Introduction

Pulmonary hypertension (PH) is associated with high morbidity and mortality. It is a disease that results in high pressure load onto the right ventricle (RV). When the RV is faced with high afterload, it hypertrophies in response to the high pressure, remodels and eventually dilates (1,2). RV failure is the most common cause of death in pulmonary arterial hypertension (PAH) and occurs when the RV can no longer adapt to the afterload (1). The definitive diagnosis of PH is made by cardiac catheterization. However, the cost and risks associated with this invasive procedure have pushed clinicians to look for other means of evaluating the RV-pulmonary artery axis that is the basis of PH pathophysiology. The result is an impressive body of literature exploring the use of non-invasive imaging in the care of PH patients. Noninvasive imaging in PH is indeed critical in the diagnosis and follow up care, and part of the guidelines recommendation for pediatric and adult PH evaluation (3-7). These include established and novel modalities: two-dimensional (2D) and three-dimensional
(3D) echocardiography for evaluation of morphology and functional assessment, cardiac magnetic resonance imaging (MRI) to quantify volumes and flows as well as to characterize the myocardium, and cardiac computed tomography (CT) for evaluation of the lung parenchyma and vasculature to rule out treatable causes of PH. This review is an update of the latest advancement in the noninvasive imaging evaluation of pediatric and adult PH.

**Echocardiography**

Transthoracic echocardiography (TTE) is the noninvasive modality of choice for the initial screening of suspected PH. The optimal scaling parameters for RV metrics that would provide body size independence has not been well established across the spectrum of pediatric and adult patients. For this reason, the assessment of pediatric PH can have reference ranges different from those in adults, which may affect the accuracy of diagnosis in children with PH (8-10). Here, we combine established key parameters with newly developed TTE variables and modalities in adult and pediatric patients to assess potential RV dysfunction in PH. Conventional TTE imaging includes hemodynamic measurements and flow assessment by using Doppler echocardiography, evaluation of RV and left ventricular (LV) function and anatomy (11-14). Advanced TTE imaging includes multiparametric approaches, analysis of RV size and function using 2D and 3D echocardiography, RV to pulmonary arterial (PA) coupling ratio and myocardial mechanics (15-17). Standard evaluation of the LV (including end-systolic and diastolic LV eccentricity index, end-systolic RV/LV diameter ratio, classical indicators of diastolic LV dysfunction) is indispensable for complete and thorough evaluation by TTE since RV dysfunction and severe PH with systemic/supra-systemic RV pressure frequently affect diastolic and systolic LV function.

**Standard imaging**

**Estimation of pulmonary artery pressure (PAP)**

The estimation of systolic PAP (PASP) is based on the peak velocity of the tricuspid regurgitation jet (TRV) usually best obtained in a modified apical view. The simplified Bernoulli equation (PASP = 4 x v^2), using continuous wave (CW)-Doppler to assess velocity within the TRV, describes the relationship of TR and right ventricular systolic pressure (RVSP) as a surrogate of PASP in the absence of an outflow tract obstruction or pulmonary artery (PA) stenosis (18). Mean right atrial pressure (RAP) is added to this value to arrive at an estimated RVSP. In adults, TRV values >3.4 m/s, usually correspond to a PASP >50 mmHg at rest, increasing the likelihood of PH. However, one must bear in mind that in cases with severe TR, PASP may be underestimated (19,20).

The mean PAP (mPAP) can be calculated from the PASP (~ RVSP) as the mPAP = 0.61 x PASP + 2 mmHg, a formula that provides a great coefficient of determination in the pulmonary system (21,22). This formula permits the use of Doppler measurements in screening for PH, applying the accepted thresholds of mPAP of 25 mmHg or more recently suggested mPAP >20 mmHg and a pulmonary vascular resistance (PVR) index >3.0 Wood Units m^-2 (7).

If pulmonary regurgitation (PR) can be interrogated with CW-Doppler in the parasternal short axis (PSAX) view, mPAP and end-diastolic PAP can be estimated from the maximum (early diastolic) and minimum (end-diastolic) PR velocity, respectively, using the simplified Bernoulli equation and taking into account mean RAP. In diagnosis and follow-up of PH, mPAP and end-diastolic PAP should be routinely used, especially when TRV is unreliable (23).

**RV longitudinal systolic function**

Reflecting the longitudinal excursion of the tricuspid annulus towards the apex, tricuspid annular plane systolic excursion (TAPSE) is typically measured with M-mode in the apical 4-chamber view (24) (Figure 1A). Using the TAPSE as a surrogate measure of systolic RV function one has to bear in mind that the TAPSE takes into account neither segmental RV dysfunction/contractility nor radial systolic function, which usually contributes significantly to RV ejection in the setting of severe RV hypertrophy (13). Physiologically, TAPSE values increase from preterm infants to healthy adolescents with age and growth. Reference values for TAPSE are available for adults and pediatric patients (25-27). A reduced TAPSE marks abnormal longitudinal function of the RV with high specificity. For every 1mm decrease in TAPSE, the unadjusted risk of death has been shown to increase by 17% in adult patients with PH (28). In children and adolescents with PH, TAPSE values diminish (z-score <−2) with increasing age compared with those in healthy age-matched controls which may indicate a decline of RV systolic function in PH due to permanent severe RV pressure overload (25,29).

**Tissue Doppler velocities**

Tissue Doppler imaging (TDI) allows a quantitative
assessment of RV systolic and diastolic function by measurement of myocardial velocities (Figure 1B). As tissue velocities vary with age and heart rate, different reference values must be applied for adult and pediatric patients (30). Various studies revealed that in patients with PH, systolic (S’) and early diastolic (E’) velocities at the tricuspid, septal, and LV wall are lower than in controls, highlighting the involvement of biventricular systolic and diastolic function (31-33). Cumulative event-free survival rate is suggested to be lower when tricuspid E’ is <8 cm/s (33). TDI is independent of chamber geometry which permits its use in any chamber configuration. In pediatric PH patients
associated with congenital heart disease, TDI measurements were able to demonstrate that the tricuspid annular peak S’ is substantially impaired, with S’ value continually decreasing with longer duration of pediatric PH (32). Of note the integral of the S wave yield a metric of TAPSE derived by the TDI.

**Presence of pericardial effusion (PE)**

In adult PAH patients, presence of PE even in small amounts applies as prognostic factor and was associated with poorer survival compared to patients without PE (34). Admittedly, in another study, the appearance of a new moderate or larger PE was associated with increased mortality in adult PH patients, whereas these findings could not have been determined in patients with only a small PE (35). In children, however, the prognostic meaning of PE in pediatric PH is unclear at this time due to lack of sufficient data. Furthermore, the definition of a significant PE in children remains controversial.

**RV fractional area change (FAC)**

The RV FAC is a measure for evaluation of RV contraction and is calculated as (diastolic RV area – systolic RV area)/diastolic RV area. RVFAC encompasses both longitudinal and radial RV function from the base to the apex although it has the disadvantage of being less reproducible than TAPSE (28). In the context of RV enlargement when incomplete depiction of the RV cavity is common, RV FAC analysis will be associated with higher inter- and intraobserver variability (28). In children, a correlation between RV FAC and indexed RV stroke work and TAPSE has been reported (36) indicating the meaningfulness of an inclusion of RV FAC analysis also in pediatric PH patients. However, one must take into account that RV FAC is dependent on loading conditions as is TAPSE or longitudinal strain (36).

**Left ventricle (LV)**

The elevated RV pressure, as part of the pathophysiology in PH, leads to flattening of the interventricular septum. Consequently, the LV appears D-shaped and is more prominent in systole (37). In PH, a reduction in LV systolic as well as in diastolic volumes can be observed, whereas the global systolic function is usually preserved when measured by ejection fraction (EF) (6,8,37). LV systolic function is often impaired when measured by LV longitudinal strain. Although not the focus of this review, evaluation of the LV, including LV systolic function (LV EF, LV fractional shortening), longitudinal systolic LV function (mitral annular plane systolic excursion), and inflow and outflow Doppler of all valves, should be an integral part of the echocardiographic assessment of PH, to assess decremental RV-LV interactions (38). Diminished LV torsion is associated with worsening RV function (39).

**LV diastolic function**

In pediatric PH patients, the LV function can be impaired furthermore through RV/LV interdependence (31,44). Adult guidelines, published by the American Society of Echocardiography (ASE), suggest for evaluating LV diastolic dysfunction a cutoff value of E/e’ ratio of >14 to potentially predict elevated left atrial pressures (45). Importantly, E/e’ ratio appears not reliable in pediatric patients with LV diastolic dysfunction. For instance, approximately 50% of children with a cardiomyopathy present with E/e’ values within normal age adapted reference ranges (46). In the future, the diagnosis of LV diastolic dysfunction will likely be improved by the development of age specific thresholds for E/e’ ratios.

**PA acceleration time**

The PA forward flow velocity profile, obtained in the right ventricular outflow tract (RVOT) with PW Doppler just proximal to the pulmonary valve, can be used to determine the PA acceleration time (PAAT) (Figure 1D). PAAT is the interval in milliseconds from the onset of ejection to peak flow velocity and is used for the estimation of RV pressure and PASP. Of note, the PAAT can be measured in 99% of patients, even when the TRV CW-Doppler
envelope is not available or incomplete (47). A linear inverse relationship between PAAT and mPAP exists in adult patients with PAAT <100 ms (48). A PAAT value of <100 msec is considered abnormal. Due to the fact that PAAT varies with heart rate, pediatric reference values and z-scores were established and could be used to identify PH children (20,49). It has also been demonstrated that similar to PAAT, the RV ejection time (RVET) is reduced in adults and in children with PH (50). A ratio of PAAT/RVET is less dependent on age, body surface area (BSA) and heart rate. Therefore, PAAT/RVET is suggested to be meaningful in pediatric patients (50,51). Furthermore, bronchopulmonary dysplasia and PH were described to have negative impact on PAAT measures, suggesting that PAAT and PAAT/RVET ratio can be used as complementary modalities to assess pulmonary hemodynamics in neonates (52).

Advanced parameters

RV base/apex ratio

The RV end-systolic base/apex (RVES b/a) ratio is a simple 2D index that takes into account the specific morphologic features of the RV (Figure 2A). It is considered of clinical value for assessment of PH in adults as well as in pediatric patients (53,54). In the four-chamber view, the RV internal diameter at the base (RVES b) is measured just apical to the tricuspid annulus and the apical internal diameter of the RV (RVES a) is measured at the level of the distal end of the moderator band. Both diameters are determined in end-systole. In children and in adults PH, the RVES b/a ratio was found to be lower than in healthy (age-matched) control subjects, mainly due to dilation of the RV base and apex (53,54).

Systolic-to-diastolic duration ratio

Patients with significant PH have prolonged contraction of the RV, despite a shorter RV ejection time (55). This permits the use of systolic (S) to diastolic (D) duration ratio as an indicator of RV dysfunction (Figure 2B). In patients with PH, the Doppler-derived S/D duration ratio is higher than in healthy controls, especially as heart rates increase (55). An S/D duration ratio >1.4 is inversely correlated with survival in pediatric PH (56). The S/D duration ratio is independent of heart size, making this parameter useful in growing children. The S/D duration ratio, however, is dependent on a clearly defined onset and ending of TR Doppler tracings, so that Doppler derivation could be more reproducible (57).

Right atrial (RA) function

Assessment of the RA function can be highly relevant in PH patients. Adults with PH who lost RA function due to atrial fibrillation, have been shown to deteriorate clinically, with poor outcomes (58). An RA active emptying fraction ≥60% and a RV FAC <25% are good predictors of clinical
worsening in pediatric PH (59). In pediatric PH patients, changes in RV diastolic function reflect worsening RA reservoir and conduit function (60). Furthermore, RA deformation indices have been shown to correlate with functional capacity, invasive hemodynamic parameters, and associated with poor outcomes in children with PH (60).

**RV enlargement**

In patients with PH, RV dilation affecting the interventricular septum leads to septal shift toward the LV in systole. In severe PH, progressive RV pressure overload results in RV failure and is related to underfilling and dysfunction of the compressed LV (61-63). RV/LV ratios measured in the PSAX were demonstrated as meaningfully higher in adult and pediatric patients with PH than in controls and correlated well with invasive hemodynamic measures (Figure 1C) (62,64). Furthermore, an RV/LV ratio in end-systole >1 is associated with adverse clinical events (64,65), highlighting the importance of ventricular interdependence in pediatric and adult PH.

**Subcostal TAPSE**

Reference values and z-scores for subcostal TAPSE (S-TAPSE) were recently provided for adult and pediatric patients (66,67). Like TAPSE values, S-TAPSE values at different ages are correlated with systolic RV function parameters. Especially in critically ill patients following cardiac surgery, evaluation of tricuspid annulus motion is often hampered in the four-chamber view due to wound dressings. In such settings, TAPSE can be determined from a subcostal view and may assist in identifying patients with reduced RV function (66,67).

**3D-echocardiography imaging**

Due to its complex geometric shape, capturing of the inflow and outflow tracts of the RV in the same time acquisition is nearly impossible using 2D echocardiography. 3D-echocardiography (Figure 3) has proved to be a feasible complementary tool in quantifying RV function, validated with cardiac MRI in adult and pediatric patients (16,17,68,69). Normal values of RV volumes and function have been reported in adults and children (69,70). PH patients usually have larger RV volumes with lower EF than controls (17,71). Furthermore, PH groups vary in RV dysfunction severity: RV functional indices were worse in idiopathic PAH patients than in PAH-congenital heart disease patients (17). The 3D RV EF in pediatric

---

*Figure 3* Three-dimensional echocardiography of right ventricular volumes and function in pulmonary hypertension patient.
PH has been correlated with hemodynamic parameters determined invasively, as well as biomarkers of severity (16). The investigation of RV arterial coupling ratio derived from 3D echocardiography by using stroke volume (SV) to end-systolic volume (ESV) in children with PH demonstrated that SV/ESV ratio is decreased in children with PH compared with controls. Furthermore, SV/ESV ratio correlated with RV strain and seemed to be a strong predictor of adverse clinical events in pediatric PH (72).

**RV strain and strain measurements**

Strain is a measure of myocardial deformation whereas strain rate is defined as rate of deformation over time (13). Deformation imaging provides unique information on regional and global RV function. Reference values of adult and pediatric RV strain measurements are available (73-75). RV global longitudinal systolic strain and strain rate are substantially lower in adults with PH than in controls (76). Furthermore, RV strain provides incremental prognostic value over echocardiographic variables and could have been demonstrated as strong predictor of survival in PH (77). While on therapy, an individual improvement of >-5% in absolute value in RV systolic strain at follow-up correlates with better pulmonary hemodynamics, improved clinical status, and less evidence of RV failure (78). Decreased longitudinal deformation, decreased or absent transverse shortening, and post-systolic shortening demonstrated RV dysfunction in children with IPAH (79). In a five year follow-up study, RV strain, as well as TAPSE measurements, was found to be worse in non-survivors compared to survivors at baseline and over 5 years (80). However, the differences observed in TAPSE between survivors and non-survivors were not as divergent as between the RV strain measurements (81). After initiation of prostacyclin analogues in pediatric PH patients, an early and sustained improvement in RV strain measures have been observed, indicating that RV strain may be sensitive marker of RV function in this population (82). Besides 2D strain variables, 3D strain parameters could have been demonstrated to show better association to RV failure hemodynamics than echocardiographic measurements in adults as well as in children (17,80). The clinical application of 3D strain needs further research.

**Exercise stress echocardiography**

Exercise stress echocardiography is no longer part of the routine diagnosis of PH (5). A disease entity of exercise PH is not recommended or used due to the lack of reliable data that define which levels of exercise-induced changes in pulmonary artery pressures or PVR are prognostic (83). There are no normal values established for pulmonary artery pressures during exercise. Therefore, it is difficult to determine if the rise in pulmonary artery pressure during normal exercise is related to physiologic increase or a true disorder of PVR. An increase in pulmonary artery pressure may occur during normal exercise without an increase in PVR (84,85). One study has shown that RV dimension is increased in adult PH patients compared to controls during exercise stress echocardiography (86). Recent studies have suggested that mean pulmonary artery pressure rises by ≥1 mmHg per liter of cardiac output in normal subjects, but pulmonary vascular disease patients have a rise of ≥3 mmHg per liter of cardiac output (7). More research is needed for exercise stress echocardiography in PH as a diagnostic tool in adult and pediatric population. The use of pulmonary arterial pressure-cardiac output output slope will likely be needed to better define PH with exercise. This needs at least three time points and assessment of both evaluation of RVSP and estimated cardiac output at rest, at peak, and in recovery.

In summary, echocardiography remains the first line imaging modality for the diagnosis and evaluation of the PH patient. 2D images measure right heart size, diastolic and systolic function, as well as give insight into the relative pressures of the right heart compared to the left. The relative ease of use and low cost make echocardiography an indispensable tool for clinical care as well as research. Often, a combination of parameters can comprehensively reflect the patient status. For example, FAC, LV eccentricity index, and RA and right ventricular sizes were all shown to improve in a study of patients initially diagnosed with severe PH receiving aggressive triple therapy (87). Color and spectral Doppler are used to access flow as a reflection of pressure and resistance. Advances in 3D echocardiography and strain will only make echocardiography more accurate and be able to pick up early changes before overt clinical deterioration. Limitation of echocardiography includes patients with poor acoustic windows in patients with lung disease.

**Magnetic resonance imaging**

The American College of Cardiology Foundation/American Heart Association (88) and the European Paediatric Pulmonary Vascular Disease Network (89) recommend both MRI and CT for prognostic information about the RV
and lung parenchyma in the care of patients at baseline and follow-up. MRI has several critical advantages over other imaging modalities. Images by MRI offer high temporal and spatial resolution without exposure to ionizing radiation (Figure 4). Multiple studies have reinforced the high reproducibility of MRI-derived ventricular measures, making it the reference standard for volumetric and functional measures. Low cardiac variability in CMR imaging in PAH allows for a strong detection of clinically relevant changes with a small trial sample size, which is a critical point for the study of a relatively rare disease (90).

In addition, the ability to cross reference flow measures, including SV and cardiac output, with multiple approaches makes MRI unique. Especially in congenital heart disease in which patients may have chest wall scarring from repeated sternotomy or in adults with large body habitus, the anterior RV may be difficult to clearly delineate by echocardiography. MRI is not limited by either. The complex geometry of the RV makes straight-forward, easy mathematical approaches, as is done in elliptical-shaped LV, impossible. MRI makes no assumption about ventricular geometry. A number of conditions may preclude subjects from undergoing MRI. Pacemakers and ICDs should be fully evaluated for MRI compatibility. Metal artifacts can notoriously obscure imaging and should be considered. Claustrophobia, which is estimated to be 4% of subjects undergoing MRI, may not be previously recognized by the patient. The presence of arrhythmia may also make imaging difficult. MRI based approach also has longer scanning time and requires multiple breath-holds. MRI has relatively higher costs compare to echo. Finally, investment in proper training of MRI technologists and post-processing analysts is critical.

Currently, standard of care is using cardiac catheterization to establish PH diagnosis and hemodynamics, including PVR and mPAP. It is important to point out that cardiac catheterization data focuses on only the vasculature, while MRI has the ability to establish data on both the ventricle and the vasculature, and thus, a more comprehensive evaluation of the pathophysiology of PH. Over the past decade, growing literature supports the use of RV parameters in combination with vascular parameters to be more predictive of outcomes in PH than PVR. The first report to show this included 64 adult PH patients and found that progressive RV dilation and decrease of LV end-diastolic volume at 1 year follow-up were the strongest predictors of mortality (91). Soon after this, multiple reports showed similar findings, with measures of volume, EF, and SV shown to be powerful predictors of survival in PH (92-95). RVEF has also gained attention. A critical paper by van de Veerendonk et al. followed 110 patients with PH, acquiring baseline and 12 months CMR and cardiac catheterization. The authors found that patients with low RVEF had significantly decreased survival compared to higher RVEF independent of the PVR. Twenty-five percent of PH patients with reduced PVR over a median 59-month follow-up still had deterioration of RV function. In fact, comparison of survivors and non-survivors showed no change in PVR, but a clear deterioration of RVEF in the non-survivors compared to survivors (95). Similarly, in children, in the largest published cohort, Moledina showed prognostic potential of RV and LV volumetric measures for predicting death or transplant (94). Thus, this shifted the conventional paradigm from relying on a single PVR to understanding the interplay of the RV-PA axis.

**Ventricular vascular coupling (VVC)**

In a healthy cardiovascular state, the ventricle adapts to changes in the arterial afterload by altering contractility in order to maintain adequate SV at the most efficient myocardial work. Response of the RV initially to rising afterload is increasing hypertrophy. At the point when this is no longer adequate, the RV becomes increasingly dilated, thus increasing diastolic volume to maintain SV. Heart rate will also increase. Both of these changes shift the system to higher myocardial consumption. Uncoupling occurs when myocardial adaptations are no longer sufficient to maintain cardiac output (1). The interaction between the RV and
the PA can be conceptualized by the VVC ratio, with the numerator being the ventricular contractility (E\(_\text{c}\)) and the denominator being the arterial elastance (E\(_\text{a}\)). Arterial elastance (E\(_\text{a}\)) represents vascular afterload. VVCR (as E\(_\text{a}/E\_\text{c}\)) is theoretically optimal at 1.5–2, but in progressive load-increasing diseases associated with failing ventricular function such as PAH, VVCR eventually decreases. VVCR is thought to decrease little in early stages of load-increasing disease such as PH, as SV is maintained by augmented RV contractility. In late stages of disease, the RV fails to adapt and a further decrease in VVCR is expected. Conventionally, VVCR is measured by cardiac catheterization, with progressively decreasing preload achieved by obstructing flow through the inferior vena cava—a time-consuming and invasive methodology, not to mention impossible in very young children secondary to the size of the catheter used. Sanz showed that VVCR can be estimated using only MRI data (VVCR\(_\text{CMR}\)) and is calculated as SV/ESV (96), and correlates to VVCR derived by the catheterization-based single beat (VVCR\(_\text{RHC}\)) method (97,98). It is important to understand that this estimation assumes zero intercept of the ventricular elastance curve which limits its accuracy. Despite this, VVCR\(_\text{CMR}\) remains a valuable tool, especially when followed longitudinally. In adults and children, both VVCR\(_\text{CMR}\) and VVCR\(_\text{RHC}\) correlated well with conventional functional and hemodynamic parameters of disease severity (96,98), as well as clinical deterioration (99) and transplant-free survival (100). Impaired right ventricular-vascular coupling in PH is likely to be a gradual and progressive feature and serial imaging could guide the clinician in this process.

**Myocardial strain**

MRI can measure myocardial strain by two technique: tissue tagging and feature tracking. The first uses a pre-pulse to nullify signals within defined parallel lines or grids. These “tags” are then followed throughout the cardiac cycle, with each tag representing small regions of the myocardium. Feature tracking is a more novel approach, and uses steady-state free precession sequences images in multiple planes to determine deformation in the longitudinal, circumferential, and radial directions by following the movement of the myocardial border over the cardiac cycle (101). This method has been validated against tagged MRI (101). In a study looking at feature tracking of the RV, RV strain correlated with RV E\(_\text{a}/E\_\text{c}\) and afterload (E\(_\text{a}\)), but not with contractility (E\(_\text{c}\)). Multivariate logistic binary regression analysis identified long-axis RV radial strain as being able to identify RV-arterial uncoupling (102). In exercise states, PH patients showed less reserve than controls, as reflected by displaying significant decrease in longitudinal RV strain despite normal EF (103). Strain has been used in several studies of therapeutic response (104,105).

Ventricular mechanics such as torsion and torsion rate can also be determined from strain analysis. Dysfunction of the RV compromises both diastolic and systolic function of the LV through ventricular-ventricular interaction. In PH patients, it is not surprising that LV torsion and torsion rate are decreased as RV EF decreases (39). In PH patients compared to normal subjects, smaller changes in RVEF lead to larger compromise of LV torsion rate (39). Hence, accurate diagnosis of the function and anatomy of both ventricles is essential in the pathophysiology of PH.

**Tissue characterization**

One of the biggest advantages of MRI is the ability to provide details about the myocardium, by bringing out differences in signals for varying tissue composition to identify myocardial pathologies. T1 relaxation time is the time for a nuclear spin to return to 63% of its magnetization in the longitudinal plane. T1-weighted imaging is used to identify scar or fatty infiltration of the myocardium. T2 reflects the presence of water, and tissue edema (106). Using T1 and T2 tissue characterization with CMR permits investigation beyond the cardiomyocyte to understanding histopathologic changes—a process not available with other imaging modalities. The failing RV in PH is associated with myocardial edema, as shown in rodents by histologic confirmation, as well as T2-weighted MRI imaging.

In recent years, the development of new MR sequences enabled the quantitative measurement of T1 and T2 magnetization properties, known as T1 and T2 mapping, which have allowed for further characterization of changes in the underlying myocardial substrate. The diffuse nature of the fibrosis, for example, makes imaging by T1-weighted approaches impractical since T1-weighted imaging identified fibrosis as a hyperintense signal compared to myocardium. T1 mapping however can identify diffuse fibrosis, in which collagen has accumulated within the myocardium leading to diastolic dysfunction. In PH patients, native T1 (performed pre-contrast) and T2 weighted images are increased in the RV insertion points, with abnormal T2 values also seen in the RV free wall (107). Extracellular volume (ECV) fraction quantification can be
determined by measuring T1 mapping before and after contrast. In PH porcine models, ECV was higher in the RV insertion point and correlated with PVR, mPAP, RVEF, and PA area (108). Studies in humans concur (109,110), with elevated native T1 and ECV seen even before the presence of late gadolinium enhancement (LGE) (110).

**LGE**

LGE is a technique by which images are acquired 5–10 minutes following gadolinium contrast administration. The presence of gadolinium within the myocardium after this time represents a breakdown of cell membrane, allowing intercalation of contrast into extracellular spaces. This is seen with fibrosis and ischemic changes. Hyperenhancement is seen in 69–83% of PH patients (111,112), and often seen in the RV insertion points, correlating to regions of highest stress. However, its prognostic value remains uncertain. The literature is conflicting as to the use of LGE in PH. The presence of LGE has been linked to more severe disease state (113), longer disease duration (114), increased RV dilation, and to worsening RV function (112). Less clear, is the relationship between LGE and RV EF, with reports indicating polar opposite findings (113-115). Two large studies looked at 162 patients (111) and 124 patients (116), neither of which found correlation with mortality or morbidity including hospitalization RV failure, initiation of parenteral prostacyclin, sustained ventricular arrhythmia, or referral for lung transplant. Differences between studies may reflect a difference in etiologic make-up of study populations and distinction between LGE in the insertion points and extended involvement into the interventricular septum.

**Vascular imaging**

CMR is now emerging as a technique for evaluating vascular function as well, including flow patterns, flow velocity, wall shear stress (WSS) (117), and recently, vorticity and helicity by 4-dimensional (4D) velocity-encoded MRI (118) (Figure 5). It can accurately and non-invasively quantify aortic velocity fields, allowing for the calculation of heterogeneously evolving WSS, as well as provide information on flow, turbulent kinetic energy, and vorticity (119). MRI also may demonstrate developmental vascular changes in PAH, which is particularly important in pediatric PH, in which disease progression is occurring simultaneously with development (120). Cross section analysis of the proximal pulmonary arteries can give accurate measures of arterial stiffness. These measures include relative area change, defined as (\(\text{Area}_{\text{systole}} - \text{Area}_{\text{diastole}}\)/\(\text{Area}_{\text{diastole}}\), compliance, and distensibility. Measures of arterial stiffness are predictive of mortality in PH (121). In fact, RAC was found to be the strongest predictor of mortality independent of pulse pressure.

Phase contrast (PC) imaging uses phase shift from flowing blood to quantitate flow and its derivatives, so that volume, velocity, and direction of flow can be determined. Pulse wave velocity (PWV), another measure of stiffness, can be measured from PC imaging from the flow-area method \((dQ/dA)\) in which flow and area change waveforms (122). PWV in PH is a potential predictor of hospitalizations, increasing symptoms, and escalation of therapy (123).

WSS, an important hemodynamic component affecting mechanotransduction (124), is the shear that is seen by the endothelium caused by moving blood and can be measured from PC MRI (123). Acute and chronic changes in WSS associated with high flow pulsatility and disturbed flow has been shown to alter downstream signalling cascade, and can lead to vasodilation or vasoconstriction, endothelium proliferation, and alter coagulation cascade (125,126). Indeed, WSS in the main and branch pulmonary arteries is decreased in PH, in which flow derangements including vortex formation have been shown by time resolved 3D (127) and 4D flow MRI (128). This correlates with increasing arterial stiffness and decreased RV function (129). In addition, oscillatory shear index (OSI), the deviation of shear stress from its principle direction, can be derived from the WSS waveform. High OSI value is seen with compromised endothelial function and atherosclerotic changes (130), and is abnormal in the pulmonary arteries of PH.

---

© Cardiovascular Diagnosis and Therapy. All rights reserved.
Like many cardiovascular diseases, flow alterations play a critical role in the progression of PH. Abnormal cardiac vorticity, for example, is involved in the early development of cardiac dysfunction (131). Thus far, flow has been understudied due to limitations of imaging modality, until the development of 4-D flow MRI. 4D MRI allows the acquisition of flow data in all three dimensions of space over time (132). This unique feature allows the visualization and quantification of complex flows that is critical to normal and diseased physiology with high spatial resolution. For example, the normal vortex formation around the mitral anterior leaflet during early filling is lost in PH. It is this vortex in the healthy state that places the volume of blood directly in the LV outflow tract to be ejected during the next systolic phase. Therefore, this loss may compromise the efficiency of the system. The loss of this vortex is hypothesized to be due to the septal shift and alteration in septal geometry in PH (133). In addition, abnormal vorticity (local spins of fluid) in the inflow of both the RV and LV is seen in PH (134,135). In COPD patients with normal LV volumetric measures and normal diastolic Doppler indices, vorticity is already reduced; thus, may be a sensitive marker for early diastolic dysfunction (134).

Pressure estimation by Doppler using the Bernoulli’s equation gives proof of concept to the ability of flow to estimate pressure. 4D MRI has not only shown decreased vorticity in the proximal pulmonary arteries, but multi-regression models using 4D data in PH by Kheyfets et al. showed that vorticity may estimate PVR using a function of vorticity in the MPA, RPA, cardiac output, and the relative area change in the MPA. The equation predicts 94% of the variability in PVR (136). A meta-analysis covering 15 studies that analyzed the estimation of PVR by MRI showed no statistically significant differences in PVR estimated by MRI and that derived by catheterization (137). Of note, this meta-analysis included multiple MRI approaches with no standard protocol. Much work is still needed, but MRI is a promising tool for providing accurate data about the myocardium and vasculature in PH patients that can be followed serially.

**Exercise MRI**

The ability to perform exercise stress to MRI imaging has been a powerful advancement for its ability to clearly define myocardial borders even with a tachypneic and tachycardic patient. Exercise testing can unmask signs and symptoms, as well as limitations that are not appreciated at rest. Recently, exercise cardiovascular MRI demonstrated that low risk versus high risk PAH patients can be differentiated; that is, MRI during exercise stratifies patients into different degrees of right ventricular inotropic reserve. Patients were evaluated based on World Health Organization Functional Classification I and II. Those in class I were able to increase SV with exercise, while those in class II did not have this ability (138). Reduced SV during exercise is a determinant of increased risk of decompensation. Understanding exercise limitation could help guide therapy towards incorporating restoration of ventricular functional reserve by intensified PAH specific treatment.

**Ventricular-ventricular interaction**

Ventricular-ventricular interaction—the phenomenon in which the performance of each ventricle affects the other—has been well described (139). In a study of isolate canine hearts, the left ventricle contributed to 68% of normal RV systolic pressure and 80% of pulmonary arterial flow (140). While the LV strain, particularly along the septum, is negatively affected in PH as shown by feature tracking (141), this in turn further reduces RV performance (Figure 6) (39).

MRI allows accurate assessment of SV and peak emptying rate (PER) during systole and early diastolic peak filling rate (PFR) for both the RV and LV. In a volume-time curve, PER is the maximal negative velocity during systole, and PFR is the maximal positive velocity in early diastole. Both were shown to correlate with SV. Thus, both can be additional surrogates for biventricular diastolic dysfunctions (142).

Another mechanism by which LV function is reduced in PH is the alteration of flow and blood volume in and out of the LV. As previously mentioned, in PH, the vortex along the anterior mitral leaflet is lost, which is thought to efficiently deliver a bolus of blood to the LVOT to be ejected with the next contraction (131).

**Vascular-vascular interaction**

While not often considered, the aorta is also adversely affected in PH. We showed that the presence of a dilated main pulmonary artery mechanically constrains the expansion of the adjacent ascending aorta, leading to decreased RAC. The lower RAC was not seen in the descending aorta (143). This apparent regional aortic stiffness is seen by the LV as increased afterload. Our data suggests that the degree of decreased RAC correlates with
lower L VEF and lower VVCR. This is yet another possible mechanism by which disease of the RV-PA axis affects both LV performance (Figure 6).

In summary, MRI allows high spatial resolution of the intracardiac structure and vascular involved in PH pathophysiology. MRI is not constrained by body habitus, which is a significant advantage in older patients, those with poor acoustic windows, and those with surgical scars. The unusual geometry of the RV does not preclude accurate measurements of RV volume and EF. Advances in MRI have allowed detailed assessment of flow through the heart and the vasculature and reflect vascular hemodynamics and vascular remodeling. Despite important limitations, MRI is a promising tool to comprehensively describe the entire RV-PA axis, as well as the involvement of the left heart.

**Computed tomography**

CT is important for identifying vascular, cardiac, parenchymal and mediastinal abnormalities (89). Details about pulmonary nodules, extracardiac shunts, bronchial and vascular anomalies are seen with a very short acquisition time by CT. Advances with multidetector (MDCT) have improved both spatial and temporal resolutions, allowing the development of high resolution CT applications (144). MDCT (also known as multislice CT) is an advancement from the conventional single slice CT, and has multiple rows of detectors, enabling more than one slice of data to be collected at any given time. High resolution CT without contrast is used for evaluation of lung parenchymal issues, especially in cases of PH suspicious for falling under Nice Classification Group 3. Parenchymal findings associated with different etiologies of PH include ground glass nodules seen in idiopathic PAH or pulmonary veno-occlusive disease interlobular thickening seen in left heart disease, pulmonary fibrosis, or sarcoidosis, ground glass opacities seen in pulmonary edema, interstitial lung disease, connective tissue disease. Enlarged bronchial arteries can be seen in CTEPH.
and Eisenmenger disease. Of note, bronchial artery dilation is often not presented in idiopathic PAH (145). Automated 3D volume segmentation is available with CTA can significantly decrease time for post-processing to as quick as 3 minutes (146).

Filling defects as seen in CTEPH is best seen with CT, along with associated findings of mosaic attenuation and enlarged bronchial arteries (145). CT can also identify PE and thickening seen in moderate to severe PH (147).

Limitations of MDCT include exposure to ionizing radiation and potentially nephrotoxic contrast, as well as being restricted to a low and stable heart rate for adequate acquisition of the myocardium. Imaging of the vasculature without need for myocardial imaging can be done with non-EKG gating.

Superior to other non-invasive imaging, CT scan image more proximal pulmonary arteries and veins. Therefore, details such as peripheral pruning can be seen. Mosaic attenuation is seen in PH and is a process by which there is dilation of vessels in areas of high attenuation and rarefaction of vessels in low areas of attenuation. This is thought to be due to regional differences in perfusion (148). Similarly, changes in segmental and subsegmental bronchi can be easily seen. A number of sentinel papers have shown reasonable sensitivity and specificity in using CTA for PH diagnosis by visualization and measurement of the pulmonary arteries (149,150). A mean PA diameter >29 mm has 87% sensitivity and 89% specificity for the diagnosis of PH. This specificity can reach 100% if there is also segmental artery:bronchus ratio >1:1 in 3 of 4 lobes (149). Another approach focusing on the ratio of main pulmonary artery diameter to the ascending aorta of >1 has a 96% positive predictive value (150,151).

In patients in whom MRI is not ideal, CT can serve as an alternative to volumetric measures. In a meta-analysis comparing CT to MRI for RV measurements, 19 studies were reviewed which had CT performed less than one month from MRI. There was no statistical difference between the two modalities in RV volumetric measurements and EF, albeit CT overestimated EDV and ESV, and underestimated RVEF and SV (152). In a small study of 30 patients with PAH, 6MWT had higher correlation to MRI-derived RV parameters compared to CT (153). Similarly, a study of 812 patients with PAH from connective tissue disorders underwent CT, MRI, and RHC within 48 hours. CT derives RV and LV ratio and RV thickness did not perform diagnostically as well as MRI -derived data. Furthermore, MRI parameters were more prognostic with 1-year survival compared to CT (154).

A brief mention about SPECT imaging in PH is warranted. Small studies have looked at the use of stress $^{99m}$Tc SPECT perfusion studies in assessing the RV in PH (155). The increased uptake of $^{99m}$Tc-labeled tracer is consistent with RV dilation as well as RV hypertrophy (156). An RV/LV ratio of 0.35 on non-attenuation corrected SPECT images had a sensitivity of 72% and specificity of 70% for identifying a pulmonary systolic pressure of >40 mmHg. This is drastically compromised when attenuation correction is used as there is a higher false positive rate for RV hypertrophy.

In summary, CT is critical for evaluation of the lung parenchyma and pulmonary vasculature as part of the evaluation for PH. In patients in whom MRI is contraindicated, CT can give accurate measurements of RV volume and EF. Chest CT can be completed within a matter of minutes, although the downsides include exposure to ionizing radiation and potential renal toxicity associated with iodinated dye used in CT.

**Conclusions**

Noninvasive imaging is important in the diagnosis and follow up of PH. Valuable information can be gained from these noninvasive measures to make clinical decisions and management of PH as invasive measurements pose risks for these patients. There is growing evidence that the deterioration of the RV function may be more predictive of PVR of outcomes. If this is indeed true, standard approach using cardiac catheterization is wholly inadequate to serially evaluate PH subjects. Rather, we must look towards noninvasive imaging as a safe and accurate means of providing comprehensive data of the ventricles, pulmonary and systemic circulation involved in PH progression. Further research will help define the most valuable parameters in determination of long-term prognosis.

**Acknowledgments**

**Funding:** None.

**Footnote**

**Provenance and Peer Review:** This article was commissioned by the Guest Editors (Martin Koestenberger, Harm-Jan Bogaard and Georg Hansmann) for the series “Right Ventricular Dysfunction” published in *Cardiovascular Diagnosis and Therapy*. 

© Cardiovascular Diagnosis and Therapy. All rights reserved.
Diagnosis and Therapy. The article was sent for external peer review organized by the Editor-in-Chief and the editorial office.

Conflicts of Interest: All authors have completed the ICMJE uniform disclosure forms (available at http://dx.doi.org/10.21037/cdt-20-272). The series “Right Ventricular Dysfunction” was commissioned by the editorial office without any funding or sponsorship. MK served as the unpaid Guest Editor of the series. UT reports grants from NIH NHLBI, from null, outside the submitted work. JC reports grants and other from Actelion, grants from Astra Zeneca, grants from United Therapeutics, outside the submitted work. DI reports grants and non-financial support from Actelion, grants from Bayer, grants from Glaxo Smith Kline, grants from Liquidia, grants and non-financial support from United Therapeutics, grants from Lilly, outside the submitted work. The other authors have no other conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

Open Access Statement: This is an Open Access article distributed in accordance with the Creative Commons Attribution-NonCommercial-NoDerivs 4.0 International License (CC BY-NC-ND 4.0), which permits the non-commercial replication and distribution of the article with the strict proviso that no changes or edits are made and the original work is properly cited (including links to both the formal publication through the relevant DOI and the license). See: https://creativecommons.org/licenses/by-nc-nd/4.0/.

References


Transthoracic echocardiography in the evaluation of pediatric pulmonary hypertension and ventricular dysfunction. Pulm Circ 2016;6:15-29.


36. Di Maria MV, Younoszai AK, Mertens L, et al. RV stroke work in children with pulmonary arterial hypertension: estimation based on invasive haemodynamic assessment


57. McCabe C, Vranesic II, Verdes MC, et al. Right


idiopathic pulmonary arterial hypertension vs those with pulmonary valve stenosis: Insights into mechanics of right ventricular dysfunction. Echocardiography 2017;34:888-97.


