INTRODUCTION

Skeletal muscles play a pivotal role in mobile and static functions, including body movement and maintenance of posture [1], and many studies have investigated the relationship of skeletal muscles with physical wellness, morbidity, and mortality [2, 3]. As a result, sarcopenia, defined as the loss of muscle mass and strength, is now formally recognized as a disease with an ICD Code. As the importance of muscle has been emphasized, a variety of...
imaging methods have been introduced for the assessment of muscle quantity. CT is considered as one of the gold-standard methods for noninvasive assessment of muscle quantity [4]. The CT measurement of muscle quantity is based on the difference in the X-ray attenuation value (measured in Hounsfield unit [HU]) of each pixel, as each component of the body (including the skeletal muscles, bone, adipose tissue, and visceral organs) has a specific attenuation threshold [5, 6], which is a prerequisite for their identification in cross-sectional CT images. Thus, additional CT scans, with resulting additional costs and radiation dose, are not needed for muscle evaluation if a CT scan was performed as a part of patient care for any other cause, including the assessment of disease and treatment response.

Recently, the European Working Group on Sarcopenia in Older People 2 revised the definition of sarcopenia by adding muscle function based on muscle quantity to the former definitions; however, muscle quantity by itself is limited in its ability to predict outcomes [7]. Muscle quality, which represents the micro- and macroscopic aspects of muscle architecture and composition, is also related to outcome: intra- and extra-myocellular fat deposition occurs with aging or disuse of muscle and leads to decreased muscle strength and function, followed by increased mortality [8]. As lipid deposition lowers the density of muscles [9], several studies have investigated the measurement of muscle quality using CT—by measuring the muscle density or stratifying the intramuscular components according to their HU distributions—and evaluated its usefulness for determining prognosis, independent of muscle quantity [10-12]. Accordingly, CT has emerged as an accurate measurement tool for the determination of muscle quantity and quality. Nevertheless, standardized parameters for image acquisition [13] have not been determined, and it is questionable whether muscle measurements remain stable if CT parameters are altered. In fact, X-ray attenuation and noise may be influenced by many factors, including tube voltage, tube current, slice thickness, and the reconstruction algorithm used. Variation in these CT parameters might worsen the accuracy of muscle quality measurement if the attenuation threshold is narrowed. Therefore, the aim of our study was to evaluate the reliability of the measurement of muscle quantity and quality under variable CT parameters.

**MATERIALS AND METHODS**

This study did not require approval from the Institutional Review Board owing to the study design and because it did not involve any humans/animals.

**Phantom**

The CIRS Model 004 CT Simulator (Computerized Imaging Reference Systems, Inc.) was used in this study. This phantom was originally designed to simulate the CT density of the lumbar vertebra, but it also simulates the size, shape, and CT density of the abdominal muscles and subcutaneous adipose tissue at the level of the 2nd to 4th lumbar region. The target CT density of the muscle compartment was 45 HU.

**CT Protocols and Image Acquisitions**

CT images of the phantom were acquired on three different systems (Somatom Definition AS+: Siemens Healthineers; Discovery CT750 HD, GE Healthcare; and Ingenuity Core 128, Philips Healthcare). Images were repeatedly obtained at the level of the 3rd lumbar vertebra, with modulation of tube voltage (120 kVp and 80 kVp) and tube current (standard mAs and low mAs), leading to three different categories of radiation dose: standard dose (STD), low dose with reduced voltage (LD-kVp), and low dose with reduced current (LD-mAs). In addition, the slice thickness (thin sections [1 or 1.25 mm], medium sections [2.5 or 3 mm], and thick sections [5 mm]) and image reconstruction algorithms (filtered back projection [FBP] and iterative reconstruction [IR]) were varied. The detailed imaging parameters and their modulations are presented in Table 1.

**Determination of the Reference Standard Muscle Compartment**

The reference standard muscle compartment area was determined according to the segmentation of known phantom compartments, including the muscle, subcutaneous fat, visceral fat, internal organs, and vertebra. Each compartment was automatically segmented according to the following processes (Fig. 1).

**Preparation: Generation of the Reference Map**

Reference maps of the compartments, including the muscle, subcutaneous fat, visceral fat, internal organs, and vertebra, were generated from known phantom compartments.
Step 1: Initial Segmentation of the Phantom Area

The area of the phantom on the CT images was separated from the background area using Otsu’s thresholding method [14]. Noise reduction using an anisotropic diffusion filter and mathematical morphology was simultaneously performed to generate an initial segmentation of the phantom area [15, 16].

Step 2: Rigid Registration

The reference map was registered to the initial segmented area (the phantom area) using a center-of-mass match and rigid transformation [17].

Step 3: Final Segmentation of the Muscle Compartment

The muscle compartment was segmented using the reference map.

Measurement of Muscle Area and Density

For the CT images acquired with different parameters, the cross-sectional areas of the muscle and its components were segmented using pre-defined HU thresholds that were deemed generally acceptable in previous studies (Fig. 2) [11]. The total muscle area was segmented using a threshold of -190 to 150 HU. The total muscle area component was categorized into skeletal muscle areas (SMA) with a threshold of -29 to 150 HU, and intermuscular adipose tissue areas (IMA) with a threshold of -190 to -30 HU. The SMA was further subcategorized into normal attenuation muscle area (NAMA; threshold, 30 to 150 HU) and low attenuation muscle area (LAMA; threshold, -29 to 29 HU) to facilitate evaluation of the quality of muscle according to CT density [11, 18]. The mean density of the reference standard muscle compartment was calculated by averaging the HU value of each pixel within the area. A web-based toolkit for automatic measurement of muscle area is available at https://iaidimage.com/app/aid-u/sarcopenia-l3.

Signal-to-Noise Ratio Analysis

The signal-to-noise ratio (SNR) was analyzed using the ImageJ program (National Institute of Health, https://imagej.nih.gov/ij). In the phantom CT images, the regions of interest were placed in the right paraspinal muscle and the air outside the phantom. The standard deviation of the air outside the phantom was regarded as background image noise, and the SNR was calculated by dividing the mean attenuation of the right paraspinal muscle by the background image noise [19].

Statistical Analysis

The mean values obtained from the three devices using the same parameters were used as representative values. The results obtained using different parameters were compared with those obtained with the standard protocol (i.e., STD with FBP reconstruction and thick sections [5 mm]) and with the pre-segmented area. In addition, the Dice similarity coefficient (DSC) [20] was used to evaluate the similarities of the cross-sectional areas of the total

Table 1. CT Image Acquisition Parameters

|                        | Somatom Definition AS+, Siemens Healthineers | Discovery CT750 HD, GE Healthcare | Ingenuity Core 128, Philips Healthcare |
|------------------------|---------------------------------------------|----------------------------------|---------------------------------------|
| Radiation dose*        | STD 120 kVp and 220 reference mAs           | STD 120 kVp and 100–400 mA       | STD 120 kVp and 321 reference mAs     |
|                        | LD-kVp 80 kVp and 220 reference mAs         | LD-kVp 80 kVp and 100–400 mA     | LD-kVp 80 kVp and 321 reference mAs   |
|                        | LD-mAs 120 kVp and 100 reference mAs        | LD-mAs 120 kVp and 10–300 mA     | LD-mAs 120 kVp and 168 reference mAs  |
| Slice thickness* (mm)  | Thin section 1                              | 1                                | 1                                     |
|                        | Medium section 3                            | 2.5                              | 3                                     |
|                        | Thick section 5                             | 5                                | 5                                     |
| Reconstruction algorithm*| IR (SAFIRE, iterative strength level 1)   | IR (ASIR 30%)                    | IR (iDose4)                           |
| Field of view (mm)     | 380                                         |                                  |                                       |
| Kernel                 | Standard                                    |                                  |                                       |

* A total of 18 images per device were obtained with the modulation of radiation dose (three conditions), slice thickness (three conditions), and reconstruction algorithm (two conditions). ASIR = adaptive statistical iterative reconstruction, FBP = filtered back projection, IR = iterative reconstruction, LD-kVp = low dose with reduced voltage, LD-mAs = low dose with reduced current, SAFIRE = sinogram affirmed iterative reconstruction, STD = standard dose
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The Wilcoxon signed-rank test was used to evaluate the relationships of the SNR and background noise to tube voltage, radiation dose, and reconstruction algorithm. The Friedman test was used to evaluate the SNR and background noise with different slice thicknesses. Statistical analysis was performed with PASW 18 (SPSS Inc.), and a \( p \) value < 0.05 was considered statistically significant.

RESULTS

A total of 18 image sets were acquired, including various combinations of imaging parameters, three radiation dose settings (STD, LD-mAs, and LD-kVp protocols), three different slice thicknesses (thin, medium, and thick sections), and two different reconstruction algorithms (FBP and IR).

Impact of CT Parameters on Segmentation Area

Compared with the reference standard muscle compartment, the segmented SMA values were higher with both the STD and the LD-mAs protocols, while they were lowest with the LD-kVp protocol, regardless of the slice thickness and reconstruction algorithm (Fig. 3). The

![Fig. 2. Segmentation of the muscle compartment and its components using HU thresholds.](image)

Fig. 2. Segmentation of the muscle compartment and its components using HU thresholds. The components are the areas segmented using HU thresholds to represent biological and clinical tissue components, including SMA (composed of NAMA and LAMA) and IMA. HU = Hounsfield unit, IMA = intermuscular adipose tissue area, LAMA = low attenuation muscle area, LD-kVp = low dose with reduced voltage, LD-mAs = low dose with reduced current, NAMA = normal attenuation muscle area, SMA = skeletal muscle area.

![Fig. 1. Determination of the reference standard muscle compartment.](image)

Fig. 1. Determination of the reference standard muscle compartment. The reference standard muscle compartment was determined using the following process: separation of the phantom area from the background area on the CT image (step 1), rigid registration of the reference map (preparation) onto the phantom area (step 2), and segmentation of the muscle compartment (step 3). The final area of muscle on the CT image was 122.2 cm².
SMAl results did not differ significantly between the thin, medium, and thick sections, or between the different reconstruction algorithms.

In all the CT protocols, SMA occupied at least 91.7% of the pre-segmented area. In contrast, NAMA was not constant across the protocols, varying between 59.7% and 81.7% of the pre-segmented area, despite the fact that the target HU of the muscle stated by the manufacturer (45 HU) was within the threshold range. Of note, the proportions of SMA and NAMA in the images generated using IR algorithm were higher than those in the images generated using FBP algorithm in all but the standard protocol, although the differences for SMA and NAMA were less than 2.7% and 4.7%, respectively.

The muscle area and SMA both showed good similarity, with the DSCs being within the range of 0.96–1.00 for the muscle area and 0.94–1.00 for the SMA. However, the DSCs of NAMA ranged from 0.74 to 0.96, showing variation that
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was dependent on the CT parameters (Fig. 4).

**Impact of CT Parameters on Density**

In all cases, the mean density of the reference standard muscle compartment was lower than the target density stated by the manufacturer (range, 39.0–44.9 HU) (Fig. 5). The mean density increased with thin slices (difference range, 0–1.3 HU) and low tube voltage (difference range, 3.4–3.5 HU), but decreased with low radiation dose (difference range, 1.01–1.8 HU) and IR usage (difference range, -0.1–1.2 HU).

**Signal-to-Noise Ratio**

The background noise increased with low tube voltage (120 kVp vs. 80 kVp: 6.55 vs. 12.75; p < 0.001) and low tube current (standard vs. low: 6.55 vs. 8.85; p < 0.001) (Table 2). In association with the changes in background noise, the SNR decreased with low tube voltage (120 kVp vs. 80 kVp: 5.92 vs. 3.21; p < 0.001) and low tube current (standard vs. low: 5.92 vs. 4.78; p < 0.001). Background noise decreased with thicker sections (1 [or 1.25] vs. 3 [or 2.5] vs. 5: 12.60 vs. 8.80 vs. 7.35; p < 0.001), while SNR increased with thicker sections (1 [or 1.25] vs. 3 [or 2.5] vs. 5: 3.41 vs. 5.04 vs. 5.53; p < 0.001). Images reconstructed with IR algorithm showed lower background noise (IR vs. FBP: 9.20 vs. 10.90; p = 0.002) and higher SNR (IR vs. FBP: 4.83 vs. 3.78; p = 0.005) than those reconstructed with FBP algorithm.

**DISCUSSION**

To the best of our knowledge, no consensus exists over which CT protocols or parameters yield the most reliable muscle measurements. Although many studies have investigated the assessment of muscle quantity and quality, they have not provided the CT acquisition parameters in sufficient detail [21], which has hampered the reproduction of their work. In this phantom study, we investigated the reliability of CT measurement of muscle quantity and quality with different CT parameters, using two popular methods for measuring muscle quantity and quality on CT: using 1) the cross-sectional area within the defined HU thresholds, and 2) the mean density. In addition, we used the DSC to evaluate similarities in cross-sectional area over different acquisitions using different parameters, relative to the standard protocol acquisitions. According to our findings, the cross-sectional total muscle areas (-190 to 150 HU) and SMA (-29 to 150 HU) with different protocols were similar to those using the standard protocol, whereas the areas of NAMA (30 to 150 HU) were not stable across the protocols. Furthermore, the mean density of the muscle compartment was inaccurate and fell short of the target attenuation stated by the manufacturer.

The most popular attenuation threshold used to measure muscle mass (or quantity) is -29 to 150 HU. In addition, the lower margin might be expanded to -190 HU to consider the IMA component. Our study revealed that the measurement of muscle quantity using these thresholds was reliable irrespective of the CT parameters used, consistent with the report of a prior study involving human subjects which reported that SMA measured with thin slice thickness and low radiation dose was constant (lower by less than 5%) [22]. In contrast, the measurement of muscle quality using the mean CT density was subject to the choice of CT parameters. Fuchs et al. [22] also reported that thin slices (2 mm vs. 5 mm thickness) and low radiation dose altered the mean density by 4.8 and 17.3 HU, respectively. In principle, the mean HU values should be constant irrespective of the radiation dose used, as the outliers of HU, caused by noise, at both ends of the range should offset each other [22]. However, this was not the case for the reference standard muscle compartment used in our study, probably because noise from neighboring structures, such as the vertebrae and adipose tissue, altered the attenuation of the affected pixels. Moreover,
in CT images with low SNR, such as those acquired with protocols using low tube current, low tube voltage, or thin slice sections, the effect could become substantial. The photoelectric effect along with the low tube voltage might have altered the CT density [23, 24].

Some studies have suggested differentiating low attenuation muscle from normal attenuation muscle on the basis of the HU threshold (usually using a cutoff of 30 HU) [11, 25]. However, according to our results, the measurement of muscle quality based on this narrower HU threshold would not be constant across different protocols. The reliability of muscle quality evaluation across protocols would most likely be worse if differentiation based on this threshold was applied to human subjects, as intra- and extra-cellular fat components actually exist within the muscle tissue of real subjects, unlike in our phantom [26]. Therefore, we suggest that the CT measurement of muscle quality should use only a limited range of CT parameter settings. However, further studies are required in future to determine which parameters should be adopted for reliable measurements.

We also investigated the usefulness of the IR algorithm for muscle measurement on dose-reduced protocols; we expected the reduced noise to better reflect the true
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Table 2. Background Image Noise and SNRs of Phantom Images with Different Protocols

| Protocol                  | Background Noise | SNR*         |
|---------------------------|------------------|--------------|
|                           | Median           | IQR          | P*   | Median | IQR          | P*   |
| Tube voltage (kVp)        |                  |              |      |        |              |      |
| 120 (n = 18)              | 6.55             | 6.05, 9.95   | < 0.001 | 5.92   | 4.24, 7.77  | < 0.001 |
| 80 (n = 18)               | 12.75            | 10.70, 19.00 |        | 3.21   | 2.48, 3.82  |        |
| Tube current (mAs)        |                  |              |      |        |              |      |
| Standard (n = 18)         | 6.55             | 6.05, 9.95   | < 0.001 | 5.92   | 4.24, 7.77  | < 0.001 |
| Low (n = 18)              | 8.85             | 7.63, 11.53  |        | 4.78   | 3.61, 5.57  |        |
| Slice thickness (mm)      |                  |              |      |        |              |      |
| 1 or 1.25 (n = 18)        | 12.60            | 11.20, 19.68 |        | 3.41   | 2.66, 3.83  |        |
| 2.5 or 3 (n = 18)         | 8.80             | 7.35, 11.33  |        | 5.04   | 3.60, 5.82  |        |
| 5 (n = 18)                | 7.35             | 6.28, 9.90   |        | 5.53   | 4.21, 7.12  |        |
| Reconstruction algorithm  |                  |              |      |        |              |      |
| FBP (n = 27)              | 10.90            | 7.30, 13.80  | 0.005 | 3.78   | 3.30, 5.66  | 0.005 |
| IR (n = 27)               | 9.20             | 7.40, 11.60  |        | 4.83   | 3.51, 5.79  |        |

*Using the Wilcoxon signed-rank test (tube voltage, radiation dose, and reconstruction algorithm) or Friedman test (slice thickness). FBP = filtered back projection, IQR = interquartile range, IR = iterative reconstruction, SNR = signal-to-noise ratio

Our study has several limitations. First, the phantom did not fully reflect the muscle components of the human body, as the HU of the muscle area was theoretically constant in the phantom. Moreover, as the phantom was primarily used to measure bone density, measurement of muscle density might have been inaccurate. Thus, the measured mean HU values of the muscle compartment were all lower than the true value (i.e., 45 HU), irrespective of the CT parameters. Nevertheless, using the phantom, we could investigate the reliability of measurements in relation to CT parameters, as we could adjust the CT parameters without any ethical considerations of radiation dose. However, additional animal or clinical studies might be required in future to confirm our results. Second, contrast agent administration would also influence the reliability of muscle measurement, and it was not possible to assess this using a phantom. In fact, many studies have reported that contrast usage and its enhancement phase alters the interpretation of muscle quantity and quality [22, 30, 31]. Lastly, we did not investigate the reliability of muscle measurement across different scanners. However, we believe that it was impractical to compare them head-to-head, as each device has innate strengths and limitations in terms of dose reduction and image reconstruction. In addition, the aim of this study was not to determine which device was superior; rather, we wanted to determine the constant effects of alterations to the protocols, irrespective of the device characteristics.

In conclusion, the measurement of muscle quantity using HU threshold is a reliable method, regardless of the CT protocol used. Conversely, the measurement of muscle quality using the mean CT density and a narrower HU threshold was inconsistent and inaccurate, with variations across the different CT protocols. Therefore, future studies are warranted to determine the optimal CT protocols for reliable measurements of muscle quality.

Conflicts of Interest
The authors have no potential conflicts of interest to disclose.

Author Contributions
Conceptualization: Kyung Won Kim. Data curation: Dong Wook Kim, Jiyeon Ha, Yousun Ko. Formal analysis: Dong
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