Comparison of Cardio–Ankle Vascular Index (CAVI) and CAVI₀ in Large Healthy and Hypertensive Populations

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Aim: The cardio–ankle vascular index (CAVI) represents the blood pressure-independent arterial stiffness from the origin of the aorta to the ankle. CAVI₀ has been proposed as a variant index. We aimed to clarify the difference between CAVI and CAVI₀ among large populations, and to explore reasons of the difference.

Methods: The subjects were 5,293 Japanese healthy and 3,338 hypertensive people. Simple and multiple regression analyses were performed using age, sex, body mass index, systolic, and diastolic blood pressure (Pd) as variables. Sub-group analysis was performed by sex and age. The CAVI values with and without adjustment by reference pressure were also compared.

Results: CAVI had a positive correlation with Pd, while CAVI₀ had a negative correlation with Pd in the healthy population. The CAVI values of the hypertensive group were higher than those of healthy group in both men and women, but the CAVI₀ values in women of the hypertensive group in the 30–39 age group was significantly lower than that of the corresponding healthy group. Differences of CAVI values with or without modification using the reference pressure were 1.09% ± 1.38% for the healthy group and 3.68% ± 1.66% for the hypertensive group.

Conclusion: CAVI showed the expected values, but CAVI₀ showed inexplicable results in the healthy and hypertensive populations. The differences were due to the strong dependency of CAVI₀ on Pd. Differences of CAVI values with or without reference pressure were negligible. These results indicate that CAVI obtained by the VaSera system is appropriate, but CAVI₀ is not.

Key words: CAVI, CAVI₀, haPWV, Bramwell–Hill's equation, Stiffness Parameter β

Introduction

It is known that arterial stiffness reflects atherosclerosis and it is related to cardiovascular events. Measurements of arterial stiffness have been attempted from the late 19th century¹. It is established that the velocity of the pulse that arises from the heart is correlated with the stiffness of the artery². Consequently, pulse wave velocity (PWV) has been used as an index of arterial stiffness³. However, PWV essentially changes according to changes in blood pressure at the measuring time. This has been shown theoretically and also experimentally⁴. Therefore, PWV is an inappropriate index for research studying the effect of blood pressure on arterial stiffness.

An attempt to establish an arterial stiffness index independent of blood pressure at measuring time was made by Hayashi et al.⁵. They found that the relationship between a change in vascular diameter and internal blood pressure showed an exponential curve,
using isolated arteries. Whereas, when blood pressure change was expressed as a natural logarithm, the relationship between diameter change and blood pressure change showed nearly a straight line. The inclination of this line, regarded as an arterial stiffness parameter which does not change according to changes in blood pressure at the measuring time, was defined as stiffness parameter $\beta$ (Supplement Seq.1). Later, Kawasaki et al. showed a method to measure $\beta$ using an ultrasonic diagnostic imaging apparatus by setting the diastolic pressure ($P_d$) in place of the reference pressure (Supplement Seq.7).

In 2004, the cardio-ankle vascular index (CAVI) was developed by applying the Bramwell–Hill equation to Kawasaki’s $\beta$ formula using PWV, as an index reflecting arterial stiffness of the arterial tree from the origin of the aorta to the ankle. The VaSera system was invented to measure CAVI. The precise methods were already published and the index was defined by eq.1.

$$CAVI=a \times \left(2\rho \times \ln \left(\frac{P_s}{P_d}\right) \times PWV^2\right) + b \quad \text{eq.1}$$

[PWV: pulse wave velocity of the arterial tree from the origin of the aorta to the ankle, $P_s$: systolic blood pressure, $P_d$: diastolic blood pressure, $\rho$: blood density, $\Delta P$: $P_s$–$P_d$, a, b: coefficients.]

Using this index, many studies have been published over the last 15 years; these studies include various valuable findings confirmed by researchers throughout the world. Though both CAVI and PWV values increase with aging, the increase is more marked with CAVI. Further, CAVI is more closely correlated to aortic distensibility than PWV. Patients with arteriosclerotic diseases, such as coronary arterial disease, cerebrovascular diseases, and chronic kidney disease show high CAVI values. Furthermore, CAVI is known to be high among those with coronary risk factors, such as hypertension, diabetes mellitus, dyslipidemia, and sleep apnea syndrome. Many studies have shown that CAVI is a predictor of cardiovascular events.

In addition, several studies on the acute changes of CAVI showed reasonable changes as a result of the administration of metoprolol, doxazosin, and nitroglycerin.

However, recently, Spronck et al. proposed a similar index, termed CAVI$_0$, which is based on the assumption that CAVI is dependent on blood pressure, whereas, CAVI$_0$ is not. The calculation formula for CAVI$_0$ is defined in eq.2.

$$CAVI_0 = 2\rho \times \frac{PWV^2}{P_d} - ln \left(\frac{P_d}{P_0}\right) \quad \text{eq.2}$$

[$P_0$: reference pressure (100 mmHg)]

According to Spronck et al., there are two reasons for the blood pressure dependence of CAVI. The first reason is that the slight difference between the $\beta$ of Hayashi and that of Kawasaki et al. is expressed by the second term of the right side of eq.2, which is related to the reference pressure, and this is the first element of blood pressure dependency. The second reason is the $\ln \frac{P_s}{P_d}$ of the CAVI expression, which is different from 1/$P_d$. Spronck et al. compared CAVI and CAVI$_0$ values based on mathematical simulations using blood pressure values from a randomized list, and showed the difference between CAVI and CAVI$_0$, as it related to the dependency on the blood pressure at measuring time.

Based on the assumption that the artery is completely monotonous and blood pressure is the same at both sides of the artery, Spronck’s equation (eq.2) could be mathematically reasonable. However, Spronck’s equation (eq.2) includes only $P_d$. This raises one concern that CAVI$_0$ might be influenced too strongly by $P_d$. The pulse transition time measured in CAVI is not just a foot-to-foot period of the pulse waves at the pressure level of $P_d$, and it is not suitable to apply the CAVI$_0$ theory simply to this condition.

To date, there has been no detailed report comparing CAVI and CAVI$_0$ with a large population; carotid–femoral PWV has been compared to CAVI and resulted in a relatively small correlation [$r^2=0.18$ or $r^2=0.31$].

**Aim**

The aim of this study was to compare and clarify the actual differences between CAVI and CAVI$_0$ in the large population. Both values were compared in a healthy group and a hypertensive group by simple and multiple regression analyses using age, sex, body mass index (BMI), $P_s$, and $P_d$ as variables. Also, the comparison was performed in subgroups by sex and age. As a result, there were differences between CAVI and CAVI$_0$. The reason for the differences is discussed from the viewpoint of the properties of heart–ankle PWV (haPWV) used in the VaSera system.

Furthermore, Spronck et al. pointed out that the arterial stiffness value should be corrected with a reference pressure, and $-\ln \frac{P_d}{P_0}$ (see eq.2) was added in the CAVI$_0$ equation. In this article, the actual differences between the CAVI values with or without reference pressure were calculated and compared among the healthy group and the hypertensive group, respectively.
Subjects
The population-based sample used in the present analysis was comprised of \( n = 8,631 \) (healthy and \( n = 3,338 \), hypertensive) Japanese subjects aged \( \geq 20 \) years residing in major cities nationwide who underwent a health check at the Japan Health Promotion Foundation with the complete data set \( n = 28,400 \) among \( n = 32,627 \) of the published data \( 32, 33 \). A flow diagram of the population is shown in Fig. 1.

Analysis Method

1. Comparison of CAVI and CAVI\( \text{I}_0 \) by Simple and Multiple Regression Analysis

Firstly, in order to overview the basic feature of CAVI and CAVI\( \text{I}_0 \) as a whole, the coefficients of correlation between CAVI and CAVI\( \text{I}_0 \) were obtained in the healthy and hypertensive groups. Then, simple and multiple regression analyses were performed in the healthy and hypertensive groups using age, sex, BMI, Ps, and Pd as variables. Dummy variables were set for sex (men: 1, women: 2). To avoid confounding by blood pressure, regression analysis was performed in separate models with Ps and with Pd. The criteria of the healthy group and the hypertensive group are shown in Fig. 1.

2. Comparison of the Differences between CAVI and CAVI\( \text{I}_0 \) among the Healthy and Hypertensive Groups in Sex and Age subgroups

To investigate the differences between CAVI and CAVI\( \text{I}_0 \) in detail, subjects were divided into 12 groups by sex and age strata. The age groups were 20–29, 30–39, 40–49, 50–59, 60–69, and \( \geq 70 \) years. The mean values of CAVI and CAVI\( \text{I}_0 \) and the statistical significance between the healthy group and the hypertensive group were calculated, respectively, in men and women.

Also, in order to investigate the relationship between CAVI and CAVI\( \text{I}_0 \) with blood pressure, Ps and Pd of the healthy group and the hypertensive group were obtained in the same manner in each age stratum in men and women.

3. Correlation between haPWV and Ps, Pd, and Mid Pressure (Pm) in the Healthy Group

Shirai et al. reported that CAVI without coefficient “\( a \)” and “\( b \)” can be described as indicated in eq.3 \( ^{34} \):

\[
\text{CAVI} = 2\rho \times \frac{\ln \left( \frac{P_s}{P_d} \right)}{\Delta P} \times \text{PWV}^2
\]

\[
= 2\rho \times \frac{\ln (Ps) - \ln (Pd)}{Ps - Pd} \times \text{PWV}^2 \quad \text{eq.3-1}
\]

\[
\text{CAVI} = 2\rho \times \frac{1}{P_{\text{m}}} \times \text{PWV}^2 \quad \text{eq.3-2}
\]

[CAVI\( \text{I}_0 \): CAVI without coefficients “\( a \)” and “\( b \),” Pm: mid blood pressure \( \left( \frac{Ps + Pd}{2} \right) \)]

The reason why CAVI is based on Pm is as follows: CAVI is calculated from haPWV, which is the PWV of the arterial tree from the origin of the aorta to the ankle. As is well known, PWV has blood pres-
sure dependency\(^{10}\). Therefore, also in the cardiac cycle, PWV takes different values depending on the point of measurement between at Pd and at Ps; Shirai et al. showed that haPWV in CAVI corresponded mostly to Pm rather than Ps and Pd\(^{30}\). It means that the formula of CAVI is substantially based on Pm, whereas, Spronck et al. assumed that Pd should be used in CAVI because PWV measured at the foot-to-foot of pulse wave corresponds to at Pd. This is the essential difference. The influence of the second term of CAVI, which is related to the reference pressure, will be described later.

In order to reconfirm this, the relationship of haPWV with Ps, Pd, and Pm were studied in a healthy group, which is considered to have less influence of vascular remodeling with hypertension, by age strata with simple regression analysis.

### 4. Analysis of the Over-Estimation of CAVI\(_0\): Comparison of Pm/Pd in Men and Women

Since in CAVI\(_0\), PWV squared is divided by incompatible Pd instead of compatible Pm, there is a concern that the CAVI\(_0\) value is overestimated by Pm/Pd, which corresponds to CAVI\(_0\)/CAVI as a whole. In order to confirm the influence of this over-estimation, the values of Pm/Pd were calculated in the healthy group and the hypertensive group in each age stratum in men and women.

### 5. Comparison of CAVI Values with and without Reference Pressure

A combination of Bramwell–Hill's equation and Stiffness Parameter \(\beta\) gives eq.4 (Supplement Seq. 15).

\[
\beta = 2\rho \times \frac{\text{PWV}^2}{P} - \ln \left( \frac{P}{P_0} \right) \tag{eq.4}
\]

When PWV corresponds to Pd, by substituting P=Pd, the CAVI\(_0\) equation of eq. 2 is derived. For CAVI, PWV corresponds to Pm, and by substituting P=Pm, eq.5 is derived.

\[
\beta = \text{CAVI\(_{ref}\)} = 2\rho \times \frac{\text{PWV}^2}{P} - \ln \left( \frac{P_m}{P_0} \right) \tag{eq.5}
\]

[CAVI\(_{ref}\): the value with reference pressure P\(_0\).]

CAVI\(_{ref}\) described in eq.3-1 is the actual measured value without the coefficients “a” and “b,” and is comparable to CAVI\(_{ref}\), the mathematical value with reference pressure described in eq.5. It has been reported that there is no discrepancy in the clinical significance between CAVI with and without the coefficients “a” and “b”\(^{30}\).

In order to evaluate the differences of values, CAVI\(_{ref}\), CAVI and CAVI\(_0\) were compared and the difference ratio to CAVI\(_{ref}\) were calculated in the healthy group and hypertensive group.

### Statistical Analysis

Unpaired Welch's t-test was used to evaluate comparisons of CAVI and CAVI\(_0\) among the groups. Results were expressed as the mean ± standard deviation, and \(p<0.05\) was considered significant.

In the regression analysis, the dependent variables were CAVI and CAVI\(_0\), and the independent variables used for regression analysis were sex, age, body mass index (BMI), Ps, and Pd. Co-linearity between independent variables was confirmed by the Durbin–Watson test and Variance Inflator Factor.

In the simple regression analysis of haPWV with Ps, Pd, and Pm, the coefficient of correlation was obtained in each age stratum of the healthy group. In the comparison of \(\beta m\), \(\beta m'\), and CAVI\(_0\), the results were expressed as the mean ± standard deviation. All statistical analyses were performed using the SPSS software package (SPSS Inc., Chicago, IL, USA).

### Results

#### 1. Comparison of CAVI and CAVI\(_0\) with Simple and Multiple Regression Analysis

The coefficients of correlation between CAVI and CAVI\(_0\) in the healthy group and the hypertensive group were 0.923 and 0.955, respectively, with \(p<0.001\) for both. Subsequently, regression analyses with age, sex, BMI, Ps, and Pd in CAVI and CAVI\(_0\) were studied. The results are shown in Table 1. In the relationship with Ps (Table 1a, b), no major discrepancy was seen between CAVI and CAVI\(_0\), except for that in CAVI; Ps was not significant in the multiple regression analysis in the healthy group.

However, in the relationship with Pd (Table 1c, d), an inexplicable result was observed for CAVI\(_0\). For CAVI, the correlation coefficient (\(r\)) in simple regression with Pd in the healthy group was 0.214 (\(p<0.001\)). In multiple regression analysis, the standardized partial regression coefficient (\(\beta\)) was 0.040 (\(p=0.001\)). In the hypertensive group, for a simple regression with Pd, the result was -0.031 (\(p=0.076\)). In the multiple regression analysis, the standardized partial regression coefficient (\(\beta\)) was 0.145 (\(p<0.001\)). From these results, a positive correlation with Pd was observed for CAVI.

For CAVI\(_0\), the correlation coefficient (\(r\)) in the simple regression analysis with Pd in the healthy group was -0.064 (\(p<0.001\)). In the multiple regression analysis, the standardized partial regression coefficient (\(\beta\)) was -0.257 (\(p<0.001\)). In the hyperten-
sive group, in the simple regression analysis with Pd, the result was $-0.166\ (p<0.001)$. In multiple regression analysis, the standardized partial regression coefficient ($\beta$) was not significant. As shown above, a negative correlation with Pd was observed for CAVI₀.

2. **Comparison of the Significant Differences of CAVI and CAVI₀ between the Healthy and the Hypertensive Groups in Sex and Age Subgroups**

As shown in Fig. 2a-d, in general, a similarity was seen between CAVI and CAVI₀, but inexplicable
results were found for CAVI\(_0\). In the analysis of CAVI, the value of the hypertensive group exceeded the value of the healthy group in both men and women, and the differences were significant, except for women at the age of 30–39 years.

However, in the analysis of CAVI\(_0\), in the group with the age of 30–39 years, the values of the hypertensive group were less than the values of the healthy group in both men and women. (10.23 ± 1.54 vs. 10.43 ± 1.33, \(p=0.21\) in men, and 9.54 ± 1.22 vs. 10.26 ± 1.25, \(p=0.024\) in women, respectively). In the group with women at the age of 30–39 years, the values of the hypertensive group were significantly less than the values of the healthy group. Thus, a contra-

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**Fig. 2.** Comparison of the significant differences of CAVI and CAVI\(_0\) between healthy group and the hypertensive group in each age stratum in men (a, b) and women (c, d)
Correlation between haPWV and Ps, Pd, and Pm in the Healthy Group

Coefficients of correlation between haPWV and Ps, Pd, and Pm in the healthy group were obtained, and the results are shown in Table 2. In all age strata, Pm showed a stronger correlation with haPWV than Ps and Pd, indicating that Pm was a stronger factor in the determination of haPWV than Ps and Pd. This result was consistent with the previous report by Shi-rai et al. 34).

Analysis of the Over-Estimation in CAVI₀; Comparison of Pm/Pd in Men and Women

The changes of Pm/Pd at each age group were studied to clarify how the contradictory results (i.e.,

Table 2. Correlation of coefficients between haPWV and Ps, Pd, and Pm in the healthy group (n=5,293)

| Age   | 20-29 | 30-39 | 40-49 | 50-59 | 60-69 | ≥ 70 | Total |
|-------|-------|-------|-------|-------|-------|------|-------|
| n     | 980   | 1,948 | 1,260 | 818   | 255   | 32   | 5,293 |
| Ps    | .565 (*** | .529 (*** | .556 (*** | .520 (*** | .464 (*** | .332 (NS) | .544 (*** |
| Pd    | .599 (*** | .587 (*** | .632 (*** | .567 (*** | .536 (*** | .082 (NS) | .624 (*** |
| Pm    | .637 (*** | .604 (*** | .641 (*** | .599 (*** | .546 (*** | .290 (NS) | .631 (*** |

*** means p<0.001 and NS means p ≥ 0.05

Abbreviations: Ps, systolic blood pressure; Pd, diastolic blood pressure; Pm, mid blood pressure.

Fig. 3. Blood pressure in each age stratum in men (a) and women (b)

Abbreviations: Ps, systolic blood pressure; Pd, diastolic blood pressure.

3. Correlation between haPWV and Ps, Pd, and Pm in the Healthy Group

Coefficients of correlation between haPWV and Ps, Pd, and Pm in the healthy group were obtained, and the results are shown in Table 2. In all age strata, Pm showed a stronger correlation with haPWV than Ps and Pd, indicating that Pm was a stronger factor in the determination of haPWV than Ps and Pd. This result was consistent with the previous report by Shi-rai et al. 34).

4. Analysis of the Over-Estimation in CAVI₀; Comparison of Pm/Pd in Men and Women

The changes of Pm/Pd at each age group were studied to clarify how the contradictory results (i.e.,
higher CAVI₀ in healthy women than in hypertensive women at the age of 30–39 years), were obtained. The results are shown in Fig. 4. Pm/Pd values in the hypertensive group were less than those aged 30–39 and 40–49 in the healthy group in both men and women.

5. Comparison between CAVI Values with and without Reference Pressure

The difference between CAVI ref (with reference pressure P₀), CAVI' (actual measured value without the coefficients “a” and “b”), and CAVI₀ are shown in Fig. 5. The difference ratio of CAVI' and CAVI ref in the healthy group and in the hypertensive group were 1.09% ± 1.39% and 3.68 ± 1.66%, respectively, while the difference ratio of CAVI₀ to CAVI ref in the healthy group and in the hypertensive group were 37.58 ± 8.66% and 34.75 ± 9.48%, respectively.

Discussion

In general, some similarity was seen between CAVI and CAVI₀ with high values more than 0.9 in coefficients of correlation both in the healthy and hypertensive groups. However, as shown in Table 1 (d), the Pd was a significantly negative contributing factor for CAVI₀ in both the healthy group and hypertensive group by simple regression analysis, and it was also a negatively contributing factor for CAVI₀ in the healthy group by multiple regression analysis. Generally, those results were unreasonable. Whereas the Pd was a significantly positive contributing factor for CAVI in the healthy group by simple regression analysis and multiple regression analysis, and there was no significance between Pd and CAVI in the hypertensive group by both simple and multiple regression analysis. These results were acceptable.

Next, CAVI and CAVI₀ values were compared in subgroups of sex and age, as shown in Fig. 2. a-d. The CAVI₀ values of the hypertensive young women group were significantly lower than the value of the healthy young women group, and the significantly higher CAVI₀ was observed age > 50 years in both men and women. Whereas, CAVI values of the hypertensive young women group were almost the same as those of the healthy young women group, and the significantly higher CAVI in the hypertensive group was observed in all age strata of men and women, except for young women in their 30s. It is unreasonable that arterial stiffness in young hypertensive women is lower than that of the young healthy women in their 30s. Therefore, we tried to clarify the reason.

At first, the equations of CAVI and CAVI₀ (see eq.1 and eq.2 as described in the introduction), were
sound to the corresponding notch of the upper brachial artery, and the time from the upper brachial artery to the ankle are measured. The former time is measured in the end-systolic period, and the pressure level is far higher than Pd. This detection system may render the strong dependency of haPWV on Pm rather than Pd. We have already reported this item\(^{34}\). Spronck et al. gave a further comment on previous data\(^{34}\) in an attempt to explain that the dependency of CAVI on Pm might be due to arterial remodeling in hypertensive patients because Pm is considered to be related to the remodeling. However, the data of Table 2 in this paper was obtained from the healthy group, and the results were consistent in all age strata, irrespective of the influence of age-related remodeling. This data was essentially consistent with the data in our previous report.

Then, it can be concluded that CAVI was obtained at Pm during the pulse cycle between Ps and Pd in the VaSera system. Furthermore, to confirm the reason why CAVI\(_0\) in healthy women was higher than in hypertensive women at the age of 30–39 years, the contribution of Pd was studied. Fig. 4 shows Pm/Pd in men and women of all ages. Pm/Pd in the hypertensive group was significantly less than those of the healthy group.

**Fig. 5.** Comparison of measured values with and without the reference pressure

CAVI\(_{\text{ref}}\) is with reference pressure and CAVI\(_{\text{r}}\) is without reference pressure. (a) values in the healthy group, (b) values in the hypertensive group, (c) difference ratio of CAVI\(_{\text{r}}\) and CAVI\(_0\) with CAVI\(_{\text{ref}}\) in the healthy group, and (d) difference ratio in the hypertensive group.
aged 30–39 years and 40–49 years in both men and women. This corresponds to the results of the comparison of the significant contradiction between CAVI and CAVI₀ among women at the age of 30–39 years in the healthy group and the hypertensive group, and is considered to be the reason why the CAVI₀ values were reversed between the young hypertensive group and the healthy group, and that there were no significant differences between them in men.

Also, as Pd increases, Pm/Pd becomes relatively small, so the CAVI₀ value decreases. As a result, CAVI₀ correlated negatively with Pd. This is considered to be the main reason why CAVI₀ showed unreasonable results.

These results suggest that CAVI₀ was strongly dependent on Pd changes and the CAVI₀ value was underestimated in the case of the elevated Pd group.

Another important difference between CAVI and CAVI₀ is that CAVI₀ included \(-\ln\left(\frac{P_d}{P_0}\right)\) to be corrected with P₀. The influence of the reference pressure on CAVI was studied using clinical data without coefficients “a” and “b.” As shown in Fig. 5, the difference ratio between CAVI and CAVI₀ was small and practically negligible in clinical usage, while the difference ratio between CAVI₀ and CAVI was large.

From the above results, the difference between CAVI and CAVI₀ is due to the application of Pm in place of Pd, rather than the difference with or without the reference pressure.

Recently, two studies comparing CAVI and CAVI₀ in pediatric subjects of Slovakia reported a slight blood pressure dependency of CAVI. One study compared between normotensive boys and those with white-coat hypertension (WCH) with higher CAVI₃⁷), and the other study compared between normotensive girls and boys with spurious systolic hypertension (SSH) with lower CAVI₃⁸). No significant differences were seen in CAVI₀ in both cases.

However, this article did not show any other standard of arterial stiffness, and it is actually difficult to decide which index is more proper. Only the values of CAVI and CAVI₀ were presented on the assumption that there was no difference in either group regarding arterial stiffness. Although it is known that WCH is associated with sympathetic hyperactivity²⁹), which can modulate arterial stiffness³⁰), and it is also known that SSH is seen in young boys with highly elastic arteries⁴¹), some possibilities remain, in that arterial stiffness of the persons with WCH is actually high, and that of boys with SSH is actually low.

Conclusion
CAVI showed reasonable values, but CAVI₀ showed unreasonable values between healthy and hypertensive populations, especially in young women. This difference was due to the strong dependency of CAVI₀ on Pd. The differences in CAVI values with or without the reference pressure were negligible. These results indicate that CAVI obtained by the VaSera system is appropriate, whereas CAVI₀ is not. We concluded that the CAVI value obtained by using the VaSera system is reasonable and appropriate as a clinical index of blood pressure-independent arterial stiffness of the arterial tree from the origin of the aorta to the ankle, and the accumulated data published for the last 15 years is reliable.

Disclosures
Kohji Shirai, Kenji Suzuki, Kazuhiro Shimizu, and Masanobu Takata had no conflict of interest concerning this article.
Shinichiro Tsuda, Tomoyuki Yamamoto, Mitsuya Maruyama, and Koji Takahashi belong to Fukuda Denshi Co., Ltd. and are involved in the development of CAVI.

References
1) Young T. The Croonian Lecture: on the functions of the heart and arteries. Phil Trans R Soc Lond, 1809; 99: 1-31
2) Moens AI. Die Pulskurve [The Pulse Curve]. Leiden, The Netherlands: E. J. Brill; 1898
3) Bramwell JC, Hill AV. The velocity of the pulse wave in man. Proc R Soc Lond B Biol Sci, 1922; 93: 298-306
4) Bramwell JC, McDowali RJS, McSwiney BA. The variation of arterial elasticity with blood pressure in man. (Part I). Proc R Soc Lond B, 1923; 94: 450-454
5) Hayashi K, Handa H, Nagasawa S, Okumura A, Moritake K. Stiffness and elastic behavior of human intracranial and extracranial arteries. J Biomech, 1980; 13: 175-184
6) Kawasaki T, Sasayama S, Yagi S, Asakawa T, Hirai T. Non-invasive assessment of the age related changes in stiffness of major branches of the human arteries. Cardiovasc Res, 1987; 21: 678-687
7) Shirai K, Utino J, Otsuka K, Takata M. A novel blood pressure-independent arterial wall stiffness parameter; cardio-ankle vascular index (CAVI). J Atheroscler Thromb, 2006; 13: 101-107
8) Takahashi K, Yamamoto T, Tsuda S, Okabe F, Shimose T, Tsuji Y, Suzuki K, Otsuka K, Takata M, Shimizu K, Uchino J, Shirai K. Coefficients in the CAVI Equation and the Comparison Between CAVI With and Without the Coefficients Using Clinical Data. J Atheroscler Thromb, 2019; 26: 465-475
9) Hayashi K, Yamamoto T, Takahara A, Shirai K. Clinical assessment of arterial stiffness with cardio-ankle vascular index: theory and applications. J Hypertens, 2015; 33: 1742-1757
10) Topouchian J, Labat C2 Gautier S, Bäck M, Achimastos
A, Blacher J, Cwynar M, de la Sierra A, Pall D, Fantin F, Farkas K, Garcia-Ortiz L, Hakobyan Z, Jankowski P, Jelkovic A, Kobalava Z, Konradi A, Kотовская Y, Kotsani M, Lazareva I, Litvin A, Milyagin V, Mintale I, Persson O, Ramos R, Rogoza A, Rylkozyte L, Scuteri A, Sirenko Y, Soulis G, Tasic N, Udovychenko M, Uralzina S, Wolff-fahrt P, Zelvian P, Benetos A, Asmar R. Effects of metabolic syndrome on arterial function in different age groups: the Advanced Approach to Arterial Stiffness study. J Hypertens, 2018; 36: 824-833

11) Boardman H, Lewandowski AJ, Lazdam M, Kenworthy Y, with obstructive sleep apnea. Am J Hypertens, 2011; 24:

12) Horinaka S, Yabe A, Yagi H, Ishimura K, Harada H, Imamura T, Matsuoka H. Comparison of atherosclerotic indicators between cardiac and peripheral vasculature. J Hypertens, 2017; 35: 513-522

13) Suzuki J, Sakakibara R, Tomaru T, Tateno F, Kishi M, Ogawa H, Kurosu T, Shirai K. Stroke and carotid-ankle vascular stiffness index. J Stroke Cerebrovasc Dis, 2013; 22: 171-175

14) Kubo-sono T, Miyama T, Ueyama K, Nagaki A, Hama-saki S, Kusano K, Kubo-sono O, Tei C. Association between arterial stiffness and estimated glomerular filtration rate in the Japanese general population. J Atheroscler Thromb, 2009; 16: 840-845

15) Okura T, Watanabe S, Kurata M, Manabe S, Koreshawa M, Irita J, Enomoto D, Miyoshi K, Fukuoka T, Higaki J. Relationship between cardiac-ankle vascular index (CAVI) and carotid atherosclerosis in patients with essential hypertension. Hypertens Res, 2007; 30: 335-340

16) Takaki A, Ogawa H, Wakeyama T, Iwami T, Kimura M, Hadano Y, Matsuwa S, Miyazaki Y, Hira-tuska A, Matsu-zaki M. Cardiac-ankle vascular index is superior to brachial-ankle pulse wave velocity as an index of arterial stiffness. Hypertens Res, 2008; 31: 1347-1355

17) Ibata J, Sasaki H, Kakimoto T, Matsuno S, Nakatani M, Irita J, Enomoto D, Miyoshi K, Fukuoka T, Higaki J. Relationship between cardiac-ankle vascular index (CAVI) and carotid atherosclerosis in patients with essential hypertension. Hypertens Res, 2007; 30: 335-340

18) Gomez-Sanchez L, et al. The Association Between the cardio-ankle vascular index from blood pressure at the time of measurement. J Hypertens, 2017; 35: 1521-1523

20) Kumagai T, Kasai T, Kato M, Naito R, Maeno K, Kasagi S, Kawana F, Ishiwata S, Narui K. Establishment of the cardiac-ankle vascular index in patients with obstructive sleep apnea. Chest, 2009; 36: 779-786

21) Kasai T, Inoue K, Kumagai T, Kato M, Kawana F, Sagara M, Ishiwata S, Ohno M, Yamaguchi T, Momomura S, Narui K. Plasma pentraxin3 and arterial stiffness in men with obstructive sleep apnea. Am J Hypertens, 2011; 24: 401-407

22) Otsuka K, Fukuda S, Shimada K, Suzuki K, Nakanishi K, Yoshiyama M, Yoshikawa J. Serial assessment of arterial stiffness by cardiac-ankle vascular index for prediction of future cardiovascular events in patients with coronary artery disease. Hypertens Res, 2014; 37: 1014-1020

23) Gohbara M, Iwahashi N, Sano Y, Akiyama E, Maejima N, Tsukahara K, Hibi K, Kosuge M, Ebina T, Umemura S, Kimura K. Clinical Impact of the Cardiac-ankle Vascular Index for Predicting Cardiovascular Events After Acute Coronary Syndrome. Circ J, 2016; 80: 1420-1426

24) Sato Y, Nagayama D, Saiki A, Watanabe R, Watanabe Y, Imamura H, Yamaguchi T, Ban N, Kawana H, Nagumo A, Ohira M, Endo K, Kurosu T, Tomaru T, Shirai K, Tatsuno I. Cardiac-ankle Vascular Index is Independently Associated with Future Cardiovascular Events in Outpatients with Metabolic Disorders. J Atheroscler Thromb, 2016; 23: 596-605

25) Hitsumo T. Clinical usefulness of the cardiac-ankle vascular index as a predictor of primary cardiovascular events in patients with chronic kidney disease. J Clin Med Res, 2018; 10: 883-890

26) Shirai K, Song M, Suzuki J, Kurosu T, Oyama T, Nagayama D, Miyashita Y, Yamamura S, Takahashi M. Contradictory effects of α-1- and α-1-adrenergic receptor blockers on cardiac-ankle vascular stiffness index (CAVI) – CAVI is independent of blood pressure-. J Atheroscler Thromb, 2011; 18: 49-55

27) Spronck B, Avolio AP, Tan I, Butlin M, Reesink KD, Del-haas T. Arterial stiffness index beta and cardiac-ankle vascular index inherently dependent on blood pressure but can be readily corrected. J Hypertens, 2017; 35: 98-104

28) Spronck B, Avolio AP, Tan I, Butlin M, Reesink KD, Delhaas T. Reply: physics cannot be disputed. J Hypertens, 2017; 35: 1523-1525

29) Spronck B, Avolio AP, Tan I, Butlin M, Reesink KD, Del-haas T. Medical science is based on facts and evidence. J Hypertens, 2018; 36: 960-962

30) Shirai K, Shimizu K, Takata M, Suzuki K. Independence of the cardiac-ankle vascular index from blood pressure at the time of measurement. J Hypertens, 2017; 35: 1521-1523

31) Gomez-Sanchez L, et al. The Association Between the Cardiac-ankle Vascular Index and Other Parameters of Vascular Structure and Function in Caucasian Adults: MARK Study. J Atheroscler Thromb, 2015; 22: 901-911

32) Suzuki K, Ishizuka N, Miyashita Y, Shirai K. Epidemiological examination about the standard value and the validity for the standardization as the examination of CAVI (cardiac-ankle vascular index) noninvasive blood pressure-independent arteriosclerosis test. Niigata J Med Technol, 2008; 48-1: 2-10; (in Japanese)

33) Namekata T, Suzuki K, Ishizuka N, Shirai K. Establishing baseline criteria of cardiac-ankle vascular index as a new indicator of arteriosclerosis: a cross-sectional study. BMC Cardiovasc Disord, 2011; 11: 51

34) Shirai K, Shimizu K, Takata M, Suzuki K. Medical science is based on evidence (answer to Spronck et al. ’s refutation: physics cannot be disputed). J Hypertens, 2018; 36: 958-960

35) Mirault T, Pernet M, Frank M, Couade M, Niarr R,
Azizi M, Emmerich J, Jeunemaître X, Fink M, Tänner M, Messas E. Carotid stiffness change over the cardiac cycle by ultrafast ultrasound imaging in healthy volunteers and vascular Ehlers-Danlos syndrome. J Hypertens, 2017; 33: 1890-1896

36) Kenyhercz WE, Rterman B, Sita V, Illapani P, Dowell J, Mo X, White RD, Kolipaka A. Quantification of aortic stiffness using magnetic resonance elastography: measurement reproducibility, pulse wave velocity comparison, changes over cardiac cycle, and relationship with age. Magn Reson Med, 2016; 75: 1920-1926

37) Mestanik M, Jurko A, Spronck B, Avolio AP, Butlin M, Jurko T, Visnovcova Z., Mestanikova A, Langer P, Tonhajzerova I. Improved assessment of arterial stiffness using corrected cardio-ankle vascular index (CAVI0) in overweight adolescents with white-coat and essential hypertension. Scand J Clin Lab Invest, 2017; 77: 665-672

38) Jurko T, Mestanik M, Jurko A Jr, Spronck B, Avolio AP, Mestanikova A, Sekaninova N, Tonhajzerova I. Pediatric reference values for arterial parameters cardio-ankle vascular index (Cavi) and Cavi0. J Am Soc Hypertens, 2018; 12: e35-e43

39) Smith PA, Graham LN, Mackintosh AF, MA, Stoker JB, Mary DA. Sympathetic neural mechanisms in white-coat hypertension. J Am Coll Cardiol, 2002; 40: 126-132

40) Nardone M, Incognito AV, Millar PJ. Evidence for pressure-independent sympathetic modulation of central pulse wave velocity. J Am Heart Assoc, 2018; 7: e007971

41) Mahmud A, Feely J. Spurious systolic hypertension of youth: fit young men with elastic arteries. Am J Hypertens, 2003; 16: 229-232
Supplement

β Described with PWV and Blood Pressure

Hayashi defied the β as blood pressure-independent parameter of arterial stiffness in Eq. 1.

\[ P = P_0 \times e^{\beta \times \left( \frac{D}{D_0} - 1 \right)} \]  \hspace{1cm} \text{Eq. 1}

(β: specific stiffness of the blood vessel, P: blood pressure, P₀: reference pressure, D: blood vessel diameter, D₀: blood vessel diameter at P₀)

By dividing both sides of Eq. 1 by P₀ and taking natural logarithm,

\[ \ln \left( \frac{P}{P_0} \right) = \beta \times \left( \frac{D - D_0}{D_0} \right) \]  \hspace{1cm} \text{Eq. 2}

Therefore,

\[ \beta = \ln \left( \frac{P}{P_0} \right) \times \frac{D_0}{D - D_0} \]  \hspace{1cm} \text{Eq. 3}

By substituting P = Pₛ and P = Pₖ for Eq. 2 and transforming,

\[ \ln (Pₛ) - \ln (P₀) = \beta \times \left( \frac{Dₛ - D₀}{D₀} \right) \]  \hspace{1cm} \text{Eq. 4}

\[ \ln (Pₖ) - \ln (P₀) = \beta \times \left( \frac{Dₖ - D₀}{D₀} \right) \]  \hspace{1cm} \text{Eq. 5}

By taking the difference between both sides of Eq. 4 and Eq. 5 and transforming,

\[ \beta = \ln \left( \frac{Pₛ}{Pₖ} \right) \times \frac{D₀}{Dₛ - Dₖ} \]  \hspace{1cm} \text{Eq. 6}

Kawasaki et al. defined the β¹ as practically measurable stiffness parameter in Eq. 7.

\[ \beta¹ = \ln \left( \frac{Pₛ}{Pₖ} \right) \times \frac{D₀}{Dₛ - Dₖ} \]  \hspace{1cm} \text{Eq. 7}

From Eq. 6 and Eq. 7,

\[ \beta = \beta¹ \times \frac{D₀}{Dₑ} \]  \hspace{1cm} \text{Eq. 8}

From Eq. 2,

\[ \frac{D₀}{D} = \frac{1}{1 + \ln \left( \frac{P}{P_0} \right)} \]  \hspace{1cm} \text{Eq. 9}

By substituting Eq. 9 for Eq. 8 with D = Dₑ

\[ \beta = \beta¹ - \ln \left( \frac{Pₖ}{P₀} \right) \]  \hspace{1cm} \text{Eq. 10}

Since in the physiological range, the second term of right side of Eq. 10 is generally small compared to the first term, β is approximated by β¹.

On the other hand, the Bramwell–Hill equation is represented by Eq. 11.

\[ \text{PWV}² = \frac{\text{dP}}{\text{dD}} \times \frac{D}{2\rho} \]  \hspace{1cm} \text{Eq. 11}

(ρ: blood density)

Here, \( \frac{\text{dP}}{\text{dD}} \times \frac{D}{2} \) is the volume elastic modulus, and Eq. 11 is a general equation expressing the relationship between the wave velocity and the volume elastic modulus. If D changes in proportion to P, the volume elastic modulus is constant and the wave velocity PWV is also constant. In blood vessels, however, D changes exponentially with respect to P, which is indicated by the Stiffness Parameter β equation of Eq. 1.

The blood pressure dependency of PWV arises from the exponential nature of this blood vessel as follows:

When both sides of Eq. 1 are differentiated with D, the exponent becomes a coefficient from the differential formula of exponential,

\[ \frac{\text{dP}}{\text{dD}} = \frac{\beta}{D₀} \times P \]  \hspace{1cm} \text{Eq. 12}

Substituting Eq. 12 for the right side of Eq. 11

\[ \text{PWV}² = \frac{\beta}{D₀} \times P \times \frac{D}{2\rho} \]  \hspace{1cm} \text{Eq. 13}

β is represented by Eq. 14 by transforming Eq. 13.

\[ \beta = \frac{2\rho \times \text{PWV}²}{P} \times \left( \frac{D₀}{D} \right) \]  \hspace{1cm} \text{Eq. 14}

Substitute Eq. 9 for Eq. 14 and organize it with β,

\[ \beta = \frac{2\rho \times \text{PWV}²}{P} - \ln \left( \frac{P}{P₀} \right) \]  \hspace{1cm} \text{Eq. 15}

Since in the physiological range, the second term of right side of Eq. 15 is generally small compared to the first term, PWV² is approximately proportional to β and P.