Single-Beat Estimation of Right Ventricular Contractility and Its Coupling to Pulmonary Arterial Load in Patients With Pulmonary Hypertension

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Background—An accurate assessment of intrinsic right ventricular (RV) contractility and its relation to pulmonary arterial load is essential for the management of pulmonary hypertension. The pressure-volume relationship with load manipulation is the gold standard assessment used for this purpose, but its clinical application has been hindered by the lack of a single-beat method that is valid for the human RV. In the present study, we sought to validate a novel single-beat method to estimate the preload recruitable stroke work (PRSW) and its derivative for ventriculoarterial coupling in the human RV.

Methods and Results—A novel single-beat slope of the PRSW relationship (Msw) was derived by calculating the mean ejection pressure when the end-systolic volume was equal to volume-axis intercept of the PRSW relationship. In addition, by using a mathematical transformation of the equation representing the linearity of the PRSW relationship, a novel index for ventriculoarterial coupling, Msw/mean ejection pressure, was developed. RV pressure-volume relationships were measured in 31 patients (including 23 patients with pulmonary hypertension) who were referred for right-sided heart catheterization. In this cohort, the single-beat Msw was strongly correlated with the multiple-beat Msw (r=0.91, P<0.0001). Moreover, a significant correlation was observed between the single- and multiple-beat Msw/mean ejection pressure (r=0.53, P=0.002), with a stronger correlation in those with greater RV systolic pressure (r=0.70, P=0.003).

Conclusions—The novel single-beat approach provided an accurate estimation of indexes for the PRSW relationship and ventriculoarterial coupling. It may be particularly useful in assessing RV adaptation to increased pressure overload. (J Am Heart Assoc. 2018;7:e007929. DOI: 10.1161/JAHA.117.007929.)

Key Words: contractility • heart failure • pressure-volume relationship • pulmonary circulation • pulmonary hypertension • pulmonary impedance

Pulmonary arterial (PA) hypertension (PAH) can lead to pathologic remodeling of the pulmonary vasculature and progressive increases in the PA load. Although the right ventricle (RV) attempts to adapt by compensatory hypertrophy and dilation, adaptation can become insufficient, leading to right-sided heart failure and ultimately death.1 Accordingly, prognosis in PAH is strongly related to RV compensation rather than to the degree of the vascular injury itself.1-3 Thus, accurate assessment of intrinsic RV contractility and its relation to the PA load is essential for refining risk stratification and optimizing treatment in these patients.1,3 However, commonly used indexes of RV contractility, such as the RV ejection fraction (EF) and tricuspid annular plane systolic excursion, are limited by considerable load dependence.4

Pressure-volume (PV) relationships provide relatively load-insensitive measures of contractility, such as end-systolic elastance (Ees), preload-recruitable stroke work (PRSW), and Ees/arterial elastance (Ea), a key metric that describes...
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load-insensitive measure of RV contractility. Compared with range of ventricular volumes, arguably provides the most work (SW) versus end-diastolic volume measured across a single beat (model SPC-570-2; Millar Instruments, Houston, TX) was advanced through the right internal jugular vein and positioned at the RV apex to measure instantaneous RV volume. The RV conductance signal was calibrated to match the RV EF, which was independently determined by same-day cardiac magnetic resonance imaging (CMR) and thermodilution cardiac output. PV loops and relationships were constructed both at baseline and during phase 2 of the Valsalva maneuver (period of preload decline), as validated previously. PV data were digitalized at 500 Hz using a custom-designed data acquisition system and stored for subsequent offline analysis. Extraction of the conventional hemodynamic parameters from the PV loops was performed using custom analysis programs (WinPVAN-3.5.10). Further analyses involving multiple-beat Ees and PRSW were conducted using R version 3.0.1.

Calculation of Ees and the PRSW Relationship Using Multiple Beats

The end-systolic points for a series of loops during the Valsalva maneuver were determined as those with maximal elastance and were fitted against the linear end-systolic PV relationship (ESPVR) using the linear least-squares method, with the use of an iterative method to calculate the multiple-beat Ees, as previously reported. Effective Ees was determined by dividing the end-systolic pressure by the stroke volume (SV), and RV-PA coupling was assessed as Ees/Ea. The PRSW relationship was also determined by a linear regression analysis of SW versus
end-diastolic volume data obtained during the Valsalva maneuver, according to the following equation:

\[ SW = M_{sw} \times (V_{ed} - V_{sw}) \]  

(1)

where \( M_{sw} \) and \( V_{sw} \) are the slope and volume-axis intercept, respectively.\(^{13}\) The PRSW coefficients determined from multiple-beat data are specifically denoted as \( M_{sw(MB)} \) and \( V_{sw(MB)} \).

**Outline of the Single-Beat Estimation of the ESPVR and the PRSW Relationship**

Our single-beat approach for estimating RV \( M_{sw} \), hereby described, involves the following: (1) an estimation of the ESPVR from the single-beat late systolic PV relation and (2) a determination of \( M_{sw} \) on the basis of a novel link between the PRSW relationship and the ESPVR. These have been validated previously for the LV in an animal study.\(^{15}\)

**1) Estimation of the curvilinear ESPVR from the single-beat late systolic PV relation**

We used a single-beat method to estimate a curvilinear ESPVR on the basis of the concept of maximum systolic myocardial stiffness, as previously proposed by Mirsky et al.\(^{15,18}\) We modified the original ESPVR formula of Mirsky et al\(^{18}\) to implement a CMR-measured wall volume into the formula, as outlined in detail in Data S1:\(^{19}\)

\[ P_{es}(V_{es}) = A \times \ln\left(\frac{V_{es}}{V_{0}}\right) \times \ln\left(1 + \frac{V_{wall}}{V_{es}}\right) \]  

(2)

where A is an amplification factor and \( V_{0} \) is the volume-axis intercept of the ESPVR, which represents the chamber volume when the fiber stress is assumed to be zero. The RV wall volume was measured using CMR or can be calculated from end-diastolic volume by assuming a constant RV wall-volume ratio when measured wall volume is not available. Figure 1 shows that the single-beat ESPVR was estimated by fitting the points between the peak systolic pressure and the end-systolic point on the signal-averaged baseline loop against equation 2 using the nonlinear least-squares method. This provides a zero-stress volume \( (V_{0}) \) on the basis of the single baseline beat, which was denoted as \( V_{0(SB)} \).

**2) Determination of \( M_{sw} \) from the estimated ESPVR**

The principle for estimating the RV \( M_{sw} \) using a single beat is based on a novel physiologic link detected between the ESPVR and the PRSW relationships.\(^{15}\) We defined the \( P_{m} \) as SW/SV:

\[ SW = P_{m} \times SV \]  

(3)

Combining equations 1 and 3 yields

\[ P_{m} \times SV = M_{sw} \times (V_{es} + SV - V_{sw}) \]  

(4)

which can be rearranged to

\[ (P_{m} - M_{sw}) = \frac{M_{sw}}{SV} \times (V_{es} - V_{sw}) \]  

(5)

More important, this equation indicates that when the end-systolic volume is equal to \( V_{sw} \), \( P_{m} \) should be equal to \( M_{sw} \). If one conceptualizes \( M_{sw} \) and \( V_{sw} \) as a PV coordinate, then the linearity of the PRSW relationship dictates that this coordinate \((V_{sw}, M_{sw})\), when placed on the PV plane, must be on the PV relationship between \( P_{m} \) and \( V_{es} \), as defined in equation 5, which is hereby referred to as the end-systolic P\(_{m}\)-volume relationship (EMPVR) (see Figure 2A). This relationship is conceptually similar to the ESPVR, but with end-systolic pressure replaced by \( P_{m} \). We assumed that the EMPVR: \( P_{m}(V_{es}) \) function has a curvilinear characteristic similar to that of the ESPVR: end-systolic pressure\( (V_{es}) \) (equation 2) with a common \( V_{0} \) but a different amplification factor, as follows:

\[ P_{m}(V_{es}) = \frac{P_{m(sb)}}{P_{es}(V_{es(sb)})} \times P_{es}(V_{es}) \]  

\[ = B \times \ln\left(\frac{V_{sw}}{V_{0(SB)}}\right) \times \ln\left(1 + \frac{V_{wall}}{V_{es}}\right) \]  

(6)

where \( P_{m(sb)} \) and \( V_{es(sb)} \), are known \( P_{m} \) values and end-systolic volumes of the baseline single beat, respectively, and B is an amplification factor.

Once the ESPVR is estimated from a single beat, the EMPVR can also be determined as in equation 6 using \( V_{0(SB)} \). As \( P_{m} \) is equal to \( M_{sw} \) when end-systolic volume is equal to \( V_{sw} \) [ie, \( P_{m}(V_{sw}) = M_{sw} \)], substituting \( V_{sw} \) for \( V_{es} \) in equation 6 yields,

\[ M_{sw} = B \times \ln\left(\frac{V_{sw}}{V_{0(SB)}}\right) \times \ln\left(1 + \frac{V_{wall}}{V_{sw}}\right) \]  

(7)

Also, from equation 1,

\[ SW_{sb} = M_{sw} \times (V_{ed(sb)} - V_{sw}) \]  

(8)

where \( SW_{sb} \) and \( V_{ed(sb)} \) are the known SW and end-diastolic volume of the baseline single beat, respectively. By solving the 2 simultaneous equations for \( M_{sw} \) and \( V_{sw} \) (equations 7 and 8), we finally obtain single-beat estimates of the PRSW coefficients \((M_{sw(SB)}\) and \( V_{sw(SB)}\)). A schematic flow chart using an example case for single-beat estimation of \( M_{sw} \) is summarized in Figure 2B. An algorithm using this outlined approach was generated in R version 3.0.1 (R Foundation) and used to calculate the \( M_{sw} \) uniformly from each signal-averaged single-beat loop.

**Proposal of a Novel Index of Venticuloarterial Coupling**

We propose the use of the ratio of contractility (ie, the \( M_{sw} \)) to \( P_{m} \) as a novel index of RV-PA coupling. By rearranging equation 5, we obtain
\[
\frac{M_{sw}}{SV} = \left( \frac{P_m - M_{sw}}{V_{es} - V_{sw}} \right) = E'_{es} \quad (9)
\]

where \((P_m - M_{sw})/(V_{es} - V_{sw})\) represents the local slope of the EMPVR and is denoted as \(E'_{es}\), as shown in Figure 3. More important, equation 9 indicates that \(E'_{es}\) is consistently equal to the \(M_{sw}\) adjusted by the SV (ie, contractility). On the other hand, pulmonary vascular impedance can be represented by the slope of a diagonal line across the rectangular PV loop, as in Figure 3, which is denoted as \(E'_a\):

\[
E'_a = \frac{P_m}{SV} = \frac{SW}{SV^2} \quad (10)
\]

As shown in Data S2, \(E'_a\) is transformed into an integrated form of vascular impedance on the basis of the concept that the external ventricular work (ie, SW) is equal to the hydraulic energy imparted to the blood.\(^1\)\(^2\)\(^9\) The pumping performance of the ventricle can be determined by \(E'_{es}/E'_a\) in a similar way to \(E_{es}/E_a\) coupling, as shown in Figure 3.\(^7\) Thus, this novel \(E'_{es}/E'_a\) coupling framework, on the basis of the EMPVR, directly relates the intrinsic contractility with the vascular impedance. Multiplying both sides of equation 9 by \(SV/P_m\) yields

\[
\frac{M_{sw}}{P_m} = \frac{(P_m - M_{sw})}{(V_{es} - V_{sw})} \frac{P_m}{SV} = E'_{es}/E'_a \quad (11)
\]

Therefore, the \(M_{sw}\) can be linked with the arterial load simply as \(M_{sw}/P_m\) to determine the RV-PA coupling \((E'_{es}/E'_a)\) in the PV plane. Single- and multiple-beat \(M_{sw}/P_m\) were calculated as \(M_{sw[SB]}/P_m\) and \(M_{sw[MB]}/P_m\) respectively.

**Statistical Analysis**

Data were presented as mean±SD. The PRSW estimates, on the basis of the single-beat approach, were compared with the multiple-beat PRSW measurements (ie, the \(V_{sw[MB]}\) and \(M_{sw[MB]}\)) using Pearson’s correlation coefficient and a linear regression analysis. Bland-Altman analysis was used to assess...
the agreement between single- and multiple-beat measurements. Statistical analyses were conducted with R version 3.0.1.21

Results

Study Population

Of the 41 patients enrolled, we evaluated data from 31 who had successful studies for PV loops and CMR (6 patients did not complete the study for safety or technical reasons, and 4 patients were excluded because of insufficient preload reduction with the Valsalva maneuver). The analyzed cohort included 23 patients with PH (idiopathic PAH, n=6; systemic sclerosis PAH, n=13; systemic sclerosis with PH attributable to heart failure with preserved EF or interstitial lung disease, n=4) and 8 patients without PH (systemic sclerosis without PH, n=6; no PH, n=2). Patient characteristics are shown in Table 1.

Multiple-beat data

Table 2 shows that the PV data were obtained from patients with variable hemodynamic statuses. The RV end-diastolic wall-to-chamber volume ratio was 0.17±0.06. During phase 2 of the Valsalva maneuver, 6.0±2.2 PV loops were obtained. Variable multiple-beat $E_{es}$ (0.14–1.9 mm Hg/mL) and $E_a$ (0.29–2.3) resulted in variable RV-PA coupling ($E_{es}/E_a$), which ranged from 0.23 to 4.5 (1.0±0.91). The PRSW relationship was highly linear ($r=0.97±0.02$), and the $M_{sw(MB)}$ ranged from 14 to 64 mm Hg. The ratio between $M_{sw(MB)}$ and $P_m$ ($M_{sw(MB)}/P_m$) established our novel index of RV-PA coupling.
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Single-beat estimation of the PRSW relationship and RV-PA coupling

An iterative algorithm yielded an average of 20 points (range, 9–56 points) on a steady-state PV loop to be used for single-beat ESPVR estimation. Figure 4 shows that the zero-stress volumes of the single- and multiple-beat ESPVRs were strongly correlated (`r=0.86, P<0.0001`). The single-beat estimation of the PRSW slope (Msw(SB)) estimated from the signal-averaged baseline PV loop was strongly correlated with the multiple-beat PRSW slope, Msw(MB) (Figure 5A, `r=0.89, P<0.0001`). Even when the RV end-diastolic wall-to-chamber volume ratio was assumed (a mean value of 0.17), the single-beat approach provided an accurate estimation of Msw(MB) without individual information on the RV wall volume (Figure 5B, `r=0.91, P<0.0001`). Bland-Altman plots showed a mild overestimation by 3.7 (SEM, 0.92) mm Hg and limits of agreement from –6.5 to 14.0 mm Hg (Figure 5C and 5D).

Although estimated and measured Ees (ie, Msw(SB)/SV and Msw(MB)/SV) strongly correlated with each other (`r=0.95, P<0.0001`, Figure S2), the correlation between estimated and measured RV-PA coupling on the basis of the Ees (ie, single-versus multiple-beat Ees'/Ea' calculated as Msw(SB)/Pm versus Msw(MB)/Pm) was only moderate (`r=0.52, P=0.002`, Figure 6A), with a stronger correlation in patients with greater RV systolic pressure (red circles; RV pressure ≥ median of 44.12 mm Hg, `r=0.68, P=0.004`). Even when the Msw(SB) on the basis of the assumed RV end-diastolic wall-to-chamber volume ratio was used to estimate RV-PA coupling, the correlation between estimated and measured RV-PA coupling was moderate (`r=0.53, P=0.002`, with a stronger correlation in those with greater RV systolic pressure (`r=0.70, P=0.003`), as shown in Figure 6B.

Validation analyses for our single-beat method

To test the validity of the basic principle of our estimation method that the point (Vsw, Msw) is on the EMPVR curve, PRSW coefficients that were determined from the multiple-beat EMPVR (as a point on the EMPVR) were compared with the actual multiple-beat PRSW coefficients (ie, Msw(MB)) and

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Table 1. Patient Characteristics

<table>
<thead>
<tr>
<th>Diagnoses</th>
<th>Value (n=31)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IPAH</td>
<td>6 (19.4)</td>
</tr>
<tr>
<td>SSc-PAH</td>
<td>13 (42.0)</td>
</tr>
<tr>
<td>SSc-PH attributable to HFpEF or ILD</td>
<td>4 (12.9)</td>
</tr>
<tr>
<td>SSc without PH</td>
<td>6 (19.4)</td>
</tr>
<tr>
<td>No PH</td>
<td>2 (6.5)</td>
</tr>
<tr>
<td>Age, y</td>
<td>61±12</td>
</tr>
<tr>
<td>Female sex, n (%)</td>
<td>28 (90)</td>
</tr>
<tr>
<td>Body surface area, m²</td>
<td>1.84±0.21</td>
</tr>
<tr>
<td>NYHA class III/IV, n (%)</td>
<td>12 (39)</td>
</tr>
<tr>
<td>Heart rate, bpm</td>
<td>74±13</td>
</tr>
<tr>
<td>Systemic arterial pressure, mm Hg</td>
<td>94±14</td>
</tr>
</tbody>
</table>

Continuous variables are shown as means±SD. Bpm indicates beats per minute; HFpEF, heart failure with preserved ejection fraction; ILD, interstitial lung disease; IPAH, idiopathic PAH; NYHA, New York Heart Association; PAH, pulmonary arterial hypertension; PH, pulmonary hypertension; SSc, systemic sclerosis.
physiologic link between the EMPVR and PRSW relationship (which identifies the $V_{sw}$ by determining the $P_m$ when the end-systolic volume is equal to $V_{sw}$). Moreover, our single-beat approach is the first to successfully link the $M_{sw}$ with the impedance of the vascular system to generate the ratio $M_{sw}/P_m$, thereby providing a simple yet reliable estimate of RV-PA coupling. These attractive features of our single-beat method should be of great clinical value for the assessment of RV contractile function and RV-PA coupling, and thus can help improve the diagnosis and treatment of diseased RV in humans.

**Single-Beat Estimation of Human RV $E_{es}$ and $M_{sw}$**

Assessment of RV function is increasingly recognized because of growing recognition that RV function is strongly related to outcomes in several cardiovascular diseases, including PH, congenital heart disease, and left-sided heart failure. However, commonly used clinical indexes of RV contractility, such as RV EF or tricuspid annular plane systolic excursion, are limited by their considerable load dependence. Both $E_{es}$ and $M_{sw}$ provide measures of an intrinsic contractile state that are largely independent of loading conditions, even for the RV, but the clinical application of these indexes has been extremely limited by the need to record multiple beats over a wide volume range. Although several single-beat methods for such indexes have been developed and validated for the LV, the application of such methods to the RV has not been demonstrated.

**Discussion**

The present study showed, for the first time, that the slope of the PRSW relationship (ie, $M_{sw}$) can be accurately estimated from a single beat in the human RV by combining the following: (1) an estimation of the curvilinear ESPVR and EMPVR from the late-systolic PV relationship and (2) a novel physiologic link between the EMPVR and PRSW relationship (which identifies the $V_{sw}$ by determining the $P_m$ when the end-systolic volume is equal to $V_{sw}$). Moreover, our single-beat approach is the first to successfully link the $M_{sw}$ with the impedance of the vascular system to generate the ratio $M_{sw}/P_m$, thereby providing a simple yet reliable estimate of RV-PA coupling. These attractive features of our single-beat method should be of great clinical value for the assessment of RV contractile function and RV-PA coupling, and thus can help improve the diagnosis and treatment of diseased RV in humans.

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necessarily been successful in the human RV. A single-beat approach for estimating Ees using the sine-wave fit, originally developed for the LV by Sunagawa et al,\textsuperscript{23,24} is believed to work for the RV. This belief is based on indirect evidence from an animal study that the maximal pressure of isovolumetric contraction (during pulmonary artery clamping) can be predicted from sine-wave fitting of the isovolumetric phase of RV pressure.\textsuperscript{8} However, this single-beat method has never been validated in the human RV and was shown to agree poorly with measured Ees in an animal study.\textsuperscript{25} Furthermore, this estimate was not found to be associated with clinical outcomes in 2 recent studies.\textsuperscript{11,12}

This is the first study to demonstrate a single-beat method that is valid for use in the human RV. We used a novel approach to estimate both the ESPVR and PRSW relationship, which has already been validated for the LV in our animal study.\textsuperscript{15} Because the ESPVR and PRSW relationships have different hemodynamic characteristics, using both relationships would be ideal to enhance clinical interpretation. The curvilinear ESPVR used in the present study is based on the concept proposed by Mirsky et al that the maximal myocardial stiffness (ie, stress/strain ratio of the ventricular wall) attained at end systole is constant throughout short-term changes in preload and afterload.\textsuperscript{18} The estimation of ESPVR from the late systolic PV relation is based on the notion that the late systolic PV relation can be approximated as a set of PV data achieving maximal myocardial stiffness (ie, ESPVR).\textsuperscript{15}

In fact, in our RV data, a near-maximal myocardial stiffness (97.7% of the maximal myocardial stiffness on average) was already attained during late systole (ie, before end systole),

![Figure 5. Estimated and measured preload recruitable stroke work slope, Msw. A, Scatterplots comparing the estimated preload recruitable stroke work (PRSW) slope (Msw(SB)) on the basis of the single-beat method using measured right ventricular wall volume to the multiple-beat PRSW slope (Msw(MB)). B, Similar analysis using the assumed right ventricular (RV) end-diastolic wall/chamber volume ratio. C, Bland-Altman plots for Msw(SB) on the basis of the single-beat method using measured RV wall volume. D, Bland-Altman plots for Msw(SB) using the assumed RV end-diastolic wall/chamber volume ratio.](image-url)
which supports the use of a late systolic PV relation to estimate ESPVR in the human RV. Although the late systolic period can be short in some patients with PH, sufficient data acquisition for curve fitting was possible in the present study. We showed that the $V_0$ of the ESPVR can be accurately estimated using 20 data points on average (range, 9–56 points). Although curvilinear ESPVR models have been shown to provide a more reliable trajectory of the ESPVR than linear ESPVR,\textsuperscript{18,26} it has not been widely acknowledged because it does not provide single numbers for contractility and for ventriculoarterial coupling, such as $E_{es}$ and $E_{es}/E_a$. However, in the present study, because of a novel link between the ESPVR and the PRSW relationship that has been validated for the first time in the RV, the curvilinear ESPVR model was able to provide another load-insensitive measure of contractility, the $M_{sw}$. Because the $M_{sw}$ has many advantages over $E_{es}$, including its strong linearity over a wide range of physiologic loads and independence of chamber size and volume signal gain, our single-beat method may be of great clinical value in assessing RV contractility in diseases of the right side of the heart.

Assessment of RV-PA Coupling Using the $M_{sw}$

Despite the attractive features of the $M_{sw}$, as previously noted, one important limitation of the $M_{sw}$ thus far has been the inability to relate it to PA loads in the PV plane to assess RV-PA coupling. In the present study, we have resolved this long-standing problem with the $M_{sw}$ by developing a novel index of $M_{sw}$-based RV-PA coupling: $M_{sw}/P_m$, which is equal to the $E_{es}/E_a$ in the PV plane. Figure 3 shows that $E_{es}$ is the local slope of the EMPVR, which is analogous to a linear approximation of the curvilinear ESPVR and reflects intrinsic contractility (ie, $M_{sw}$ adjusted by SV). More important, the afterload ($E_a$) that is coupled with the $E_{es}$ represents an integrated form of vascular impedance (Data S2), which is known to provide a comprehensive description of both steady and pulsatile afterloads (including PA resistance and compliance). Thus, the $E_{es}/E_a$ framework is physically meaningful because it is derived mathematically from well-established measures of contractility and vascular load. This is in contrast to the $E_{es}/E_a$ framework, wherein $E_a$ is only indirectly related to vascular impedance.\textsuperscript{27} By considering $P_m$, rather than end-systolic pressure (ie, at EMPVR rather than at ESPVR), one can assess RV-PA coupling on the basis of $E_{es}/E_a$, in a similar way to $E_{es}/E_a$. Although a minor difference between end-systolic pressure and $P_m$ is expected in the LV, where the pulsatile component of the external ventricular power is far less than the steady component, a significant difference would exist for the RV, where the pulsatile power is reported to account for as much as 23% of the total ventricular power.\textsuperscript{20} This was clearly shown by only a moderate correlation ($r=0.53$) and a
considerable difference in the range of change between multiple-beat $E_{es}/E_a$ and $E_{es}/E_a'$ (ie, $E_{es}/E_{a[MB]}$ and $M_{sw}/P_m$ [MB]), as shown in Figure S1. We speculate that the optimal value for the novel index of RV-PA coupling is $\approx 0.8$ to $1.0$, which was attained by those without PH (Figure S1). Although $E_{es}/E_a$ is an established marker of RV-PA coupling, $E_{es}/E_a'$ potentially provides an even better characterization of RV-PA coupling, considering the significantly pulsatile PA load caused by enhanced PA wall stiffness and its substantial impact on survival in PAH.\(^\text{24}\) Because survival in PAH is closely related to RV adaptation to the increased pressure overload (ie, RV-PA coupling),\(^1\) whether the PRSW-based RV-PA coupling index (ie, $M_{sw}/P_m$ or $E_{es}/E_a'$) predicts clinical outcomes better than the traditional RV-PA coupling indexes, such as RV EF or an ESPVR-based index ($E_{es}/E_a$), warrants future investigations.

In the present study, although the agreement between estimated and measured $E_{es}'$ on the basis of our single-beat method was excellent, the correlation between measured and estimated RV-PA coupling on the basis of the $E_{es}'$ was only moderate for the whole population. This was mainly because the estimation of RV-PA coupling was less accurate in those with low afterload, wherein a small estimation error in contractility would result in a relatively large estimation error in RV-PA coupling (ie, ratio of contractility/afterload). However, RV-PA coupling is of critical issue for those with elevated RV pressure in the setting of PH. For such patients, our single-beat method provided a better estimation of RV-PA coupling ($r=0.70$).

Study Limitations

First, this was a retrospective single-center study that enrolled a comparatively small number of patients. The sensitivity of our single-beat $M_{sw}$ to short-term changes in inotropic status was not investigated in the present study, which needs to be elucidated in future investigations. Second, the formula used for modeling the curvilinear ESPVR in the present study was developed under the assumption of the ventricular shape to be prolate spheroid, and thus, the applicability of such a formula to the RV was not guaranteed.\(^1\)\(^5\) However, our single-beat approach was capable of estimating the $M_{sw}$ much more accurately than those based on a generally used linear ESPVR model, which provides strong evidence for the validity of our model. In addition, our single-beat method has an advantage of allowing selection of any other ESPVR formulas for further refinement. Third, the algorithm of our single-beat method may be somewhat complicated compared with that of the conventional single-beat method for $E_{es}$ using sine-wave fitting. However, when the conventional sine-wave fit was applied to our data,\(^8\)^\(^\text{22}\) the correlation between single- and multiple-beat $E_{es}$ was low ($r=0.41$), as shown in Figure S4, which would preclude a reliable assessment of contractility. A program to automate the data processing of our single-beat method is provided in Data S3 for the readers’ wide use. Finally, our single-beat approach still requires measurement of instantaneous PV data. A combination of the pressure recording of the normal right-sided heart catheterization and flow/volume data on CMR would possibly provide a good approximation of the late systolic PV relation as well as the baseline SW necessary for our single-beat approach. This needs further study.

Conclusions

A load-insensitive measure of contractility, $M_{sw}$, for the human RV and its coupling to PA afterloads can be accurately estimated using our single-beat approach. This approach would help in precisely assessing RV adaptation to increased pressure overload and thereby help improve the management of patients with PH.

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Disclosures

None.

References


SUPPLEMENTAL MATERIAL
Data S1. End-systolic pressure volume relationship based on maximum myocardial stiffness

Mirsky et al. demonstrated that the maximal stress–strain ratio (i.e., myocardial stiffness) attained during end-systole is constant throughout acute changes in preload, and that the end-systolic pressure volume relationship (ESPVR) derived from the myocardial stress-strain relationship is curvilinear, representing a more physiologic ESPVR than linear ESPVR. The average fiber stress (σ) can be defined by
\[ \sigma = \frac{3}{2} \times P \times \frac{V_m}{V_{wall}}, \]
where P is the left ventricular pressure, and V_m and V_wall are the midwall and wall volumes, respectively. The midwall volume is defined as the logarithmic mean of the chamber volume (V) and the outer volume (V_out),
\[ V_m = V_{wall} / \ln(1 + \frac{V_{wall}}{V}). \] [equation S1] Therefore,
\[ V_m = \frac{V_{wall}}{\ln(1 + \frac{V_{wall}}{V})}. \]

The midwall natural strain (\(\varepsilon_n\)) can be defined by
\[ \varepsilon_n = \frac{1}{3} \ln(V_m / V_{m0}), \]
where V_{m0} is the midwall volume at zero stress. We set the systolic zero-stress volume as the reference distension, as described by Mirsky et al. The average fiber strain can be calculated as
\[ \varepsilon = K_m \varepsilon_n, \]
where K_m is the constant value determined by the assumed geometry of the ventricle. The average systolic myocardial stiffness (E_av) is defined as
\[ E_{av} = \frac{\sigma}{\varepsilon}, \]
and end-systole was defined as the latest time at which the systolic myocardial stiffness reached its maximum value (max E_{av}). The end-systolic, stress–strain relationship (\(\sigma_{es}\) versus \(\varepsilon_{es}\)) based on the maximal stiffness concept can be represented in the form:
MaxE\text{av} = \frac{\sigma_{es}}{\varepsilon_{es}} = \left(\frac{9}{2K_m}\right) \times P_{es} \times V_{mes}/(V_{wall} \times \ln(V_{mes}/V_{m0})). \quad \text{[equation S2]}

where $P_{es}$ and $V_{mes}$ are the left ventricular pressure and midwall volume at end-systole, respectively.

Rearranging equation S2 results in the following equation:

$$P_{es} = \left(\frac{2K_m}{9}\right) \times \text{maxE}_{av} \times \ln\left(\frac{V_{mes}}{V_{m0}}\right) \times \frac{V_{wall}}{V_{mes}} \quad \text{[equation S3]}$$

Converting the midwall volume to the chamber volume using equation S1 yields,

$$P_{es} = \left[\left(\frac{2K_m}{9}\right) \times \text{maxE}_{av}\right] \times \ln\left(\frac{\ln(1+\frac{V_{wall}}{V_0})}{\ln(1+\frac{V_{wall}}{V_{es}})}\right) \times \ln(1 + \frac{V_{wall}}{V_{es}}). \quad \text{[equation S4]}$$

Therefore, ESPVR can be expressed in the form:

$$P_{es}(V_{es}) = a \times \ln\left(\frac{V_{mes}}{V_{m0}}\right) \times \ln(1 + \frac{V_{wall}}{V_{es}}), \quad \text{[equation S5]}$$

where $a$ is an amplification factor. Equation S5 can be approximated using the following simpler formula:

$$P_{es}(V_{es}) = A \times \ln\left(\frac{V_{es}}{V_0}\right) \times \ln(1 + \frac{V_{wall}}{V_{es}}), \quad \text{[equation S6]}$$

where $A$ is an amplification factor.
Data S2. Relationship between $\dot{E}_a$ and pulmonary impedance

Milnor et al. described that the mean external ventricular work per time ($W$) can be expressed as a sum of the steady and pulsatile components:\(^3\)

$$W = P_0 Q_0 + \frac{1}{2} \sum_{n=1}^{N} Q_n^2 Z_n \cos \theta_n,$$  \hspace{1cm} [equation A1]

using the mean values of pressure and volume flow $P_0$ and $Q_0$, harmonic components $Q_n$ of the ventricular ejection wave, input impedance $Z_n$, and phase $\theta_n$. At any given rate, the ratio of each harmonic amplitude to mean flow ($Q_n/Q_0$) was found to be fairly consistent.\(^3\)

Therefore,

$$Q_n = C_n \times Q_0,$$ \hspace{1cm} [equation A2]

where $C_n$ is a function of heart rate. Combined with equation A1,

$$W = P_0 Q_0 + \frac{1}{2} Q_0^2 \sum_{n=1}^{N} C_n^2 Z_n \cos \theta_n,$$  \hspace{1cm} [equation A3]

Stroke work (SW) and stroke volume (SV) can be expressed as

$$SW = W \times T_0,$$ \hspace{1cm} [equation A4]

$$SV = Q_0 \times T_0,$$ \hspace{1cm} [equation A5]

where $T_0$ is cycle length. We defined $\dot{E}_a$ as $P_m/SV$, which is equal to $SW/SV^2$. Combined with equations A3-A5,

$$\dot{E}_a = \frac{SW}{SV^2} = \frac{1}{T_0} \left( \frac{P_0}{Q_0} + \frac{1}{2} \sum_{n=1}^{N} C_n^2 Z_n \cos \theta_n \right).$$ \hspace{1cm} [equation A6]

Therefore, $\dot{E}_a$ is a global marker of vascular impedance, which inherently accounts for both steady and pulsatile afterloads.
**Data S3. R code for our single-beat method**

#Pres, pressure data of the signal-averaged pressure–volume loop  
#Vol, volume data of the signal-averaged pressure–volume loop  
#MRIwallv, right ventricular wall volume as measured by cardiac magnetic resonance imaging  
#sESV, end-systolic volume of the signal-averaged pressure–volume loop  
#sEDV, end-diastolic volume of the signal-averaged pressure–volume loop  
#sSW, stroke work of the signal-averaged pressure–volume loop

library(nleqslv)
pp<-min(c(1:length(Pres))[Pres== max(Pres,na.rm=T)][1])
ep<-min(c(1:length(Pres))[Vol== min(Vol, na.rm=T),na.rm=T)
esp<-NA
pVo<-NA
pa<-NA
pVo[1]<- 0.1*sEDV
pa[1]<-100
for(i in 1:100){ 
  Tc<-function(x){log(x/pVo[i])*log(1+(MRIwallv/x))}
  maxeav<-max(Pres[pp:ep]/Tc(Vol[pp:ep]),na.rm=T)
  esp[i]<-max(c(pp:ep)[Pres[pp:ep]/Tc(Vol[pp:ep])]>0.99*maxeav,na.rm=T)
  Xs<-Vol[pp: esp[i]]
  Ys<-Pres[pp: esp[i]]
  fits<-nls(Ys~ a*log(Xs/c)*log(1+(MRIwallv/Xs)),start=c(c=pVo[i],a=pa[i]),trace=T)
  pVo[i+1]<-summary(fits)$coefficient[1]
  pa[i+1]<-summary(fits)$coefficient[2]
  Vo<-pVo[length(pVo)]
  A<-pa[length(pa)]
  tPm<-function(x){A*log(x/Vo)*log(1+(MRIwallv/x))}
  A2<-sSW/(sEDV-sESV)/(log(sESV/ Vo)*log(1+(MRIwallv/sESV))
  Pm<-function(x){A2*log(x/ Vo)*log(1+(MRIwallv/x))}
  Est<-function(Vsw){Msw<-sSW/(sEDV-Vsw)
  y<-Pm (Vsw)-Msw}
  Vsw<-nleqslv(0.5*sESV,Est)$x
  Msw<-sSW/(sEDV-Vsw)
  Msw
Figure S1. Relationship between conventional and novel indices of right ventricular–pulmonary arterial coupling

Scatterplots comparing multiple-beat $E_{es}/E_{a}$ ($E_{es}/E_{a(MB)}$) and multiple-beat $M_{sw}/P_{m}$ ($M_{sw(MB)}/P_{m}$). The right ventricular–pulmonary arterial coupling ratio in patients with pulmonary hypertension (black points) was significantly lower than that in those without pulmonary hypertension (red points) ($p=0.03$ for $E_{es(MB)}/E_{a}$ and $p=0.003$ for $M_{sw(MB)}/P_{m}$). Multiple-beat $M_{sw}/P_{m}$ had limited values around 0.80–1.0 in those without PH (red points), whereas multiple-beat $E_{es}/E_{a}$ ranged much more widely.
Figure S2. Estimated and measured $E'_{es}$

A) Scatterplots comparing the estimated and measured $E'_{es}$ (i.e., $M_{sw(SB)}$/stroke volume and $M_{sw(MB)}$/stroke volume) based on measured right ventricular wall volume.

B) Bland–Altman plots for the estimation of $E'_{es}$ based on measured right ventricular wall volume, which shows a mild overestimation by 0.05 (standard error: 0.013) mmHg/mL and limits of agreement from -0.10 to 0.20 mmHg/mL.
Figure S3. Consistency assessment within multiple-beat data.

Scatterplots comparing the preload recruitable stroke work (PRSW) coefficients estimated as a point on the end-systolic mean ejection pressure–volume relationship (EMPVR) using multiple-beat EMPVR versus the actual multiple-beat PRSW coefficients calculated based on the stroke work-volume plane (i.e., EMPVR-based versus actual $M_{sw(MB)}$ and $V_{sw(MB)}$). These graphs validate the basic principle of the estimation method that point $(V_{sw}, M_{sw})$ is on the EMPVR curve.
Figure S4. Conventional single-beat estimation of end-systolic elastance ($E_{es}$) based on sine-wave fit.

Scatterplots comparing single-beat end-systolic elastance ($E_{es}$) based on the conventional sine-wave method and multiple-beat $E_{es}$ ($E_{es(SB)}$ versus $E_{es(MB)}$).
Figure S5. Iterative method used for single-beat estimation of the end-systolic pressure–volume relationship.

(A) The first curve fitting was performed on late systolic points (red open points) based on a randomly selected end-systolic point. The zero-stress volume ($V_0$, red closed point) was obtained from the estimated end-systolic pressure–volume relationship (ESPVR, red curve), which was used to calculate myocardial stiffness (stress–strain ratio).

(B) Myocardial stiffness was calculated based on the first $V_0$. The last point attaining the maximal myocardial stiffness (black closed point) was used as the end-systolic point for the second fitting. The second end-systolic point is different from the first randomly selected end-systolic point.

(C) The second curve fitting was performed for the late systolic points determined based on myocardial stiffness (blue open points). The second $V_0$ (blue closed point) was obtained from the second fitting and was used to calculate myocardial stiffness again. This process was repeated until we identified the end-systolic point that attained the maximal myocardial stiffness, as shown in Figure 1B.
Supplemental References:

