweight and/or obese populations [12].

Melton III et al. [16] developed the TMMI, which was used in four studies [16, 18, 22, 41]. This index also shows important differences in prevalence due to the same factors that explain the variation in prevalence with AMMI. As TMMI identifies the total muscle content, the prevalence of sarcopenia could be higher in comparison with AMMI since appendicular muscle mass represents $75 \%$ of total body
muscle mass [16, 43]. However, prevalence measured by TMMI was lower than that calculated by AMMI and is justified by errors in the estimation of total muscle mass, such as a potential overstatement of water or fat contents, which limit the usefulness of TMMI [43].

Although DXA is precise and is recommended, muscle mass was validated through other measurements in order to enable operationalization and applicability to clinical settings, such as electrical impedance (BIA) and anthropometric measurements. Starting from BIA, Janssen et al. [32] proposed that SMI be adjusted both for height and for weight. Sarcopenia prevalence according to this method also presented significant differences, [33-35] attributed to the different characteristics of study populations and cutoff point references, as well as to the inherent limitations of BIA, which presents a standard error of 9\% [32] in the estimation of muscle mass. The increase in total body water, particularly extracellular water, may result in underestimation of fat body mass and overestimation of lean body mass [3].

Less frequently $(n=3)$ the anthropometric measurement was also utilized to evaluate muscle mass and diagnose sarcopenia, due to the low cost, noninvasive character, and


Figure 3: Sarcopenia prevalence and CI $95 \%$ in men. (A: DEXA; A1: appendicular skeletal muscle mass index (AMMI); A2: total skeletal muscle mass index (AMMT); A3: residuals method; B: BIA; B1: skeletal muscle index/height; B2: skeletal muscle index/weight; C: calf circumference).
basic training requirements. Prevalence found through this method was significantly lower than that obtained with DXA or BIA $[39,40]$.

From the 28 articles selected for review, four types of cutoff points for sarcopenia diagnosis were identified, being two standard deviations below the mean for a young reference population, the most used cutoff point ( $n=17$ ), despite its limitations. Only one cohort study defined sarcopenia as a loss of 3\% of baseline AMM, based on the coefficient of variation for the measurement of AMM using DXA, which was 2$3 \%$ [21]. Visser [44] points out that most definitions include a cutoff point for low muscle mass, but not for loss of muscle mass. The statement of sarcopenia refers to a relative deficiency in muscle mass and does not specify loss [44]. At this point it is discussed that the comparison with elderly population of the same population, noninstitutionalized and with
high life quality standards, could reflect with greater precision the deficiency of muscle mass instead of the comparison with young population. The affirmation is supported by studies that show that after the age of 30 , the musculoskeletal system starts to undergo a progressive loss, with a $1-2 \%$ decrease in muscle mass starting at the age of 50 , which becomes more pronounced after the age of 60 [45]. Caution must be exercised when making comparisons with a young population, as young people have not been exposed to the same factors that older people have experienced throughout their lives, besides the progressive loss of muscle mass that is characteristic of the natural aging process. Thus, studies [46] on the causes of sarcopenia evaluate a wide variety of conditions that go beyond known risk factors, such as sedentary lifestyles, dietary intake, influence of hormones, and cytokine levels, supporting the definition of sarcopenia as a geriatric syndrome.

Despite the differences encountered between the studies, regarding methods and definitions for estimating muscle mass, the present study demonstrates that a substantial proportion of the elderly population has sarcopenia, even in healthy populations. It is questioned, however, what the acceptable progression of loss of muscle mass is as a consequence of the aging progress and what values can identify a harmful loss; that is, that could place the elderly at risk of falling, dependence, and frailty syndrome. These are questions that can direct future research and therefore prospective studies are required and recommended to delineate the natural progression of sarcopenia and its predispositional factors.

The evaluation of sarcopenia, as proposed by the first definitions and by the EWGSOP and IWGS consensus, has been performed in research, even with its inherent limitations. The use of DXA for the estimation of muscle mass guarantees higher reliability and must be the method chosen to evaluate muscle mass in research and for patients of higher clinical complexity. DXA, however, is of difficult operationalization and access in the health service routine, due to elevated cost and specialized professional requirements. It is recommended that other methods, such as BIA and CC, be developed and validated by research devoted to the tracking and consequently to the screening of sarcopenia in health services, due to easiness of application and low cost.

In conclusion, more than one operational definition, it is necessary that the current methods are applied in clinical practice, because sarcopenia presents low visibility in the health services and has not achieved the same space in clinical settings as in research. Therefore, propagation among geriatrics and gerontology healthcare professionals is important and must be included in the context of public health politics.

## Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

## References

[1] I. H. Rosenberg, "Summary comments", The American Journal of Clinical Nutrition, vol. 50, no. 5, pp. 1231-1233, 1989.
[2] S. B. Heymsfield, R. Smith, M. Aulet et al., "Appendicular skeletal muscle mass: measurement by dual-photon absorptiometry," American Journal of Clinical Nutrition, vol. 52, no. 2, pp. 214-218, 1990.
[3] I. Janssen, S. B. Heymsfield, R. N. Baumgartner, and R. Ross, "Estimation of skeletal muscle mass by bioelectrical impedance analysis," Journal of Applied Physiology, vol. 89, no. 2, pp. 465471, 2000.
[4] R. N. Baumgartner, K. M. Koehler, D. Gallagher et al., "Epidemiology of sarcopenia among the elderly in New Mexico," American Journal of Epidemiology, vol. 147, no. 8, pp. 755-763, 1998.
[5] M. Muscaritoli, S. D. Anker, J. Argilés et al., "Consensus definition of sarcopenia, cachexia and pre-cachexia: joint document elaborated by Special Interest Groups (SIG) "cachexia-anorexia in chronic wasting disease" and "nutrition in geriatrics"," Clinical Nutrition, vol. 29, no. 2, pp. 154-159, 2010.
[6] A. J. Cruz-Jentoft, J. P. Baeyens, J. M. Bauer et al., "Sarcopenia: European consensus on definition and diagnosis," Age and Ageing, vol. 39, no. 4, pp. 412-423, 2010.
[7] R. A. Fielding, B. Vellas, W. J. Evans et al., "Sarcopenia : an undiagnosed condition in older adults. Current consensus definition: prevelence, etiology, and consequences. International Working Group on Sarcopenia," Journal of the American Medical Directors Association, vol. 12, no. 4, pp. 249-256, 2011.
[8] J. E. Morley, A. M. Abbatecola, J. M. Argiles et al., "Sarcopenia with limited mobility: an international consensus," Journal of the American Medical Directors Association, vol. 12, no. 6, pp. 403-409, 2011.
[9] I. Janssen, R. N. Baumgartner, R. Ross, I. H. Rosenberg, and R. Roubenoff, "Skeletal muscle cutpoints associated with elevated physical disability risk in older men and women," The American Journal of Epidemiology, vol. 159, no. 4, pp. 413-421, 2004.
[10] F. Landi, R. Liperoti, D. Fusco et al., "Sarcopenia and mortality among older nursing home residents," Journal of the American Medical Directors Association, vol. 13, no. 2, pp. 121-126, 2012.
[11] F. Landi, A. J. Cruz-Jentoft, R. Liperoti et al., "Sarcopenia and mortality risk in frail older persons aged 80 years and older: results from ilSIRENTE study," Age Ageing, vol. 42, no. 2, pp. 203-209, 2013.
[12] A. B. Newman, V. Kupelian, M. Visser et al., "Sarcopenia: alternative definitions and associat ions with lower extremity function," Journal of the American Geriatrics Society, vol. 51, no. 11, pp. 1602-1609, 2003.
[13] D. Moher, A. Liberati, J. Tetzlaff, and D. G. Altman, "Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement," Annals of Internal Medicine, vol. 151, no. 4, pp. 264-269, 2009.
[14] S. H. Downs and N. Black, "The feasibility of creating a checklist for the assessment of the methodological quality both of randomised and non-randomised studies of health care interventions," Journal of Epidemiology and Community Health, vol. 52, no. 6, pp. 377-384, 1998.
[15] P. O. A. Monteiro and C. G. Victora, "Rapid growth in infancy and childhood and obesity in later life-a systematic review," Obesity Reviews, vol. 6, no. 2, pp. 143-154, 2005.
[16] L. J. Melton III, S. Khosla, C. S. Crowson, M. K. O'Connor, W. M. O'Fallon, and B. L. Riggs, "Epidemiology of sarcopenia," Journal of the American Geriatrics Society, vol. 48, no. 6, pp. 625-630, 2000.
[17] L. B. Tankó, L. Movsesyan, U. Mouritzen, C. Christiansen, and O. L. Svendsen, "Appendicular lean tissue mass and the prevalence of sarcopenia among healthy women," Metabolism: Clinical and Experimental, vol. 51, no. 1, pp. 69-74, 2002.
[18] M. Iannuzzi-Sucich, K. M. Prestwood, and A. M. Kenny, "Prevalence of sarcopenia and predictors of skeletal muscle mass in healthy, older men and women," The Journals of Gerontology Series A: Biological Sciences and Medical Sciences, vol. 57, no. 12, pp. M772-M777, 2002.
[19] A. M. Kenny, L. Dawson, A. Kleppinger, M. Iannuzzi-Sucich, and J. O. Judge, "Prevalence of sarcopenia and predictors of skeletal muscle mass in nonobese women who are long-term users of estrogen-replacement therapy," The Journals of Gerontology, Series A: Biological Sciences and Medical Sciences, vol. 58, no. 5, pp. 436-440, 2003.
[20] S. Gillette-Guyonnet, F. Nourhashemi, S. Andrieu et al., "Body composition in French women 75+ years of age: the EPIDOS study," Mechanisms of Ageing and Development, vol. 124, no. 3, pp. 311-316, 2003.
[21] M. Visser, D. J. H. Deeg, and P. Lips, "Low vitamin D and high parathyroid hormone levels as determinants of loss of muscle strength and muscle mass ( sarcopenia ): the longitudinal Aging Study Amsterdam," The Journal of Clinical Endocrinology and Metabolism, vol. 88, no. 12, pp. 5766-5772, 2003.
[22] E. M. C. Lau, H. S. H. Lynn, J. W. Woo, T. C. Y. Kwok, and L. J. Melton III, "Prevalence of and risk factors for sarcopenia in elderly Chinese men and women," Journals of Gerontology A. Biological Sciences and Medical Sciences, vol. 60, no. 2, pp. 213216, 2005.
[23] M. J. Delmonico, T. B. Harris, J.-S. Lee et al., "Alternative definitions of sarcopenia, lower extremity performance, and functional impairment with aging in older men and women," Journal of the American Geriatrics Society, vol. 55, no. 5, pp. 769-774, 2007.
[24] J. S. W. Lee, T. W. Auyeung, T. Kwok, E. M. C. Lau, P. C. Leung, and J. Woo, "Associated factors and health impact of sarcopenia in older Chinese men and women: a cross-sectional study," Gerontology, vol. 53, no. 6, pp. 404-410, 2008.
[25] T. N. Kim, M. S. Park, S. J. Yang et al., "Prevalence and determinant factors sarcopenia in patients with type 2 diabetes: the Korean Sarcopenic Obesity Study (KSOS)," Diabetes Care, vol. 33, no. 7, pp. 1497-1499, 2010.
[26] J. Woo, J. Leung, A. Sham, and T. Kwok, "Defining sarcopenia in terms of risk of physical limitations: a 5-year follow-up study of 3,153 Chinese men and women," Journal of the American Geriatrics Society, vol. 57, no. 12, pp. 2224-2231, 2009.
[27] Y. Rolland, V. Lauwers-Cances, C. Cristini et al., "Difficulties with physical function associated with obesity, sarcopenia, and sarcopenic-obesity in community-dwelling elderly women: the EPIDOS (EPIDemiologie de l'OSteoporose) Study," The American Journal of Clinical Nutrition, vol. 89, no. 6, pp. 1895-1900, 2009.
[28] K. Sanada, M. Miyachi, M. Tanimoto et al., "A cross-sectional study of sarcopenia in Japanese men and women: reference values and association with cardiovascular risk factors," European Journal of Applied Physiology, vol. 110, no. 1, pp. 57-65, 2010.
[29] X. Wen, M. Wang, C. M. Jiang, and Y. M. Zhang, "Are current definitions of sarcopenia applicable for older Chinese adults?" Journal of Nutrition, Health \& Aging, vol. 15, no. 10, pp. 847-851, 2011.
[30] Y.-S. Kim, Y. Lee, Y.-S. Chung et al., "Prevalence of sarcopenia and sarcopenic obesity in the Korean population based on the Fourth Korean National Health and Nutritional Examination Surveys," The Journals of Gerontology, Series A: Biological Sciences and Medical Sciences, vol. 67, no. 10, pp. 1107-1113, 2012.
[31] D. S. Domiciano, C. P. Figueiredo, J. B. Lopes et al., "Discriminating sarcopenia in community-dwelling older women with high frequency of overweight/obesity: the São Paulo Ageing \& Health Study (SPAH)," Osteoporosis International, vol. 24, no. 2, pp. 595-603, 2013.
[32] I. Janssen, S. B. Heymsfield, and R. Ross, "Low relative skeletal muscle mass (sarcopenia) in older persons is associated with functional impairment and physical disability," Journal of the American Geriatrics Society, vol. 50, no. 5, pp. 889-896, 2002.
[33] E. M. Castillo, D. Goodman-Gruen, D. Kritz-Silverstein, D. J. Morton, D. L. Wingard, and E. Barrett-Connor, "Sarcopenia in elderly men and women: the Rancho Bernardo study", American Journal of Preventive Medicine, vol. 25, no. 3, pp. 226-231, 2003.
[34] M. Y. Chien, T. Y. Huang, and Y. T. Wu, "Prevalence of sarcopenia estimated using a bioelectrical impedance analysis prediction equation in community-dwelling elderly people in

Taiwan," Journal of the American Geriatrics Society, vol. 56, no. 9, pp. 1710-1715, 2008.
[35] J. Tichet, S. Vol, D. Goxe, A. Salle, G. Berrut, and P. Ritz, "Prevalence of sarcopenia in the French senior population," Journal of Nutrition, Health and Aging, vol. 12, no. 3, pp. 202206, 2008.
[36] G. Bahat, B. Saka, F. Tufan et al., "Prevalence of sarcopenia and its association with functional and nutritional status among male residents in a nursing home in Turkey," Aging Male, vol. 13, no. 3, pp. 211-214, 2010.
[37] F. Landi, R. Liperoti, D. Fusco et al., "Prevalence and risk factors of sarcopenia among nursing home older residents," Journals of Gerontology-Series A Biological Sciences and Medical Sciences, vol. 67, no. 1, pp. 48-55, 2012.
[38] F. Masanes, A. Culla, M. Navarro-Gonzalez et al., "Prevalence of sarcopenia in healthy community-dwelling elderly in an urban area of Barcelona (Spain)," Journal of Nutrition, Health and Aging, vol. 16, no. 2, pp. 184-187, 2012.
[39] Y. Rolland, V. Lauwers-Cances, M. Cournot et al., "Sarcopenia, calf circumference, and physical function of elderly women: a cross-sectional study," Journal of the American Geriatrics Society, vol. 51, no. 8, pp. 1120-1124, 2003.
[40] L. P. Bastiaanse, T. I. M. Hilgenkamp, M. A. Echteld, and H. M. Evenhuis, "Prevalence and associated factors of sarcopenia in older adults with intellectual disabilities," Research in Developmental Disabilities, vol. 33, no. 6, pp. 2004-2012, 2012.
[41] X. Wen, M. Wang, C.-M. Jiang, and Y.-M. Zhang, "Are current definitions of sarcopenia applicable for older Chinese adults?" Journal of Nutrition, Health and Aging, vol. 15, no. 10, pp. 847851, 2011.
[42] M. Di Monaco, C. Castiglioni, F. Vallero, R. Di Monaco, and R. Tappero, "Sarcopenia is more prevalent in men than in women after hip fracture: a cross-sectional study of 591 inpatients," Archives of Gerontology and Geriatrics, vol. 55, no. 2, pp. e48e52, 2012.
[43] D. N. Proctor, P. C. O'Brien, E. J. Atkinson, and K. S. Nair, "Comparison of techniques to estimate total body skeletal muscle mass in people of different age groups," American Journal of Physiology-Endocrinology and Metabolism, vol. 277, no. 3, pp. E489-E495, 1999.
[44] M. Visser, "Towards a definition of sarcopenia-results from epidemiologic studies," The Journal of Nutrition Health and Aging, vol. 13, no. 8, pp. 713-716, 2009.
[45] R. J. Zacker, "Health-related implications and management of sarcopenia," Journal of the American Academy of Physician Assistants, vol. 19, no. 10, pp. 24-29, 2006.
[46] Y. Rolland, S. Czerwinski, G. A. Van Kan et al., "Sarcopenia: its assessment, etiology, pathogenesis, consequences and future perspectives," Journal of Nutrition, Health and Aging, vol. 12, no. 7, pp. 433-450, 2008.


[^0]:    * Bastiaanse et al. 2012 [40]: EWGSOP criteria. The prevalence using only muscle mass estimated by CC was $9.1 \%$ in both sexes.

