ORIGINAL ARTICLE

RV-pulmonary arterial coupling predicts outcome in patients referred for pulmonary hypertension

Rebecca R Vanderpool, Michael R Pinsky, Robert Naeije, Christopher Deible, Vijaya Kosaraju, Cheryl Bunner, Michael A Mathier, Joan Lacomis, Hunter C Champion, Marc A Simon

ABSTRACT

Objective  Prognosis in pulmonary hypertension (PH) is largely determined by RV function. However, uncertainty remains about what metrics of RV function might be most clinically relevant. The purpose of this study was to assess the clinical relevance of metrics of RV functional adaptation to increased afterload.

Methods  Patients referred for PH underwent right heart catheterisation and RV volumetric assessment within 48 h. A RV maximum pressure (Pmax) was calculated from the RV pressure curve. The adequacy of RV systolic functional adaptation to increased afterload was estimated either by a stroke volume (SV)/end-systolic volume (ESV) ratio, or by EF (RVEF). Diastolic function of the RV was estimated by a diastolic elastance coefficient β. Survival analysis was via Cox proportional hazard analysis.

Results  Patients (n=50; age 58±13 yrs) covered a range of mPAP (13–79 mm Hg) with an average RVEF of 39±17% and ESV of 143±89 mL. Average estimates of the ratio of end-systolic ventricular to arterial elastance were 0.79±0.67 (SV/ESV) and 2.3±0.65 (Pmax/mPAP-1). Transplantation-free survival was predicted by right atrial pressure, mPAP, pulmonary vascular resistance, β, SV, ESV, SV/ESV and RVEF, but after controlling for right atrial pressure, mPAP, and SV, SV/ESV was the only independent predictor.

Conclusions  The adequacy of RV functional adaptation to afterload predicts survival in patients referred for PH. Whether this can simply be evaluated using RV volumetric imaging will require additional confirmation.

BACKGROUND

It has been better appreciated in recent years that RV function is a major determinant of functional state, exercise capacity and survival in patients with severe pulmonary hypertension (PH). However, how to measure RV function and what parameters might be most clinically relevant remains debated.

The normal RV is a thin-walled crescent shape flow generator unable to cope with brisk increases in pulmonary artery pressures (PAP). However, in the presence of chronically increased PAP the RV adapts by a progressive increase in contractility allowing for maintained output. When eventually this ‘homeometric adaptation’ fails, filling pressures and dimensions of the RV increase, resulting in maintenance of flow output through a ‘heterometric adaptation’, as described by Starling’s law of the heart. Therefore, gold standard measurements of RV function in patients with PH necessarily include estimations of contractility and afterload.

RV contractility is quantified by maximum elastance, or the maximum value of RV pressure-volume relationships, usually estimated by an end-systolic elastance (Ees) or ratio of end-systolic pressure (ESP) to end-systolic volume (ESV). An acceptable reference measure for RV afterload is arterial elastance (Ea), or ESP divided by stroke volume (SV). Importantly, the optimal RV-arterial coupling, which allows for flow output at minimal energy cost, corresponds to an Ees/Ea ratio of approximately 1.5 to 2.

A limited number of studies in patients with severe PH have reported an increase in Ees in proportion to increased pulmonary vascular resistance (PVR) with preserved or decreased Ees/Ea. These studies used either single beat or multiple beat measurements of Ees, or attempted a simplified approach by exclusive volume measurements. Different methods result in variable values for the Ees/Ea ratio. Furthermore, the prognostic relevance of any representation of Ees/Ea as a measure of RV-arterial coupling in patients with or the suspicion of PH has never been studied.

In the present study, we evaluated the prognostic impact of different methods to estimate Ees/Ea from right heart catheterisation (RHC) and imaging in patients with a spectrum of severities of PH. Because systolic function impacts on diastolic function, we also calculated an index of diastolic stiffness β recently shown to be significantly increased in patients with pulmonary arterial hypertension (PAH).

METHODS

Study setting  Patients referred to the University of Pittsburgh Comprehensive Pulmonary Hypertension Program, and undergoing RHC for evaluation of PH, gave an informed consent to the study which was approved by the University of Pittsburgh Institutional Review Board. The study was prospective. Inclusion criteria were: (1) age ≥18 years; (2) clinically indicated RHC and cardiac imaging (either multislice CT (MSCT) of the chest, or cardiac MRI (CMR)). Exclusion criteria were: (1) international normalized ratio (INR)>1.6; (2) platelet count<50 000; (3) unable to lay flat for imaging.
procedures; (4) known allergy to intravenous contrast; (5) serum creatinine greater than 2.0; (6) known arrhythmia precluding adequate gating for cardiac imaging.

**RV volumetric assessment**

RV volume was measured by either CMR (n=6) or gated cardiac MSCT (n=44). The cardiac MSCT scanning protocol was developed specifically for this study as previously reported. Patients were scanned on a GE Lightspeed 16-slice scanner (n=13), or GE VCT 64-slice scanner (n=31) (General Electric Healthcare; Milwaukee, Wisconsin, USA). Multiplanar reformatted images and three-dimensional (3D) and 4D whole heart volume renderings were performed on a free-standing workstation (GE ADW V4.3, GE Healthcare, Milwaukee, Wisconsin, USA). RV function was measured using standard summation of slab volume technique as has been used in evaluating left ventricular volumes. The endoluminal borders of the RV in each slice were manually traced from apex to the tricuspid valve plane in both end-systole and end-diastole. CMR was performed by dedicated CMR technologists with a 1.5 T Siemens Magnetom Espree (Siemens Medical Solutions, Erlangen, Germany) and a 32-channel phased array cardiovascular coil. The CMR scans included standard breath-held segmented cine imaging with steady-state free precession and prospective ECG gating. RV volumes were measured using Simpson’s rule without geometric assumptions from short axis stacks of end diastolic and end systolic cine frames (slices 6 mm thick, 4 mm gap, 30 frames per cardiac cycle) by physicians experienced in CMR.10

**Doppler echocardiography**

In a subset of patients, Doppler RV outflow tract flow velocity measurements obtained simultaneously with RHC were sampled for RV-arterial coupling measurements using the single beat approach, as indicated below.

**Right heart catheterisation**

RHC was performed within 48 h of MSCT or CMR using a balloon-tipped thermolitldation 7 F Swan-Ganz catheter for the measurements of right atrial pressure (RAP), PAB occluded PAP, RV pressure and cardiac output (CO). Screen captures of the RV pressure tracings stored in the WITT system were later analysed and semiautomatically redigitised using a custom programme in Matlab.11

**Pressure-volume analysis**

To calculate Ees, the single beat method was employed.12 Briefly, a maximum pressure, RV maximum pressure (Pmax), was calculated from the non-linear extrapolation of the early systolic and diastolic portions of the RV pressure curve. ESP was approximated by mean PAP (mPAP).13 Ees was then calculated as the slope of end-systolic pressure-volume (P-V) relationship: (Pmax–mPAP) divided by SV.14 Ea was estimated by the ratio of mPAP to SV, assuming RV pressure at end-diastolic volume (EDV) equal to zero.15 Since the Ees/Ea ratio can be simplified by omitting either mPAP or SV as common terms, Ees/Ea was also estimated by either the ratio of SV/ESV defined as the ‘volume method’ or (Pmax/mPAP–1) defined as the ‘pressure method’ (figure 1). Because the shape of RV pressure-volume loops may suggest that ESP is closer to systolic RV pressure (sRVP) than to mPAP, Ees/Ea was also calculated by the pressure method with sRVP instead of mPAP.

Diastolic stiffness was calculated by fitting a non-linear exponential curve to the diastolic portion of the single-beat pressure-volume loops as recently published.8 Specifically, the diastolic stiffness β and a curve-fitting parameter α were determined by fitting the non-linear exponential, \( P = \alpha (e^{\beta t} - 1) \) using three points from the pressure-volume loop (figure 1). The first point is the pressure at the beginning of diastole following the isovolumetric relaxation when the RV is at the ESV. The second point is at the pressure and volume before the isovolumetric contraction. The third point is assumed to go through the origin (0,0).

All single beat pressure-volume loop analyses of RV systolic and diastolic function were performed in Matlab (Matlab 2013a programme The MathWorks, Natick, Massachusetts, USA).

**Study protocol**

All patients had a RHC and either MSCT or CMR within 48 h. Additionally, the single beat method with Doppler echocardiographic measurement of pulmonary flow was applied in a subset of three patients. Instantaneous changes in RV volume were estimated by integration of the volumetric flow rate calculated from the product of the Doppler velocities and the mean cross-sectional area of the pulmonary artery determined from thermodilution (where mean cross-sectional area=CO/mean velocity). The single beat method allows for the definition of ESP a line drawn from Pmax tangent to the RV pressure-volume loop with relative change in volume calculated from the integration of pulmonary flow.12

**Statistics**

Data are expressed as mean±SD. Statistical significance was defined as two-sided p<0.05. Primary outcome of survival was defined as freedom from lung transplantation or death. Difference between estimation methods were calculated using a paired Student t test. In order to determine factors prognostic of survival, Cox regression analyses were conducted on the whole population. Univariate (significance set at p<0.05 cut-off), and multivariate (backward selection set at p<0.1 cut-off, including all variables having a p<0.1 in the univariate analysis) Cox analyses were employed. Receiver operator characteristic curves (ROC) were plotted for variables significant in the multivariate Cox models to determine area under the curve, specificity and sensitivity of the optimal cut-offs. Optimal cut-offs were defined as the point on the ROC curve nearest the upper left corner of the ROC graph. Kaplan–Meier survival curves were constructed and the log-rank test was used to compare survival in patients stratified by cut-off determined by ROC. Statistical calculations were made using SPSS for Windows (V20, SPSS, Chicago, Illinois, USA) or R (R Project for Statistical Computing, http://www.r-project.org/) for the survival analysis.

**RESULTS**

A total of 50 patients (34 women) were enrolled, aged 58±13 years, with a New York Heart Association (NYHA) functional class of 2.5±0.8 and a distance walked in 6 min of 295±125 m. Nine of them had no PH as defined by a mPAP<25 mm Hg. Their mPAP was 17±4 (range: 10–21) mm Hg. They had been referred for a suspicion of PH because of unexplained dyspnoea and a question of increased maximum velocity of tricuspid regurgitation at Doppler echocardiography. The diagnoses in the remaining 41 patients were idiopathic PAH in 11, connective disease associated PAH in 12, chronic thromboembolic PH in 3, sarcoidosis-associated PH in 3, portal hypertension-associated PH in one, PH secondary to idiopathic lung fibrosis in 7, and left ventricular diastolic dysfunction in 4. LVEF was 62±6%. Treatments of PH consisted in

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**References**

phosphodiesterase-5 inhibitor in 12 patients, endothelin receptor antagonists in 5, Prostacyclins in 5, calcium channel blockers in 11, warfarin in 11, diuretics in 29 and digitalis in 3.

The haemodynamic measurements are summarised in Table 1. There was a wide range of pulmonary haemodynamics and RV adaptation to afterload as assessed by RVEF and N-terminal pro b-type natriuretic peptide (NT-pro-BNP) levels.

The measurements of Ees and Ea, and Ees/Ea ratio by the volume and pressure methods are shown in Table 2. Ees was increased, but Ees/Ea, by either the volume or the pressure method, was decreased in proportion to increased PAP (Figure 2). Ees and Ees/Ea were the highest with the mPAP pressure method and lowest with the volume method, with the sRVP pressure method in between (Table 2). The diastolic stiffness coefficient β was increased in proportion to increased Ees (pressure method) and decreased Ees/Ea (SV/ESV) (Figure 3). β was not significantly associated with Ees by volume method (mPAP/ESV; R=0.016; p=0.91) and Ees/Ea by pressure method (Pmax/mPAP-1; R=0.34; p=0.34).

The average follow-up time was 2.1±1.9 years with a maximum follow-up of 5.9 years with 16 events (11 deaths and 5 lung transplants). Unadjusted univariate Cox proportional hazards ratio to predict the primary outcome of time to death or lung transplant was performed on the following variables: RAP, mPAP, PVR, SV, ESV, EF, SV/ESV, β and Pmax/mPAP-1. Higher mPAP, PVR, ESV and β, and lower SV, RVEF and Ees/Ea by the volume method (SV/ESV) were all predictive of outcome by univariate Cox proportional hazards ratio (Figure 4) in the entire cohort. Ees/Ea estimated by the pressure method with either mPAP or sRVP to estimate ESP did not predict outcome in the entire cohort. In the multivariate cox model of the entire cohort controlling for RAP, mPAP and SV, SV/ESV emerged as the only independent predictor of transplant-free survival (OR 0.030, 95% CI 0.003 to 0.324, p=0.004, overall model p=0.003). This remained the case after controlling for diastolic stiffness (β), RAP, and SV in another model (OR 0.066, 95% CI 0.005 to 0.873, p=0.039, overall model p=0.002). But in a model controlling for sex, age and WHO functional class, SV/ESV was the only significant predictor (OR 0.24, 95% CI 0.002 to 0.391, p=0.009, overall model p<0.001). When the cohort was limited to those with PH (mPAP>25 mm Hg), SV, ESV, RVEF, β and SV/ESV were the only univariate predictors of

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**Table 1** Right heart catheterisation and RV volumetric measurements

<table>
<thead>
<tr>
<th></th>
<th>Mean±SD</th>
<th>Range</th>
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<tbody>
<tr>
<td>Heart rate (bpm)</td>
<td>75±13</td>
<td>45–98</td>
</tr>
<tr>
<td>Right atrial pressure (mm Hg)</td>
<td>9±5</td>
<td>0–20</td>
</tr>
<tr>
<td>Mean pulmonary artery pressure (mm Hg)</td>
<td>40±16</td>
<td>10–79</td>
</tr>
<tr>
<td>Occluded pulmonary artery pressure (mm Hg)</td>
<td>13±5</td>
<td>3–28</td>
</tr>
<tr>
<td>Pulmonary vascular resistance (wood units)</td>
<td>6.7±6.3</td>
<td>0.6–24.9</td>
</tr>
<tr>
<td>Cardiac output (L/min)</td>
<td>5.2±1.8</td>
<td>2.4–9.3</td>
</tr>
<tr>
<td>RV end-diastolic volume (mL)</td>
<td>219±91</td>
<td>93–630</td>
</tr>
<tr>
<td>RV end-systolic volume (mL)</td>
<td>143±89</td>
<td>25–540</td>
</tr>
<tr>
<td>RVEF (%)</td>
<td>39±17</td>
<td>7–78</td>
</tr>
<tr>
<td>NT pro-BNP</td>
<td>1030±1870</td>
<td>4–9863</td>
</tr>
</tbody>
</table>

**Table 2** End-systolic elastance (Ees), arterial elastance (Ea) and Ees/Ea in 50 patients referred for pulmonary hypertension

<table>
<thead>
<tr>
<th></th>
<th>Mean±SD</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pressure method (ESP=mPAP)</td>
<td>Ees</td>
<td>1.38±0.88</td>
</tr>
<tr>
<td></td>
<td>1.1±0.7*</td>
<td>0.37±0.33*</td>
</tr>
<tr>
<td></td>
<td>Ea</td>
<td>0.64±0.46</td>
</tr>
<tr>
<td></td>
<td>1.0±0.7*</td>
<td>0.64±0.46</td>
</tr>
<tr>
<td></td>
<td>Ees/Ea</td>
<td>2.30±0.65</td>
</tr>
<tr>
<td></td>
<td>1.4±0.88*</td>
<td>0.79±0.67*</td>
</tr>
</tbody>
</table>

Means±SD values of Ees calculated as mPAP/ESV, (Pmax-mPAP)/SV, or (Pmax-sRVP)/SV, of Ea calculated as mPAP/SV or sRVP/SV. The RV-arterial coupling ratio was reduced to SV/ESV in the volume method and (Pmax/mPAP-1) or (Pmax/sRVP-1) in the pressure methods. The lowest Ees/Ea was calculated with the volume method and the highest with the pressure method (with mPAP to estimate ESP).

*p Value<0.05 compared to pressure method with ESP=mPAP.

ESP, end-systolic pressure; SV, end-systolic volume; mPAP, mean pulmonary artery pressure; sRVP, peak systolic RV pressure; SV, stroke volume.
outcomes. In this cohort limited to mPAP ≥ 25 mm Hg, SV/ESV remained an independent predictor of transplant-free survival by multivariate analysis controlling for RAP, mPAP and SV (OR 0.035, 95% CI 0.003 to 0.379, p=0.006, overall model p=0.015), and this was a much stronger predictor than RVEF in a similar model (OR 0.928, 95% CI 0.883 to 0.974, p=0.003, overall model p=0.002). Patients stratified by a ROC-derived cut-off value of ≤ 0.515 for Ees/Ea showed highly significant differences in transplant-free survival in the associated Kaplan-Meier curve (figure 5).

Because assumptions about the shape of the pressure-volume loop significantly impact the measured Ees calculated from non-simultaneous measures of pressure and volume, we used simultaneous RV pressure and Doppler pulmonary flow velocity in three patients to construct representative instantaneous pressure-volume loops. Relative changes in RV volume during a cardiac cycle were calculated from the Doppler flow velocities (figure 6). The patients, respectively, had a normal mPAP (17 mm Hg) and CO (5.0 L/min), or elevated mPAP (62 and 47 mm Hg), and reduced or normal CO (B: 2.4 L/min and C: 8.3 L/min). Ees calculated from the line drawn from Pmax and tangent to the P-V curve was close to Ees calculated by the sRVP pressure method. Ees slopes calculated with mPAP were slightly higher.

DISCUSSION

The present results show that the coupling of RV systolic function to afterload is an independent predictor of transplantation-free survival in patients with PH of variable severities and origins. In this population of patients referred for PH, the simplest metric of RV-arterial coupling based on imaging of end-systolic and EDVs was prognostically superior to invasive measurements provided by a RHC.

The adequacy of RV functional adaptation to afterload in patients with PH is usually evaluated in clinical practice by a variety of invasive and non-invasive measurements. Of those, RAP and CO (RHC), tricuspid annular plane excursion, RV performance index, atrial size, strain and pericardial effusion (echo-cardiography), SV, ESV, EDV and RVEF (CMR) and brain natriuretic peptides have been demonstrated to be of prognostic relevance. However, available data do not offer a hierarchy allowing clinicians to select pertinent and practical ones depending on patients and available resources. This is why in the present study we explored the predictive value of measurements more closely pertaining to gold standards of systolic and diastolic function of the RV.

Systolic ventricular function, or contractility, is defined by a maximum ventricular elastance, or Ees. Similarly, ventricular afterload is best defined as Ea. Both can be measured on the same PV loop. An Ees/Ea higher than 1, normally in the range of 1.5–2, allows for RV flow output at minimal energy cost, and thus reflects optimal ventriculo-arterial coupling. Several recent studies have reported on Ees/Ea in patients with severe forms of PH such as PAH or chronic thromboembolic PH, showing adaptive increase in Ees with more or less preserved Ees/Ea ratios. However, the predictive value of RV-arterial coupling measured by Ees/Ea has not yet been determined.

In the present study, we looked for the best combination of physiologically sound and clinically feasible prognostic methods.

Figure 2 Ratio of end-systolic elastance (Ees)/elastance (Ea) calculated by pressure method (A) or volume method (B) plotted versus mean pulmonary arterial pressure (mPAP). Ees/Ea is greater when calculated by the pressure method than when calculated by the volume measurement, however, inverse relationship to mPAP is consistent between methods.

Figure 3 Diastolic stiffness increased linearly with RV contractility (A). However, there is an inverse relationship between diastolic stiffness and the RV-arterial coupling ratio, stroke volume (SV)/end-systolic volume (ESV).
The volume method is an Ees/Ea ratio simplified for pressure. In doing so, there is an inherent assumption that V₀, or the RV volume at zero filling pressure, would be equal to zero, which is unrealistic. The end-systolic P-V ratio measured experimentally on a non-ejecting ventricle, or on a family of P-V loops from an ejecting ventricle, is actually slightly curvilinear but with a positive V₀. Therefore, SV/ESV underestimates Ees/Ea as compared with measurements taking ESP into account. The pressure method calculates Ees/Ea from Pmax, mPAP and SV, resulting in a pressure ratio (Pmax/mPAP−1). This method assumes preload-independency of ESV and mPAP as a valid estimate of ESP. Both assumptions suffer from several limitations. However, the pressure method may provide more accurate measurements of Ees/Ea than the volume method, as relative preload-independency is more likely over the range of reported measurements than a zero value for V₀. This was confirmed by Ees/Ea measurements calculated via the single beat methodology in three patients. It is therefore surprising that the volume measurement of RV-arterial coupling outperformed other measurements of the adequacy of the coupling of RV systolic function to afterload in predicting outcomes. The reasons for this are not entirely clear. The practical relevance of the finding is that prognostically relevant RV-arterial coupling can be measured using an entirely non-invasive approach. This will require confirmation on a larger patient population.

The PV loop of a normal RV is of triangular shape, with ejection continuing after maximum elastance, in relation to the low pulmonary vascular impedance. The normal left ventricular P-V loop is of square shape with maximum Ea at end-systole, and thus Ees and Emax coinciding at the upper left corner of the PV loop. In severe PH as PVR progresses, the PV loop of the RV transitions to a square shape such that the difference between Ees and Emax becomes insignificant, or may take a triangular shape with early peaking, and ESP getting close to peak systolic pressure of the RV as also illustrated in figure 6. Therefore, at a low PVR, ESV measured by any imaging technique underestimates RV volume at Emax, but this error is minimised at high PVR. On the other hand, the pressure method assuming mPAP close to RV pressure at Emax overestimates Ees/Ea, so that sRVP to estimate ESP may be preferable. However, in the present study, while the pressure method with sRVP instead of mPAP led to slightly lower Ees and Ees/Ea, as expected (figure 6) this was not associated with an improved prognostic capability.

Recent studies focusing on RVEF as a measure of RV adaptation to afterload in patients with PAH disclosed the importance of changes over time along with the natural evolution of the disease and pharmacological interventions. However, RVEF is a markedly load-dependent measure of Ees/Ea. Whereas, SV/ESV should be more load-independent than RVEF because ESV changes proportionally less than EDV at any given change in venous return. Due to this, SV/ESV is a surrogate of the RV-arterial coupling ratio and why the optimal measurement of maximal systolic Ea relies on a family of pressure-volume loops at variable venous return. How changes over time of more robust measurements of RV-arterial coupling improve outcome prediction in severe PH remains to be explored.

Diastolic function is defined by a diastolic elastance curve determined by a family of pressure-volume loops at variable loading. Several formulas have been proposed. Most recently, Rain et al reported on 21 patients with PAH in whom RV diastolic stiffness was estimated by fitting a non-linear exponential curve through the diastolic pressure-volume relationships, with the formula P=α(eβV−1). In that study, the diastolic stiffness constant β was closely associated with disease severity. The pathogenesis of RV diastolic dysfunction was related to...
increased RV collagen content (ie, fibrosis) and stiffness of the RV sarcomeres, in turn due to reduced phosphorylation of titin, a key protein regulating myocyte stiffness. Accordingly, we calculated $\beta$ via this method and found it to be predictive of transplant-free survival, though not independently.

In the present study, the Ees/Ea ratio and $\beta$ were highly correlated. This is not surprising as cardiomyocyte relaxation is dependent on afterload, calcium handling and contractile protein function, all affected by systolic functional adaptation to loading conditions, so that ventricular relaxation can be considered as an integral part of a continuous contraction-relaxation cycle of the heart.

There are several limitations to the present findings. The first is the small size of the patient population and inhomogeneity of severity and type of PH which precluded valid subgroup analyses. Therefore, it is not known if the present study’s conclusions are generally applicable. The second is that only prediction for initial measurements was investigated, not changes with progression of the disease which will hopefully be addressed in future studies. The third is that SV was measured as the difference between EDV and ESV which could have caused underestimation of its true value in case of tricuspid regurgitation. However, SV calculated from CO and HR was 72 ±29 mL, and the difference between EDV and ESV 76±28 mL, indicating absence of significant tricuspid regurgitation in the present study population. Additionally, prospective ECG gating during CMR could have missed true EDV and underestimate EDV. The fourth is in the absence of a true gold standard for RV-arterial coupling, based on the measurements of Emax, Ea and $\beta$ on families of pressure volume loops generated at variable venous return. While the Valsalva manoeuvre has been reported for this purpose, this has only been in one study and not found to be generally feasible. On the other hand, inserting an inferior vena cava balloon to manipulate venous return was felt to be excessively invasive and, therefore, ethically problematic.

In conclusion, this study shows prognostic relevance of RV-arterial coupling evaluated by the imaging of ESV and EDV in patients referred for PH. This will require confirmation in larger PH patient populations.

**Figure 6** Pressure-volume loops from instantaneous RV pressure and pulmonary artery Doppler flow (A) compared to the simplified pressure-volume loops with peak systolic RV pressure (sRVP) to estimate end-systolic pressure (B) for three subjects. The corresponding RV pressure curves are also displayed (left, inset). Slopes of end-systolic elastance (Ees) in A were 0.91 mm Hg/mL (subject A), 1.02 mm Hg/mL (subject B), and 1.21 mm Hg/mL (subject C). Slopes of Ees in B were respectively of 0.83 mm Hg/mL, 1.19 mm Hg/mL, and C: 1.53 mm Hg/mL. Slopes of Ees with mean pulmonary artery pressure (mPAP) instead of sRVP (not shown) were respectively of 1.03 mm Hg/mL, 1.46 mm Hg/mL and 1.93 mm Hg/mL.

**Key messages**

**What is already known on this subject?**
RV function correlates with prognosis in severe pulmonary hypertension, but which metric of it is most clinically relevant is still uncertain. Clinical methods to estimate RV function from simplified pressure volume loops correlate with disease severity, but the clinical relevance has not been assessed.

**What might this study add?**
This study investigated the ability of the RV-pulmonary arterial coupling parameters stroke volume/end-systolic volume ratio (SV/ESV), a RV maximum isovolumetric pressure/mean pulmonary artery pressure ratio, EF, and by diastolic function of the RV to predict clinical outcomes. The adequacy of RV functional adaptation to afterload in patients referred for pulmonary hypertension predicts survival, and can be evaluated using imaging with the ratio of SV/ESV.

**How might this impact on clinical practice?**
The ability to relate quantitative metrics of RV function in pulmonary hypertension to clinical outcomes can provide a powerful tool for management. Such metrics could also be used in the future as surrogate endpoints for outcomes and evaluation of response to therapies.

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Contributors MAS, MAM, JL and HCC contributed to the conception and design of the study. CD, VK and JL contributed to the analysis and interpretation of the CT data. CB contributed to the analysis and sand interpretation of clinical data. RR, RN, MRP, MAM, HCC and MAS contributed to the analysis and interpretation of data. RR, RN, MRP, JL, HCC and MAS drafted the manuscript and all authors contributed to revising it critically. All authors have read and had final approval of the manuscript.

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REFERENCES


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