

A model for estimating body shape biological age based on clinical parameters associated with body composition

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Background: To date, no studies have attempted to estimate body shape biological age using clinical parameters associated with body composition for the purposes of examining a person's body shape based on their age.

Objective: We examined the relations between clinical parameters associated with body composition and chronological age, and proposed a model for estimating the body shape biological age.

Methods: The study was conducted in 243,778 subjects aged between 20 and 90 years who received a general medical checkup at health promotion centers at university and community hospitals in Korea from 2004 to 2011.

Results: In men, the clinical parameters with the highest correlation to age included the waist-to-hip ratio ($r = 0.786$, $P < 0.001$), hip circumference ($r = -0.448$, $P < 0.001$), and height ($r = -0.377$, $P < 0.001$). In women, the clinical parameters with the highest correlation to age include the waist-to-hip ratio ($r = 0.859$, $P < 0.001$), waist circumference ($r = 0.580$, $P < 0.001$), and hip circumference ($r = 0.520$, $P < 0.001$). To estimate the optimal body shape biological age based on clinical parameters associated with body composition, we performed a multiple regression analysis. In a model estimating the body shape biological age, the coefficient of determination (R^2) was 0.71 in men and 0.76 in women.

Conclusion: Our model for estimating body shape biological age might be a novel approach to variation in body shape that is due to aging. We assume that our estimation model would be used as an adjunctive measure in easily predicting differences in body shape with the use of clinical parameters that are commonly used to assess the status of obesity in a clinical setting.

Keywords: chronological age, body shape, biological age

Introduction

Substantial changes in body composition occur over the aging process. Many studies have suggested that over the course of the aging process, the amount of fat tends to increase and is preferentially stored in the abdominal area, while skeletal muscle mass and bone mineral density decline.¹⁻⁴ Understanding how changes in body composition, especially fat distribution, affect the risk for many chronic diseases such as cardiovascular diseases and type 2 diabetes mellitus, is a primary concern.

It is well known that body mass index (BMI) is one of the most popular clinical parameters for estimating body composition, and it is a reliable indicator for body fat mass percentage (BFM%). It is also an obesity index whose causal relationship with morbidity and mortality has been well established.⁵⁻⁷ Because it can be efficiently used to determine obesity in a large-sized population, BMI is frequently used for the assessment of obesity in a clinical setting.

It has been reported, however, that the distribution of body fat rather than its amount is more likely to be associated with health-related risks,⁸ and that abdominal obesity is an independent risk factor for raising the morbidity and mortality as such.^{9–11} This has led to an emphasis on the importance of other parameters, including the abdominal obesity index in particular, than BMI in assessing obesity.

Measures for abdominal obesity include waist circumference (WC), waist-to-hip ratio (WHR), and abdominal sagittal diameter. In addition, abdominal obesity is also evaluated by measuring the amount of body fat using ultrasonography, computed tomography, and magnetic resonance imaging. Of the various types of measures that are used to assess obesity, the amount of visceral body fat measured on computed tomography scans is known to best indicate abdominal obesity;^{12–14} however, its use is somewhat restricted because of the cost and exposure to irradiation. Accordingly, in an actual clinical setting, WC and WHR are mainly used to assess abdominal obesity. According to several previous studies examining various parameters that are commonly used to assess abdominal obesity in a clinical setting, WC was a more reliable parameter than WHR in making a diagnosis of abdominal obesity.¹⁵ In addition, these studies have also shown that WC was also the most reliable indicator in predicting the amount of visceral fat.^{10,16,17}

Body fat is redistributed with increasing age, as more of it becomes located in the abdominal cavity.^{14,18} The reason for this redistribution is not really known, but declining testosterone and growth hormone levels in combination with declining rates of lipolysis of visceral fat associated with aging may play a role in men.¹⁹ Low testosterone levels have been shown to be associated with increased visceral fat mass in men.²⁰ In women, the declining estrogen levels after menopause may be a critical factor. While several studies have investigated the relationship between weight change, obesity, and visceral fat with age and the risk of developing lifestyle-related diseases,^{21–24} the effect of maintaining one's youthful body shape (in terms of appearance) on reducing such risks has been given significantly less attention.

Several recent studies have reported the difference in body shape between men and women, and they have examined age-related differences in body shape using three-dimensional body scans. It has become more and more clinically important to assess body shape based on the pattern of body fat distribution and redistribution, rather than to assess obesity simply based on an increased amount of body fat. Meanwhile, several approaches have been used to objectively assess body shape from this context.^{25–30} Presumably, the assessment of body shape based on three-dimensional body scans might

be unavailable in a clinical setting. Therefore, it might also be mandatory to conduct studies to assess the status of body shape associated with age using clinical parameters that are currently used to assess obesity in a clinical setting.

It remains problematic that in general, anthropometric measures such as BMI or WHR are insufficient for the assessment of this redistribution.³¹ In addition, anthropometric measures have been mainly used to assess the status of obesity based on body composition up until the present.

Age after birth is a measure of the degree of aging in individual, this is termed as chronological age or calendar age, which is a definition of aging as a function of time. Aging is characterized by a time-related decline in physiological functions and changes in morphology, and is usually assessed by the chronological age, which is defined as time elapsed since birth. Chronological age fails to provide an accurate indicator of the aging process. Because tissues age at different rates and because diseases vary enormously among individuals, humans become increasingly different from one another with age. But, biological age estimates the functional status of an individual in reference to his or her chronological peers on the basis of how well he or she functions in comparison with others of the same chronological age.^{32,33} Biological age is estimated based on aging biomarkers that are subject to change depending on one's calendar age. Overall, biological age is a more reliable indicator of the degree of aging and one's health status as compared to an individual's calendar age. To date, a variety of models for estimating biological age have therefore been proposed to meet relevant study objectives. To estimate the physical, biochemical, and hormonal biological ages, we have also previously proposed the models for estimating biological ages using relevant biomarkers;³⁴ however, we did not propose a model for estimating body shape biological age using the clinical parameters associated with body composition.

Given this background, we conducted the current study to examine the correlations of age with BMI, BFM%, lean body mass percentage (LBM%), WC, and the WHR using clinical parameters associated with body composition. In addition, by using these parameters, we also proposed a model of body shape biological age that estimated a person's body shape based on their age. Thus, our estimation model will help people understand the above correlations, and it will promote bidirectional communication between physicians and patients.

Methods

Subjects

The current study was conducted in 243,778 subjects aged between 20 and 90 years; the subjects comprised

men and women who received a general medical checkup for health promotion and disease prevention at health promotion centers at university and community hospitals during a period ranging from June 2004 to January 2011. In the current study, exclusion criteria included the presence of uncontrolled hypertension and diabetes, and the subjects taking current medications for cardiovascular, pulmonary, hepatic, pancreatic, or renal diseases. Subjects were also excluded if their reported clinical parameters exceeded the standard deviations of the normal range.

Clinical parameters

To analyze the differences in clinical parameters due to aging, and to minimize the potential confounding effects of diseases or abnormal measurements, we set inclusion criteria considering both the standard deviations of parameters that were collected and the normal range which was established by the American Medical Association. Of these clinical parameters, we excluded those which were not in agreement with the inclusion criteria. Only the screened data were used for the statistical analysis.

For the clinical parameters associated with obesity, we collected data such as height, weight, WC, hip circumference (HC), LBM%, and BFM%. Height and weight were measured using height-measuring equipment and an electronic scale while subjects wore light clothes. As for WC, the thinnest area between the inferior part of the lowest rib and the iliac crest was measured as a unit of cm when subjects were in an upright position. As for HC, the location of the greater trochanter or the widest circumference was measured as a unit of cm. This was followed by the measurement of LBM% and BFM% using multifrequency segmental bioelectrical impedance (Inbody®, Biospace Co, Ltd, Seoul, Korea). BMI was calculated using height and weight as shown below.

Then, using the measurements of WC and HC, the WHR was calculated:

$$\text{BMI (kg/m}^2\text{)} = \text{Weight (kg)}/\text{Height}^2 \text{ (m}^2\text{)} \quad (1)$$

Statistical analysis

To identify the correlation between the clinical parameters and chronological age, a Pearson's correlation analysis was performed. To propose a model for estimating body shape biological age, we performed a stepwise method in multiple regression analysis between the age and clinical parameters that had been selected by the correlation analysis. We employed multiple regression analysis with assigning variance inflation factor to evaluate multicollinearity among the parameters. Statistical analysis was completed using SPSS 12.0 (SPSS Inc, Chicago, IL, USA). Statistical significance was set at $P < 0.05$.

Results

Baseline characteristics

A total of 243,778 subjects participated in the current study, of which there were 132,336 men and 110,042 women. All the clinical parameters associated with obesity are represented in Table 1. Mean age was 47.17 ± 10.98 years in men and 46.94 ± 11.96 years in women. In men, mean height and weight were 170.57 ± 5.96 cm and 71.16 ± 9.23 kg, respectively. In women, these values were 157.91 ± 5.63 cm and 56.92 ± 7.64 kg, respectively.

The correlation between age and the clinical parameters associated with body composition

In men, the clinical parameters with the highest correlation with the age included the WHR ($r = 0.786$, $P < 0.001$), HC

Table 1 Clinical characteristics of study subjects by gender

Parameters	Males (n = 132,336)		Females (n = 110,042)	
	Mean \pm SD	Range	Mean \pm SD	Range
Age (years)	47.17 \pm 10.98	20.00–90.00	46.94 \pm 11.95	20.00–90.00
Ht (cm)	170.57 \pm 5.96	141.00–190.00	157.91 \pm 5.63	140.00–189.00
Wt (kg)	71.16 \pm 9.23	37.50–100.00	56.92 \pm 7.64	31.00–99.80
BMI (kg/m ²)	24.25 \pm 2.45	15.01–30.01	22.65 \pm 2.72	15.01–30.01
LBM%	72.26 \pm 4.11	52.39–85.00	65.51 \pm 4.64	52.01–85.00
BFM%	23.30 \pm 4.58	10.00–42.98	29.85 \pm 4.82	10.00–43.00
WC (cm)	84.83 \pm 4.78	60.00–105.00	77.07 \pm 6.08	60.00–105.00
HC (cm)	95.51 \pm 1.79	76.50–110.00	92.74 \pm 1.96	75.00–110.00
WHR	0.90 \pm 0.03	0.67–1.19	0.83 \pm 0.04	0.66–1.18

Abbreviations: SD, standard deviation; Ht, height; Wt, weight; BMI, body mass index; LBM%, lean body mass percentage; BFM%, body fat mass percentage; WC, waist circumference; HC, hip circumference; WHR, waist-to-hip ratio.

($r = -0.448, P < 0.001$), and height ($r = -0.377, P < 0.001$). In women, the clinical parameters with the highest correlation with age included the WHR ($r = 0.859, P < 0.001$), WC ($r = 0.580, P < 0.001$), and HC ($r = 0.520, P < 0.001$). All of the collected data surrounding the clinical parameters had a significant correlation with age. In men, however, only the clinical parameters associated with obesity other than BMI ($r = 0.005, P = 0.191$) had a significant correlation with age (Tables 2 and 3).

Estimation of body shape biological age based on clinical parameters associated with body composition

To estimate the optimal body shape biological age based on the clinical parameters associated with body composition, we performed a multiple regression analysis. In a model for estimating body shape biological age in men, the coefficient of determination (R^2) was 0.71, and the relevant clinical parameters included the WHR, WC, height, LBM%, and weight. In a model for estimating the body shape biological age in women, the coefficient of determination (R^2) was 0.76, and the relevant clinical parameters included the WHR, height, WC, LBM%, and weight (Tables 4 and 5). Figures 1 and 2 illustrate the correlations between biological age and chronological age in a model for estimating body shape biological age in men and women.

Discussion

As shown in the current study, a model for estimating body shape biological age was effective in predicting the age-related changes that occur in body shape based on clinical parameters associated with body composition (WHR, WC, height, LBM%, weight, and HC) in approximately 71% of men and approximately 76% of women. Regarding the correlation between age and the clinical parameters associ-

Table 3 Correlation coefficients between age and clinical parameters in females

Parameters	Females (n = 110,042)	P-value
Ht	-0.46	<0.001
Wt	0.14	<0.001
BMI	0.41	<0.001
LBM%	-0.43	<0.001
BFM%	0.41	<0.001
WC	0.58	<0.001
HC	0.52	<0.001
WHR	0.86	<0.001

Abbreviations: Ht, height; Wt, weight; BMI, body mass index; LBM%, lean body mass percentage; BFM%, body fat mass percentage; WC, waist circumference; HC, hip circumference; WHR, hip-to-waist ratio.

ated with body composition, seven parameters other than BMI for men, and all of the eight parameters for women had a significant correlation with age. Of the clinical parameters that had a significant correlation with age. Of the clinical parameters that had a significant correlation with age in men, height, weight, LBM%, and HC had a negative correlation with age, while BFM%, WC, and the WHR had a positive correlation. In women, height and LBM% had a negative correlation with age, while weight, BMI, BFM%, WC, HC, and the WHR had a positive correlation.

Previous studies have shown that both men and women have experienced a decrease in height due to increased age, and this was more notable in women.³⁵⁻⁴³ A decrease in height due to increased age mainly originates from changes in the vertebrae (eg, vertebral fracture, weakening of the paravertebral muscles, changes in posture, degenerative changes in the intervertebral disc, vertebral deformity, and kyphosis). In addition, its onset begins at the age of 45 years or so, and then it continues with increased age.^{36,37,43,44} It is therefore known that a decrease in height due to increased age is associated with osteoporotic fractures, and it is also an indicator of the prediction of vertebral fractures and risks of developing femoral fractures in the elderly.^{45,46} In the current study, there was also a significant correlation

Table 2 Correlation coefficients between age and clinical parameters in males

Parameters	Males (n = 132,336)	P-value
Ht	-0.38	<0.001
Wt	-0.21	<0.001
BMI	0.01	0.191
LBM%	-0.14	<0.001
BFM%	0.15	<0.001
WC	0.16	<0.001
HC	-0.45	<0.001
WHR	0.79	<0.001

Abbreviations: Ht, height; Wt, weight; BMI, body mass index; LBM%, lean body mass percentage; BFM%, body fat mass percentage; WC, waist circumference; HC, hip circumference; WHR, waist-to-hip ratio.

Table 4 Body shape biological age prediction model in males ($R^2 = 0.705$)

Parameters	PRC ± SEM	Standard PRC	t	P-value
WHR	281.419 ± 0.922	0.692	305.362	0.000
HC	-1.595 ± 0.014	-0.260	-114.282	0.000
Ht	-0.149 ± 0.005	-0.081	-27.987	0.000
LBM%	-0.112 ± 0.007	-0.042	-15.500	0.000
Wt	-0.052 ± 0.004	-0.043	-13.681	0.000

Abbreviations: PRC, partial regression coefficients; SEM, standard error of the mean; WHR, waist-to-hip ratio; HC, hip circumference; Ht, height; LBM%, lean body mass percentage; Wt, weight.

Table 5 Body shape biological age prediction model in females ($R^2 = 0.763$)

Parameters	PRC \pm SEM	Standard PRC	t	P-value
WHR	224.964 \pm 0.922	0.768	244.049	0.000
Ht	-0.245 \pm 0.005	-0.115	-49.171	0.000
HC	-0.631 \pm 0.016	0.103	40.325	0.000
LBM%	-0.200 \pm 0.006	-0.077	-32.074	0.000
WC	-0.123 \pm 0.006	-0.063	-20.244	0.000

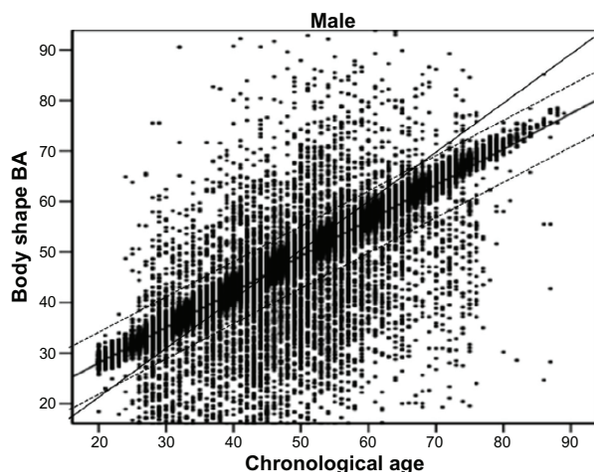
Abbreviations: PRC, partial regression coefficients; SEM, standard error of the mean; WHR, waist-to-hip ratio; Ht, height; HC, hip circumference; LBM%, lean body mass percentage; WC, waist circumference.

between a decrease in height and increased age. In a model for estimating body shape biological age, the degree of the correlation between the two parameters was significantly higher in women than men.

BMI is widely used for nutritional assessment, obesity classification, and as a prognostic variable for mortality; however, there are many limitations. First, BMI could potentially produce an inaccurate diagnosis of “overweight” and “obese” in some special populations such as athletes, body builders, and elderly patients. Second, BMI-associated mortality in specific situations is controversial.⁴⁷

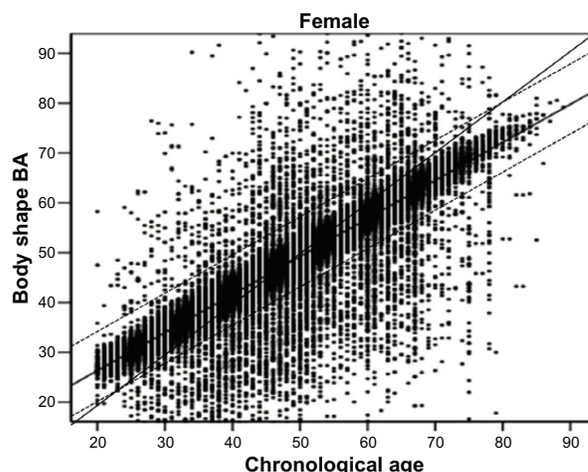
In a model for estimating the body shape biological age, we included the height and weight of men and only the height of women rather than the BMI. In addition, the correlation between BMI and age reached statistical significance only in women.

As shown in the current results, both men and women demonstrated a positive correlation between BFM% and

**Figure 1** The body shape BA prediction model in males.

Notes: The 45° line represents the theoretical normal aging line in males. The experimental line is the predicted regression line on body shape BA, and both dotted lines mean the range of ± 1 standard deviation (67%). The equation: body shape BA: $-16.958 + 281.419$ (WHR) $- 1.595$ (HC) $- 0.149$ (Ht) $- 0.112$ (LBM%) $- 0.052$ (Wt).

Abbreviations: BA, biological age; WHR, waist-to-hip ratio; HC, hip circumference; Ht, height; LBM%, lean body mass percentage; Wt, weight.

**Figure 2** The body shape and BA prediction model in females.

Notes: The 45° line represents the theoretical normal aging line in females. The experimental line is the predicted regression line on body shape BA and both dotted lines mean the range of 67%. The equation: body shape BA: $-137.388 + 224.964$ (WHR) $- 0.245$ (Ht) $+ 0.631$ (HC) $- 0.2$ (LBM%) $- 0.123$ (WC).

Abbreviations: BA, biological age; WHR, waist-to-hip ratio; Ht, height; HC, hip circumference; LBM%, lean body mass percentage; WC, waist circumference.

age, and a negative correlation between age and LBM%. In addition, we included only the LBM% in a model for estimating biological age. Previous studies have established that BFM% was increased and LBM% was decreased with increased age.^{48,49} An increase in BFM% due to increased age originated from a decrease in LBM%, rather than from an increase in BFM% in men. It has been reported that an increase in BFM% rather than a decrease in LBM% might play a key role in causing an increase body fat due to aging in women.^{50,51}

Regarding the distribution of body fat depending on body shape, it has been reported that abdominal obesity is a notable, independent risk factor for raising morbidity and mortality rates.⁹⁻¹¹ Therefore, the clinical significance of weight, BMI, and the BFM% would greatly vary depending on the types of abdominal obesity, (ie, android or gynoid type). Previous studies have shown that WC is the most reliable indicator for reflecting the amount of visceral fat in the assessment of abdominal obesity.^{10,15-17} As shown in the current results, both men and women had a positive correlation between increased age and both WC and the WHR; however, HC had a negative correlation with age in men and a positive correlation with in women. According to previous studies, the WHR was increased until the age of 55–64 years, and then it decreased thereafter in men, though it continually increased with increased age in women.^{52,53} These authors noted that these findings occurred because of the redistribution of body fat due to changes in sex hormones in both men and women aged between 45 and 54 years.^{52,53} Other studies have also

shown that hormone therapy reduced the WHR, thus providing more evidence for the redistribution of body fat because of hormonal changes.^{54,55} As shown in the current results, the WHR was included in a model for estimating body shape biological age in both men and women. In addition, HC was included in this model for men, and both HC and WC were included for women. In particular, in estimating the body shape biological age, both men and women showed that the WHR was the most powerful factor. Previous studies have shown that WC is a more reliable indicator of predicting abdominal obesity. From a perspective that considers the differences in body shape due to aging, however, the WHR is a more powerful factor, and it better reflects differences in body shape.

In cross-sectional studies, the prevalence of high body weight or obesity ($BMI > 30 \text{ kg/m}^2$) increases with age up to about age 60 and then declines.⁵⁶ Although this does not indicate that the excessive accumulation of body fat rarely occurs in the elderly, this does suggest that visceral obesity increasingly occurs because of the redistribution of body fat due to increased age, and this leads to changes in body shape.^{25–30,56} Accordingly, considering variation in body shape due to aging, it would be more appropriate for clinical evaluation to be based on body shape due to the increased occurrence of visceral obesity, rather than the consistent application of clinical parameters for assessing obesity.

We propose that a model for estimating body shape biological age be appropriate for assessing differences in body shape due to increased age with the application of clinical parameters associated with body composition. Moreover, by presenting the clinical parameters for assessing obesity that are commonly used in a clinical setting – and thereby comparing the results with those obtained from age-matched controls – our estimation model would be used to perform an overall analysis of the differences in body shape due to aging. Furthermore, by comparing one's current body shape biological age with their calendar age (thereby expressing whether the body shape biological age is older or younger than one's calendar age), our estimation model would be easily accepted by people, and it would also be of help for facilitating communication between physicians and patients. In addition, it would also be of help for providing the motives to improve the status of obesity, and for setting personal goals. In a model for assessing body shape biological age, however, we did not include the factors that might affect the clinical parameters associated with obesity, including the amount of physical activity, eating habits

such as nutritional intake, and socioeconomic status, and we excluded the data of some subjects who were suspected of having diseases or conditions based on certain criteria. There were also limitations in the current study in that we did not perform a detailed analysis of the participants' medical history such as a present illness, a past history, and a family history.

Conclusion

Our model for estimating body shape biological age might be a novel approach to variation in body shape that is due to aging despite several limitations. We assume that our estimation model would be used as an adjunctive measure in easily predicting differences in body shape with the use of clinical parameters that are commonly used to assess the status of obesity in a clinical setting. Our results will provide baseline data for further studies to examine not only the effects of changes in body shape due to the aging process on the degree of aging in individuals, the rate of aging progression, and life expectancy, but they will also provide information surrounding correlations between body shape, disease, morbidity, and mortality.

Disclosure

The authors report no conflicts of interest in this work.

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