Wearable Seismocardiography-Based Assessment of Stroke Volume in Congenital Heart Disease

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BACKGROUND: Patients with congenital heart disease (CHD) are at risk for the development of low cardiac output and other physiologic derangements, which could be detected early through continuous stroke volume (SV) measurement. Unfortunately, existing SV measurement methods are limited in the clinic because of their invasiveness (eg, thermodilution), location (eg, cardiac magnetic resonance imaging), or unreliability (eg, bioimpedance). Multimodal wearable sensing, leveraging the seismocardiogram, a sternal vibration signal associated with cardiomechanical activity, offers a means to monitoring SV conveniently, affordably, and continuously. However, it has not been evaluated in a population with significant anatomical and physiological differences (ie, children with CHD) or compared against a true gold standard (ie, cardiac magnetic resonance). Here, we present the feasibility of wearable estimation of SV in a diverse CHD population (N=45 patients).

METHODS AND RESULTS: We used our chest-worn wearable biosensor to measure baseline ECG and seismocardiogram signals from patients with CHD before and after their routine cardiovascular magnetic resonance imaging, and derived features from the measured signals, predominantly systolic time intervals, to estimate SV using ridge regression. Wearable signal features achieved acceptable SV estimation (28% error with respect to cardiovascular magnetic resonance imaging) in a held-out test set, per cardiac output measurement guidelines, with a root-mean-square error of 11.48 mL and $R^2$ of 0.76. Additionally, we observed that using a combination of electrical and cardiomechanical features surpassed the performance of either modality alone.

CONCLUSIONS: A convenient wearable biosensor that estimates SV enables remote monitoring of cardiac function and may potentially help identify decompensation in patients with CHD.

Key Words: cardiac output ■ machine learning ■ multimodal ■ noninvasive ■ pediatrics

Congenital heart disease (CHD) affects approximately 40,000 births per year in the United States alone, a quarter of whom suffer from critical cases that require surgery or other interventions in the first year of life. Even more disturbingly, only 69% of those presenting with these critical types of CHD reach adulthood.1,2 Although advancements in cardiac care and surgery have significantly improved the CHD survival rate from a few decades ago, these treatments are not curative by nature. Hence, the growing population of patients with CHD are at high risk of clinical deterioration, either sudden or gradual. Specifically, low cardiac output syndrome is the leading cause of post-CHD surgery death, whereas the development of heart failure is the leading cause of mortality among adult patients with CHD.3,4 Fortunately, routine assessment of key hemodynamic

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parameters, such as ejection fraction, stroke volume (SV), and cardiac output, which is the percentage and volume of blood pumped out by the heart per heartbeat and volume per minute, has been shown to inform prognosis and guide interventions, reducing overall mortality. Because these parameters assess the ability of the heart to pump blood effectively to meet oxygen demand, they are hallmark indicators of left ventricular dysfunction when depressed and a strong predictor of major adverse cardiac events. Thus, the development of technologies allowing continuous, noninvasive, and inexpensive measurement of SV and cardiac output represents a critical need in CHD; such technologies could be used in both inpatient and outpatient settings to improve outcomes.

Existing SV and cardiac output measurement methods are suboptimal, especially in children and those with CHD. Thermodynamics-based pulmonary artery catheterization is accurate, but is not commonly used in children because of the large size of catheters and physiological differences, such as those present in single-ventricle patients, affect the waveform morphology. Furthermore, the ability of baseline seismocardiogram features to accurately assess diagnostic differences in absolute SV across different people has not been examined. Finally, seismocardiogram-based SV estimation has only been studied in comparison with the transesophageal Doppler echocardiogram and therefore never with respect to an unequivocal gold-standard measurement such as CMR.

We sought to evaluate the use of seismocardiography to measure SV in a unique population of patients with CHD using our convenient wearable biosensor. In this exploratory work, simple, intuitive, physiologically inspired ECG and seismocardiogram features derived from this wearable biosensor were used, along with machine learning, to estimate the baseline SV of patients with CHD undergoing clinically indicated CMR. Inaccuracies in patients with shunts. Transesophageal Doppler echocardiography is less invasive but cannot often be used continuously, angle dependent and therefore less accurate, bulky, and requires a trained professional. Cardiovascular magnetic resonance imaging (CMR), a noninvasive technique widely considered as the gold standard in children and those with CHD because of high accuracy and excellent reproducibility, is not feasible for continuous SV monitoring because of the inability to be performed at bedside or in the outpatient setting. Therefore, noninvasive continuous cardiac output monitoring (NICCOM) technologies have been developed that estimate SV from models using demographic information combined with either the impedance cardiogram or the finger arterial pressure waveform, obtained through bioimpedance and the vascular unloading technique, respectively. However, these approaches are obtrusive, require strict placement of multiple electrodes or cuff sizes, have low accuracy in critically ill patients, and are rarely tested in children, so their practicality remains in doubt when used for monitoring SV in patients with CHD.

Seismocardiography is a promising method for NICCOM that uses a low-noise accelerometer placed on the chest to capture the seismocardiogram, which provides cardiomechanical information unobtainable by other NICCOM methods. When combined with the ECG, the seismocardiogram allows for the calculation of systolic time intervals, such as the pre-ejection period (PEP) and ventricular ejection time (VET). In recent years, groups have demonstrated that ECG and seismocardiogram signals acquired from wearable devices can accurately estimate SV, heart failure clinical status, and underlying events in the cardiac cycle using echocardiography and CMR in people with structurally normal hearts. However, seismocardiogram signals have not been evaluated in patients with CHD, resulting in a lack of understanding in how major anatomical and physiological differences, such as those present in single-ventricle patients, affect the waveform morphology. Furthermore, the ability of baseline seismocardiogram features to accurately assess diagnostic differences in absolute SV across different people has not been examined. Finally, seismocardiogram-based SV estimation has only been studied in comparison with the transesophageal Doppler echocardiogram and therefore never with respect to an unequivocal gold-standard measurement such as CMR.
addition, this preliminary work provides greater insight into how cardiomechanical signals, such as the seis-
mocardiogram, are modulated in patients with CHD
with severe anatomical and physiological differences. Ultimately, it will yield a framework, along with the
pertinent features necessary, for estimating SV using
wearable ECG and seismocardiogram signals toward
noninvasive, continuous, and ubiquitous monitoring of
this vital hemodynamic parameter.

METHODS
The data presented in this study are available from the
corresponding author upon reasonable request.

Study Protocol
Detailed overviews of the study design and patient de-
mographics are provided in Figure 1 and Table 1, re-
spectively. This study was conducted under institutional
review board protocol STU2019-1280 at the University
of Texas Southwestern Medical Center. We approached
consecutive patients with CHD undergoing clinically in-
dicated CMR and obtained written consent and assent
as appropriate. In the preoperative area, chest-worn
biosensor data were collected in a supine position for
3 minutes, unless unable to do so because of COVID-
19–related anesthesia restrictions. If the patient was
undergoing the scan under anesthesia, the device was
placed 10 minutes after induction of anesthesia to allow
for them to reach physiological equilibrium. During
the clinical CMR scan, left ventricular and right ventric-
ular SV, and if recommended, aortic valve forward flow,
were collected per clinical protocols. After the CMR
scan, we obtained another 3 minutes of chest-worn bi-
osensor data, although because of space restrictions,
most of these data were obtained in a sitting position.
To maintain consistency with body positioning, only ei-
ther pre-CMR or post-CMR supine data were examined
in this work, with most data taken from the pre-CMR
measurement, unless unavailable. Therefore, the simi-
larity between pre-CMR and post-CMR measurements
was not analyzed. Furthermore, matching the supine
posture also used during CMR allowed us to account
for the known impacts that changes in venous return
because of posture can have on both SV and wearable
biosignals. The aortic forward flow measurement
was prioritized over the volumetric one, if acquired.

Figure 1. Concept overview.
Study design showing wearable biosensor placement when supine and asynchronous reference cardiovascular magnetic resonance
imaging (CMR) measurement. Seismocardiogram (SCG) mechanistic overview detailing modulation due to cardiac physiology,
acquisition with an accelerometer, and sensing axes for ECG (negative, positive, and right-leg-drive [RLD] electrodes), and triaxial
SCG signals. Analysis pipeline, from sensor input to model estimation of stroke volume, for wearable (blue), demographic (green), and
CMR (purple) data. H indicates the transfer function between the input internal sources of cardiomechanical vibration and the output
SCG waveform measured on the surface of the torso; MRI, magnetic resonance imaging; and SCG_dv, dorso-ventral SCG.
variables, where applicable, was computed using Fisher exact test.

**Table 1. Overview of Patient Demographics and Cardiac Function Clinical Parameters for Study Participants**

| Demographics and clinical parameters | Training set, n=36 | Held-out test set, n=9 | P value |
|--------------------------------------|--------------------|-----------------------|---------|
| Sex, n (%)                           |                    |                       | 1.00*   |
| Men                                  | 22 (61)            | 6 (67)                |         |
| Women                                | 14 (39)            | 3 (33)                |         |
| Height, cm, mean (SD)                | 151.3 (26.6)       | 163.1 (16.0)          | 0.21†   |
| Weight, kg, mean (SD)                | 56.3 (26.0)        | 59.2 (19.7)           | 0.76†   |
| Body surface area, m², mean (SD)     | 1.52 (0.48)        | 1.63 (0.34)           | 0.52†   |
| Age, y, mean (SD)                    | 15.0 (7.9)         | 14.7 (2.6)            | 0.90†   |
| Stroke volume, mL, mean (SD)         | 68.8 (32.34)       | 78.19 (24.81)         | 0.42†   |
| Cardiac output, L/min, mean (SD)     | 4.94 (1.88)        | 5.01 (0.84)           | 0.92†   |
| Ejection fraction, %, mean (SD)      | 0.57 (0.08)        | 0.59 (0.12)           | 0.51†   |
| Reference heart rate, bpm, mean (SD) | 76.1 (14.2)        | 67.7 (14.8)           | 0.12†   |
| Single ventricle, n (%)              | 5 (14)             | 3 (33)                | 0.33*   |
| Systemic ventricle, n (%)            |                    |                       | 0.65*   |
| Right ventricle                      | 6 (17)             | 2 (22)                |         |
| Left ventricle                       | 30 (83)            | 7 (78)                |         |

*Statistical significance between training and testing sets in categorical variables, where applicable, was computed using Fisher exact test.
†Statistical significance between training and testing sets in values, where applicable, was computed using an unpaired t test.
‡Values for systemic ventricle data are shown.

The systemic ventricle, which is connected to the aorta and responsible for cardiac output, was labeled by the cardiologist, and SV data from this ventricle along with the instantaneous heart rate (HR) taken from the corresponding CMR measurement, for either the volumetric or aortic forward flow measurement, were used as the reference SV and HR measurement, respectively. The systemic ventricle measurements were used for the target SV based on the assumption that features from the wearable signals would correspond more closely with the systemic ventricle responsible for ejecting the systemic SV. Additionally, patients were not separated by ventricular morphology to train ventricle-specific SV estimation models because of an already small sample size for ridge regression models and literature that suggests that SV estimates are not based on differences in systemic ventricles.18

**Multimodal Hardware Design**

The electronic hardware used in this study, shown in Figure 2, is an updated and miniaturized version to that previously described in detail.19 Updates focused on decreasing the overall device size to accommodate a pediatric population. From a sensing standpoint, identical sensors and analog front ends were used in this study to acquire the ECG (ADS1291; Texas Instruments, Dallas, TX) and seismocardiogram (ADXL355; Analog Devices, Norwood, MA) signals across all versions of the hardware, and the same 3-dimensional printing filament, polyactic acid, was used to manufacture the device housing. Data were saved locally on an internal secure digital card and downloaded over universal serial bus using the custom software application previously mentioned.19 Further descriptions of the hardware used are provided in Data S1.

**Signal Processing**

A signal processing block diagram is depicted in Figure S1. Preprocessing consisted of bandpass filtering the ECG and the dorso-ventral axis of the seismocardiogram signal between 5 and 30 Hz, and 0.8 and 30 Hz to remove their respective out-of-band noise. The seismocardiogram was also filtered into a higher frequency bandwidth 30 to 125 Hz to produce a signal hereafter referred to as high-frequency seismocardiogram, which is more closely representative of the phonocardiogram. In turn, this wider bandwidth accelerometer signal is capable of picking up on higher frequency vibrations, coupled to the acoustics of the phonocardiogram, thus offering a more reliable timeframe to estimate aortic valve opening and closing events,20 a common technical challenge in seismocardiogram processing.19 After filtering, all signals were resampled to 1 kHz. The R peaks of the ECG were used to segment the seismocardiogram and high-frequency seismocardiogram signals into different heartbeats. The seismocardiogram and high-frequency seismocardiogram heartbeats were ensemble averaged using 30 heartbeat windows with 50% overlap to reduce 0 mean noise, account for respiratory-induced variability in seismocardiogram signals, and improve the consistency of amplitude features before selecting the highest signal-to-noise ratio beat later used to extract the features shown in Table 2. A detailed description of the calculation and meaning of the features used is provided in Data S1.

**Machine-Learning Regression Analysis**

Ridge regression was used to estimate SV because of its ability to handle multicollinearity, a trait common among systolic time intervals, as well as provide feature importance with reduced complexity simply from its weights as a linear model. To avoid data leakage, an 80% to 20% fixed training–testing scheme (ie, 36 patients for training, 9 for testing) was determined using a true random number generator (RANDOM.ORG, Dublin, Ireland). A 10-fold cross-validation on the training set was used to perform the grid search necessary for hyperparameter optimization. Forward feature selection, on the training set, was used to reduce the feature set from 14 down to 9 features by examining the coefficient
of determination, through simple linear regression between ECG and seismocardiogram features and SV. Different ridge regression models, trained on a combination of feature sets, each with their respective optimized $\lambda$ hyperparameter from the 10-fold cross-validation on the training set, were used to estimate SV. Specifically, we began by examining a model that was merely trained on demographic features alone (ie, body surface area and age) because of their well-known correlation to cardiac output, which assessed the ability to quantify SV without the use of a wearable biosensor. Next, we tested using ECG features, namely HR, to assess the estimation accuracy when using only a conventional metric that is readily, remotely, and continuously available through Holter monitors. Then, we tested our novel approach by adding seismocardiogram features to the ECG model to provide for a holistic evaluation of both the electrical and mechanical aspects of cardiac health. Finally, we have provided various combinations of these wearable biosensor and demographic feature sets to determine whether the easily accessible demographic information can augment model estimation.

Statistical Analysis
To assess these model performances, we computed the root-mean-square error (RMSE) and the coefficient of determination ($R^2$) between the estimated SV and true CMR SV in the held-out test set that was unseen to the machine-learning algorithm until final testing. Percent error was calculated for the highest performing
model given the guidelines for cardiac output measurement devices as the limits of agreement (ie, 1.96 times the SD of the bias) divided by the mean SV from the CMR and wearable × 100, with a percent error <30% regarded as acceptable.22 Importance of the features was derived from the magnitude of the weights from ridge regression, and importance of the permutations, iterated 1000 times, were also computed and are provided in Figure S2. Specifics on model selection, train-test splitting, cross-validation, and feature selection are provided in Data S1.

RESULTS

We enrolled 57 patients and successfully acquired both supine wearable and CMR data from 45 patients. Twelve patients were excluded because they either did not contain accompanying CMR, a supine wearable measurement because of COVID-19 anesthesia area restrictions, usable wearable data because of system malfunction, or withdrew from the study. Their detailed patient demographics are presented in Table 1.

The regression model performance, the coefficient of determination and RMSE, for the training (ie, 10-fold CV) and testing set for all feature sets is shown in Table 3.

The test set performance, when combining features from both the ECG and seismocardiogram modalities, improved upon that of the ECG model alone (R², 0.76 and RMSE, 11.48 mL versus R², 0.69 and RMSE, 13.05 mL) and substantially upon the demographic one (R², −0.10 and RMSE, 24.56 mL). However, the ECG-only model still outperformed the seismocardiogram only model (R², 0.20 and RMSE, 20.51 mL). To put the significance of these low RMSEs into context, the dynamic range of measured SV in the training and test set were 146.5 and 74.8 mL, respectively. The regression and Bland-Altman plot for the highest performing model, combining ECG and seismocardiogram features to estimate SV, are shown in Figure 3. The 95% CIs for the highest performing model train and test sets are −44.63 to 44.63 mL and −23.82 to 20.8 mL, respectively. The percent error, the metric that is used by cardiac output measurement guidelines, for this highest performing model was 28%, within the acceptable criteria of 30%.22 By contrast, the percent error for the ECG-only model was outside the threshold for acceptability at 31%.

The demographic feature model has a stark difference between training (R², 0.72) and testing (R², −0.10) performance, a clear sign of overfitting. Therefore, combining the demographic information with ECG and seismocardiogram features improved their training set performance (R², 0.88), but not the test set performance (R², 0.27). However, as presented in Table 1, there was no significant difference between the demographics of the training and testing set.

The importance of the features based on the magnitude of the weights from the ridge regression model is shown in Figure 4. Other than HR, the most important features are either the PEP, VET, or a ratio derived from the combination of them. The importance of the permutations, given in Figure S2, was comparable to the magnitude of the weights from ridge regression, but with VET and PEP having opposite ordering in importance. In addition, the Pearson correlation coefficients between the physiological features and SV, provided in Table S1, demonstrate that HR, aortic valve closure, and VET have the greatest correlation to SV while also giving indication of their relationship to SV.

DISCUSSION

In this preliminary study, we show that a combination of seismocardiogram and ECG parameters, obtained noninvasively, can be acceptable estimators of SV in patients with CHD undergoing CMR per cardiac output measurement devices.
measurement guidelines. This exploratory work represents a necessary advancement toward both more holistic wearable SV estimation for diagnostics and remote monitoring for patients with CHD. Furthermore, our study is novel in using seismocardiography in children and those with CHD; other studies have focused on structurally normal hearts. Specifically, although it has been shown previously that noninvasive ECG and seismocardiogram measurements contain information that can be used to estimate SV, this relationship had not been examined in a diverse population of patients with congenital heart defects and compared against a true gold-standard measurement. Here, we evaluated the future usefulness of a convenient method for estimating SV and observed that there was a strong correlation between simple, highly interpretable seismocardiogram features and SV, across a wide range of CHD diagnoses, ages, and anatomical differences. We further achieved acceptable estimation of SV in a completely held-out test set by using a regression model that combined both electrical and cardiomechanical wearable features, producing performance superior to that of either feature subset alone.

**Wearable Multimodal Signal Features Are the First Acceptable Estimator of Baseline SV in a Completely Held-Out Test Set of Patients With CHD**

Wearable features were strongly correlated to baseline gold-standard CMR SV in a heterogenous population of patients with CHD. Furthermore, with an overall 28% error, we were able to achieve an acceptable estimation of SV, based on the limits set forth by cardiac output measurement guidelines, in a completely randomized held-out test set. Meanwhile, the training set, for which the model was an unacceptable estimator of SV based on the guidelines, had a wide dynamic range of nearly double that of the test set of 146.5 mL, which may explain the lower performance therein and the discrepancy observed in this random set. Overall, this SV estimation model trained on multimodal wearable signal features and tested on unseen data is of greater use than a purely correlation-based result. Additionally, for the Bland-Altman analysis, all estimations were within the limits of agreement indicative of high model precision. Meanwhile, existing inconvenient methods, although more comprehensively evaluated, such as transesophageal Doppler and NICCOM methods,
have estimation errors of >40%. In the field of physiological research and biomedical signal processing, to be able to use baseline measurements to estimate absolute values in clinical parameters across a subject population is extremely difficult and rarely performed. Typically, a perturbation or intervention is leveraged to modulate physiological properties, in this case hemodynamics, which allows for a greater dynamic range and ability to track subject-specific changes in waveform morphology, usually resulting in higher accuracy. In this study, the comprehensive age range and diagnoses, representative of higher-risk children and adults with CHD who would be undergoing CMR, not only adds difficulty in hardware design, but also contributes to high intersubject variability. Eventually, if the device were to be used to obtain continuous, noninvasive measurements, perhaps in the case of monitoring patients with CHD after surgery for low cardiac output syndrome, then tracking subject-specific changes in seismocardiogram signals would suffice to monitor status, predict exacerbation, and offer personalized health care without a specific gold-standard baseline assessment. Generally, subject-specific models will outperform globalized models, although the reduced complexity of the latter may prove to be beneficial in certain scenarios.

Our low RMSE for the test set of 11.48 mL, underscored by the wide dynamic range in SV of 74.8 mL, within, demonstrates that our estimation is relatively robust to outliers. Hence, the inability to explain these outliers with a demographics-only model, which performed worse than our overall model, suggests that the acceptable estimation was not driven primarily by patient size (eg, body surface area). From the residual errors in the regression plot shown in Figure 3, our model also had comparable estimation of high and low SVs: a well-known limitation of machine-learning approaches that cleverly estimate the mean to reduce error. However, there was a slightly better performance at lower SV, most likely because of a larger number of data points with similar target values. Additionally, in our training set, there were fewer data points near the highest SVs as well, which lead to the greatest error when estimating them and potentially explains the difference between the training and testing performance. Therefore, increasing the number of data points with an emphasis on those exhibiting the boundaries of SV should improve estimation performance and overall model robustness.

Nevertheless, using wearable signal features as an acceptable estimator of SV across a heterogeneous population of patients with CHD, suggests that eventually seismocardiogram signals can assist in overall diagnostics, which was previously not demonstrated in seismocardiography literature.

Cardiomechanical Seismocardiogram Features Improve Model Estimation

As shown in Table 2, adding the cardiomechanical seismocardiogram features to the purely electrical ECG model resulted in a modest improvement in performance that reduced the percent error from 31% to 28%, which was sufficient to achieve acceptable SV estimation per cardiac output measurement guidelines. Although the heart is electrically activated, it remains a mechanical pump, and therefore assessing these other aspects of cardiac health, although traditionally ignored by NICCOM methods, are essential to quantifying its mechanical function. Specifically, although HR is well known to exhibit a strong correlation to SV, the other most important features essential to achieving a good estimation were the PEP, VET, and PEP/VET. The PEP, a combination of the intrinsic electromechanical delay and isovolumic contraction time, was our second most important feature. This is in accordance with established knowledge that the PEP can vary based on age, between infancy and puberty, and differences in contractility and preload, both captured in our data set. Similarly, the VET, the time it takes to eject the SV of blood out the aorta, is related to SV. In addition, several NICCOM technologies use the impedance cardiogram to estimate SV through mathematical formulas, which are grounded in well-understood relationships between bioimpedance and cardiac output, and leverage the VET as a strong correlate to SV. However, bioimpedance inconveniently requires multiple electrodes to be placed on the body, whereas seismocardiography can capture the same VET in a significantly more convenient manner simply through an accelerometer placed on the chest. Additionally, their ratio (ie, PEP/VET) has been demonstrated to be inversely related to contractility and helpful in determining heart failure. This relationship is understood to be because of the greater amount of time required for the failing heart to build up the pressure necessary for ejection, related to PEP, and the smaller SV ejected during a shorter VET.

Though the ECG features had a more significant independent contribution in estimating baseline SV, the seismocardiogram signal has been shown to better capture longitudinal changes in ventricular function by assessing the mechanical aspects of cardiac health. Nonetheless, given the complex determinants of SV, it is not surprising that combining features from multiple sensing modalities was necessary to create the holistic model that had the greatest performance.

Demographic Based Correlations to SV Do Not Necessarily Generalize

Demographic feature models (ie, those using age and body surface area) did not generalize well to our
SV Estimation Is Robust Against Anatomical Differences

Largely, our data suggest that acceptable SV estimation can be achieved regardless of the unique anatomies and physiologies in patients with CHD. Although there are considerable anatomical modifications between single-ventricle and 2-ventricle patients, it has previously been shown that there are no distinguishable differences in their SV estimation before and after the hemi-Fontan operation when using magnetic resonance imaging. In addition, because of their wide-ranging diagnoses and demographics, the intersubject variability in the single-ventricle patients may have overshadowed their population variability with respect to the 2-ventricle patients. In future studies, to fully determine whether any anatomically induced modulation in seismocardiogram morphology exists, data should be taken from a larger and more homogenous single-ventricle population, for instance, reducing the age gap to only neonates with single and 2 ventricles.

Eventually, remote monitoring of the growing population of older children and adults with heart defects appears to be more feasible because of their uncoupled characteristics with respect to this novel sensing modality and key hemodynamic parameters of ventricular function such as SV.

Future Work

In this novel, exploratory work relating wearable signals to SV in patients with CHD, we present technology that can be refined to improve accuracy through multiple means from developments in the wearable device, data collection, and algorithms used herein. Specifically, plethysmogram signals were not used in this work; however, features from the plethysmogram have been shown to be correlated to SV and offer a measure of peripheral hemodynamics that is uncaptured by the seismocardiogram and ECG. This alternative modality when fused with the seismocardiogram and ECG can likely help improve performance by creating a more holistic model. Next, when using machine-learning methods, increasing data size can have significant influence on determining true model accuracy and generalizability. Therefore, further studies should aim to collect data on more patients with similar diagnoses for longer periods of time, perhaps with a continuous SV reference measurement, and analyze differences between either single- and 2-ventricle patients or those with a systemic right versus systemic left ventricle, separately. Using longitudinal data to train subject-specific models will likely result in increased accuracy when compared with non-subject-specific models. Finally, with an increase in data from both another modality measured by the device and sample size, more complex machine-learning algorithms can be used to better estimate SV.

STUDY LIMITATIONS

This study has limitations related to both the technologies used and the patient population studied. The wearable biosensor is not magnetic resonance imaging safe, and thus, simultaneous measurement of CMR flows and volumetrics with ECG and seismocardiogram was not possible. Thus, to mitigate this limitation, we obtained the wearable measurements as soon as possible before and after the CMR scan and after the patient was under anesthesia for those getting the CMR with anesthesia. However, it is possible that the patients were in different physiologic states during the CMR compared with when the wearable measurements were taken. Furthermore, the small population studied herein, especially considering the application of machine learning to estimate SV, requires the model to be further validated in patients with CHD to ensure generalizability. To mitigate this in this first study, we used a small subset of highly interpretable features along with a ridge regression model that offers a regularization penalty to reduce model overfitting and used a fixed train-test split. Regardless, this study represents exploratory work, because the accuracy of machine-learning models with small sample sizes cannot be truly determined until more data are collected in multiple settings. Overall seismocardiogram and ECG metrics from a chest-worn wearable biosensor correlate well to SV, but further longitudinal studies in larger
and more diverse populations and multiple settings are needed for seismocardiography to realize its potential as a continuous, noninvasive tracker of SV for those with congenital and functional heart disease.

CONCLUSIONS

We demonstrated that a multimodal wearable biosensor that measures both seismocardiogram and ECG signals can serve as an acceptable estimator of SV in patients with CHD based on the cardiac output measurement guidelines. In the future, this exploratory work could be expanded to monitor patients conveniently and longitudinally either after surgery or from the comfort of their homes. Noninvasive, continuous monitoring of SV using a wearable biosensor equips clinicians with the tools necessary to track their patients longitudinally, not currently captured by any clinical program and seldom studied, which is essential to comprehend the lifetime complications facing this growing population. Eventually, advanced machine-learning algorithms may even be capable of predicting the periodic decompensations of patients with CHD. In addition, it is well known that there are racial, socioeconomic, and geographic factors that contribute to disturbing health disparities in CHD mortality. Ultimately, following further studies in a larger population, an inexpensive ECG and seismocardiogram wearable biosensor may provide accurate low-user-input SV monitoring in a noninvasive, continuous, and affordable manner for patients in out-of-office settings in low-resource settings.

ARTICLE INFORMATION

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Supplemental Material
Appendix S1
Table S1
Figures S1–S2
References [32–36]

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Supplemental Material
Data S1

Supplemental Methods

Multimodal hardware design

Overall, the key upgrades from the hardware used in\textsuperscript{19} include the addition of a flexible connector, two main sensing printed circuit boards as opposed to three, eventually the use of gel-electrode ECG, a separate photoplethysmogram sensor board with newer discretized photodiodes and light-emitting-diodes, and a foam-based spring backing mechanism for improved photoplethysmogram sensing. However, the photoplethysmogram signals were not explored in this work and therefore—to prevent detracting from the focus of this work—the specific details of that hardware will not be expanded on further. The sample rate of the ECG was 1 kHz and the SCG either 500 Hz or 2 kHz depending on the prototype version. Specifically, in the newer version, shown in Figure 2a, the sample rate of the SCG was increased to 2 kHz to provide a bandwidth of 500 Hz; the SCG sampling frequency was adjusted to capture higher frequency sounds that may eventually be utilized to monitor patients with heart murmurs—a subsection of the CHD population at a greater risk of decline. Unfortunately, as with a proof-of-concept study, the hardware required few—mostly device housing—modifications at different stages of the study before reaching the current prototype pictured in Figure 2a. Most importantly, the earlier version of the hardware utilized in this study featured the use of a dry electrode ECG, using stainless steel tape, for which the device was pressed against the chest of the patient to acquire the biosignals, while the later version used standard infant AgCl gel electrodes (Kendall HP69, Medtronic PLC, Dublin, Ireland) to adhere to the chest, eliminating the need for an extra contact force. To help mitigate any issues from differences in contact pressure with the dry electrode
version, in addition to having the same group of few clinicians collect all data, only segments of the signals where the dry electrode acquired ECG—which is susceptible to variations in contact pressure due to changes in skin-electrode-impedance—had a consistent amplitude were analyzed. Devices with both versions of the ECG featured a firmware modification which leveraged the lead-on detect feature of the ECG chip and would toggle a light-emitting diode facing the clinician between red and green for when ECG lead was detected as off or on, respectively. This also removed the possibility of accidentally applying an excessive amount of pressure without knowing whether a signal was being acquired.

In future work, though high-fidelity wearable measurements were acquired and only few minutes of data collection were necessary for this study, the wearable biosensor still needs further miniaturization to be used in future longitudinal studies in a pediatric population. However, given the considerably smaller footprint of the internal essential sensing elements, the hardware could readily be miniaturized and exploit the advent of flexible electronics which can offer a low-profile, less obtrusive solution for even greater convenience when performing longitudinal monitoring.

Signal processing and feature extraction

All signal processing and feature extraction was carried out in MATLAB 2018a (MathWorks, Natick, Massachusetts, USA) and entirely automated. A high-frequency SCG signal more closely related to the phonocardiogram was extracted for this analysis. The phonocardiogram, typically acquired from digital stethoscopes, is a wide bandwidth, high-frequency acoustic signal that captures heart sounds (i.e., $S_1$ and $S_2$) and obtains information of valve closures when placed at specific auscultation sites. Although, the phonocardiogram should be acquired using a wide-bandwidth, piezoelectric accelerometer (i.e., a contact microphone)
rather than the capacitive, direct current micro-electro-mechanical systems accelerometer used herein, the sampling rate of the accelerometer was increased to provide this bandwidth. First, the R-peaks of the ECG—marking ventricular depolarization—were found using Pan-Tompkins’s algorithm and used to determine the wearable HR. Then the SCG and high-frequency SCG signals were segmented into different heartbeats using and beginning with the detected R-peaks of the ECG. Due to the large differences in HR in this dataset, all of the heartbeats were zero-padded to a fixed length of 1300 ms, based on the slowest HR in the dataset. Next, the SCG and high-frequency SCG heartbeats were ensemble averaged using 30 heartbeat windows with 50% overlap—to reduce zero-mean noise, remove respiratory induced variability, and improve the consistency of amplitude features—before selecting the highest SNR beat—calculated using the algorithm in\textsuperscript{34}. First the envelope of the high-frequency SCG was computed which provided the profiles for the conventional heart sounds S\textsubscript{1} and S\textsubscript{2}. The algorithm for detecting the aortic opening point on the max SNR SCG beat was the same as that used in\textsuperscript{20}, where the aortic opening was detected by finding the nearest zero-crossing after the peak of the high-frequency SCG envelope between 0 and 150 ms; the aortic closing point was determined by finding the most consistent peak of the high-frequency SCG itself between 250 ms to the end of the beat. The aortic opening point resembles the PEP with the difference between that and the aortic closing point being the VET. Two other reciprocal features, PEP/VET and VET/PEP—systolic marker robust to differences in HR—are the quotient of the PEP and VET. Two interpretable systolic amplitude features were calculated as the RMS amplitude of the SCG during the PEP and during the VET. In total 9 systolic features were extracted from the wearable signals. Note that the HR from the CMR was added as a feature, due to both the inability to acquire continuous measurements with the wearable patch during the CMR—because of magnetic interference and injury— and due to expected high
accuracy in HR estimation when using wearable ECG during baseline measurements, as a closer measure of the HR during the reference measurement.

Leveraging surrounding physiological information can contextualize and improve the estimation accuracy of wearable measurements. The SV measurement from the CMR, is computed from a composition of several images which are obtained a relatively slow sampling rate. Therefore, due to respiratory induced variability in SV readings—stemming from changes in venous return, preload, and HR—clinicians typically ask patients to hold their breath. However, as imaginable, for younger children this is obviously not possible. Instead, multiple scans are taken, are the resulting images are averaged before computing SV from the averaged image. Similarly, when using wearable measurements to accurately estimate SV compared to CMR readings should also factor in respiratory variability by averaging over a larger timespan—such as the 30 heartbeats employed in this analysis.

**Machine learning**

All machine learning and cross validation was performed in Python 3.0 using scikit-learn ridge regression and grid search packages, respectively. Despite a considerable sample size with respect to other SCG literature—especially given the diversity of demographics and diagnoses in such a diseased population—due to a small number of overall datapoints for a machine learning problem, multi-variate ridge regression was chosen as a less complex linear, and more interpretable, model to estimate SV. Ridge regression is similar to multiple-linear regression but with a regularization penalty—commonly referred to as lambda—that penalizes the model to prevent overfitting to the training data, thereby hopefully improving model generalizability.

10-fold cross validation was chosen as a commonly regarded robust method for optimizing hyperparameters and given that each subject had only one datapoint there would be
no overlap of subject-specific data in each fold. The completely randomized, held-out test set was determined by utilizing a true random number generator (RANDOM.ORG, Dublin, Ireland). To approximately balance and have a representative number of the number of single-ventricle patients in the training and testing set based on their size, originally during the data collection we again randomly split them into groups of four and three, respectively. However, after the final data was collected, a last single-ventricle patient was added to the training set to achieve a perfect 80%-20% split, hence a slight imbalance.

When selecting features for biomedical machine learning problem with a small dataset size there is a greater importance placed on not only selecting a few features that can explain a lot of the variance but also ones that can be clinically understood. Therefore, the original feature set of predictor variables consisted of 14 features that were chosen based on those with strong overlap between commonly used SCG features in existing literature and those that are simple and intuitive to cardiologists. Using forward feature selection on the training set, we decreased the feature set from 14 down to nine features based on the simple linear regression coefficient of determination between ECG and SCG features and SV. The Pearson’s correlation coefficient values between these final nine features and SV, for the training set, are shown in Table S1.

We compared different ridge regression models trained on unique feature sets in our work. Specifically, we sought to compare models leveraging different combinations of ECG, SCG, and demographic (i.e., age and body surface area) features. Both training and testing set features were normalized based on the training set mean and standard deviation. The hyperparameter lambda for the highest performing model, which combined ECG and SCG features, came out to the maximum regularization penalty of 1.0.
Table S1. Training Set Correlation Coefficients Between Physiological Features and Stroke Volume.

| Physiological Feature | Training Set Pearson’s r |
|-----------------------|--------------------------|
| HR<sub>CMR</sub>      | -0.65                    |
| HR<sub>Wearable</sub> | -0.63                    |
| VET                   | 0.39                     |
| PEP                   | -0.02                    |
| PEP/VET               | 0.01                     |
| VET/PEP               | -0.06                    |
| AC                    | 0.45                     |
| RMS<sub>PEP</sub>     | -0.20                    |
| RMS<sub>VET</sub>     | -0.21                    |

HR indicates heart rate; CMR, cardiac magnetic resonance imaging; VET, ventricular ejection time; PEP, pre-ejection period; AC, timing of aortic valve closure; RMS, root-mean-square power.
Figure S1. Signal processing pipeline. Block diagram of signal processing overview showing interpolation of electrocardiogram (ECG) and seismocardiogram (SCG) signals acquired from the wearable before bandpass filtering, R-peak detection, heartbeat windowing, and signal
quality assessment using the signal-to-noise ratio (SNR). Illustration of the custom high-frequency SCG (HF-SCG)—indicative of valve closures—assisted feature selection algorithm, helping to locate key fiducial points such as the aortic valve opening (AO) and aortic valve closure (AC) on the SCG—used to compute the pre-ejection period (PEP), ventricular ejection time (VET), and the AC. Additionally, the search radius for the AO (green) and AC (red) algorithm as well as their candidate points are shown.

Figure S2. Permutation feature importances for stroke volume (SV) estimation model.

Permutation feature importances for wearable system with features randomly shuffled 1000 times, ranked in order from top to bottom, and color-coded by wearable sensing modality—electrocardiogram (ECG) and seismocardiogram (SCG) signals.