




Application of ultrasound for muscle assessment in sarcopenia: towards standardized measurements

Stany Perkisas^{1,19}  · Stéphane Baudry^{2,19} · Jürgen Bauer³ · David Beckwée^{4,5} · Anne-Marie De Cock¹ · Hans Hobbelen^{6,7,20} · Harriët Jager-Wittenaar^{6,7,21} · Agnieszka Kasiukiewicz⁸ · Francesco Landi⁹ · Ester Marco^{10,11,12} · Ana Merello¹³ · Karolina Piotrowicz¹⁴ · Elisabet Sanchez¹³ · Dolores Sanchez-Rodriguez^{12,15,16,17} · Aldo Scafoglieri^{18,19} · Alfonso Cruz-Jentoft¹³ · Maurits Vandewoude^{1,19}

Received: 15 June 2018 / Accepted: 29 August 2018
© European Geriatric Medicine Society 2018

Abstract

Purpose Measurement of muscle mass is paramount in the screening and diagnosis of sarcopenia. Besides muscle quantity however, also quality assessment is important. Ultrasonography (US) has the advantage over dual-energy X-ray absorptiometry (DEXA) and bio-impedance analysis (BIA) to give both quantitative and qualitative information on muscle. However, before its use in clinical practice, several methodological aspects still need to be addressed. Both standardization in measurement techniques and the availability of reference values are currently lacking. This review aims to provide an evidence-based standardization of assessing appendicular muscle with the use of US.

Methods A systematic review was performed for ultrasonography to assess muscle in older people. Pubmed, SCOPUS and Web of Sciences were searched. All articles regarding the use of US in assessing appendicular muscle were used. Description of US-specific parameters and localization of the measurement were retrieved.

Results Through this process, five items of muscle assessment were identified in the evaluated articles: thickness, cross-sectional area, echogenicity, fascicle length and pennation angle. Different techniques for measurement and location of measurement used were noted, as also the different muscles in which this was evaluated. Then, a translation for a clinical setting in a standardized way was proposed.

Conclusions The results of this review provide thus an evidence base for an ultrasound protocol in the assessment of skeletal muscle. This standardization of measurements is the first step in creating conditions to further test the applicability of US for use on a large scale as a routine assessment and follow-up tool for appendicular muscle.

Keywords Sarcopenia · Ultrasound · Muscle assessment

Introduction

Sarcopenia is becoming one of the biggest health care challenges that arises together with increasing age expectations [1]. It is associated with mortality, functional decline, disability, a higher rate of falls, a higher incidence

of hospitalizations, increased health care costs and a lower quality of life [2]. Therefore, clear cutoffs are desirable for an early diagnosis. However, there is still some debate about a universal definition. The currently most used operational definition is the age-related decline of muscle mass, together with the decline of strength and/or function [3]. This definition can still be regarded as largely heterogeneous. This debate is continued in the use of specific cutoff points for muscle mass, as is clearly seen in the many criteria that are proposed by various international organizations [3–8].

When applying these criteria, it is evident that in diagnosing sarcopenia, muscle mass still holds the most weight, more than strength or function. This also finds its way in a case finding algorithm, published by the European Working Group for Sarcopenia in Older People [3], where there can be no diagnosis of sarcopenia without the assessment

The authors are part of the SARCUS working group on behalf of the Sarcopenia Special Interest Group of the European Geriatric Medicine Society.

Electronic supplementary material The online version of this article (<https://doi.org/10.1007/s41999-018-0104-9>) contains supplementary material, which is available to authorized users.

✉ Stany Perkisas
Stany.perkisas@zna.be

Extended author information available on the last page of the article

of muscle mass. This focus on muscle mass, however, gives way to a few practical implications, as there seems to be a large cleft between guidelines and clinical practice.

First of all, the prevalence of sarcopenia is highly dependent on the diagnostic method used [9]. As stated, there are many different cutoff points for diagnosing sarcopenia through muscle mass. All are either based on dual-energy X-ray absorptiometry (DXA) or bio-impedancemetry (BIA); currently, there are no cutoff values for either CT or MRI. DXA cutoff points are based on skeletal muscle mass index (SMI), which is the sum of the all the appendicular muscle mass, divided by height. For DXA SMI, three groups of cutoff points are used [10–12]. BIA also uses SMI, with one cutoff point proposition based on BIA-predicted skeletal muscle mass [13] and the other based on absolute muscle mass [14, 15]. It is unclear in how far both DXA and BIA are available in clinical practice.

Secondly, it has become clear that measuring muscle quantity, i.e., muscle mass, in itself is only part of the problem of age-related muscle degradation. Also the muscle architectural qualities—here used synonymously as muscle quality—need to be assessed, as volume in itself has no linear relation with either strength or function. Muscle quality is a broad term and is in current literature used in two meanings. On one hand, it can mean the muscle strength or muscle power per unit of muscle mass [16–18]. On the other hand—as in this study—it can be used as description of the components of the muscle described [19]. The neural component in the process of sarcopenia could possibly amount up to 50% [19], but it is clear that shear volume alone does not explain the rest. Measuring the components of this volume is vital [20, 21]. The assessment of these components cannot be done either by DXA or BIA [22], although there is some evidence that multi-frequency bio-impedance and segmental bioelectrical impedance spectroscopy could measure muscle composition to a certain degree and give information regarding skeletal muscle physiology [23, 24]. For assessing muscle quality, computerized tomography (CT) and magnetic resonance imaging (MRI) are considered ‘gold standards’ [25], in as far this is possible, because many different scanning techniques exist: CT muscle attenuation [26], diffusion tensor MRI [27], Dixon MRI [28], proton magnetic resonance spectroscopy (MRS) [29], ¹³C-MRS [30] and ³¹P-MRS [31]. For both CT and MRI, whole body scanning can be done, as also single/multiple slice scanning [32–34]. The former are considered as superior, but adequate data are present to use the latter as a convenient alternative [3]. In clinical practice, however, CT and MRI are not practical, due to the limited availability, lack of portability, high cost and radiation exposure (CT). Also, currently there are no cutoff values for either CT or MRI. This way, current guidelines and

criteria fall short, in the way that they are often not applicable in a clinical routine setting due to practical reasons as stated above, and do not yet have incorporated the quality aspect of muscle mass. There is an important need for an instrument that can give information about both muscle quantity and quality, that is cheap and easily available in routine practice, and that can be used in large population-based screenings.

Ultrasonography or ultrasound (US) could fill this gap.

US is a well-studied technique that has already proven its worth in the detection of neuromuscular pathology with positive predictive values of up to 90% [35] and in the assessment of muscle–tendon interactions [36]. US is a portable, inexpensive, non-invasive technique without using ionising radiation that also has a high repeatability. Regarding muscle mass assessment, US-based measurements have a strong positive correlation with DEXA [37–40], CT [41] and MRI-based measurements [42–44]. For the estimation of muscle quality and quantity, US is a valid and accurate technique [45, 46]. The validity of ultrasound to discern architectural properties has also been demonstrated in cadaver validation studies [47–51]. It has good intra- and interrater reliability [45], as well as test–retest reliability [52], both in an elderly people [45] as in a younger population [53]. However, it is unclear which anatomical site is best to be used for specific outcomes, e.g., prediction of total skeletal muscle mass [54]. It must be taken into account that not all peripheral muscles decline alike [55, 56]. In this regard, prediction equations for total skeletal muscle mass need further validation [45]. Also, there are hardly any reference data currently available, as there is no standardization of the measurement technique. To our knowledge, there are only four studies giving limited reference data, for muscle thickness and echo intensity of selected muscles [57–60].

This review aims to provide a standardization of ultrasound measurements for assessing muscle mass in the assessment of sarcopenia. To the best of our knowledge, there are no hard clinical data to prefer assessing one muscle or set of muscles over another, or to choose one type of measurement and disregards the others. Therefore, an overview will be given of how appendicular muscles are measured in the literature so far, followed by a proposition of how to do a muscle assessment in a standardized way. These propositions will be based on the literature and consensus within the review group. Among these propositions, there will be a list summarizing which information needs to be minimally included in a protocol when planning future studies. This way, we hope to advance the study of the application of ultrasound in sarcopenia assessment in different settings [61, 62].

Methods

Registration

The protocol for this systematic review has been registered at PROSPERO (Registration number CRD42018085587).

Search strategy

The search strategy was set up based on three main components: elderly (population) [63], ultrasound (exposure), and muscle (outcome). For this, a modified PECO model for clinical questions was used. The search was performed in Pubmed, Web of Science and Cochrane Library, up until the 20th of January 2018. All eligible studies in English, German, French and Dutch were screened for their applicability. Studies regarding the use of ultrasound in the assessment of muscle mass were considered for this review. Bibliographic lists of included papers were hand-searched for additional studies. Animal studies, studies using cadaver specimens, studies assessing non-appendicular muscle, case reports, letters to the editor, editorials and (systematic) reviews were excluded.

Search structure for PubMed was as follows: (((Elderly[tiab] OR community-dwelling[tiab] OR geriatric[tiab] OR Frailty[tiab] OR Ageing[tiab] OR elders[tiab] OR Frail[tiab] OR “postmenopausal women”[tiab] OR aging[tiab] OR older[tiab] OR residents[tiab] OR “old people”[tiab] OR nursing homes[mh] OR aging[mh] OR frail elderly[mh] OR homes for the aged[mh] OR aged, 80 and over[mh]))) AND (((((((Ultrasonography[Mesh] OR Ultrasound) OR Echograph* OR Ultrasonograph*) OR Ultrasonic) OR Echotomograph* OR Sonograph*)) AND (“Muscles”[Mesh] OR “Lean tissue” OR “Lean mass” OR “Lean body mass” OR muscle OR “Fat free mass”))).

Search structure for SCOPUS was as follows: ((TITLE-ABS-KEY (elder*)) OR (TITLE-ABS-KEY (community-dwelling)) OR (TITLE-ABS-KEY (geriatric)) OR (TITLE-ABS-KEY (frail*)) OR (TITLE-ABS-KEY (ag*ing)) OR (TITLE-ABS-KEY (“postmenopausal women”)) OR (TITLE-ABS-KEY (old*)) OR (TITLE-ABS-KEY (resident*)) OR (TITLE-ABS-KEY (“nursing homes”))) AND ((TITLE-ABS-KEY (ultrasound)) OR (TITLE-ABS-KEY (echograph*)) OR (TITLE-ABS-KEY (ultrasonograph*)) OR (TITLE-ABS-KEY (ultrasonic)) OR (TITLE-ABS-KEY (echotomograph*)) OR (TITLE-ABS-KEY (sonograph*)) AND ((TITLE-ABS-KEY (muscle*)) OR (TITLE-ABS-KEY (“Lean tissue”)) OR (TITLE-ABS-KEY (“Lean mass”)) OR (TITLE-ABS-KEY (“Lean body mass”)) OR (TITLE-ABS-KEY (“Fat free mass”))).

Search structure for Web of Science was as follows: Elder* OR community-dwelling OR geriatric OR Frail* OR Ageing OR Frail OR “postmenopausal women” OR old* OR resident* OR “nursing homes”. Ultrasound OR Echograph* OR Ultrasonograph* OR Ultrasonic OR Echotomograph* OR Sonograph*. Muscle* OR “Lean tissue” OR “Lean mass” OR “Lean body mass” OR “Fat free mass”.

An overview of the study selection process is shown in Fig. 1. After deleting duplicates, abstracts were gathered. Abstracts were divided among 12 independent reviewers, experienced in either geriatrics, physical therapy, radiology or body composition. One other reviewer (SP) screened all the abstracts. Title and abstract of all manuscripts were screened for eligibility, reviewers being blinded from each other, using the Rayyan web-based software [64]. Disagreements were resolved by consensus within the group. All review articles, case reports, letters to the editor, and editorials were excluded. Then, the selected articles were used for full-text reading. Again, the full-text articles were divided among the same 12 independent reviewers as in the previous step. However, for the reviewers, the full texts to be reviewed were not the same as the abstracts that were reviewed. In other words, each reviewer reviewed two different sets of publications. Reasons for exclusion of an article after full-text assessment were: absence of clear description of location of measuring point, absence of clear description of muscle measured, content being article outside the scope of the manuscript, or referencing to another article regarding measurement technique/location. For the last category, the article to which was referred for measurement technique/location, was checked for inclusion. If not already included, it was added.

Results

Search strategy

The initial search yielded 17.579 abstracts (PubMed = 5565, SCOPUS = 7255, Web of Science = 4759). There was one additional record identified through other sources. After deleting duplicates ($n = 2751$), 14.829 abstracts were screened. Of these, 359 articles were withheld. After full-text assessment, 76 articles were used in this review. See also Fig. 1 for details about the study selection process.

All articles included in this paper are detailed in a supplemental volume (Table S1), with cohort size, age (median or range) and ethnicity.

Mean cohort size was 58 (median 44, range 9–347). One study did not mention the cohort size. Two papers did not have cohorts as they were suggestions for US protocols.

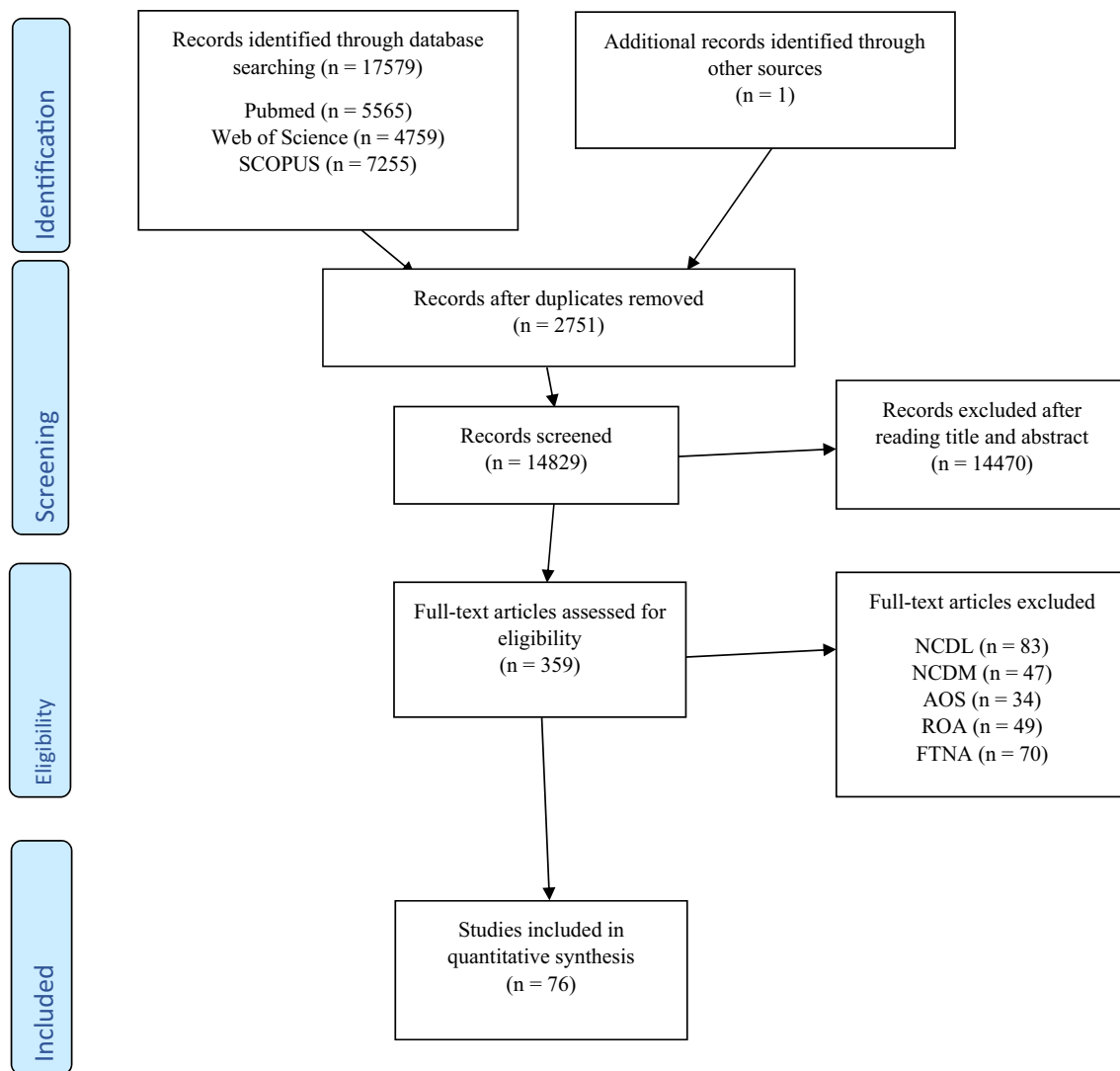


Fig. 1 Overview of the study selection process using the PRISMA 2009 flow chart [166]. *NCDL* no clear description of location of measuring point, *NCDM* no clear description of muscle measured,

AOS article outside of scope of manuscript, *ROA* referencing to other articles for measurement technique, *FTNA* full-text not available

Age was clearly described in 38 studies. In 33 studies, only ranges were given. Three studies did not mention age. In the two protocol studies, no patients were used so no age was noted.

Ethnicity was very poorly described explicitly. Eight studies mentioned Asian ethnicity (one Chinese, seven Japanese), one study mentioned Caucasian ethnicity, and one study mentioned white/Indian ethnicity. One study mentioned different ethnicities: Caucasian 87%, Asian 29%, native American 3%, African 1%. In two studies (protocol), ethnicity was not applicable.

The sex of study subjects was not included in this review.

As there is no gold standard in US muscle assessment, it was difficult to have a good quality assessment of the

included articles. Although a multitude of research questions were present, it must be noted that no studies compared different US techniques in the assessment of muscle.

Patient positioning pre-investigation

Only seven studies stated clearly which subject side was assessed. In five studies, the dominant side was used [65–69]. Only one study specified using the non-dominant side [70]. One study used only the right side, not mentioning if this was the dominant side or not [71].

Regarding the state of the muscle assessed, one article mentioned letting the test subject perform three maximum contractions of the muscle assessed before measurement

Table 1 Overview of different patient positioning used in muscle assessment

	Supine				Lying on non-dominant side, knees 10°	Prone	Sitting			Standing	
	Full extension	Knees at 10°	Semi-Fowler position	Legs at 50°			Knee angle 115° and hip angle 140°	Sitting			
								Hips 90°, knees 90°	Hips 90°, knees 60°		Hips 85°, knees 0°
Lower limb											
Rectus femoris	X	X	X	X				X	X	X	
Vastus lateralis	X	X		X	X			X			
Vastus medialis	X										
Vastus intermedius	X		X					X	X	X	
Quadriceps (all 4 muscles)	X							X	X		
Biceps femoris						X					
Tibialis anterior	X										
Gastrocnemius medialis						X		X		X	
Soleus						X		X			
Upper limb											
Biceps brachii	X										
Triceps brachii										X	

[65]. Two studies had the test subjects being relaxed for 15 min, one to allow fluid shifts to stabilize [72], the other to avoid muscle contraction-induced fluid shifts and muscle blood flow during the measurements [73]. The latter, therefore, also specified to have done all ultrasound measurement before any functional testing. Two groups refrained the study subjects from rigorous exercises, one only for the same day [74], one for the last 48 h [75].

The position of the study subject was dependent on the specific muscle investigated, but also herein there were some differences. For an overview of the different positions used in selected muscles, see Table 1. In the supplemental material, a table is included that indicates which study assessed which muscle at a certain position (Table S2).

For supine and prone positions, study subject is regarded as having hips and shoulders in neutral position, knees and elbows in full extension, ankles at 90°, unless specified otherwise. It has to be mentioned that it is important to correctly describe the positioning, as the angle of joints is not standardized in one direction, e.g., knees in 150° or 30° could mean the same. In none of the studies, it was described whether the supine position included the feet being of the table or not. One study, assessing the gastrocnemius medialis, specified having the tibiotalar joint angle at 115°, for which they had a cast specially made for all participants [69].

Upper leg muscles

- Rectus femoris was assessed in supine position in full extension [65, 68, 72, 76–86], with knees at 10° [87–89], in semi-Fowler position [90–92]. It was assessed in sitting position with hips and knees at 90° [66, 93] and with hips at 90° and knees at 60° [94]. Lastly, it was also assessed standing [95, 96].
- Vastus lateralis was assessed in supine position in full extension [72, 78, 84, 85, 89, 97–101], with knees at 10° [88], with legs at 50° [71] and with knee angle at 115° and hip angle at 140° [102]. It was assessed lying in the non-dominant side with knees at 10° lateralis [87]. Lastly, it was assessed sitting with hips and knees at 90° [103].
- Vastus medialis was assessed in supine position in full extension [72, 84, 104].
- Vastus intermedius was assessed in supine position in full extension [72, 76–79, 84] and in semi-Fowler position [90]. It was assessed in sitting position with hips at 90° and knees at 60° [94]. Lastly, it was also assessed standing [96].
- Quadriceps—comprising all 4 muscles: rectus femoris, vastus medialis, vastus intermedius and vastus lateralis—were assessed in supine position in full extension [105].

It was also assessed in sitting position with hips at 90° and knees at 60° [106].

- Biceps femoris was only assessed in prone position [78].

Lower leg muscles

- Tibialis anterior was only assessed in full extension [97].
- Gastrocnemius medialis was assessed in prone position [78, 82, 97, 102, 107–111] and in sitting position with hips and knees at 90° [69, 103, 112].
- Soleus was assessed in prone position [78] and in sitting position with hips and knees at 90° [112].

Upper arm muscles

- Biceps brachii was only assessed in full extension [57, 97, 113].
- Triceps brachii was assessed standing [114, 115]. One study that assessed triceps brachii did not mention the patient position [116].

System and system settings

Many different types of ultrasound machines were used, from various manufacturers (SonoSite, GE Healthcare, Siemens, Mindray, Philips, Toshiba, Hitachi Aloka Medical, Esaote, Fukuda Denshi, Hewlett-Packard, Telemed). A summary of the different types used is beyond the scope of this article.

Every research group used B-mode ultrasound, and all but one used a linear transducer probe. The only article using a curved transducer had as preposition the validation of the use of a curved versus a linear transducer [91].

The length of the transducer varied from 3.8 to 7 cm. The frequency used was described in 63 cases, and ranged from 3 to 15 MHz.

In 13 cases, the bandwidth instead of the exact frequency was described: 10–15 MHz ($n=1$), 7–12 MHz ($n=3$), 6–13 MHz ($n=1$), 5–12 MHz ($n=2$), 5–10 MHz ($n=3$), 3–13 MHz ($n=1$), 3–12 MHz ($n=1$), 3–11 MHz ($n=1$).

In 50 articles, a clear frequency was described. The most common used frequency was 7.5 MHz ($n=23$); further frequencies used were 15 MHz ($n=1$), 13.6 MHz ($n=1$), 12 MHz ($n=7$), 10 MHz ($n=6$), 9 MHz ($n=3$), 8 MHz ($n=7$), 5 MHz (linear array, $n=6$), 5 MHz (curved array, $n=1$).

No information on the inclination of the probe was noted.

Some additional system settings were described. Most were on image depth (focus point) and general gain.

Image depth was described in 7 cases: 45 mm (mm) [71], 50 mm [87, 88, 117], 60 mm [101, 118] and 70 mm [72].

General gain (in decibels, dB) was described in 15 cases: was either set to 50 dB [82, 83, 87, 117, 119], 58 dB [120], 68 dB [71, 108, 111, 121], 85 dB [94], 86 dB [122] or 90 dB [72, 84, 123].

Time gain compensation was described as being in a neutral position in two cases [72, 124].

Dynamic range was described in five cases: being set to 40 [121], set to 69 [120], or set to 72 [83, 117, 119].

Compression—which is altering the display of the range of echo intensities to end up with a lower amount of different shades of gray—was described in one study as set to 70 dB [122].

No further system settings were described.

For image post-processing and measurement, 35 of the 75 articles mentioned the use of additional software. These included Image J [125] ($n=26$), Matlab (The MathWorks, Inc., Natick, Massachusetts, United States) ($n=2$) and Photoshop (Adobe, Adobe Systems Incorporated, San Jose, California, United States) ($n=7$).

Components and measuring points

Five main components were distilled from the literature: muscle thickness (MT), pennation angle (PA), fascicle

length (Lf), echo intensity (EI) and cross-sectional area (CSA).

Four studies mentioned that all measurements were done three times [66, 67, 91, 97], and after that, the mean value was taken. Two other studies mentioned doing all measurements five times [89, 97]. One of the latter two discarded the highest and lowest value, and then took the mean value of the remaining three [89].

Muscle thickness was defined as the distance between deep and superficial aponeurosis [97, 109]. It can be expressed either in centimetres (cm) or in millimetres (mm).

Cross-sectional area was divided into two: anatomical cross-sectional area (ACSA) and physiological cross-sectional area (PCSA). Anatomical cross-sectional area was defined as the area of cross-section of a muscle perpendicular to its longitudinal axis. Physiological cross-sectional area was defined as the area of cross-section of a muscle perpendicular to its fibers. In non-pennate muscle, ACSA and PCSA are the same; in pennate muscles they are not. ACSA underestimates the number of total fibers in a pennated muscle. Muscle strength is more correlated with PCSA than with ACSA because the former represents the maximal number of acto-myosin crossbridges that can be activated in parallel during contraction [126]. Therefore, when studying muscle strength, it is not advised to only measure ACSA.

Table 2 Measuring points and components of quadriceps muscles used in assessment

Proximal point	Distal point	Distance	MT	CSA	ACSA	PCSA	FL	PA	EI
Greater trochanter	Popliteal crease	60–70%	RF						
	Lateral condyle	30%	VM, VL	RF			VL	VM, VL	VM, VL
		50%	RF, VL, VI	RF, VL	VL		VL	RF, VL, VI	RF, VL, VI
		2/3	RF, VL	VL			VL	VL	VL
		50%	RF, VI						
	Articular cleft of knee	50%			RF	RF			
Proximal border of patella	50%								
Anterior superior iliac spine	Lateral condyle	2/3	RF, VI						RF, VI
	Proximal border of patella	50%	RF, VI, VL						RF, VI
		60%	RF, VL						
		2/3	RF, VL, VI						VL
		3/5		RF					
	75%				RF				
	Midpoint of patella	50%	RF, VI						
Distal border of patella	50%	RF		RF					
Anterior inferior iliac spine	Proximal border of patella	50%	RF, VI	RF	RF		RF	RF	RF, VI
	Proximal border of patella	10 cm proximal to distal point	RF						
–	–	15 cm proximal to distal point		RF	RF				

MT muscle thickness, CSA cross-sectional area, ACSA anatomical cross-sectional area, PCSA physiological cross-sectional area, FL fascicle length, PA pennation angle, EI echo intensity, RF rectus femoris, VL vastus lateralis, VM vastus medialis, VI vastus intermedius

Measurements of CSA were usually presented in square centimeter. Some studies mentioned to measure the CSA by manually drawing the circumference of the muscle with a cursor [66, 91, 127]. One study normalized the CSA to body mass to represent it as a relative measure of quadriceps muscle [71].

The pennation angle (Ap) was defined as the angle of insertion of muscle fiber fascicles into the deep aponeurosis [69, 97]. The angle at which fibers in a pennate arrangement are oriented relative to the longitudinal axis varies from muscle to muscle [128]. The pennation angle is proportional to the number of sarcomeres packed in parallel along the aponeurosis and closely related to the force-generating capacity of the muscle [109].

Fascicle length (Lf) was defined as the length of the fascicular path between the insertions of the fascicle into the superficial and deep aponeuroses. In the cases where the fascicle extended outside of the acquired ultrasound image, the length of the missing portion of the fascicle was estimated by extrapolating linearly both the fascicular path, visible in the image, and the aponeurosis [65, 69, 97, 129, 130]. Another method of estimating the fascicle length is using a formula: multiplying the muscle thickness times the hypotenuse of the pennation angle inversed [131] or, stated differently, dividing the muscle thickness by the hypotenuse of the pennation angle [99]. These formulae do not account for fascicle curvature [73]. Fascicle length is proportional to the number of sarcomeres arranged in series and the excursion range of the muscle fiber [109].

Echo intensity was defined as the brightness of the image acquired through ultrasound. It is expressed in gray scales (0–255). Some studies used a gray scale analysis for determining the echo intensity [72, 119]. Different programs were used for this analysis, which came down to post-producing the images.

Components and specific measuring points will be given per individual muscle.

Upper leg muscles

The largest part of information on anatomical landmarks is found on the four bellies of the quadriceps muscle (rectus femoris, vastus lateralis, vastus medialis, vastus intermedius). As common landmarks can be used for the identification of these four muscles, the data gathered are taken together. All measuring points and components of quadriceps muscles are also represented in Table 2. Besides the quadriceps muscle, the only other upper leg muscle assessed was the biceps femoris (data in the text, not in the table).

At a distance of 60–70% between the greater trochanter and the popliteal crease, rectus femoris was assessed for muscle thickness [65].

At a distance of 30% between the greater trochanter and the lateral condyle, vastus medialis was assessed for muscle thickness [84, 131, 132], pennation angle [131] and echo intensity [84, 123]. Vastus lateralis was assessed for muscle thickness [85, 132, 133], fascicle length [85, 133], pennation angle [85] and echo intensity [133]. Rectus femoris was assessed for cross-sectional area [85].

At a distance of 50% between the greater trochanter and the lateral condyle, rectus femoris was assessed for muscle thickness [41, 75, 84, 89, 95, 131, 134], cross-sectional area [71, 97], pennation angle [89, 131] and echo intensity [71, 84, 123]. Vastus lateralis was assessed for muscle thickness [70, 80, 84, 88, 89, 99–101, 103, 115, 131, 135], cross-sectional area [88], anatomical cross-sectional area [71, 87, 98, 119, 135], fascicle length [70, 88, 99–101, 103, 115], pennation angle [70, 88, 89, 99–101, 103, 115] and echo intensity [71, 84, 88, 119, 135, 136]. Vastus intermedius was assessed for muscle thickness [84, 131, 134], pennation angle [131] and echo intensity [84, 136]. Quadriceps (all 4 bellies) were assessed for muscle thickness [137].

At a two-third distance between the greater trochanter and the lateral condyle, rectus femoris was assessed for muscle thickness [67]. Vastus lateralis was assessed for muscle thickness [86], cross-sectional area [86], fascicle length [86], pennation angle [86] and echo intensity [86].

At a distance of 50% between the greater trochanter and the articular cleft of the knee, rectus femoris and vastus intermedius were assessed for muscle thickness [96].

At a distance of 50% between the greater trochanter and the proximal border of the patella, rectus femoris was assessed for anatomical cross-sectional area [138] and physiological cross-sectional area [127].

At a two-third distance between the anterior superior iliac spine and the lateral condyle, both rectus femoris and vastus intermedius were assessed for muscle thickness and echo intensity [90].

At a distance of 50% between the anterior superior iliac spine and the proximal border of the patella, rectus femoris was assessed for muscle thickness [43, 78, 79, 82, 117, 118, 120–122, 139, 140] and echo intensity [68, 83, 118, 120, 122]. Vastus intermedius was assessed for muscle thickness [78, 79, 120–122, 139] and echo intensity [120, 122]. Vastus lateralis was assessed for muscle thickness [140].

At a distance of 60% between the anterior superior iliac spine and the proximal border of the patella, rectus femoris was assessed for muscle thickness [92] as was the vastus lateralis [78].

At a two-third distance between the anterior superior iliac spine and the proximal border of the patella, rectus femoris was assessed for muscle thickness [43, 77, 141]. Vastus lateralis was assessed for muscle thickness [118] and echo intensity [118]. Vastus intermedius was assessed for muscle thickness [77, 141].

At a three-fifth distance between the anterior superior iliac spine and the proximal border of the patella, rectus femoris was assessed for cross-sectional area [81].

At a distance of 75% between the anterior superior iliac spine and the proximal border of the patella, rectus femoris was assessed for physiological cross-sectional area [91].

At a distance of 50% between the anterior superior iliac spine and the midpoint of the patella, rectus femoris was assessed for muscle thickness [76], as was the vastus intermedius [76].

At a distance of 50% between the anterior superior iliac spine and the distal border of the patella, rectus femoris was assessed for muscle thickness and anatomical cross-sectional area [93].

At a distance of 50% between the anterior inferior iliac spine and the proximal border of the patella, rectus femoris was assessed for muscle thickness [88, 94], cross-sectional area [87, 88], anatomical cross-sectional area [119], fascicle length [88], pennation angle [88] and echo intensity [88, 94, 119]. Vastus intermedius was assessed for muscle thickness [94] and echo intensity [94].

At 10 cm proximal of the proximal border of the patella, rectus femoris was assessed for muscle thickness [142].

At 15 cm proximal of the proximal border of the patella, rectus femoris was assessed for muscle thickness [74] and anatomical cross-sectional area [143].

At 50% between the ischial tuberosity and the lateral condyle of the tibia, biceps femoris was assessed for muscle thickness [78].

Lower leg muscles

The lower leg muscles assessed were the gastrocnemius medialis, the soleus, and the tibialis anterior muscle. One study mentioned that as a bipennate muscle, both the deep and superficial part of the tibialis anterior was measured [97].

At 30% proximal between the medial condyle of the tibia and the medial malleolus of the fibula, the gastrocnemius medialis was assessed for muscle thickness [78, 103, 115], cross-sectional area [111], anatomical cross-sectional area [108], pennation angle [108, 115], fascicle length [108, 115] and echo intensity [108, 111].

Further measurements use landmarks from the muscle itself.

At 50% between the proximal and distal tendon insertion of the muscle, gastrocnemius medialis was assessed for muscle thickness [82, 102, 109], fascicle length [69, 97, 102, 110] and pennation angle [69, 97, 102, 107, 110].

At the most bulky area of the leg, gastrocnemius medialis was assessed for muscle thickness [70, 80, 112, 144],

cross-sectional area [145], fascicle length [70, 145] and pennation angle [70, 145].

No studies assessed the gastrocnemius lateralis.

At 30% proximal between the medial condyle of the tibia and the medial malleolus of the fibula, the soleus was assessed for muscle thickness [78].

At 50% between the proximal and distal tendon insertion of the muscle, soleus was assessed for pennation angle [146].

At the most bulky area of the leg, soleus was assessed for muscle thickness [112].

For the tibialis anterior, there was only one study, assessing pennation angle and fascicle length at 50% of muscle length, without giving clear anatomical landmarks [97].

Upper arm muscles

The upper arm muscles assessed were the biceps brachii and triceps brachii. Less studies used clear anatomical landmarks for the upper arm muscles than for the upper leg muscles.

At 50% between the acromion and the cubital fossa with the elbows flexed at 90 degrees, biceps brachii was assessed for muscle thickness [113].

At two-thirds between the acromion and the antecubital crease with the arms fully stretched, biceps brachii was assessed for muscle thickness [57].

At 50% of muscle length (not defined how this is determined), biceps brachii was assessed for fascicle length [97] and pennation angle [97]. This article also mentions that since the fascicles were almost parallel to the superficial aponeurosis, no fascicle length was measured as too much extrapolation was needed [97].

At maximal girth of the upper arm, biceps brachii was assessed for anatomical cross-sectional area [143].

At 40% between the acromion process of the scapula and the lateral epicondyle of the humerus (starting at the lateral epicondyle), triceps brachii was assessed for muscle thickness [114] and pennation angle [114].

At 40% between the acromion process of the scapula and the lateral epicondyle of the humerus (starting at the lateral epicondyle), triceps brachii was assessed for muscle thickness [115] and pennation angle [115].

At 50% between the posterior crista of the acromion and the olecranon triceps brachii was assessed for muscle thickness [116] and pennation angle [116].

Discussion

It is clear from the multitude of different measuring points that there is little consistency in the current ultrasonographic muscle assessment. To advance US as a routine technique

Table 3 Consensus proposition, shortcomings in knowledge and protocol listings for patient positioning pre-investigation

Consensus proposition:
No exercise 30 min before investigation
Preferably minimum 30 min (maximum 60 min) in the same position before investigation, for measurements in recumbent position
Muscle should be assessed in a relaxed state
If the patient is placed in a recumbent position, it is recommended to use the full extension position (either supine or prone)
Shortcomings in knowledge:
Exact influence of (minor) muscle exercise on measurements
To be mentioned in the protocol:
Preparations in advance of the investigation (amount of minutes rest, in which position)
State of muscle being investigated (relaxed, contracted)
Which position the patient is placed in, including the angles of the relevant joints, clearly describing which angle is meant exactly
Whether left/right side was taken and whether this was the dominant/non-dominant side
Sex and age of patient

to be used for muscle quality and quantity assessment in old age, standardization is paramount. In the following paragraphs, the possible points of discussion will be addressed regarding the different aspects of measurement. A consensus proposition will be given at the end of each paragraph. Shortcomings in knowledge will also be addressed, as more studies need to be done before able to give recommendations about this subject. We will also propose what information should minimally be mentioned in a study protocol.

Patient positioning pre-investigation

Since there is a clear difference between relaxed and contracted muscle [147], one state should be chosen. As it is easier to keep a muscle in a relaxed state than in a certain degree of contraction, it seems logical to choose for the former. For the record, it must be said that there is some evidence that contracted muscle correlates better with muscle function than muscle in a relaxed state [147]. Ideally, this should be studied for better intra- and interrater reliability and test–retest reliability. Also the state of contraction before an examination could possibly have an influence on the component measurements. Refraining study subjects from exercises for 24–48 h is only practical in a purely scientific setting [74, 75], not a clinical one, where one would want its patients to exercise as much as possible. Also letting patients performing an amount of maximal contractions [65] could possibly influence measurements. An older study has shown that intensive exercise can give a 15% increase in water content in a given muscle bulk [148], as also giving way to a possible large measuring error. As it is unclear to what degree minor contractions and light-to-moderate exercises influence muscle volume, it is advised not to let the study subjects do exercises before taking measurements. As there

is no clear data on a timeframe, a period of minimal 30 min is proposed. Ideally, this period is spent laying down.

There are little data on how long patients should be in the same position in order to allow fluid shifts to stabilize [72], or to avoid muscle contraction-induced fluid shifts and muscle blood flow during the measurements [73]. Differences in measurements made in the standing and recumbent positions for example may be due to postural or positional forces acting on muscle shape (for instance joint angle), or due to physiological changes [149]. When going from standing to recumbent position, the most significant changes in thigh muscle size occurred within the first 15–20 min of recumbency, with a stabilization after 60 min of bed rest [150, 151]. When measuring a subject in a recumbent position, a minimum of 30 min and a maximum of 60 min in the same position prior to the measurements are thus advisable.

The patient positioning in itself is less important for measuring the five components, because depending on what one wants to measure, the positioning could change. However, it must be mentioned that muscle components are significantly different if measured in a standing or in a recumbent position. This is seen more in CSA than in muscle thickness or pennation angle [152]. If the patient is placed in a recumbent position, it is recommended to place the patient in full extension (described above), as in this position, most muscles are relaxed maximally. Mentioning the position in a protocol is thus strongly advised. Regarding dominant/non-dominant side, there is some discussion about the relevance of functional asymmetry and strength differences in dominant versus non-dominant side [153]. In non-athletic populations, inter-limb differences in strength are possibly more related to neural factors than pure muscle-related factors [86]. Since this is not clear in literature, it is advised to clearly indicate whether the dominant or non-dominant side is assessed.

Table 4 Consensus proposition, shortcomings in knowledge and protocol listings for system and system settings

Consensus proposition:

All types of ultrasound machine can be used, as long as B-mode is present

Extended field of view is not necessary but recommended

A linear transducer probe is recommended. A minimum length of 5 cm is advised

Inclination of the probe should be neutral, which is perpendicular to the skin

Using a generous amount of transmission gel is recommended

Maintaining the most minimal pressure possible between transducer and skin is recommended

Shortcomings in knowledge:

Exact influence of different system settings on measurements of echo intensity

To be mentioned in the protocol:

Manufacturer and type of US machine

Type of probe, including length of probe

Frequency of beam (other system setting, see “Components and measuring points: echo intensity”)

Any additional software used in post-production of images

Consensus proposition, shortcomings in knowledge and protocol listings for patient positioning pre-investigation are addressed in Table 3.

System and system settings

The brand of ultrasonographic machine used is of no relevance for most of the measurements.

Standard B-mode is applied in all studies to visualize the different muscle components and is available on most machines.

Although a curved probe can be used [91], a linear transducer probe is more adapted to assess muscle anatomy.

The length of the transducer is less important for measuring the five main components. However, for assessing cross-sectional area and echo intensity, larger (longer) probes can potentially visualize more tissue, which can be helpful if a large muscle bulk is present. Therefore, a minimum length of 5 cm seems advisable. Extended field of view techniques could help solve this problem.

As no information on inclination of the probe is at hand, it is advised to keep the probe as much perpendicular to the skin as possible.

When applying the probe to the skin, it is important to avoid compression of dermal surface and distortion of muscle surface [72, 104, 154, 155]. Dupont et al. found that applying strong pressure with the ultrasound transducer could flatten the deltoid muscle by 50% or more and because this error is proportional to muscle size, the absolute error in muscle thickness measurement would be even greater for larger muscles [155]. To minimize both beam loss/scatter and need for dermal/muscle compression, the use of a transmission gel is standard in ultrasonographic investigations. There is no defined standard amount of gel to be used. It is advised to use a generous amount and maintain the minimal

pressure possible/necessary between transducer and the skin [155].

The frequency of the transducer beam is less relevant for measuring components, except for assessing echo intensity. However, the higher the frequency, the better is the visualization of anatomical structures. Using ultrasound, there is therefore a constant compromise between image resolution and depth of penetration of the sound waves. Higher frequency transducers provide better spatial resolution, but these transducers have a shallower depth of penetration than a lower frequency transducer [156].

To our knowledge, other system settings do not seem to be relevant for measuring the components, except for assessing echo intensity. Depth focus, general gain, time-gain compensation, dynamic range, etc. can be set in order to have the best possible view of the muscle that is to be assessed.

The use of software for post-processing and measurement is less relevant for measuring the components, except for assessing echo intensity. Most modern machines have software that allows measurements to be directly done during the investigation; otherwise, all other picture post-processing software can be used.

Measuring echo intensity is dependent on many factors. These are discussed under the paragraphs “Components” and “Measuring points”.

Consensus proposition, shortcomings in knowledge and protocol listings for system and system settings are addressed in Table 4.

Components

The five components that can be easily measured when assessing muscle components are already mentioned: muscle thickness, pennation angle, fascicle length, echo intensity and cross-sectional area. The technique to measure these

items is relative easy. The most difficult part is to define where to do the measurements. As there is no information available on the most ideal location within the muscle, there exists a multitude of measuring points. Since no scientific substantiated “best” point can be defined, consensus locations will be provided for each muscle described in literature.

It seems logical to use the thickest zone of a muscle when wanting to assess muscle thickness, as this will be the place that the muscle will generate the most contractive power. However, there are no studies to our knowledge that have looked at the evolution of muscle thickness throughout a specific muscle. Therefore, we advise different approaches for different kinds of muscles.

Theoretically, muscles like the four bellies from the quadriceps are the thickest at 50% of their length, measured from tendon to tendon [157]. Therefore, we propose to mark and use the point at 50% of the muscle’s length (50% rule).

Unfortunately, not every muscle is shaped like the bellies of the quadriceps, e.g., the gastrocnemius medialis. This poses a certain difficulty, and a possible reason for measurement bias. In the literature, in this type of muscle the “maximal bulk” was most often noted as measuring point. We propose that in more asymmetrical muscles, the point of 50% length between tendons is visualized, and then 4 additional points are checked: at 30, 40, 60 and 70% length between

Table 5 Consensus proposition, shortcomings in knowledge and protocol listings for components

Consensus proposition:

- Five components can be measured: muscle thickness, pennation angle, fascicle length, echo intensity and cross-sectional area
- Measurements are ideally done at maximal muscle bulk
- Depending on muscle anatomy, different techniques are advised for determining maximal muscle bulk
- Panoramic vision and extended-field-of-view software are not absolutely necessary but recommended
- In pennate muscles, measuring physiological CSA rather than anatomical CSA is recommended
- When the fascicle length cannot be directly measured, it can be calculated using the standard formula
- When measuring echo intensity, all system settings need to be kept the same. Currently, no proposition for specific system settings based upon literature can be done for echo intensity

Shortcomings in knowledge:

- Exact point of maximal muscle thickness for each muscle
- Changes of the main components (MT, CSA, FL, PA, EI) throughout the muscle bulk
- A good measure for comparing echo intensity between different US machines/systems

To be mentioned in the protocol:

- The muscle that is assessed, with inclusion of the anatomical landmarks that are used and the exact point in between the landmarks. If not the midpoint, clearly describe whether the proximal or distal end is meant
- The components that are measured. If CSA is measured, define if anatomical or physiological CSA is meant
- Total length of muscle (to calculate relative muscle thickness values)
- The technique that is used to determine the position of maximal bulk

Table 6 Proposed anatomical landmarks for each muscle discussed

	Proximal landmark	Distal landmark	Asymmetry
Lower limb			
Rectus femoris	Greater trochanter	Proximal border of patella	Minimal
Vastus lateralis	Greater trochanter	Proximal border of patella	Minimal
Vastus medialis	Greater trochanter	Proximal border of patella	Minimal
Vastus intermedius	Greater trochanter	Proximal border of patella	Minimal
Biceps femoris (long head)	Ischial tuberosity	Proximal head of fibula	Minimal
Tibialis anterior	Lateral condyle (anterior) of tibia	US-measurement dependant	Minimal
Gastrocnemius (medialis)	Medial condyle (posterior) of the femur	US-measurement dependant	Minimal
Gastrocnemius (lateralis)	Medial condyle (posterior) of the femur	US-measurement dependant	Minimal
Soleus	Proximal head of fibula (posterior part)	Posterior superior part of calcaneus	Yes
Upper limb			
Biceps brachii	Anterior part of acromion process (acromio-clavicular joint)	Elbow crease where tendon can be palpated	Yes
Triceps brachii	Most lateral distal part of acromion	Tip of olecranon	Yes

tendons. Then, the maximal muscle thickness can be chosen from these measurements. Without having large studies that provide reference data per muscle, this will have to be done per muscle, per patient. In muscles with severe asymmetry, e.g., biceps femoris, it is best to specify which part of the muscle is assessed, e.g., biceps femoris—long head.

Another variety is the muscle with a long tendon, which has no clear anatomical landmarks to indicate where the muscle bulk ends, e.g., tibialis anterior. In these muscles, it is advised to locate the proximal and distal border of the muscle through ultrasound before referring to the 50% rule.

Noting down the 100% length of the muscle from tendon to tendon is advised, as a longer muscle can potentially generate more power. It is not known whether absolute muscle thickness or relative muscle thickness (= muscle thickness/length of the muscle) is more representative [158]. To be complete, there could be other reasons to choose for a different measurement site in specific muscles. Ticinesi et al. [86] suggested to use the distal point of 65% of the length of the vastus lateralis, as this site is most free of vessels and muscle biopsies can be easily made, avoiding major vessels and nerves. However, without specific arguments for clinical correlates, this is not recommended. For this, studies are needed.

For measuring the maximal thickness of a muscle, it is advised not only to take the midpoint of the muscle in between the tendons, but at this point of the longitudinal axis, to use the point at 50% between the medial and lateral border of the muscle bulk. This will also have to be visualized through ultrasound and marked for easy Ref. [86].

The argument of using the thickest zone of the muscle bulk is also valid for measuring the cross-sectional area. One of the disadvantages of this point could be that in some cases it will be difficult to get a complete image of the CSA, in the case of a large muscle bulk. Panoramic vision and extended-field-of-view software that are almost standard on most US machines can nullify this problem. Also, since it is unknown how the muscle volume diminishes exactly towards the tendons, submaximal measurements do not weigh up to the maximal CSA. For this, studies are needed.

As said, cross-sectional area can be divided into anatomical CSA and physiological CSA. The former underestimates the number of total fibers in a pennated muscle, so in these types of muscles the use of the PCSA is advised. In non-pennate muscles, the ASCA is the same as the PCSA.

As there is no hard information available to our knowledge about the pattern of echo intensity or pennation angle throughout a given muscle—homogenous versus heterogenous [159]—the proposition is to take all measurements (including the fascicle length) at the point of maximal bulk, as discussed for muscle thickness and CSA.

There is no doubt that echo intensity is an important parameter. As a parameter of fatty infiltration of the muscle

(myosteatorsis), it helps unravel an important aspect of the process of sarcopenia. Myosteatorsis is linked with increased mortality in specific populations [20, 160–162], with a strong need for further clinical investigation. However, the standardisation and comparability of measurements of echo intensity between different US system brands are appalling. Defined as the brightness of the image and expressed in gray scales, there are many factors (system relates) that influence the image. A small difference in beam frequency or gain—or many other settings—can give completely different results, and comparing system settings between different manufacturers is currently impossible. No good calibration model exists to date. Even the age of a probe can influence results by influencing the strength of the beam emitted. Also, although the analysis of muscle tissue acquired via biopsy suggests that echogenicity is more strongly associated with intramuscular adipose tissue rather than fibrosis [163, 164], biopsies are still needed to determine the percentage both components [165]. More studies are needed on both the comparability between systems and the differentiation of fat and fibrosis. Ideally, a universally accepted calibration dummy will be developed in the near future.

Consensus proposition, shortcomings in knowledge and protocol listings for components are addressed in Table 5.

Measuring points

For the appendicular muscle described, we refer to Table 6 for an overview of the proposed anatomical landmarks. These landmarks were selected in view of the discussion above, and with regard to the possible use of good-identifiable anatomical landmarks. As discussed in the paragraph about muscle thickness, some muscles show an asymmetry, either in length, width, or both. Table 6 also indicates whether a certain asymmetry exists and, thus, caution has to be taken in use of the 50% rule. When there are multiple origins, or if there is a larger area of insertion, the most identifiable/representative landmark is chosen. Future studies will have to confirm the usability of these landmarks.

Method of measurement

As a short overview, the proposed procedure of measurement will be shortly addressed. This procedure is based on the discussion above. It will be represented schematically as a sort of checklist.

- Place the patient in the desired position, preferably 30 min before investigation.
- Select the muscle that is to be assessed, and check for the desired technique to locate the maximal muscle bulk.

- Use the appropriate technique and the anatomical landmarks provided to locate the longitudinal measuring point. Mark this with a dermatographic pencil.
- Locate the medial and lateral side of the muscle and mark these with a dermatographic pencil. Use the middle point of these marks. Now the correct measuring point is found.
- Keeping the transducer probe in a longitudinal direction in line with the muscle fiber fascicles. At this position, measure muscle thickness (from aponeurosis to aponeurosis), pennation angle (angle of muscle fiber fascicles into the deep aponeurosis) and fascicle length.
- Turn the transducer probe 90°. At this position, measure cross-sectional area. As this will probably not be a perfect ellipse, the circumference of the muscle can be manually drawn with a cursor. Use this maximal area to also measure echo intensity.
- Repeat all measurements three times and use the mean value of these measurements.

Limitations

There are some limitations to this review. Although the search strategy was very broad, we have focused on appendicular muscle mass, and have not included facial, thoracic, abdominal and pelvic muscle. Also the incidental reports from smaller muscle groups from the hand were not included, because too little information was available per muscle. The muscle groups that were not included in this review could be the subject of future investigations.

Also some ultrasound-based techniques were not included in this review, such as elastography. This could also be the subject of a review in itself.

Future direction

The ultimate goal would be to have not only a standardized way to assess muscle characteristics, but also to have a workable algorithm to diagnose sarcopenia using ultrasound. For this, some barriers have to be taken.

First of all more insights should be gained into the age-related evolution of the different muscle characteristics. It seems reasonable to assume that muscle thickness and cross-sectional area for instance will decrease with age, but perhaps the pennation angle will start shifting before any thickness or volume dimensions change. This is important because making a diagnosis as early as possible also means that treatment can start earlier.

Secondly, reference values of a ‘normal population’ are lacking. In 2016, Minetto et al. [82] have already done some work on this, using muscle thickness values 2 standard deviations below the gender-specific means of a sample of

younger subjects to diagnose sarcopenia. In this regard, one could discuss about using one set of cutoff points derived from a younger population, or if age-related reference values should be used. It is also clear that all reference points should be specific for each muscle. This means that every muscle should be examined separately, because cutoff points for, e.g., the vastus lateralis are different than those for the gastrocnemius.

Thirdly, it is too early to say if diagnosing sarcopenia with ultrasound should be done by a composite score or by a single measurement. In this stage, it seems advisable to measure all muscle characteristics and link this with diagnostic criteria that are currently used (e.g., BIA, DEXA). In a later stage, perhaps it will become clear that certain ultrasound-based characteristics are more important than others. For this, more studies have to be done.

Fourthly, the measurements proposed in this manuscript are not exhaustive. Perhaps other parameters are equal or more important, e.g., elastography of the muscle or tendon. Although an area that should certainly be explored, too little is known of this in the context of sarcopenia to currently make any statements.

In summary, there is certainly a dynamic towards using ultrasound for diagnosing sarcopenia, but more studies need to be done, preferentially starting with the creation of reference values.

Conclusion

To compare studies and advance the use of ultrasound in the assessment of sarcopenia, certain standardization has to be done. However, it is not yet clear which measuring points are more relevant than others. Therefore, the recommendations proposed in this review should not be regarded as set in stone. They are rather intended as a reference point or as a guideline, from which comparative studies can be initiated. Future directions can certainly involve changing some of the recommendations, if there is new evidence to support these changes. The limitations of this review are clear since there are a lot of unknown factors: the exact spreading pattern and evolution of the different architectural components throughout the muscle, which muscles or muscle measurements are most clinically relevant, influence of position of the patient, influence of pre-investigation activity, etcetera. Therefore, this consensus approach is used as a starting point. Hopefully, this way future studies will have an extra support to build upon. Also, this way interested researchers can collaborate towards an ultrasonographic diagnosis of sarcopenia.

In conclusion, this review offers a guideline for investigators wanting to set up a study using ultrasound in muscle assessment. Studies in clinical settings are needed to validate the effectiveness of these propositions.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

Ethical approval This article does not contain any studies with human participants performed by any of the authors.

References


- Beaudart C et al (2014) Sarcopenia: burden and challenges for public health. *Arch Public Health* 72(1):45
- Beaudart C et al (2017) Health outcomes of sarcopenia: a systematic review and meta-analysis. *PLoS ONE* 12(1):e0169548
- Cruz-Jentoft AJ et al (2010) Sarcopenia: European consensus on definition and diagnosis: report of the European Working Group on sarcopenia in older people. *Age Ageing* 39(4):412–423
- Muscaritoli M et al (2010) Consensus definition of sarcopenia, cachexia and pre-cachexia: joint document elaborated by Special Interest Groups (SIG) “cachexia-anorexia in chronic wasting diseases” and “nutrition in geriatrics”. *Clin Nutr* 29(2):154–159
- Fielding RA et al (2011) Sarcopenia: an undiagnosed condition in older adults. Current consensus definition: prevalence, etiology, and consequences. International working group on sarcopenia. *J Am Med Dir Assoc* 12(4):249–256
- Morley JE et al (2011) Sarcopenia with limited mobility: an international consensus. *J Am Med Dir Assoc* 12(6):403–409
- Dam TT et al (2014) An evidence-based comparison of operational criteria for the presence of sarcopenia. *J Gerontol A Biol Sci Med Sci* 69(5):584–590
- Chen LK et al (2014) Sarcopenia in Asia: consensus report of the Asian Working Group for Sarcopenia. *J Am Med Dir Assoc* 15(2):95–101
- Beaudart C et al (2015) Estimation of sarcopenia prevalence using various assessment tools. *Exp Gerontol* 61:31–37
- Baumgartner RN et al (1998) Epidemiology of sarcopenia among the elderly in New Mexico. *Am J Epidemiol* 147(8):755–763
- Delmonico MJ et al (2007) Alternative definitions of sarcopenia, lower extremity performance, and functional impairment with aging in older men and women. *J Am Geriatr Soc* 55(5):769–774
- Newman AB et al (2003) Sarcopenia: alternative definitions and associations with lower extremity function. *J Am Geriatr Soc* 51(11):1602–1609
- Chien MY, Huang TY, Wu YT (2008) Prevalence of sarcopenia estimated using a bioelectrical impedance analysis prediction equation in community-dwelling elderly people in Taiwan. *J Am Geriatr Soc* 56(9):1710–1715
- Janssen I et al (2004) Skeletal muscle cutpoints associated with elevated physical disability risk in older men and women. *Am J Epidemiol* 159(4):413–421
- Janssen I, Heymsfield SB, Ross R (2002) Low relative skeletal muscle mass (sarcopenia) in older persons is associated with functional impairment and physical disability. *J Am Geriatr Soc* 50(5):889–896
- Barbat-Artigas S et al (2012) How to assess functional status: a new muscle quality index. *J Nutr Health Aging* 16(1):67–77
- Newman AB et al (2003) Strength and muscle quality in a well-functioning cohort of older adults: the Health, Aging and Body Composition Study. *J Am Geriatr Soc* 51(3):323–330
- Goodpaster BH et al (2006) The loss of skeletal muscle strength, mass, and quality in older adults: the health, aging and body composition study. *J Gerontol A Biol Sci Med Sci* 61(10):1059–1064
- Perkisas S et al (2016) Physiological and architectural changes in the ageing muscle and their relation to strength and function in sarcopenia. *Eur Geriatr Med* 7(3):201–206. <https://doi.org/10.1016/j.eurger.2015.12.016>
- Perkisas S, De Cock AM, Verhoeven V, Vandewoude M (2017) Intramuscular adipose tissue and the functional components of sarcopenia in hospitalized geriatric patients. *Geriatrics* 2(1):11. <https://doi.org/10.3390/geriatrics2010011>
- Reinders I et al (2016) Muscle quality and myosteatosis: novel associations with mortality risk: the Age, Gene/Environment Susceptibility (AGES)-Reykjavik Study. *Am J Epidemiol* 183(1):53–60
- Buckinx F et al (2018) Pitfalls in the measurement of muscle mass: a need for a reference standard. *J Cachexia Sarcopenia Muscle* 9(2):269–278
- Yamada Y et al (2017) Electrical properties assessed by bioelectrical impedance spectroscopy as biomarkers of age-related loss of skeletal muscle quantity and quality. *J Gerontol A Biol Sci Med Sci* 72(9):1180–1186
- Bartels EM, Sorensen ER, Harrison AP (2015) Multi-frequency bioimpedance in human muscle assessment. *Physiol Rep* 3(4):e12354
- Beaudart C et al (2016) Sarcopenia in daily practice: assessment and management. *BMC Geriatr* 16(1):170
- Goodpaster BH et al (2000) Skeletal muscle attenuation determined by computed tomography is associated with skeletal muscle lipid content. *J Appl Physiol* (1985) 89(1):104–110
- Heymsfield SB et al (2014) Assessing skeletal muscle mass: historical overview and state of the art. *J Cachexia Sarcopenia Muscle* 5(1):9–18
- Fischer MA et al (2014) Dixon-based MRI for assessment of muscle-fat content in phantoms, healthy volunteers and patients with achillobodynia: comparison to visual assessment of calf muscle quality. *Eur Radiol* 24(6):1366–1375
- Pfirrmann CW et al (2004) Assessment of fat content in supraspinatus muscle with proton MR spectroscopy in asymptomatic volunteers and patients with supraspinatus tendon lesions. *Radiology* 232(3):709–715
- Prompers JJ et al (2006) Dynamic MRS and MRI of skeletal muscle function and biomechanics. *NMR Biomed* 19(7):927–953
- Valkovic L, Chmelik M, Krssak M (2017) In-vivo(31)P-MRS of skeletal muscle and liver: a way for non-invasive assessment of their metabolism. *Anal Biochem* 529:193–215
- Mourtzakis M et al (2008) A practical and precise approach to quantification of body composition in cancer patients using computed tomography images acquired during routine care. *Appl Physiol Nutr Metab* 33(5):997–1006
- Shen W et al (2004) Total body skeletal muscle and adipose tissue volumes: estimation from a single abdominal cross-sectional image. *J Appl Physiol* (1985) 97(6):2333–2338
- Janssen I et al (2000) Skeletal muscle mass and distribution in 468 men and women aged 18–88 yr. *J Appl Physiol* (1985) 89(1):81–88
- Pillen S, van Alfen N (2011) Skeletal muscle ultrasound. *Neurol Res* 33(10):1016–1024
- Passmore E, Lai A, Sangeux M et al (2017) Application of ultrasound imaging to subject-specific modelling of the human musculoskeletal system. *Meccanica* 52:665. <https://doi.org/10.1007/s11012-016-0478-z>
- Abe T et al (2016) Ultrasound-derived forearm muscle thickness is a powerful predictor for estimating DXA-derived appendicular lean mass in Japanese older adults. *Ultrasound Med Biol* 42(9):2341–2344
- Takai Y et al (2014) Applicability of ultrasound muscle thickness measurements for predicting fat-free mass in elderly population. *J Nutr Health Aging* 18(6):579–585

39. Takai Y et al (2013) Validity of ultrasound muscle thickness measurements for predicting leg skeletal muscle mass in healthy Japanese middle-aged and older individuals. *J Physiol Anthropol* 32:12
40. Abe T et al (2015) Validity of ultrasound prediction equations for total and regional muscularity in middle-aged and older men and women. *Ultrasound Med Biol* 41(2):557–564
41. Thomaes T et al (2012) Reliability and validity of the ultrasound technique to measure the rectus femoris muscle diameter in older CAD-patients. *BMC Med Imaging* 12:7
42. Sanada K et al (2006) Prediction and validation of total and regional skeletal muscle mass by ultrasound in Japanese adults. *Eur J Appl Physiol* 96(1):24–31
43. Tandon P et al (2016) A model to identify sarcopenia in patients with cirrhosis. *Clin Gastroenterol Hepatol* 14(10):1473–1480 e3
44. Reeves ND, Maganaris CN, Narici MV (2004) Ultrasonographic assessment of human skeletal muscle size. *Eur J Appl Physiol* 91:116–118
45. Nijholt W et al (2017) The reliability and validity of ultrasound to quantify muscles in older adults: a systematic review. *J Cachexia Sarcopenia Muscle* 8(5):702–712
46. Mourtzakis M, Wischmeyer P (2014) Bedside ultrasound measurement of skeletal muscle. *Curr Opin Clin Nutr Metab Care* 17(5):389–395
47. Haberfehlner H et al (2016) Freehand three-dimensional ultrasound to assess semitendinosus muscle morphology. *J Anat* 229:591–599
48. Martin DC et al (2001) Comparing human skeletal muscle architectural parameters of cadavers with in vivo ultrasonographic measurements. *J Anat* 199:429–434
49. Tosovic D et al (2016) Anatomy of the long head of biceps femoris: an ultrasound study. *Clin Anat* 29(6):738–745
50. Kellis E et al (2009) Validity of architectural properties of the hamstring muscles: correlation of ultrasound findings with cadaveric dissection. *J Biomech* 42(15):2549–2554
51. Infantolino BW, Challis JH (2016) Evaluation of a simple method for determining muscle volume in vivo. *J Biomech* 49(9):1973–1975
52. Tillquist M et al (2014) Bedside ultrasound is a practical and reliable measurement tool for assessing quadriceps muscle layer thickness. *J Parenter Enter Nutr* 38(7):886–890
53. English C, Fisher L, Thoirs K (2012) Reliability of real-time ultrasound for measuring skeletal muscle size in human limbs in vivo: a systematic review. *Clin Rehabil* 26(10):934–944
54. Miyatani M et al (2004) The accuracy of volume estimates using ultrasound muscle thickness measurements in different muscle groups. *Eur J Appl Physiol* 91(2):264–272
55. Nijholt W et al (2018) Response to: “The use of ultrasound for the estimation of muscle mass: one site fits most?”. *J Cachexia Sarcopenia Muscle* 9:627–628
56. Reimers CD, Harder T, Saxe H (1998) Age-related muscle atrophy does not affect all muscles and can partly be compensated by physical activity: an ultrasound study. *J Neurol Sci* 159(1):60–66
57. Arts IMP et al (2007) Rise and fall of skeletal muscle size over the entire life span. *J Am Geriatr Soc* 55:1150–1152
58. Arts IMP et al (2010) Normal values for quantitative muscle ultrasonography in adults. *Muscle Nerve* 41(1):32–41
59. Verhulst FV et al (2011) Quantitative ultrasound of lower leg and foot muscles: feasibility and reference values. *Foot Ankle Surg Off J Eur Soc Foot Ankle Surg* 17(3):145–149
60. Maurits NM et al (2002) Quantitative muscle ultrasound analysis in healthy adults: weight- and age-related normal values. *Neuromuscular Disord* 12(7):776
61. Ticinesi A et al (2017) Muscle ultrasound and sarcopenia in older individuals: a clinical perspective. *J Am Med Dir Assoc* 18(4):290–300
62. Welch C et al (2018) Acute sarcopenia secondary to hospitalisation—an emerging condition affecting older adults. *Aging Dis* 9(1):151–164
63. van de Glind EM et al (2012) Search filters to identify geriatric medicine in Medline. *J Am Med Inform Assoc* 19(3):468–472
64. Ouzzani M et al (2016) Rayyan—a web and mobile app for systematic reviews. *Syst Rev* 5(1):210
65. Ema R et al (2013) In vivo measurement of human rectus femoris architecture by ultrasonography: validity and applicability. *Clin Physiol Funct Imaging* 33:267–273
66. de Bruin PF et al (1997) Size and strength of the respiratory and quadriceps muscles in patients with chronic asthma. *Eur Respir J* 10:59–64
67. Correa CS et al (2016) Effects of strength training, detraining and retraining in muscle strength, hypertrophy and functional tasks in older female adults. *Clin Physiol Funct Imaging* 36(4):306–310
68. Harris-Love MO et al (2016) Ultrasound estimates of muscle quality in older adults: reliability and comparison of Photoshop and ImageJ for the grayscale analysis of muscle echogenicity. *PeerJ* 4:e1721
69. Narici MV et al (2003) Effect of aging on human muscle architecture. *J Appl Physiol (Bethesda, Md.: 1985)* 95(6):2229–2234
70. Kaya A et al (2013) Ultrasonographic evaluation of the muscle architecture in patients with systemic lupus erythematosus. *Clin Rheumatol* 32(8):1155–1160
71. Kleinberg CR et al (2016) Influence of lower extremity muscle size and quality on stair-climb performance in career firefighters. *J Strength Cond Res* 30(6):1613–1618
72. da Silva RP et al (2017) Effect of strength training on sleep apnea severity in the elderly: study protocol for a randomized controlled trial. *Trials* 18(1):489
73. Akima H et al (2017) Relationship between quadriceps echo intensity and functional and morphological characteristics in older men and women. *Arch Gerontol Geriatr* 70:105–111
74. Bemben DA, Langdon DB (2002) Relationship between estrogen use and musculoskeletal function in postmenopausal women. *Maturitas* 42:119–127
75. Baroni BM et al (2013) Time course of neuromuscular adaptations to knee extensor eccentric training. *Int J Sports Med* 34(10):904–911
76. Gerovasilis V et al (2009) Electrical muscle stimulation preserves the muscle mass of critically ill patients: a randomized study. *Crit Care* 13:R161
77. Guerreiro AC et al (2017) Bedside ultrasound of quadriceps to predict rehospitalization and functional decline in hospitalized elders. *Front Med* 4:122
78. Ikezoe T et al (2011) Age-related muscle atrophy in the lower extremities and daily physical activity in elderly women. *Arch Gerontol Geriatr* 53(2):e153–e157
79. Ikezoe T et al (2012) Associations of muscle stiffness and thickness with muscle strength and muscle power in elderly women. *Geriatr Gerontol Int* 12:86–92
80. Kara M et al (2015) Does vitamin D affect muscle strength and architecture? An isokinetic and ultrasonographic study. *Asia Pac J Clin Nutr* 26(1):85–88
81. Seymour JM et al (2009) Ultrasound measurement of rectus femoris cross-sectional area and the relationship with quadriceps strength in COPD. *Thorax* 64:418–423
82. Minetto MA et al (2016) Ultrasound-based detection of low muscle mass for diagnosis of sarcopenia in older adults. *PM&R J Inj Funct Rehabil* 8(5):453–462

83. Mota JA, Stock MS (2017) Rectus femoris echo intensity correlates with muscle strength, but not endurance, in younger and older men. *Ultrasound Med Biol* 43(8):1651–1657
84. Rech A et al (2014) Echo intensity is negatively associated with functional capacity in older women. *Age* 36:9708
85. Rieder F et al (2015) Alpine Skiing With total knee ArthroPlasty (ASWAP): muscular adaptations. *Scand J Med Sci Sports* 25:26–32
86. Ticinesi A, Narici MV, Lauretani F et al (2018) Assessing sarcopenia with vastus lateralis muscle ultrasound: an operative protocol. *Aging Clin Exp Res*. <https://doi.org/10.1007/s40520-018-0958-1>
87. Scanlon, T.C., et al., Muscle architecture and strength: adaptations to short-term resistance training in older adults. Vol. 49. 2014. 584-92
88. Mangine GT et al (2014) Bilateral differences in muscle architecture and increased rate of injury in national basketball association players. *J Athl Train* 49(6):794–799
89. Marin PJ et al (2013) Effects of whole-body vibration on muscle architecture, muscle strength, and balance in stroke patients a randomized controlled trial. *Am J Phys Med Rehabil* 92(10):881–888
90. Cruz-Montecinos C et al (2016) Sonographic measurement of the quadriceps muscle in patients with chronic obstructive pulmonary disease: functional and clinical implications. *J Ultrasound Med* 35(11):2405–2412
91. Hammond K et al (2014) Validity and reliability of rectus femoris ultrasound measurements: comparison of curved-array and linear-array transducers. *J Rehabil Res Dev* 51:1155–1164
92. Mueller N et al (2016) Can sarcopenia quantified by ultrasound of the rectus femoris muscle predict adverse outcome of surgical intensive care unit patients as well as frailty? A prospective, observational cohort study. *Ann Surg* 264:1116–1124
93. Berger J et al (2015) Rectus femoris (RF) ultrasound for the assessment of muscle mass in older people. *Arch Gerontol Geriatr* 61:33–38
94. Fukumoto Y et al (2012) Skeletal muscle quality assessed from echo intensity is associated with muscle strength of middle-aged and elderly persons. *Eur J Appl Physiol* 112(4):1519–1525
95. Shimizu S et al (2006) The relationship between cardiac muscle, skeletal muscle mass, and vessel structure in elderly women. *Jpn J Phys Fit Sports Med* 55:213–216
96. Nakatani M et al (2016) Validity of muscle thickness-based prediction equation for quadriceps femoris volume in middle-aged and older men and women. *Eur J Appl Physiol* 116(11):2125–2133
97. de Boer MD et al (2008) Effect of 5 weeks horizontal bed rest on human muscle thickness and architecture of weight bearing and non-weight bearing muscles. *Eur J Appl Physiol* 104(2):401–407
98. Lixandrão ME et al (2016) Time course of resistance training-induced muscle hypertrophy in the elderly. *J Strength Cond Res* 30:159–163
99. Alegre LM et al (2006) Effects of dynamic resistance training on fascicle length and isometric strength. *J Sports Sci* 24(5):501–508
100. Malas FÜ et al (2013) Effects of different strength training on muscle architecture: clinical and ultrasonographic evaluation in knee osteoarthritis. *PM&R* 5:655–662
101. de Oliveira Melo M et al (2015) Effects of neuromuscular electrical stimulation and low-level laser therapy on the muscle architecture and functional capacity in elderly patients with knee osteoarthritis: a randomized controlled trial. *Clin Rehabil* 29:570–580
102. Karamanidis K, Arampatzis A (2005) Mechanical and morphological properties of different muscle-tendon units in the lower extremity and running mechanics: effect of aging and physical activity. *J Exp Biol* 208:3907–3923
103. Fukutani A, Kurihara T (2015) Comparison of the muscle fascicle length between resistance-trained and untrained individuals: cross-sectional observation. *Springerplus* 4:341
104. Engelina S et al (2014) Ultrasound investigation of vastus medialis oblique muscle architecture: an in vivo study. *Clin Anat (New York, NY)* 27(7):1076–1084
105. Akazawa N et al (2017) Relationships between intramuscular fat, muscle strength and gait independence in older women: a cross-sectional study. *Geriatr Gerontol Int* 17(10):1683–1688
106. Selva Raj I, Bird SR, Shield AJ (2017) Ultrasound measurements of skeletal muscle architecture are associated with strength and functional capacity in older adults. *Ultrasound Med Biol* 43:586–594
107. Erskine RM et al (2017) The individual and combined effects of obesity- and ageing-induced systemic inflammation on human skeletal muscle properties. *Int J Obes* (2005) 41(1):102–111
108. Gerstner GR et al (2017) Neural and muscular contributions to the age-related reductions in rapid strength. *Med Sci Sports Exerc* 49(7):1331–1339
109. Atkinson RA et al (2010) Effects of testosterone on skeletal muscle architecture in intermediate-frail and frail elderly men. *J Gerontol Ser A Biomed Sci Med Sci* 65:1215–1219
110. Morse CI et al (2005) In vivo physiological cross-sectional area and specific force are reduced in the gastrocnemius of elderly men. *J Appl Physiol* (1985) 99(3):1050–1055
111. Rosenberg JG et al (2014) Reliability of panoramic ultrasound imaging to simultaneously examine muscle size and quality of the medial gastrocnemius. *Muscle Nerve* 49(5):736–740
112. Fujiwara K et al (2010) Regular heel-raise training focused on the soleus for the elderly: evaluation of muscle thickness by ultrasound. *J Physiol Anthropol* 29(1):23–28
113. Ohata K et al (2006) Measurement of muscle thickness as quantitative muscle evaluation for adults with severe cerebral palsy. *Phys Ther* 86(9):1231–1239
114. Kawakami Y, Abe T, Fukunaga T (1993) Muscle-fiber pennation angles are greater in hypertrophied than in normal muscles. *J Appl Physiol* 74:2740–2744
115. Kubo K et al (2003) Muscle architectural characteristics in women aged 20–79 years. *Med Sci Sports Exerc* 35:39–44
116. Ojanen T, Rauhala T, Häkkinen K (2007) Strength and power profiles of the lower and upper extremities in master throwers at different ages. *J Strength Cond Res* 21(1):216–222
117. Stock MS et al (2016) Evidence of muscular adaptations within four weeks of barbell training in women. *Human Mov Sci* 45:7–22
118. Nijboer-Oosterveld J, Van Alfen N, Pillen S (2011) New normal values for quantitative muscle ultrasound: obesity increases muscle echo intensity. *Muscle Nerve* 43:142–143
119. Fragala MS et al (2014) Biomarkers of muscle quality: N-terminal propeptide of type III procollagen and C-terminal agrin fragment responses to resistance exercise training in older adults. *J Cachexia Sarcopenia Muscle* 5(2):139–148
120. Taniguchi M et al (2017) Increase in echo intensity and extracellular-to-intracellular water ratio is independently associated with muscle weakness in elderly women. *Eur J Appl Physiol* 117(10):2001–2007
121. Yamada M et al (2017) Differential characteristics of skeletal muscle in community-dwelling older adults. *J Am Med Dir Assoc* 18(9):807.e9–807.e16
122. Ye X, Wang MJ, Xiao H (2017) Echo intensity of the rectus femoris in stable COPD patients. *Int J Chronic Obstr Pulm Dis* 12:3007–3015

123. Wilhelm EN et al (2014) Concurrent strength and endurance training exercise sequence does not affect neuromuscular adaptations in older men. *Exp Gerontol* 60:207–214
124. Fukumoto Y et al (2015) Age-related ultrasound changes in muscle quantity and quality in women. *Ultrasound Med Biol* 41(11):3013–3017
125. Schneider CA, Rasband WS, Eliceiri KW (2012) NIH Image to ImageJ: 25 years of image analysis. *Nat Methods* 9(7):671–675
126. Aagaard P et al (2001) A mechanism for increased contractile strength of human pennate muscle in response to strength training: changes in muscle architecture. *J Physiol* 534(Pt. 2):613–623
127. Greening NJ et al (2015) Bedside assessment of quadriceps muscle by ultrasound after admission for acute exacerbations of chronic respiratory disease. *Am J Respir Crit Care Med* 192:810–816
128. Lieber RL, Friden J (2002) Morphologic and mechanical basis of delayed-onset muscle soreness. *J Am Acad Orthop Surg* 10(1):67–73
129. Austin N, Nilwik R, Herzog W (2010) In vivo operational fascicle lengths of vastus lateralis during sub-maximal and maximal cycling. *J Biomech* 43(12):2394–2399
130. Erskine RM et al (2009) In vivo specific tension of the human quadriceps femoris muscle. *Eur J Appl Physiol* 106(6):827–838
131. Korhonen MT et al (2009) Biomechanical and skeletal muscle determinants of maximum running speed with aging. *Med Sci Sports Exerc* 41:844–856
132. Cadore EL et al (2012) Echo intensity is associated with skeletal muscle power and cardiovascular performance in elderly men. *Exp Gerontol* 47:473–478
133. Wu R et al (2016) Effects of age and sex on neuromuscular-mechanical determinants of muscle strength. *Age* 38:57
134. Muraki S, Fukumoto K, Fukuda O (2013) Prediction of the muscle strength by the muscle thickness and hardness using ultrasound muscle hardness meter. *Springerplus* 2:457
135. Varanoske AN et al (2017) Scanning plane comparison of ultrasound-derived morphological characteristics of the vastus lateralis. *Clin Anat* 30(4):533–542
136. Wilhelm EN et al (2014) Relationship between quadriceps femoris echo intensity, muscle power, and functional capacity of older men. *Age* 36:9625
137. Tang ZW et al (2015) Size of quadriceps femoris may contribute to thyrotoxic periodic paralysis. *Med Hypotheses* 85(6):749–753
138. Watson EL et al (2015) Progressive resistance exercise training in CKD: a feasibility study. *Am J Kidney Dis* 66(2):249–257
139. Kawai H et al (2018) Morphological and qualitative characteristics of the quadriceps muscle of community-dwelling older adults based on ultrasound imaging: classification using latent class analysis. *Aging Clin Exp Res* 30(4):283–291
140. Caresio C et al (2017) Fully automated muscle ultrasound analysis (MUSA): robust and accurate muscle thickness measurement. *Ultrasound Med Biol* 43(1):195–205
141. Welch D et al. (2018) Thigh muscle and subcutaneous tissue thickness measured using ultrasound imaging in older females living in extended care: a preliminary study. *Aging Clin Exp Res* 30(5):463–469
142. Shim DG, Kwon TY, Lee KB (2017) Rectus femoris muscle atrophy and recovery caused by preoperative pretibial traction in femoral shaft fractures-comparison between traction period. *Orthop Traumatol Surg Res* 103(5):691–695
143. Bembem MG (2002) Use of diagnostic ultrasound for assessing muscle size. *J Strength Cond Res* 16:103–108
144. Tok F et al (2011) Effects of botulinum toxin-A on the muscle architecture of stroke patients: an ultrasonographic study. *J Rehabil Med* 43(11):1016–1019
145. Tomlinson DJ et al (2014) The impact of obesity on skeletal muscle architecture in untrained young vs. old women. *J Anat* 225(6):675–684
146. Padhiar N et al (2008) Pennation angle of the soleus in patients with unilateral Achilles tendinopathy. *Disabil Rehabil* 30(20):1640–1645
147. Nagano Y, Higashihara A, Edama M (2015) Change in muscle thickness under contracting conditions following return to sports after a hamstring muscle strain injury—a pilot study. *Asia Pac J Sports Med Arthrosc Rehabil Technol* 2(2):63–67
148. Sjogaard G, Saltin B (1982) Extra- and intracellular water spaces in muscles of man at rest and with dynamic exercise. *Am J Physiol* 243(3):R271–R280
149. Thoirs K, English C (2009) Ultrasound measures of muscle thickness: intra-examiner reliability and influence of body position. *Clin Physiol Funct Imaging* 29(6):440–446
150. Berg HE, Tedner B, Tesch PA (1993) Changes in lower limb muscle cross-sectional area and tissue fluid volume after transition from standing to supine. *Acta Physiol Scand* 148(4):379–385
151. Cerniglia LM et al (2007) Effects of acute supine rest on mid-thigh cross-sectional area as measured by computed tomography. *Clin Physiol Funct Imaging* 27(4):249–253
152. Wagle JP et al (2017) Comparison of the relationship between lying and standing ultrasonography measures of muscle morphology with isometric and dynamic force production capabilities. *Sports* 5(4):88
153. Koda H et al (2018) Relationship between muscle strength asymmetry and body sway in older adults. *J Aging Phys Act* 26(3):457–461
154. Heckmatt JZ, Pier N, Dubowitz V (1988) Assessment of quadriceps femoris muscle atrophy and hypertrophy in neuromuscular disease in children. *J Clin Ultrasound* 16(3):177–181
155. Dupont AC et al (2001) Real-time sonography to estimate muscle thickness: comparison with MRI and CT. *J Clin Ultrasound* 29(4):230–236
156. Backhaus M et al (2001) Guidelines for musculoskeletal ultrasound in rheumatology. *Ann Rheum Dis* 60(7):641–649
157. Narici MV et al (1989) Changes in force, cross-sectional area and neural activation during strength training and detraining of the human quadriceps. *Eur J Appl Physiol Occup Physiol* 59(4):310–319
158. Kubo K et al (2003) Muscle architectural characteristics in young and elderly men and women. *Int J Sports Med* 24(2):125–130
159. Reeves ND, Narici MV, Maganaris CN (2004) In vivo human muscle structure and function: adaptations to resistance training in old age. *Exp Physiol* 89:675–689
160. Miljkovic I et al (2015) Greater skeletal muscle fat infiltration is associated with higher all-cause and cardiovascular mortality in older men. *J Gerontol A Biol Sci Med Sci* 70(9):1133–1140
161. Rollins KE et al (2016) The impact of sarcopenia and myosteatosis on outcomes of unresectable pancreatic cancer or distal cholangiocarcinoma. *Clin Nutr* 35(5):1103–1109
162. Montano-Loza AJ et al (2016) Sarcopenic obesity and myosteatosis are associated with higher mortality in patients with cirrhosis. *J Cachexia Sarcopenia Muscle* 7(2):126–135
163. Reimers K et al (1993) Skeletal muscle sonography: a correlative study of echogenicity and morphology. *J Ultrasound Med* 12(2):73–77
164. Ismail C et al (2015) Diagnostic ultrasound estimates of muscle mass and muscle quality discriminate between women with and without sarcopenia. *Front Physiol* 6:302
165. Pillen S et al (2009) Skeletal muscle ultrasound: correlation between fibrous tissue and echo intensity. *Ultrasound Med Biol* 35(3):443–446
166. Moher D et al (2009) Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. *J Clin Epidemiol* 62(10):1006–1012

Affiliations

Stany Perkisas^{1,19}  · **Stéphane Baudry**^{2,19} · **Jürgen Bauer**³ · **David Beckwée**^{4,5} · **Anne-Marie De Cock**¹ · **Hans Hobbelen**^{6,7,20} · **Harriët Jager-Wittenaar**^{6,7,21} · **Agnieszka Kasiukiewicz**⁸ · **Francesco Landi**⁹ · **Ester Marco**^{10,11,12} · **Ana Merello**¹³ · **Karolina Piotrowicz**¹⁴ · **Elisabet Sanchez**¹³ · **Dolores Sanchez-Rodriguez**^{12,15,16,17} · **Aldo Scafoglieri**^{18,19} · **Alfonso Cruz-Jentoft**¹³ · **Maurits Vandewoude**^{1,19}

¹ University of Antwerp—University Geriatric Center, Leopoldstraat 26, 2000 Antwerp, Belgium

² Laboratory of Applied Biology and Neurophysiologie, Université libre de Bruxelles, Brussels, Belgium

³ Department of Geriatric Medicine, Klinikum, Carl von Ossietzky University, Oldenburg, Germany

⁴ Rehabilitation Sciences Research Department, Vrije Universiteit Brussel, Brussels, Belgium

⁵ Department of Rehabilitation Sciences and Physiotherapy, Faculty of Medicine and Health Sciences, University of Antwerp, Antwerp, Belgium

⁶ Hanze University of Applied Sciences, Groningen, The Netherlands

⁷ Research Group Healthy Ageing, Allied Health Care and Nursing, University of Groningen, Groningen, The Netherlands

⁸ Department of Geriatrics, Medical University in Białystok, Białystok, Poland

⁹ Department of Geriatrics, Neuroscience and Orthopedics, Teaching Hospital “Agostino Gemelli”, Catholic University of the Sacred Heart School of Medicine, Rome, Italy

¹⁰ Physical and Rehabilitation Medicine, Parc de Salut Mar, Barcelona, Spain

¹¹ Rehabilitation Research Group, Hospital del Mar Research Institute, Barcelona, Spain

¹² Universitat Autònoma de Barcelona, Barcelona, Spain

¹³ University Hospital Ramón y Cajal, Madrid, Spain

¹⁴ Department of Internal Medicine and Gerontology, Faculty of Medicine, Jagiellonian University Medical College, Kraków, Poland

¹⁵ Geriatrics Department, Parc Salut Mar, Barcelona, Spain

¹⁶ Institut Hospital Del Mar d’Investigacions Mèdiques (IMIM), Barcelona, Spain

¹⁷ Universitat Pompeu Fabra, Barcelona, Spain

¹⁸ Faculty of Medicine and Pharmacy, Vrije Universiteit Brussel, Brussels, Belgium

¹⁹ Belgian Ageing Muscle Society, Liege, Belgium

²⁰ Department of General Practice and Elderly Care Medicine, University Medical Centre, University of Groningen, Groningen, The Netherlands

²¹ Department of Maxillofacial Surgery, University Medical Centre Groningen, Groningen, The Netherlands