Aquatic exercise is an attractive form of exercise that utilizes the various properties of water to improve physical health, including arterial stiffness. However, it is unclear whether regular head-out aquatic exercise affects aortic hemodynamics, the emerging risk factors for future cardiovascular disease. The purpose of this study was to investigate whether head-out aquatic exercise training improves aortic hemodynamics in middle-aged and elderly people. In addition, to shed light on the underlying mechanisms, we determined the contribution of change in arterial stiffness to the hypothesized changes in aortic hemodynamics. Twenty-three middle-aged and elderly subjects (62 ± 9 years) underwent a weekly aquatic exercise course for 15 weeks. Aortic hemodynamics were evaluated by pulse wave analysis via the general transfer function method. Using a polar coordinate description, companion metrics of aortic pulse pressure ($PPC = \sqrt{(systolic~blood~pressure)^2 + (diastolic~blood~pressure)^2}$) and augmentation index ($AIxC = \sqrt{(augmentation~pressure)^2 + (pulse~pressure)^2}$) were calculated as measures of arterial load. Brachial-ankle (baPWV, reflecting stiffness of the abdominal aorta and leg artery) and heart-ankle (haPWV, reflecting stiffness of the whole aortic and leg artery) pulse wave velocities were also measured. The rate of participation in the aquatic training program was 83.5 ± 13.0%. Aortic systolic blood pressure, pulse pressure, PPC, AIxC, baPWV, and haPWV decreased after the training ($P < 0.05$ for all), whereas augmentation index remained unchanged. Changes in aortic SBP were correlated with changes in haPWV ($r = 0.613, P = 0.002$) but not baPWV ($r = 0.296, P = 0.170$). These findings suggest that head-out aquatic exercise training may improve aortic hemodynamics in middle-aged and elderly people, with the particular benefits for reducing aortic SBP which is associated with proximal aortic stiffness.

Keywords: aquatic exercise, aortic blood pressure, reflection wave, Windkessel function, polar coordinate description
INTRODUCTION

The advancing age is the major risk factor for cardiovascular diseases (CVD), and this is attributable in part to stiffening of large elastic arteries (1). Central arterial stiffening leads to the augmented wave reflection which pre-maturely return during early systole and thus increases aortic systolic blood pressure (SBP), pulse pressure (PP), and augmentation index (AIx). These hemodynamic characteristics could be independent predictors of future cardiovascular events and all-cause mortality (2).

Regular aerobic exercise is a recommended lifestyle for preventing CVD and risk factors (3, 4). Particularly, aquatic exercise (e.g., swimming and head-out aquatic exercise) has less stress on joints and lower risk of falling due to buoyancy and water resistance, and it is performed by a wide range of populations such as overweight and obese individuals (5, 6), the elderly people (7), and patients with osteoarthritis (8). Several intervention studies demonstrated that swimming exercise training improved arterial stiffness and aortic hemodynamics (8–12). On the other hand, we (13) and others (14, 15) showed head-out aquatic exercise training-induced favorable reduction in arterial stiffness, while the influence on aortic hemodynamics remained fully unknown.

Recently, there is a gaining recognition about the limitations of using dimensionless ratio-based metrics in cardiovascular medicine such as AIx. Indices based on ratios may not sufficiently reflect information about their constituents. As shown in the top panel in Figure 1, although points A, B, and C have different magnitudes of wave reflection from the periphery (i.e., augmented pressure, AP), the same values of AIx are obtained for the corresponding aortic PP values. After calculating AIx, the physiological units of AP and PP are canceled out, which results in a dimensionless number that no longer characterizes the unique properties of cardiac load. To overcome such simplification, polar coordinate description has recently been proposed as an alternative method (16). Using this method, the ratio can be presented in the pressure domain using a hypotenuse via the Pythagorean theorem as the “companion” of AIx (AIxC) (17). Likewise, the “companion” of PP (PPC) can also be calculated by the Pythagorean theorem. In the bottom panel of Figure 1, points D and E have different PPC for the same PP (i.e., 50 mmHg). Thus, the polar coordinate description may offer a clue to the physiologically relevant interpretation without need for additional measurement.

The purpose of this study was to investigate the hypothesis that head-out aquatic exercise training improves aortic hemodynamics in middle-aged and elderly people. Combining the traditional and newly described companion metrics (i.e., AIxC and PPC), we characterized changes in aortic hemodynamics comprehensively. In addition, to shed light on the underlying mechanisms, we determined the contribution of change in arterial stiffness to the hypothesized changes in aortic hemodynamics.

MATERIALS AND METHODS

Subjects

A total of 59 individuals joined the city-operated aquatic exercise training class. Of all, 38 volunteers (10 men, 4 pre-menopausal, and 24 post-menopausal women, age range: 39–79 years) gave their written informed consent after the explanation of all potential risks and procedures of the present study. No one engaged in regular intense endurance training such as joining masters’ competition. All subjects were non-smokers. No women were taking hormone replacement therapy. This study was reviewed and approved by the Institutional Research Board.
Aquatic Exercise Training Program
The exercise program consisted of 15 training sessions (once a week) from July to October 2017. Each exercise session consisted of a 5-min warm-up (i.e., stretching) on land and 50 min of aquatic exercise in indoor pool (31.5°C of water temperature; 1.05–1.15 m of water depth), including 10 min of walking, 30 min of resistance and stepping exercise, and 10 min of stretching and relaxation (18). The last 5 min served as the cooling down on land. Based on the subjects’ age, self-reported physical activity level, and disease status (based on the most recent health checkup), each subject was assigned to the moderate (9 of 23 subjects) or the low-intensity (14 of 23 subjects) exercise classes. Five subjects in the moderate-intensity exercise class and 3 subjects in the low-intensity exercise class had exercise habits before enrolling in this intervention study. None of them had strength training or water exercise within the past 6 months. Aside from exercise training intervention, all subjects were instructed not to change daily activities, including their exercise habits. An experienced instructor supervised and provided typical water-based movements (19). Participants reported on a 6–20 scale of perceived exertion scale (RPE) during training after every class.

Measurements
Hemodynamic variables and arterial stiffness measurement were performed after 3 h of fasting and abstinence from caffeine intake. Each subject was asked not to drink alcohol and engage in vigorous physical activity within 24 h before the experiment. Individuals with prescribed medications (hyperlipidemia: n = 2; diabetes mellitus: n = 2) were instructed to take them as usual. After 5 min of supine rest, heart rate, limb BP, heart sound, and two segments of PWV [i.e., brachial to ankle PWV (baPWV), aortic (heart) to ankle PWV (haPWV)] were measured using a semiautomatic cardiovascular screening device (Form PWV/ABI, Omron-Colin, Kyoto, Japan), as previously reported (20). Aortic BP was estimated based on the previous study (21). The brachial arterial pressure waveform was resampled at 256 Hz with data acquisition and analysis software (AcqKnowledge; BIOPAC Systems). Subsequently, the arterial waveform data were fed into the SphygmoCor Excel software (AtCor Medical, Sydney, Australia), and a generalized transfer function was applied to estimate aortic BP. Aortic BP was calibrated by brachial BP. Schema of aortic pulse wave analysis was illustrated in Figure 2.

To quantify the magnitude of wave reflection from the periphery to the heart, aortic Alx was calculated as AP divided by aortic PP (22). In addition, the ratio-based measure of Alx was transformed to the pressure domain measure AlxC via the Pythagorean theorem (16).

\[ \text{AlxC (mmHg)} = \sqrt{\text{AP}^2 + \text{PP}^2} \]

Furthermore, the “companion” of PP was also calculated as follows.

\[ \text{PPC (mmHg)} = \sqrt{\text{SBP}^2 - \text{DBP}^2} \]

Figure 2 is the graphic representation of these aortic hemodynamics. In our pilot experiment (n = 5), the day-to-day reproducibility (i.e., coefficient of variation) of aortic SBP, PP, PPC, Alx, AlxC, baPWV, and haPWV was 1.2, 5.9, 1.3, 10.6, 7.2, 2.4, and 3.0%, respectively.

Data Analysis
Exclusion criteria were (1) pre-menopause (n = 4) (via the questionnaire), (2) suspected peripheral arterial disease [ABI < 0.9 (23)] (n = 2), (3) antihypertensive medication (n = 6), (4) inability to measure heart sound (n = 1), and (5) outlier of baPWV (n = 2) which was evaluated by Smirnov-Grubbs test. A total of 23 subjects (6 men and 17 post-menopausal women) including 6 untreated hypertensives were included in statistical analysis.

Statistic
Data are expressed as mean ± SD. Intrigoup comparisons of the data obtained before and after the intervention period were performed using a paired t-test. Analysis of covariance (ANCOVA) was used to examine the contribution of the correlated variables (changes in haPWV and AlxC) to change in aortic SBP. Pearson’s product-moment correlation was used to determine the relationship between variables of interest. Statistical significance was set a priori at P < 0.05 for all comparisons.

RESULTS
The subjects analyzed had mean age of 62 ± 9 years and a height of 159.0 ± 7.4 cm. The mean RPE for all 15 sessions recorded...
after each class was 12.5 ± 2.0 (low-intensity class: 12.2 ± 0.6, moderate-intensity class: 12.8 ± 1.5). The rate of participation in exercise training sessions was 83.5 ± 13.0% (low-intensity class: 83.3%, moderate-intensity class: 83.7%, range 7–15 sessions).

After the intervention, body weight significantly decreased (from 43 ± 9.4 kg, P = 0.015). Heart rate and ABI did not change significantly, whereas brachial SBP, MAP, PP, baPWV, and haPWV significantly decreased after the exercise intervention (Table 1). Figure 3 depicts changes in aortic hemodynamics.

Aortic SBP (from 119 ± 15 to 115 ± 14 mmHg, P = 0.005), PP (from 43 ± 9 to 41 ± 8 mmHg, P = 0.020), and PPC (from 141 ± 18 to 136 ± 17 mmHg, P = 0.007), decreased significantly after aquatic exercise training. AlxC decreased after exercise training intervention (from 47 ± 9 to 45 ± 8 mmHg, P = 0.029) although AP (from 19 ± 5 to 18 ± 5 mmHg, P = 0.210) and Alx (from 44 ± 10 to 43 ± 11 mmHg, P = 0.726) remained unchanged.

Change in aortic SBP was correlated with change in haPWV (r = 0.613, P = 0.002) but not with change in baPWV (r = 0.296, P = 0.170) (Figure 4). Besides, change in aortic SBP was correlated with change in AlxC (r = 0.632, P = 0.001) but not with change in Alx (r = 0.056, P = 0.800) (Figure 5). When ANCOVA was performed with either change in haPWV or AlxC as the covariate, the change in aortic SBP was no longer statistically significant (P = 0.371 and P = 0.653, respectively). Other correlation coefficients between variables of interest were summarized in Supplementary Table 1. Changes in aortic SBP, PP, PPC, and AlxC were not correlated with either participation rate or change in body weight. Change in aortic SBP was not correlated with the baseline AoSBP (r = −0.350, P = 0.102).

DISCUSSION

The main findings of this study are as follows. First, short-term, head-out aquatic exercise training improve aortic hemodynamics in middle-aged and elderly subjects. Second, changes in aortic SBP were significantly associated with haPWV but not with baPWV.

To the best of our knowledge, this is the first study to provide the favorable effects of head-out aquatic exercise on aortic hemodynamics, the emerging risk factors for future CV events and all-cause mortality (2). Reductions in aortic SBP and PP were only 4 mmHg and 3 mmHg. However, these changes might have pathophysiological implication because the 10 mmHg increases in aortic SBP and PP corresponded to the increasing risk for total CVD events by 8.8 and 13.7%, respectively (2). Also, it should be emphasized the low frequency (once per week) of our training regimen. Although one 60-min training session per week is typical as a city-operated aquatic exercise program, the frequency of the training is remarkably lower than those in previous studies that demonstrated the positive effect of aquatic exercise training on cardiovascular health (8–12, 14). Therefore, it is needed to examine whether the increasing frequency of the training reinforces the training effect on aortic hemodynamics.

We can only speculate the underlying mechanisms of decreased aortic SBP. The change in aortic SBP with aquatic exercise training intervention was not associated with the change in baPWV, but it was associated with the change in haPWV. Currently, baPWV has become a popular modality of arterial stiffness measurement in population-based studies because of the procedural advantage of being simple to use (i.e., only wrapping cuff-sensors around four extremities) (24). Previous studies have shown that baPWV is correlated with carotid-to-femoral PWV, the gold standard measure of aortic stiffness (25–27). baPWV may provide qualitatively similar information to those derived from central arterial stiffness, although some portions of baPWV may be determined by peripheral arterial stiffness (25). However, in the measurement of baPWV, brachial arterial pressure waveforms are used as substitutes for pressure waves in the proximal aortic site, which could not be recorded easily. Thus, the segment from the ascending aorta to the upper arm is omitted from the effective path length for baPWV, as the ascending aorta is not included in the path length for cPWV (27). On contrary, the arterial path of haPWV includes the ascending aorta due to the use of the second heart sound, indicating the timing of the aortic valve closure (i.e., dicrotic notch of the brachial pressure wave). Therefore, the significant correlation between changes in aortic SBP and haPWV but not baPWV suggests that the proximal aortic destiffening is associated with the reduction of aortic SBP. However, the causality cannot be clarified by this study design.

The proximal aorta is passively stretched by the ejection flow and stores about 50% of the blood during systole. In diastole, the stretched wall recoils back and releases its potential energy to deliver blood flow to the coronary and peripheral arteries (28). The aortic wall movement is called the Windkessel function. This function avoids the excessive rise in ejection wave and pulse pressure and maintains the coronary perfusion. Histologically, this hemodynamic function is made capable by the effect of elastin, an elastic protein that is a major component of the proximal aortic wall. However, aging causes the proximal aortic stiffening characterized by severe structural remodeling of the arterial wall. Previous studies have reported that unlike the improvement of peripheral arterial function contributing to vascular stiffness, structural modification of the aortic wall is
difficult to recover when exercise training was initiated in later life (29). However, in animal studies, repeated immersion has been reported to improve the aortic Windkessel and left ventricular functions (30). These authors described that water immersion leads to increased venous return due to elevations in hydrostatic pressure, cardiac output, and vascular wall stretch and shear stress. Another study also reported that elastin gene expression is enhanced by vascular wall stretching (31). In a human study, we found that lifelong Japanese female pearl divers who performed ∼100–150 breath-holding free dives for up to 2 min at a time had lower aortic SBP, AIx, and heart-to-brachial PWV, which partly reflects the proximal aortic stiffness, when compared to inactive women of the same age (32). Likewise, in the present study, short-term aquatic exercise training in middle-aged and elderly subjects
resulted in significant decreases in aortic PP and PPC, indicating the attenuated aortic pulsatility and the improvement of the proximal aortic Windkessel function. Collectively, increased venous return during water immersion may have a positive effect on the structure and function of the proximal aorta. On the other hand, Sherlock et al. reported that aortic SBP did not change significantly after a 10-week “shallow-water” aquatic exercise training (14). Although precise depth was not described, the lack of change in aortic SBP might be attributed to lower hydrostatic pressure which results in smaller increases in venous return and stroke volume compared with the present study.

Focusing on a single number as reflected by the ratio implies that information is lost. To overcome these simplifications, it is required to formulate alternative routes of data representation (17). Thus, in this study, we calculated PPC and AIxC by the polar coordinate description. Of note, AIxC decreased significantly after the intervention, although AIx did not change significantly. Parallel changes in aortic AP and PP might have masked the attenuated wave reflection with the aquatic exercise intervention as illustrated in Figure 1. Besides, change in aortic SBP was correlated with the corresponding change in AIxC but not AIx.

The present study argues that these changes may not be identified by the conventional ratio-based indices. In this context, the newly introduced companion metric may provide better insight into underlying physiology. However, careful attention should be paid to the interpretation of the results of this index, as it has not yet been validated under a variety of physiological conditions.

**Study Limitation**

Several methodological limitations should be mentioned. First, since aortic BP was estimated by a general transfer function from the brachial artery to the aorta, estimation error might not be equal among subjects. However, intra-individual change in estimates could be evaluated reliably. Second, arterial stiffness measurement using PWV may be prone to error due to inaccuracy of the arterial path length measurement. A sophisticated methodology such as MRI is required for evaluating the local proximal aortic stiffness response to the training intervention. Third, because of the nature of the field study, we could not control the characteristics of subjects (e.g., age, sex, and disease status). Furthermore, we did not have a sedentary control group, although we confirmed the excellent day-to-day reproducibility of key measurements. In addition, physiological
exercise intensity (i.e., heart rate, oxygen consumption) during exercise training was not evaluated. These drawbacks should be overcome by future studies.

CONCLUSION

In this study, we investigated whether head-out aquatic exercise training improves central aortic hemodynamics in middle-aged and elderly people. We found that head-out aquatic exercise training even at a low training frequency (once per week) may improve aortic hemodynamics. In particular, the reduction in aortic SBP was associated with the reduction in proximal aortic stiffness.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by National Institute of Advanced Industrial Science and Technology Research Board. The patients/participants provided their written informed consent to participate in this study.

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AUTHOR CONTRIBUTIONS

JS and MF decided on the conception, design of research, drafted the manuscript, and papered figures. MF, DH, TH, and JS performed the experiments, analyzed the data, and interpreted the results of the experiments. All authors edited and revised the manuscript and read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

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