Assessment of Right Ventricular Function in the Research Setting: Knowledge Gaps and Pathways Forward
An Official American Thoracic Society Research Statement


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Background: Right ventricular (RV) adaptation to acute and chronic pulmonary hypertensive syndromes is a significant determinant of short- and long-term outcomes. Although remarkable progress has been made in the understanding of RV function and failure since the meeting of the NIH Working Group on Cellular and Molecular Mechanisms of Right Heart Failure in 2005, significant gaps remain at many levels in the understanding of cellular and molecular mechanisms of RV responses to pressure and volume overload, in the validation of diagnostic modalities, and in the development of evidence-based therapies.

Methods: A multidisciplinary working group of 20 international experts from the American Thoracic Society Assemblies on Pulmonary Circulation and Critical Care, as well as external content experts, reviewed the literature, identified important knowledge gaps, and provided recommendations.

Results: This document reviews the knowledge in the field of RV failure, identifies and prioritizes the most pertinent research gaps, and provides a prioritized pathway for addressing these preclinical and clinical questions. The group identified knowledge gaps and research opportunities in three major topic areas: 1) optimizing the methodology to assess RV function in acute and chronic conditions in preclinical models, human studies, and clinical trials; 2) analyzing advanced RV hemodynamic parameters at rest and in response to exercise; and 3) deciphering the underlying molecular and pathogenic mechanisms of RV function and failure in diverse pulmonary hypertension syndromes.

Conclusions: This statement provides a roadmap to further advance the state of knowledge, with the ultimate goal of developing RV-targeted therapies for patients with RV failure of any etiology.

Keywords: right ventricle; pulmonary hypertension; pulmonary embolism; acute respiratory distress syndrome; pulmonary circulation

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This article has an online supplement, which is accessible from this issue’s table of contents at www.atsjournals.org.
Overview

Right ventricular (RV) function is the single most important prognostic determinant of survival in various forms of pulmonary hypertension (PH). Significant progress has been made since improving the understanding of RV function and failure was identified in 2005 as a major area of need in the field; however, many knowledge gaps remain. A better understanding of the mechanisms of RV adaptation and maladaptation to pulmonary hypertensive syndromes will advance the field toward the goals of defining physiological and pathophysiological RV responses, identifying biomarkers of RV function, and developing RV-directed therapies. This document lays the framework for addressing knowledge gaps in clinical and basic RV research and for identifying research opportunities and pathways forward.

- The task force defined RV failure (RVF) as a complex clinical syndrome characterized by insufficient delivery of blood from the RV in the setting of elevated systemic venous pressure at rest or exercise.
- RVF is a heterogeneous syndrome. Multiple RVF phenotypes exist.
- The task force identified multiple knowledge gaps and research opportunities in the field. The four major research priorities that should be addressed in the next 5 years are:
  1. Enhancing mechanistic understandings of adaptive versus maladaptive RV responses to pulmonary vascular load. This includes, but is not limited to, understanding the roles of fibrosis, angiogenesis, inflammation and metabolic shifts. Genetic and epigenetic contributions to successful adaptation need to be identified.
  2. Deriving and validating a series of plasma and/or imaging biomarkers (by echocardiography and cardiac magnetic resonance imaging) for accurate evaluation of RV function, perfusion, and RV–pulmonary vascular coupling and to serve as endpoints in clinical trials.
  3. Developing and validating novel animal models and improving existing models of RVF in the setting of increased pulmonary vascular load to provide a platform for better understanding mechanisms of RVF and for testing novel RV-directed therapies.
  4. Developing novel therapies aimed at targeting RV myocardial contractility (e.g., calcium sensitizing agents) as a critical area of need that would be expected to improve patient outcomes.
- Increased availability to and methods to study human RV tissue will be critical in addressing these research priorities and in advancing the field.

Introduction

RV failure (sometimes also referred to as “right heart failure”) is increasingly recognized in acute and chronic pulmonary hypertensive syndromes, and RV adaptation to these disease states is a significant determinant of short- and long-term outcomes (1–5). An estimated 70 million individuals in the United States may have abnormal RV function (3, 4, 6–10).

Thirteen years have passed since the initial meeting of the NIH Working Group on Cellular and Molecular Mechanisms of Right Heart Failure (1). Although remarkable progress has been made in the intervening period, significant gaps remain at many levels in the understanding of cellular and molecular mechanisms of RV responses to pressure and volume overload, in the validation of diagnostic modalities, and in the development of evidence-based therapies (1, 2, 11, 12). RV function is the single most important prognostic determinant of survival in pulmonary arterial hypertension (PAH) (3) as well as in multiple highly prevalent chronic heart and lung diseases (4, 6, 7, 10); however, no consensus exists regarding the definition of RV failure (RVF). This deficiency reflects in part the current paucity of clinically available, specific biomarkers reflecting RV function. In addition, targeted therapy for RVF remains elusive. Most importantly, factors determining transition from adaptive (or compensated) to maladaptive (or decompensated) RV remodeling and predictors of RVF remain unknown.

The purpose of this document is to:

1. Comprehensively elucidate the current understanding of RV function and its assessment in healthy individuals and those with preclinical, acute, and chronic diseases;
2. Identify major knowledge gaps; and
3. Outline specific strategies for addressing these gaps.

We sought to address these objectives across three topic domains:

Topic Domain 1
Optimizing the methodology to assess RV function in acute and chronic conditions in pre-clinical models, human studies, and clinical trials. This provides the basic framework for optimally assessing RV morphology, function, and failure in animal models of disease and in humans.
Topic Domain 2
Analyzing advanced RV hemodynamic parameters at rest and in response to exercise. Recommendations are provided for improving the understanding of RV contractile and diastolic function as well as RV–pulmonary vascular coupling in response to load and exercise in both preclinical and clinical models. This will further our understanding of RV dynamic function adaptation and functional reserve and will establish optimal measures to be used in clinical assessment of the RV.

Topic Domain 3
Deciphering the underlying molecular and pathogenic mechanisms of RV function and failure in diverse PH syndromes. Identifying pathophysiologically relevant adaptive and maladaptive processes (e.g., angiogenesis, inflammation, fibrosis) in preclinical models will set the stage for targeted therapy to be tested in clinical trials.

Findings and recommendations generated from these topic domains address the overarching questions and cross-cutting themes listed in Table 1.

The intended audience of this report includes basic, translational, and clinical researchers in the fields of pulmonary circulation, cardiovascular medicine, respiratory and critical care medicine, research funding organizations, health policy experts, and pharmaceutical and device industries.

Methods
A working group of international experts from the American Thoracic Society (ATS) Assemblies on Pulmonary Circulation and Critical Care (18 ATS members and two external content experts) was convened, which included basic, translational, and clinical researchers with expertise in a broad range of areas within adult pulmonary and critical care medicine, cardiovascular medicine, cardiothoracic surgery, physiology, and pathology. The scope of the project was reviewed and approved by the ATS. Potential conflicts of interest were disclosed and managed in accordance with the policies and procedures of the ATS. No unresolved conflict of interest was identified.

This task force met twice in person (ATS 2015 for a full-day workshop and ATS 2016 for a 2-hour summation) and multiple times in between by teleconference. Major focus areas were developed by semistructured systematic review of the literature, expert opinion, and consultation. During the initial workshop, goals and ground rules guiding the work group were established. Subjects to be discussed were suggested by the steering committee and agreed on by the group. Presentations by content experts were made in each of the three topic domains, followed by group discussion aimed at defining key areas of consensus and disagreement. Current knowledge state summaries were then developed and rigorously analyzed by the entire workgroup and in focus area subgroups. Structured knowledge gap analyses were conducted in all three topic domains.

The draft was circulated to the task force members with revisions at each step. Consensus was achieved through moderated discussion. Data cloud document management was used to coordinate version control and integrate components from the subgroups. The final report was peer reviewed, approved by the leadership of each of the sponsoring Assemblies, and ultimately approved by the ATS Board of Directors.

Definition, Development, and Staging of RVF
The task force defined RVF as a complex clinical syndrome characterized by insufficient delivery of blood from the RV and elevated systemic venous pressure at rest or exercise.

RVF occurs as a consequence of 1) alterations in preload; 2) changes in RV mechanics, lusitropy, and/or contractility; or 3) increases in afterload (Figure 1A). Similar to left ventricular (LV) failure, chronic RVF occurs in four distinct stages (Figure 1B). Some investigators use the term “RV dysfunction” to indicate structural RV changes in the absence of functional alterations. In this statement, we avoid this term, because such a constellation would qualify as “RV failure” according to the definition provided above. Although frequently progressive in character, with appropriate intervention(s), progression may stop at any of these stages or even be reversible. On the other hand, acute compensation may develop at any of the four RVF stages. Once acute RVF develops, patients may deteriorate or stabilize (Figure 1C). Several clinically relevant modifiers of RV function exist (Figure 2A). PH is a syndrome commonly associated with RVF and is a prime example of how the RV responds to chronically elevated afterload (Figure 2B). With increases in pulmonary artery pressure (PAP), the RV exhibits compensatory mechanisms that include homeometric and heterometric adaptation as well as neurohormonal activation (2, 13). Adaptive RV hypertrophy (RVH) develops. It is believed, although not fully proven, that once the RV’s compensatory mechanisms are exhausted (purportedly as a result of transition from adaptive to maladaptive RVH), RVF develops. Genetic predispositions and early life events are putative modifiers of the RV compensatory response (Figure 2B). Figure 3 highlights key aspects of RVF pathophysiology addressed in this research statement.

Topic Domain 1: Optimizing the Methodology to Assess RV Function in Acute and Chronic Conditions in Preclinical Models, Human Studies, and Clinical Trials

Normative Indices of RV Morphology and Function

Rationale: Despite the high prevalence of RVF in the United States and throughout the world (3, 4, 6–10), there are no
Figure 1. Etiologies and stages of right ventricular failure (RVF). (A) RVF occurs as a consequence of alterations in preload, changes in mechanics and/or decreases in contractility, or increases in afterload. (B) Classification of RVF according to stages of development in keeping with the recent classification of the American College of Cardiology Foundation/American Heart Association Task Force for Heart Failure (298, 299). Note that stages are not static and that stages C and D are potentially reversible with normalization or significant decrease in pulmonary vascular resistance (e.g., after pulmonary endarterectomy or lung transplantation) (300, 301). Symptomatic RVF (stage C) is usually managed pharmacologically, whereas refractory RVF (stage D) often requires specialized interventional or surgical measures. Preventive measures may be applied at any of the different stages of RVF. Decompensation may occur at any stage. (C) Cycle of acute on chronic decompensation of RVF. Acute decompensation is usually provoked by one or more precipitating factors (e.g., infection, pulmonary embolism, bleeding disorders) or by progression of the underlying disease (112). Patients frequently deteriorate and enter a vicious cycle of hypotension, ischemia, and decreased pump function or may stabilize and revert to more or less stable chronic RVF. However, mortality after hospital discharge remains high (35% at 12 mo [112]). PAH = pulmonary arterial hypertension; RHF = right heart failure.
well-established biologic or clinical determinants of RV structure and function. Thus, there remains significant scientific and clinical justification for studying the structure and function of the RV in greater detail during health. Establishing normative values will enhance our understanding of subclinical changes in disease and may allow for early/preventative treatment. Unlike the LV, the thin-walled, compliant RV has difficulty accommodating increases in afterload, and even small increments or fluctuations in afterload may lead to adverse RV sequelae (13, 14), particularly if the increase in PAP is rapid. There is, however, great variability in the clinical trajectory of patients, who often present at later stages of disease, when RVF is overt (Figure 1B). Furthermore, observations from population-based cohorts may provide insight into important mechanistic pathways generalizable to RVF, regardless of etiology. Finally, targeting patients before the development of significant RVF may be more successful than instituting therapy after advanced disease is established.

**Current state of knowledge.** An ideal RV metric would be sensitive enough to capture subtle cardiac structural changes, reliable, reproducible, inexpensive, and easily obtained. It would be validated across settings and in different populations. Echocardiography can qualitatively assess RV function and can be used to measure ventricular volumes and wall motion. However, quantification requires geometric assumptions, and these are hindered by the crescentic, thin-walled shape of the normal RV and also by subject characteristics (e.g., body habitus), which could introduce bias. In contrast, cardiac magnetic resonance imaging (CMR) is a fundamentally safe and reliable modality from which quantitative data regarding RV size, function, and remodeling are acquired rapidly with limited artifact. However, it is relatively expensive in some settings and not universally available.

Measurement of RV ejection fraction (RVEF) and RV mass and volumes by CMR is highly accurate and reliable in normal subjects and in those with RV or LV failure (15–18). Thus, CMR is the standard of measuring imaging technique for RV assessment (although echocardiography is more practical in the unstable patient because it can be performed at the bedside). RVEF has been identified as a key determinant of outcome in RVF regardless of etiology, including PH due to LV failure (group 2 PH) or advanced lung disease (group 3 PH), and PAH (4, 6, 7) (group 1 PH). RVEF predicts outcome in PAH, and changes in RVEF mediate improvements in survival with disease-specific therapy, making it currently the most reliable surrogate RV endpoint (3, 19–22). Several other imaging techniques have been shown to measure RV function and predict outcome (e.g., tricuspid annular plane systolic excursion [TAPSE], speckle-tracking for strain imaging, three-dimensional [3D] echocardiography, fluorodeoxyglucose–positron emission tomography [PET]) but are less well validated, especially in health or early disease (23–27).

**Current knowledge gaps.** Normative values for RV size and function have been derived from the MESA-RV (Multi-Ethnic Study of Atherosclerosis–Right Ventricle)
Study, the Framingham Heart Study, and other cohorts (28–35). However, these observations need to be validated in health and disease, longitudinally, and in other populations. Racial, ethnic and socioeconomic differences need further exploration (28, 36, 37). Normal values for RV function are needed for physiological modifiers of the normal RV in pregnant subjects and athletes, and the effects of processes such as puberty, menopause, and aging merit study. The temporal evolution with early life events (e.g., prematurity) also requires further study (38–40). The definition and relevance of subclinical RV changes, and their role in the natural history of RVF, are also incompletely elucidated.

Recommendations/pathway for progress. Several knowledge gaps exist, which include a better understanding of the normative indices of RV morphology and function, the impact of key demographic characteristics (age, sex, race/ethnicity, and socioeconomic status) on the healthy RV, RV development throughout the life span, subclinical RV abnormalities, cardiopulmonary interactions under stress, and, finally, the impact of biventricular interactions. Such studies should include neonates and prematurely born babies, especially because the latter are at risk for developing RV abnormalities later in life (38).

Optimization of RV Assessment in Clinical Research

Rationale. Trials to detect differences in clinical worsening or survival generally require a substantial sample size, which can be difficult to complete in rare diseases such as PAH, especially when the intervention being studied is added to background therapy. A well-chosen RV surrogate endpoint in clinical trials may facilitate trial design, and ultimately drug approval, because RVF is the leading cause of death in PAH (3). An endpoint must be both accurate and precise, as measurement error can lead to bias and, therefore, erroneous conclusions.

Current state of knowledge. A given endpoint needs to satisfy a number of criteria to serve as a surrogate for “hard” clinical outcomes before incorporation into clinical trials (41–43). It should be reliable, integral to the disease causal pathway, and targeted by the intervention of interest, and
be linked with a clinically important outcome, such as survival. Finally, a significant proportion of the treatment effect (50–75%) on clinical outcome should be explained by the effect of the intervention on the surrogate endpoint. The requirements for a disease surrogate are much more stringent than for a correlate, and in fact several long-held endpoints in PAH, such as 6-minute walk distance (6MWD) and hemodynamics, are inadequate surrogates for short-term events (44–48).

Current knowledge gaps. A number of RV endpoints have been proposed, including biomarkers (e.g., BNP [brain natriuretic peptide] and troponin) as well as metrics obtained from traditional and novel imaging techniques (23–27, 49–52). Although there is currently no consensus on how to best measure RV function in clinical trials, there is some evidence that current PAH treatments may have unique effects on the RV (criterion #2 for a surrogate; as demonstrated in a limited way with sildenafil and simvastatin and RV mass, and with bosentan and RV stroke volume [53–55]). Up-front combination therapy with ambrisentan and tadalafil reduced RV mass and NT-pro-BNP levels and increased RV contractility as measured by TAPSE or speckle-tracking echocardiography in an open-label trial in scleroderma-associated PAH (56, 57). Small observational studies demonstrated that worsening RV performance is associated with poor outcomes independent of pulmonary vascular resistance (3, 53, 58). These observations, along with those that demonstrated classic endpoints (e.g., 6MWD and hemodynamics) to be inadequate, suggest we may not fully understand how currently approved PAH treatments work (e.g., by targeting the pulmonary circulation or the RV or through other systemic effects) (59). The study of patients with PH with a severe RVF phenotype independent of the etiology (i.e., WHO groups 1–5) may allow for larger and faster trials; however, such an approach has not yet been tested. CMR is accurate, reproducible, and allows for detecting small changes in RV EF (~3%) in sample sizes that are modest. It is considered the standard of measure for RV morphology and function and has been shown to be more cost saving in PAH drug trials than echocardiography, as a result of its lower measurement variability (60). Although some investigators advocate that CMR should be routinely incorporated into clinical trials, wide-scale incorporation into trials and ultimately clinical practice may be limited in some centers by cost and technical expertise.

Recommendations/pathway for progress. Measurements of RV function should be routinely incorporated into observational studies and clinical trials in pulmonary vascular disease, even if not as the primary endpoint. Longer, time-to-event clinical trials are now being conducted in PAH using composite morbidity and mortality endpoints (61, 62), which is the ideal setting in which to validate potential RV-based surrogate endpoints. The validation of an RV surrogate in this setting would contribute greatly to the feasibility of smaller phase II trials to identify potentially promising therapies from ineffective ones. CMR has greater sensitivity and reproducibility than ultrasound and thus can rapidly detect an efficacy signal with a small sample size in a short period of time, potentially minimizing cost and avoiding trial futility (60). However, its routine use in clinical trials needs further validation.

Optimization of Animal Models of RVF

Rationale. Animal models of disease are critical to understanding the physiology and structural and molecular underpinnings of the human conditions they attempt to recapitulate. They facilitate therapy discovery and safety evaluation. For RVF, an animal model that recapitulates the mechanisms and prognostic implications of RVF in humans would minimize requirements for human tissues, and new understanding of molecular processes driving failure would accelerate the discovery and preclinical testing of potential drugs for RVF.

Current state of knowledge. An ideal RVF animal model would be reproducible by individual investigators over time, as well as reproducible across institutions. Such a model would also be inexpensive, feature relatively brisk disease development for rapid knowledge acquisition, and, most importantly, accurately represent physiologic and molecular characteristics of the human RVF phenotypes. Given the varied clinical conditions resulting in primary or secondary RVF, a single animal model fulfilling all these criteria is unlikely (63). Presently, there are several commonly used animal models of RVF (Table 2), all with limitations and none considered the single “best” model. With the exception of exposure to hypoxia and pulmonary artery banding (PAB), which can be done in mice, rats have been used predominantly, as they most reliably reproduce the vascular lesions characteristic of human PAH (64). Pulmonary vascular lesions generated in different models vary (64).

Current knowledge gaps. Whether the development of RVF in these preclinical models is purely a function of hemodynamic overload or specific lung and systemic vascular lesions in various models is unknown. PAB can be applied to both rats and mice and does not elicit pulmonary vascular disease, but it has technical limitations and variable effects on the RV. The comparison of pure RV pressure overload models (such as PAB) versus pressure overload syndromes due to pulmonary vascular disease (as induced by monocrotaline or Sugen plus hypoxia) suggests that the former model manifests adaptive RVH and may be relevant to congenital heart diseases (such as pulmonic stenosis), whereas the latter may be more reflective of the maladaptive RVH seen in most patients with WHO group 1 PH (65). However, it is believed, although not fully proven, that decreased cardiac output (CO) and maladaptive RVH can develop in this model as well, depending on the degree and duration of the banding (66). A need exists for models of pure RV volume overload. The role of alternative models of RVF, including large animal models (67, 68) and transgenic mouse models (69), needs to be defined further.

Recommendations/pathway for progress. We recommend matching the animal model to the experimental question asked. For instance, when the experimental question involves the etiology of RV fibrosis, monocrotaline or Sugen plus hypoxia may be relevant. Alternatively, when studying molecular mechanisms behind the progression from compensated RVH to RVF, PAB with various degrees and/or durations of banding is preferable. Corroborations of results in more than one model strengthens experimental findings (70). The RV Langendorff model is a useful ex vivo model to assess RV inotropy and lusitropy (71–73) and a useful adjunct to in vivo assessments of the RV, which are largely load dependent. Large animal models of RV pressure and volume overload (with and without pulmonary vascular disease) will likely
be very helpful in the latter stages of preclinical testing of drugs and devices, before use in humans. RV volume overload could be generated by inducing compromising tricuspid valve insufficiency. The design of novel, translationally relevant models is dependent on a better molecular understanding of the unique determinants of human RVF. A better molecular understanding of the basis for RVF may lead to improved transgenic animal models or preference for one of the existing models with greater understanding of its implications. In the absence of these data, we recommend:

1. Correlation of findings from animal models with human tissue.
2. Consideration of sex and age in animal models of RVF.
3. Incorporation of right heart catheterization (RHC) (thermodilution and/or pressure–volume analyses using high-fidelity conductance catheters), echocardiography, and/or CMR to measure CO, a marker of RV function.
4. Combine CO measurements with other functional measurements (e.g., TAPSE and RV dilatation with echocardiography), RV volumes (CMR), and/or ventriculoarterial coupling (using pressure–volume loops).

Table 2. Commonly Used Animal Models of Right Ventricular Failure

<table>
<thead>
<tr>
<th>Monocrotaline Hypoxia</th>
<th>Sugen/Hypoxia</th>
<th>PA Banding</th>
</tr>
</thead>
<tbody>
<tr>
<td>Method</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Subcutaneous injection of alkaloid monocrotaline (40–60 mg/kg; lower dose results in milder phenotype)</td>
<td>Exposure to FiO2 10% (or 0.5 atmospheric pressure) for 3–5 wk</td>
<td>Subcutaneous injection of VEGFR2 antagonist Su5416 + hypoxia (3 wk) + room air (≥4 wk)</td>
</tr>
<tr>
<td>RVSP, mm Hg</td>
<td></td>
<td></td>
</tr>
<tr>
<td>40–60</td>
<td>30–40</td>
<td>60–80</td>
</tr>
<tr>
<td>Cardiac output</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Depressed</td>
<td>Maintained or slightly depressed</td>
<td>Severely depressed</td>
</tr>
<tr>
<td>RV histology</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fibrosis, cardiomyocyte hypertrophy, possibly biventricular myocarditis (conflicting data on the latter)</td>
<td>Fibrosis (less than other models), cardiomyocyte hypertrophy</td>
<td>Fibrosis, cardiomyocyte hypertrophy, apoptosis</td>
</tr>
<tr>
<td>RV vascular effects</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Vascular inflammation, decreased capillary density</td>
<td>Capillary proliferation</td>
<td>No angiogenesis, decreased capillary volume and density</td>
</tr>
<tr>
<td>Type of RV remodeling</td>
<td>Maladaptive</td>
<td>Adaptive</td>
</tr>
<tr>
<td>Pulmonary vascular effects</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Systemic effects</td>
<td>Yes (less PH in females)</td>
<td>Yes (less PH in females)</td>
</tr>
<tr>
<td>Sex of animal tied to experimental phenotype</td>
<td>Yes (less PH in females)</td>
<td>Yes (better RV function in females)</td>
</tr>
<tr>
<td>What is being modeled?</td>
<td>Inflammatory PAH and RV dysfunction</td>
<td>Chronic hypoxia, high altitude, some aspects of chronic lung disease</td>
</tr>
<tr>
<td>Modifications of model</td>
<td>Combination with pneumonectomy or aortocaval shunting to result in pulmonary vascular overflow and more pronounced remodeling</td>
<td>Extension of room air exposure associated with progressive pulmonary vascular remodeling and afterload increase</td>
</tr>
</tbody>
</table>

Definition of abbreviations: PA = pulmonary artery; PAH = pulmonary arterial hypertension; PH = pulmonary hypertension; RV = right ventricle; RVSP = right ventricular systolic pressure; VEGFR2 = vascular endothelial growth factor receptor 2.

PH development is usually more pronounced in younger animals. Data from References 64, 65, 81, 264–279.
Although optimization of animal models of RVF is of utmost importance in the field, these efforts should be paralleled by the development of new in vitro technologies, such as a “heart on a chip” approach and/or cardiomyocytes or cardiac endothelial cells differentiated from human-PH induced pluripotent stem cells. Such models could be used to study RV cardiomyocyte–endothelial cell interactions, screen drugs, and identify novel signaling pathways.

**Animal Models for RVF in Group 2 and 3 PH**

**Rationale.** The majority of PH cases worldwide are due to diseases such as chronic LV and lung disease, sleep-disordered breathing, or high altitude exposure (group 2 and 3 PH) (74–76). Treating the underlying condition is typically the goal; however, a significant number of patients may develop RVF, and there has been considerable interest in the development of animal models of RVF in affected individuals. Therefore, there is a need for adequate animal models of RVF resulting from group 2 and 3 PH.

**Current state of knowledge.** RVF generally does not develop in rodent models of hypoxia exposure but is present in the hypoxic newborn calf (77). In patients with chronic lung disease, the development of PH and/or RVF is not strictly linked to chronic hypoxia but may also develop because of hypercapnia, neurohumoral alterations, comorbid cardiovascular disease, and pulmonary vascular obstruction and obliteration (78). Rodents with Sugen- or cigarette smoke–induced emphysema as well as bleomycin- or adenoviral TGF-β–induced pulmonary fibrosis may exhibit RVH but do not exhibit significant RVF (79–83).

In models of LV disease, RVF may develop as an integral part of a global cardiac dysfunction (e.g., doxorubicin-induced biventricular failure) (84). In addition, isolated damage to the LV can directly impair RV function (85), even in the absence of PH (86). Development of secondary PH in models of LV failure induced by transverse aortic constriction is little studied. Indeed, a significant degree of lung vascular remodeling, PH, and RVF can develop as soon as 4 weeks after transverse aortic constriction in mice (87), suggesting this may be a model of combined pre- and postcapillary PH. Combined pre- and postcapillary PH and RVF in the setting of mitral stenosis or heart failure with preserved ejection fraction may be more aptly represented by a model of pulmonary vein banding (reported in swine [88], although with technical limitations for mechanistic studies). A model of PH due to LV diastolic dysfunction using Su5416 administration in obese ZSF1 rats has recently been published; this model exhibits moderate PH, but no overt RVF (89). Similarly, AKR/J mice fed with a high-fat diet develop increased LV end-diastolic pressure and biventricular hypertrophy, but it is unknown if they develop RVF (90). As in patients, animal models may require substantial time to develop RVF; this feature may be overlooked because of cost and time constraints, as well as arbitrary requirements by animal care committees that animals be killed if they become ill. Clinically relevant surrogate endpoints of failure to thrive (e.g., weight loss or decreased oral intake) may occur before overt RVF develops in these models.

**Current knowledge gaps.** Mechanisms of RVF development in group 2 and 3 PH are multifactorial, but the relative contributions of the various causes have not yet been fully elucidated and are difficult to mimic in small or large animals. Whether treating RVF would translate to improved clinical outcomes is currently unknown, particularly as PAH medications have largely been found to be ineffective or harmful in patients with group 2 and 3 PH.

**Recommendations/pathway for progress.** A better understanding of RV function in patients with group 2 or 3 PH will be critical for understanding the human disease and for optimizing animal models. Therefore, clinical studies should carefully phenotype patients with RVF from these syndromes, and studies should explore whether treating RVF improves clinically relevant outcomes. “Multiple hits” rather than a “single hit” likely contribute to human disease, and such multifactorial etiologies should be considered in animal models (89). Ultimately, even though the “perfect” animal model may not ever exist, the development of animal models mimicking group 2 and 3 PH with significant RVF would be a major milestone. Isolated cardiomyocytes may allow for mechanistic investigations of specific pathways. Animal studies of group 2 or 3 PH should be of sufficient duration to detect RVF. Ideally, more sophisticated RV endpoints including BNP levels, CO (measured by RHC and Doppler), and formal assessment of RV size and function should be incorporated. Studies in humans should focus on deep phenotyping and ideally involve the study of RV tissues.

**RV Responses to Pulmonary Embolism and Acute Respiratory Distress Syndrome**

**Rationale.** Both pulmonary embolism (PE) and acute respiratory distress syndrome (ARDS) cause relatively acute PH and RV injury; the ensuing RVF results in significant morbidity and mortality in both conditions. Causes of RV injury in both scenarios include increased afterload from PH and impaired subendocardial RV perfusion (91–94).

**Current state of knowledge.** Rapid increases in PAP and PVR, as occur in PE and ARDS, are poorly tolerated, and the RV will fail and develop conduction abnormalities that would not occur if the pressure increased slowly, allowing time for the development of compensatory RVH. Both large PE and severe ARDS result in uncoupling of the RV from the pulmonary circulation (defined and discussed in **Assessment of RV–PA Coupling**), and subsequent inefficient forward flow. The primary insult with PE comes from macrovessel occlusion, whereas ARDS results from microvessel occlusion with widespread but heterogeneous alveolar consolidation and elements of acute hypoxic pulmonary vasoconstriction. Both conditions shift the pulmonary vasconstrictor–dilator balance toward an increase in afterload and result in chemokine and cytokine release, recruitment of inflammatory cells, and production of reactive oxygen species (95–101). LV diastolic dysfunction (i.e., stiffness) may arise after acute PE and persist despite PH resolution (102–104) but likely is not a main contributor to symptoms. Approximately 4% of acute PE survivors develop chronic thromboembolic PH (CTEPH) (105), making this one of the most common forms of PH. Uniquely, CTEPH has a high index of recovery after intervention.

**Current knowledge gaps.** Although RVF is an independent cause of increased mortality in ARDS (106), the impact of RVF prevention or treatment on prognosis in...
ARDS is unknown (107). Prone positioning improves survival and RV function in severe ARDS (108, 109); however, the optimal methods and modes of ventilation, as well as the roles of inotropic support, varying levels of sedation, and paralysis to protect the RV and limit RVF in patients with ARDS all remain poorly studied. Whether prone positioning is indicated in cases of RVF independent of decreases in PaO₂/FiO₂ remains to be evaluated as well. Despite extensive research on the mechanisms promoting lung injury and pulmonary vascular dysfunction in ARDS, no specific and effective targeted therapies have emerged. For PE, the role of catheter-directed fibrinolysis (vs. standard anticoagulation) for intermediate-risk PE requires large-scale randomized controlled trial investigation (110). Significant gaps exist in understanding the best ways to support the failing RV during these relatively acute RVF syndromes. The optimal intravenous inotrope for clinical use has not been identified. In animal models of chronic RVH, dobutamine appears superior for acute RV support (111), but it is unclear if this also pertains to acute RVF. Although use of inotropes in RVF in the face of PAH identifies a patient cohort at high risk (46%) of in-hospital mortality, it is unclear if this is a mere reflection of the severity of the underlying condition rather than an inotrope effect per se (112). It is also uncertain how best to prevent secondary inflammatory RV injury from PE and ARDS. Modulating LV stiffness in resuscitation remains unstudied for both conditions (102–104). Knowledge about vasoconstrictor–vasodilator balance, platelet hyperreactivity, and coagulation–fibrinolytic balance has not been adequately translated into early-phase clinical trials. Mechanisms of development of CTEPH after acute PE are incompletely understood and need further study. Recently developed rat and piglet models (113–115) will facilitate such studies.

**Recommendations/pathway for progress.** We recommend developing robust and translationally applicable animal models, designing RV-focused outcome studies to assign risk in large cohorts, and performing RV-focused randomized controlled trials, particularly at the phase I stage. Strategies aimed at understanding common mechanisms of lung injury caused by precapillary pulmonary vascular occlusion associated with PE and/or ARDS need to be identified. Goals should include identifying optimal mechanisms of enhancing CO while supporting systemic vascular tone. This should include studies modulating the β-adrenergic receptor in RV cardiomyocytes as well as studies of mechanical devices. In addition, this should include the unraveling of cause and effect interplay between small vessel occlusion by cells and vasospasm and inflammatory processes that can cause RV injury directly and indirectly. The roles of disordered coagulation, platelet activation, neutrophil-derived networks, and regulation of fibrinolysis need to be defined. For both PE and ARDS, existing evidence raises the hypothesis of a secondary inflammatory hit to the RV, initiated by either direct injury (e.g., shear stress or ischemia) or as collateral damage from systemic inflammation (116). The importance and magnitude of effect of this secondary hit require quantitative study. Priorities should focus on producing translatable knowledge in the laboratory and in clinical trials to enhance the pipeline of new therapies for both conditions. In addition, better data are required to determine the optimal duration of therapy. Guidelines should be developed to assess whether and when RV function should be assessed in patients who recover from acute PE or ARDS. Investigating whether CTEPH surveillance in survivors of a documented PE improves outcomes is therefore indicated. Mechanistic and therapeutic preclinical studies in recently developed CTEPH animal models will provide further knowledge that can be harnessed to improve outcomes of patients with CTEPH.

**Differences between Acute and Chronic RV Responses**

**Rationale.** Although identifying cellular and molecular mechanisms that underlie successful adaptation of the RV to chronic stress can lead to efforts to prevent RVF in patients with chronic RV pressure overload (RVPO), this may also lead to strategies that improve outcomes of acute PH. Similarly, a better understanding of mechanisms of acute RVF may lead to treatment strategies for chronic RVF. For example, a detailed knowledge of the mechanisms leading to—or protecting against—myocardial apoptosis in the acute setting could improve the management of patients suffering from both acute and chronic RVF.

**Current state of knowledge.** Acute RVPO causes wall stress, exhaustion of cardiomyocyte energy resources, impaired calcium handling, release of proinflammatory chemokines and cytokines, inflammatory cell infiltration, and generation of toxic reactive oxygen and nitrogen species, all of which contribute to cardiomyocyte death within hours or days and loss of pump function (96, 98, 100, 101). In contrast, chronic RVPO triggers a slower and more or less successful adaptation to pressure overload (96, 117), including complex alterations in energy metabolism (generally away from fatty acid oxidation [FAO] toward uncoupled glycolysis and carbohydrate oxidation) as well as hypoxemia, ischemia, inflammation, and oxidative stress (1, 2, 13, 118, 119). These changes are accompanied by macroscopic and microscopic structural remodeling of the myocardium and cardiac microcirculation. The remodeling of chronic RVPO includes cardiomyocyte hypertrophy and, in case of unsuccessful adaptation, fibrosis and capillary rarefaction (65). Failure of compensatory mechanisms is believed to trigger transition from adaptive to maladaptive RVH, with ischemia considered a major contributor (65, 120, 121).

**Current knowledge gaps.** It is likely that changes in metabolism vary among forms of RVF. For example, FAO is increased in PAB (and inhibiting FAO is beneficial to RV function) (122); conversely, in fawn hooded rats (which develop spontaneous PH), FAO is decreased (123). It is likely this disease heterogeneity, which is relatively chamber specific and primarily affecting the RV, is also seen in humans; however, this has not been systematically assessed. Other metabolic pathways are induced in RVH as well, including the de novo appearance of cardiomyocyte glutaminolysis (although this finding requires confirmation) (124). It is unknown if a “point of no return” exists in the transition from adaptive to maladaptive RV remodeling and whether certain failure components are more important than others (117). For example, if impaired FAO in cardiomyocytes is the root cause of RVF, other processes (e.g., apoptosis, inflammation, fibrosis) would be downstream consequences, and therapeutic interventions would need to aim at reversing impaired FAO. Furthermore, it is unclear whether inflammation, oxidative stress, neurohormonal activation, or metabolic alterations equally affect RV function in the
acute versus chronic setting and whether targeting these processes has the same outcome in acute versus chronic RVF. The role of hematological factors (e.g., hypercoagulation, platelet activation, hypofibrinolysis) in initiating or propagating acute (and possibly even chronic) RVF is unknown. Furthermore, the contribution of pulmonary vasoconstriction to inducing and maintaining acute or chronic RVPO has not been well defined. Last, the role of the LV in the setting of RVPO needs further definition. Even with acute PH, the LV becomes compressed and stiffened by an enlarged RV, thereby contributing to reduced CO (102, 103).

**Recommendations/pathway for progress.** Targets and pathways important in acute RVF should be evaluated in chronic RVF (and vice versa), allowing for identification of common as well as unique targets and pathways in either setting. For example, in acute PE, hemolysis increases shear forces in the RV (125), and multiple sites in the coagulation cascade are believed to contribute to acute RVF development (126, 127). If these processes are identified as being unique to acute RVF, their study may lead to novel treatment targets. If a role in chronic RVF is established, these concepts could then be expanded to treating the chronically failing RV. Comparing acute with chronic RVF will allow for a better understanding of the transition from compensated RV remodeling. Phenotypic, genetic, and epigenetic determinants of adaptation in the acute or chronic setting should be studied. Comprehensive and integrated systems biology/omics approaches are most likely to yield novel insights regarding the shared and divergent pathways and mechanisms between acute and chronic RVF.

**Regional Differences in RV Responses**

**Rationale.** Recent data indicate that RV responses to acute or chronic afterload stress vary between different regions of the RV. Uncovering these regional differences will help identify mechanisms of transition from adaptive to maladaptive RVH. Please see the online supplement for further details. LV involvement in RV disease and ventricular interdependence are covered in Assessment Of RV–LV Interactions and Ventricular Interdependence.

Table 3 lists key knowledge gaps and solution approaches for topic domain 1.

**Topic Domain 2: Analyzing Advanced RV Hemodynamic Parameters at Rest and in Response to Exercise**

**Rationale**

Identification of knowledge gaps and solution approaches in hemodynamic phenotyping and prognostication requires a detailed understanding of the various tools used. This section, therefore, reviews strengths and weaknesses of commonly used tools, identification of knowledge gaps, and pathways for progress.

**Invasive and Noninvasive Phenotyping of RV Function**

**Current state of knowledge and knowledge gaps.** Comprehensive assessment of the RV requires a combination of RHC and RV imaging via echocardiography or CMR. Biomarkers and exercise testing serve as additional tools. RHC allows for a thorough (though not complete) description of RV function, with right atrial pressure to estimate RV end-diastolic volume, or preload, PAP, or PVR to estimate afterload, and stroke volume (SV) to reflect contractility. Imaging by echocardiography and/or CMR offers additional information. Both provide accurate (although sometimes imprecise) estimates of systolic, mean, and diastolic PAP, left atrial pressure (LAP), and CO, and derived calculations of PVR and pulmonary arterial (PA) compliance. More importantly, both techniques provide indices of RV systolic function, diastolic function and filling pressures, planar and volumetric estimations of dimensions, and quantifications of dyssynchrony (interregional inhomogeneity of contraction) and asynchrony (interventricular inhomogeneity of contraction) and RV volumes.

**Recommendations/pathway for progress.** RV function is often reported in a cursory fashion and frequently limited to an EF measurement. A physiologically sound definition of RVF (see Definition, Development, and Staging of RVF) should be implemented in biological and clinical studies, to avoid ambiguity in bench-to-bedside translation and epidemiology. Clinical studies can achieve this through an integrated imaging and invasive approach (e.g., RHC, imaging by echocardiography and/or CMR, and possibly PET [128]); preclinical investigations should use comprehensive hemodynamics and RV imaging studies (70).

**Prognostic Relevance of Measurements of RV Function**

**Current state of knowledge and knowledge gaps.** Table E1 in the online supplement provides an overview of the prognostic capabilities of RHC-, echocardiography-, and CMR-derived variables in various types of severe PH. Various combinations of these parameters have been used in prediction models (129, 130). RHC-derived predictors of outcomes include CO (measured or calculated from mixed venous blood oxygenation) (131–135), right atrial pressure (132–134, 136), PVR (135–137), and PAP (133). Echocardiographic predictors include pericardial effusion (138–141), right heart dimensions (139, 140, 142–144), estimated RV diastolic pressure (140, 145), tricuspid regurgitation (142), TAPSE (23, 140, 143, 146), maximum tissue velocity of isovolumic contraction (147), dP/dt (146), strain (evaluated by variable and evolving methodologies) (141, 148, 149), asynchrony and regional inhomogeneity of RV contraction (150), myocardial performance index (151, 152), RV fractional area change (130), right atrial size index (130), and RV contractile reserve (defined as the exercise-induced increase in the maximum velocity of tricuspid regurgitation) (153). These data suggest that echocardiography offers prognostication via measurements of RV systolic function (TAPSE, velocity of isovolumic contraction, dP/dt, strain, myocardial performance index, contractile reserve), dimensions, and estimates of filling pressures. CMR-derived predictors include SV (21, 154), end-diastolic volume (21, 155), RVEF (3, 154, 156), end-systolic volume (ESV) (129, 157), SV/ESV (3), and late gadolinium enhancement (158). The latter, however, is not uniformly confirmed (159). CMR therefore offers prognostication from measurements of systolic function (ESV, EF, SV/ESV) and right heart dimensions. A recent CMR-based study used machine learning of patterns of RV motion for outcome prediction in PH (160). Because of their noninvasive nature, there has been particular interest in using imaging
parameters to assess RV function. The current knowledge, knowledge gaps, and potential future roles of commonly used RV imaging parameters are listed in Table 4. As of now, CMR remains the standard of measurement, and RVEF is the most robustly validated parameter. Head-to-head comparisons of other imaging parameters with regard to superiority are lacking, and the appropriateness for use of one specific parameter rather than others in special clinical scenarios has not been well described. Please see the online supplement for further discussion of caveats pertinent to the use of RV function markers as prognosticators.

Recommendations/pathway for progress. Imaging or invasive measurements of RV function, identified as prognostic indicators in PAH, should be evaluated together in the framework of a multicenter trial and prediction scores constructed from rigorously identified independent predictors. A hierarchy of imaging parameters and their appropriateness for specific scenarios should be determined.

Assessment of RV Afterload

Current state of knowledge and knowledge gaps. With low-normal PVR, the contribution of the RV to pulmonary blood flow is modest at rest (evidenced after a Fontan operation). However, RV pump function is required in cases of increased afterload (e.g., in acute or chronic PH). RV afterload is affected by PVR, but there are several other equally valid contributors (161): 1) Maximum wall tension (an impractical parameter because of the irregular shape of the RV and regional inhomogeneities in contraction); 2) hydraulic power (W$_{TOT}$) calculated from the integration of instantaneous pressure and flow waves, which is the sum of oscillatory power [W$_{osc}$] and steady power [W$_{st}$ = mPAP × CO]; 3) arterial elastance (Ea; calculated from an RV pressure–volume loop as end-systolic pressure [ESP]/SV), corresponding to a measurement of afterload as “seen” by the ventricle; and 4) pulmonary vascular impedance, which represents the most comprehensive assessment of RV afterload (although least accessible to the clinician) (162).

RV pressure–volume loops with preload reduction allow for identification of the end-systolic elastance (Ees; the standard of measure for load-independent contractility in vivo). Direct measurements of Ees and Ea allow for calculation of the Ees/Ea ratio as a measurement of the coupling of the RV to the pulmonary circulation (2, 161) (Figure 4). This concept was initially developed for the LV (163) but soon thereafter was shown to be applicable to the RV (164). Sagawa and colleagues demonstrated that optimal efficiency of ventriculoarterial coupling (allowing for flow output at minimal amount of energy cost) is achieved in either ventricle when maximal or end-systolic elastance is 1.5 to 2 times greater than Ea (165). In the RV, the amount of uncoupling associated with decreased maximum cardiac output and increased RV size with high filling pressures and systemic congestion is not exactly known (128). Although a coupling ratio of 1 at rest may still be associated with preserved RV dimensions, at this level of Ees/Ea, there is impaired contractile reserve, and exercise is associated with uncoupling and increased dimensions (166, 167). Furthermore, in experimental animal models of embolic PH, the RV starts to dilate when Ees/Ea is decreased to 0.7 to 1.0 (168, 169). Most investigators therefore consider “optimal” mechanical RV–arterial coupling to correspond to an Ees/Ea of 1 to 2. Interestingly, for a similar afterload, patients with scleroderma-PAH demonstrate lower Ees/Ea than patients with idiopathic PAH (1.0 vs. 2.1) (170), suggesting significant uncoupling in scleroderma-PAH.

Ea and W$_{TOT}$ are metrics of RV afterload, so it is understandable that these measurements of the pulmonary circulation emerge as prognostic markers of RVF in severe PH (171, 172). For example, increased stiffness of the proximal PA, mediated by excessive collagen accumulation, increases the pulsatile component of RV afterload (quantified by measurement of impedance [173]) and thus decreases RV–PA coupling and RV contractile reserve (defined as the ability of the RV to increase contractile function during exercise) (2, 153, 174).
Table 4. Current State of Knowledge, Knowledge Gaps, and Potential Future Roles of Commonly Used Right Ventricular Imaging Parameters

<table>
<thead>
<tr>
<th>Imaging Parameter</th>
<th>Current State</th>
<th>Disadvantages and Knowledge Gaps</th>
<th>Future Role</th>
</tr>
</thead>
<tbody>
<tr>
<td>TAPSE</td>
<td>Easy and quick assessment</td>
<td>Less suitable for serial assessment (284, 285)</td>
<td>2D measures will still be used for assessment of RV function</td>
</tr>
<tr>
<td></td>
<td>Cheap and widely available</td>
<td>2D measures are less suitable for serial assessment owing to the complex geometry of the RV</td>
<td></td>
</tr>
<tr>
<td></td>
<td>High reproducibility (280, 281)</td>
<td>Prognostic value at baseline (23, 141, 282, 283)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Prognostic value at baseline (282, 283)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RVFAC</td>
<td>Easy and quick assessment</td>
<td>Less reproducible than TAPSE (280, 281, 285)</td>
<td>See TAPSE</td>
</tr>
<tr>
<td></td>
<td>Good relation with RVEF (280, 285, 286)</td>
<td>Less suitable for serial assessment (285)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Prognostic value at baseline in PAH (141, 282, 283)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Eccentricity index</td>
<td>Easy and quick assessment</td>
<td>Lack of information on serial assessment</td>
<td>See TAPSE</td>
</tr>
<tr>
<td></td>
<td>Prognostic value at baseline in PAH (282, 283)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Strain/strain rate</td>
<td>Prognostic value in PAH (141, 283, 287)</td>
<td>Strain assessment possible with echo (2D speckle tracking) and CMR (tagging)</td>
<td>Shows potential to assess RV dysfunction in a very early stage</td>
</tr>
<tr>
<td></td>
<td>Early detection of RV dysfunction; detects differences in RV function when other traditional measurements, including TAPSE, fail to do so (288)</td>
<td>Tagging analyses time-consuming</td>
<td></td>
</tr>
<tr>
<td>Myocardial performance index (Tei index)</td>
<td>No need to make geometric assumptions</td>
<td>Doppler echocardiography only</td>
<td>See TAPSE</td>
</tr>
<tr>
<td></td>
<td>Prognostic value in PAH (141, 282)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RVEF</td>
<td>Gold standard parameter for assessment of RV systolic function</td>
<td>Assessment is time-consuming</td>
<td>Likely to be useful in assessment of prognosis and treatment response</td>
</tr>
<tr>
<td></td>
<td>RVEF is reproducible (289)</td>
<td>Presence of tricuspid regurgitation can overestimate RVEF</td>
<td>Automated assessment of RV volumes with CMR needed</td>
</tr>
<tr>
<td></td>
<td>Prognostic value at baseline and during follow-up (3, 21)</td>
<td>Minimal clinical important difference unknown</td>
<td>3D echocardiography may enable accurate assessment; however, currently 3D echocardiography RV underestimates volumes and is less reproducible (290–294)</td>
</tr>
<tr>
<td>RVS/RVESV</td>
<td>Prognostic value at baseline (129, 184)</td>
<td>Assumption time-consuming</td>
<td>May be an important parameter, although the measure shows a great similarity to RVEF</td>
</tr>
<tr>
<td></td>
<td>Minimal clinical important difference unknown</td>
<td>Prognostic value of the change over time unknown</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Reproducibility unknown</td>
<td>Indexing to body size may be necessary but is not consistently done</td>
<td></td>
</tr>
</tbody>
</table>

(Continued)
Consequently, PA compliance is an important predictor of mortality in PAH (2, 172, 174, 175).

The most comprehensive measure of RV afterload is pulmonary vascular impedance, which represents the opposition of the vascular bed to pulsatile flow generated by friction in small-diameter vessels, stiff vessels that do not accommodate pulsations, and branching and tapering vessels that generate wave reflections (162).

Recommendations/pathway for progress. Contributions of the various components of RV afterload to RV function and RV adaptive and maladaptive processes need better definition. Effects of PAH-targeted therapies on components of RV afterload require further study. Easy-to-obtain bedside measures of RV afterload would be of value.

Assessment of RV–PA Coupling

Current state of knowledge and knowledge gaps. The most accurate measurement of RV–PA coupling is Ees/Ea. However, the complex geometry of the RV makes functional evaluations with measurement of instantaneous volume changes technically difficult. In addition, manipulating systemic venous return to alter RV preload clinically is not practical. Accordingly, a single-beat method has been developed, allowing for determining Ees and Ea from instantaneous ventricular pressure and flow output (176) or volume (129) measurements (Figure 4). However, this method is controversial, as it relies on multiple assumptions (176).

RV–PA coupling has been assessed in several studies in patients with PAH or CTEPH, and in one case report of a patient with a systemic RV (129, 167, 170, 177–179). As shown in Table E2, Ees/Ea was either maintained or decreased at rest but consistently decreased at exercise. In aggregate, these results agree with the notion of homeometric adaptation of RV function to afterload, even though the results do not allow identification of critical levels of decoupling associated with onset of heterometric adaptation and congestion.

Recommendations/pathway for progress. RV–PA coupling measurement should be implemented in studies investigating the mechanisms of action of targeted therapies in PAH (which decrease RV afterload but may or may not have intrinsic myocardial effects), even if not as the primary endpoint. It is controversial...
whether Ees/Ea can be measured on a single-beat pressure–volume loop versus a family of such loops generated by a manipulation of venous return. A study comparing these methods in patients with various types of PH is urgently needed. Simplified surrogate measurements of Ees/Ea, such as SV/ESP (156, 180) and (RV maximum pressure/mPAP – 1) (181) (Figure 4), should be validated.

Assessment of RV Diastolic Function

Current state of knowledge and knowledge gaps. Coupling of RV function to afterload has an inevitable diastolic component (2, 161, 182). Diastolic stiffness may be a predictor of outcomes in severe PH (183, 184). Please see the online supplement for a summary of RV diastolic function assessment.

Figure 4. Methods used to estimate right ventricle–pulmonary artery (RV–PA) coupling and diastolic stiffness. In both (A) the volume method, and (B) the pressure method, arterial elastance (Ea) is calculated from the ratio of end-systolic pressure (ESP) to stroke volume (SV). End-systolic elastance (Ees) as an approximation of maximum elastance in the volume method is estimated by the ratio of ESP to end-systolic volume (ESV), which results in a simplified Ees/Ea of SV/ESV. In the pressure method, Pmax is estimated from the nonlinear extrapolation of the early systolic and diastolic portions of the RV pressure curve. Ees is then defined by a tangent from Pmax determined from the extrapolation of early and late isovolumic portions of an RV pressure curve and synchronized absolute or relative volume measurements. Ees is then defined by a tangent from Pmax to the pressure–volume relationship, and Ea is defined by a line drawn from the Eos point to end-diastolic volume (EDV) (at zero pressure). (D) Diastolic stiffness (β) is calculated by fitting the nonlinear exponential $P = a e^{b \cdot V} - 1$ to pressure and volume measured at the beginning of diastole (BDP, ESV) and the end of diastole (EDP, EDV). Adapted by permission from Reference 184. BDP = beginning diastolic pressure; ESP = end-diastolic pressure; EVol = Ees estimated by the volume method; mPAP = mean pulmonary arterial pressure; Pmax = RV maximum pressure; sRVP = peak systolic RV pressure.

RV diastolic function has not been specifically explored by imaging studies. Its description is generally limited to the isovolumic relaxation time, the ratio of trans-tricuspid flow E wave to tricuspid annulus tissue Doppler imaging tricuspid e’; the deceleration of the E wave, estimated RV filling pressure from the inferior vena cava dimension and inspiratory collapse, and end-diastolic area or volume. However, these measurements have not been systematically confirmed with diastolic pressure–volume curves or, with the exception of occasional parameters (Table E1), been considered as prognostic markers.

Recommendations/pathway for progress. The relevance of RV diastolic elastance or its surrogate, the end-diastolic pressure–volume ratio, should be further assessed. Novel, noninvasive measurement techniques using echocardiographic or other imaging methods are needed.

Assessment of RV–LV Interactions and Ventricular Interdependence

Current state of knowledge and knowledge gaps. RV function needs to be understood in the context of its direct and indirect interactions with LV function. Direct interaction, or ventricular interdependence, is defined as the forces that are transmitted from one ventricle to the other through the myocardium, perivalvular fibrous annulus, and pericardium, independent of neural, humoral, or circulatory effects (185). Diastolic ventricular interaction refers to the competition for space within the indistensible pericardium when the RV dilates. RHC and imaging studies demonstrated that in patients with severe PH, mPAP and LV peak filling rate are altered in proportion to decreased RVEF (186). Systolic interaction refers to positive interaction between RV and LV contractions. For example, aortic constriction and the subsequent increase in LV contraction markedly improve RV function in animals with PAB (187). Similarly, in electrically isolated ventricular preparations, LV contraction contributes significantly (~30%) to both RV contraction and pulmonary flow (187). This is explained by a mechanical entrainment effect, but also by LV systolic function determining systemic blood pressure (an essential determinant of RV coronary perfusion). This is important because increased RV filling pressures and excessive decrease in
blood pressure have been linked to RV ischemia and decreased contractility (188).

An additional cause of negative ventricular interaction is regional and interventricular asynchrony with postsystolic contraction or “shortening.” This develops in parallel with increased PAP and contributes to altered RV systolic function and LV underfilling (189). RV regional asynchrony can be identified and quantified by echocardiography using two-dimensional (2D) or 3D speckle tracking (149, 156) (see Regional Differences in RV Responses for further details). Electrophysiologically, this phenomenon is reflected by increased QRS width and prolonged duration of the monophasic action potential of RV myocytes secondary to downregulation of repolarizing potassium channels (72). Interestingly, this is amenable to metabolic therapies (72). Likewise, QTc duration increases in patients with PAH and serves as a prognostic indicator that appears to be related to RV mass (190). LV abnormalities in patients with RVF are not limited to functional changes. In fact, fibrotic and inflammatory changes in LVs from patients with PAH were recently reported (191).

Recommendations/pathway for progress. Further insight into the functional significance and clinical relevance of dysynchrony and asynchrony is needed. Because interventricular interdependence evolves in both acute and chronic RVF, studies in relevant preclinical models are needed to determine the relative contribution of this pathogenic mechanism in those conditions. Mechanisms of structural, biochemical and molecular LV abnormalities in patients with RVF require further study.

Emerging and Alternative Approaches
Please see the online supplement for a detailed discussion of this topic.

Table 5 lists key knowledge gaps and solution approaches for topic domain 2.

### Topic Domain 3: Deciphering the Underlying Molecular and Pathogenic Mechanisms of RV Function and Failure

#### Assessment of Hypertrophy and Angiogenesis

**Rationale.** Adaptive myocardial remodeling requires coordination of myocyte hypertrophy and angiogenesis to preserve ventricular function. Myocyte hypertrophy associated with a disproportional number of capillaries and/or endothelial cell (EC) dysfunction may ultimately promote RVF. Regulation of angiogenesis is complex (reviewed in Reference 192) and involves molecular cross-talk between myocytes and ECs (193, 194). Insight regarding molecular mechanisms linking RV myocyte hypertrophy and angiogenesis is limited. Further investigation is likely to identify new paradigms and therapeutic strategies.

**Current state of knowledge.** In the LV, angiogenesis is necessary (195, 196) and sufficient (197, 198) for myocyte hypertrophy. The interplay between angiogenesis and myocyte hypertrophy is poorly defined in the RV, which has unique mechanisms to augment myocyte perfusion during increased myocardial demand (199). A detailed review of the topic was recently published (200). RV angiogenesis has been evaluated in animal models of PH, although most studies relied on counting capillary profiles in 2D sections. Some authors believe that this approach may lead to underestimation of effective capillary length and surface area (201) and unintentional sampling bias (201, 202) that may lead to inconsistent findings, even within the same model (203–208). Analysis of 2D sections in PH models associated with RVF demonstrated reduced RV capillary density (65, 124, 207), potentially reflecting capillary rarefaction. In the rat monocrotaline model, early microvascular expansion was reversed as RVF ensued (209). These data, obtained from several laboratories and consistent with data reporting vascular rarefaction in LV hypertrophy, suggest that a combination of cardiomyocyte hypertrophy (causing increased demand) and capillary rarefaction leads to clinically relevant RV ischemia in PAH (210). However, a recent analysis of human RV tissue by a stereological approach noted an increase in total vascular length in PAH.
RVs versus controls (211). Although the role of microvascular ischemia is still controversial, the role of macrovascular ischemia (reduced coronary perfusion pressure due to elevated RV end-diastolic pressure) is established (188).

Molecular regulators of RV angiogenesis and myocyte hypertrophy have received limited attention. In addition, whether molecular regulation is similar between the failing LV and RV is also incompletely understood. Animal models support potential roles for hypoxia-inducible factor and vascular endothelial growth factor (65, 207, 209), although these pathways may be impaired in RVF (64, 119). Recent studies identified additional potential proangiogenic regulators in the LV (e.g., PIGF, NF-kB [nuclear factor-kB], extracellular ubiquitin, STAT3, Nrf2, and Ang-2/Tie-2) (197, 212–216), but these have not been studied in the RV. A potential regulatory role for the proangiogenic microRNA miR-126 in RV capillary rarefaction was recently shown (217). Although several angiostatic factors were recently identified as unique biomarkers in PAH (218, 219), their role in regulating RV angiogenesis has not been explored.

Current knowledge gaps. Pivotal questions are whether RV angiogenesis is necessary for adaptive myocardial remodeling and whether EC dysfunction, insufficient angiogenesis, and/or capillary rarefaction lead to maladaptive remodeling and RVF. The quantitative and temporal relationship between myocyte hypertrophy and angiogenesis required to preserve RV function in PH has not been established. Better understanding of the molecular regulators of RVH and angiogenesis is needed to develop studies designed to investigate whether hypertrophy and successful adaptation depend on angiogenesis. The potential role of angiostatic/antiangiogenic factors (217, 218, 220) in counteracting initial angiogenic responses is incompletely understood. The applicability of any single animal model to human RV angiogenesis and hypertrophy in PH is uncertain, and differences between models require further characterization. Angiogenesis data from patients with PAH are extremely limited, and current noninvasive imaging modalities lack sufficient resolution.

Recommendations/pathway for progress. Studies focused on RV myocyte hypertrophy and angiogenesis that use standardized, quantitative metrics avoid unintentional sampling bias and produce physiologically relevant endpoints (stereological methods [221]) that will improve precision and significance of morphometric studies. Mechanistic studies, including both unbiased screens for angiogenic and angiostatic mediators and hypothesis-driven assessment of specific regulatory candidates during multiple stages of RV remodeling, will advance the field. Additional insight is likely to be gained by more advanced imaging techniques (micro–computed tomography, microsphere injection), although these will require comprehensive assessment of the entire RV or a rigorous, unbiased sampling strategy.

Table 6 lists several key recommendations for the study of molecular and pathogenic mechanisms of RV function, which are relevant for the study of RVH and angiogenesis but are also critical for the study of the other mechanisms listed in topic domain 3. RV vascularization should be measured in an unbiased fashion; however, there currently is no agreement on whether stereology is the only acceptable technique for this. At the very least, further studies are needed that compare stereological versus other techniques (e.g., two-photon confocal imaging with 3D reconstructions). We strongly recommend the inclusion of human samples for morphological studies.

Table 6. Key Knowledge Gaps and Solution Approaches for Deciphering the Underlying Molecular and Pathogenic Mechanisms of Right Ventricular Function and Failure

<table>
<thead>
<tr>
<th>Key knowledge gaps</th>
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<tbody>
<tr>
<td>Is angiogenesis and/or capillary maintenance necessary to preserve RV function during remodeling, and does capillary rarefaction cause maladaptive RV remodeling?</td>
</tr>
<tr>
<td>Does myocardial fibrosis promote RVF, and can it be reversed?</td>
</tr>
<tr>
<td>Are metabolic alterations necessary for transition from adaptive to maladaptive RV remodeling? If so, which changes are the most important ones in this process? Which alterations in myocardial metabolic pathways should be targeted therapeutically, and how?</td>
</tr>
<tr>
<td>Are apoptosis and other forms of cellular death necessary for transition from adaptive to maladaptive RV remodeling?</td>
</tr>
<tr>
<td>Are myocardial inflammation and oxidative stress causes or consequences of RVF?</td>
</tr>
<tr>
<td>What is the role of circulating and resident progenitor cells in modulating RV hypertrophy, angiogenesis, fibrosis, metabolism, cell death, inflammation, oxidative stress, and contractile signaling?</td>
</tr>
<tr>
<td>How do established and novel PAH therapies modulate RV hypertrophy, angiogenesis, fibrosis, metabolism, cell death, inflammation, oxidative stress, and contractile signaling?</td>
</tr>
<tr>
<td>Are potential short-term detrimental effects on RV contractile function with neurohormonal modulators (e.g., β-adrenergic receptor antagonists) offset by long-term beneficial effects on other signaling pathways?</td>
</tr>
</tbody>
</table>

Solution approaches/pathways forward
- More human studies focused on identifying mechanisms of RVF and facilitated through RV biopsy samples, tissue repositories, and development of advanced imaging techniques. In order to enhance scientific impact, human subjects need to be well phenotyped
- Identification and validation of biomarkers of RV remodeling and dysfunction to facilitate human studies in PH-induced RV dysfunction
- Use of stereological methods in all morphometric analyses, including both human tissue and animal models
- More studies investigating the roles of sex and age as modifiers of RV function
- More studies investigating the roles of circulating and resident progenitor cells as modifiers of RV function
- Next-generation sequencing techniques in humans and animals to identify genetic contributors to RV adaptation and maladaptation
- Correlation of findings from cellular and animal studies with human phenotypes
- Identification/clarification of important mechanistic differences between preclinical models of RV adaptation and RVF
- Expanded use of RV working heart models and in vitro analyses of diverse myocardial cell types to evaluate specific regulatory pathways

Definition of abbreviations: PAH = pulmonary arterial hypertension; PH = pulmonary hypertension; RV = right ventricular; RVF = right ventricular failure.
Assessment of Fibrosis

**Rationale.** Fibrosis is a common end result of various pathogenic processes (2, 119), resulting in contractile dysfunction, diastolic dysfunction, and arrhythmias (2, 119). Cardiac fibrosis can be detected using CMR; patients with PAH with late gadolinium uptake (indicating fibrosis) have a poor prognosis (158). Strategies aimed at decreasing fibrosis may attenuate RVF. The routine use of gadolinium-enhanced CMR (already part of the monitoring algorithm in many centers) should be considered as a potential gold standard for noninvasive detection and quantification of RV fibrosis. Identification and validation of potential biomarkers of RV fibrosis would move the field forward. The study of isolated fibroblasts from normal and diseased RVs would provide novel mechanistic insights in fibrosis progression and regression. It will be important to know whether disorders found in pulmonary vascular cells and RV cardiomyocytes are also detected in RV cardiac fibroblasts. *In vitro* studies of PH-RV fibroblasts should confirm that these cells retain their disease phenotype in cell culture. Table 6 lists further recommendations regarding the study of human samples, stereology, biomarkers, progenitor cells, consideration of sex and age, and use of next-generation sequencing techniques.

**Assessment of Cardiomyocyte Metabolism**

**Rationale.** The normal adult RV relies primarily on primary FAO for ATP production (230). However, metabolic plasticity in the RV is seen at the time of birth and with disease states inducing RVH. RV myocytes engage in several maladaptive forms of metabolism during hypertrophy, including uncoupled glycolysis, dysregulated FAO, and glutaminolysis (reviewed in Reference 229). Because these alterations contribute to cardiomyocyte dysfunction (69, 72, 122, 231, 232), further insight into mechanisms of metabolic dysregulation is likely to identify new paradigms and new targets for therapeutic intervention.

**Current state of knowledge.** RV myocardial energetics are implicated in the regulation of RV function in experimental and clinical PAH, and metabolic therapies are well validated in preclinical models (65, 69, 72, 122, 231, 232). Metabolic changes may be (in part) elicited by ischemia and potentially genetic and epigenetic alterations (69, 233, 234). Epigenetic changes may be mediated by histone deacetylation, methylation of CpG islands (promoter regions with a high frequency of C-phosphate-G nucleotides), and miRNAs. Fluorodeoxyglucose F 18–PET studies have shed light on metabolism in humans, suggesting PAH-associated RVH is associated with increased glucose uptake. There are early, conflicting data regarding changes in FAO in the PAH-RV (234–239). Clinical trials are currently evaluating whether correction of metabolic alterations, such as reducing FAO or enhancing glucose oxidation, attenuates RVF (Clinicaltrials.gov identifiers NCT02102672 and NCT01083524).

**Current knowledge gaps.** Gaps exist in understanding the contribution of metabolic dysfunction to RVF, in particular the metabolic processes that drive transition from RV compensation to failure. Defining which metabolic changes are considered adaptive versus maladaptive will assist in testing metabolic therapies to prolong RV compensation and improve outcomes. Triggers of altered metabolism in the human RV, and whether correction of ischemia would reverse RV metabolic dysfunction, remain unknown. Moreover, it is unclear whether RV ECs or fibroblasts exhibit similar metabolic abnormalities to myocytes.

**Recommendations/pathway for progress.** Additional study of metabolic dysregulation in humans and human tissues is needed. Use of microperimetry would allow for direct measurement of metabolism in freshly isolated RV myocytes (reviewed in Reference 229). Such direct *ex vivo* studies, as well as use of *in vivo* confocal microscopy, would be important to avoid
pharmacological inference from *in vivo* experiments as a basis for defining changes in the RV in PH. Metabolic imaging, using PET with conventional and novel radiotracers and MR spectroscopy, holds promise to enhance understanding of the *in vivo* metabolic RV phenotypes and defining the transition from compensation to decompensation. In preclinical studies, use of Langendorff and working heart models allows more definitive attribution of metabolic therapies to changes in RV function. *In vivo*, the use of high-fidelity catheters to measure pressure–volume relationships can help establish effects of metabolic therapies on the RV (240). Metabolic studies in skeletal muscle may help establish effects of metabolic therapies to changes in RV function. In particular, more mechanistic studies in animal models and isolated RV cells focusing on initiators and modifiers of cell death of cardiomyocytes and other relevant cell types (e.g., cardiac ECs) and correlation of findings with human phenotypes will provide a better understanding of mechanisms of RV cell death. Studies of isolated cardiomyocytes and ECs (ideally from animal models of severe RVF and/or well-phenotyped human samples) would provide novel mechanistic insights in RV cell death initiation and progression. The contribution of endoplasmic reticulum stress, mitochondrial dysfunction and mitophagy (249), autophagy (245–247), and nonapoptotic types of cell death to RVF development requires further study. Identification and validation of cardiac troponins as markers of cardiomyocyte death (rather than damage) would move the field forward. Identification of novel targets and probes for apoptosis and cell death imaging will help study cell death in animals and in humans. Strategies and regulators listed in Table 6 also apply to the study of RV cell death.

**Assessment of Inflammation**

**Rationale.** Inflammation is an integral contributor to the development of both acute and chronic RVF (2, 95, 119, 229, 250). Strategies aimed at decreasing RV inflammation may prevent or delay the progression from adaptive to maladaptive RV remodeling and result in improved outcomes. Monitoring RV inflammation may serve as a “biomarker” of RV function and/or treatment responses.

**Current state of knowledge.** Inflammation, among other processes, likely contributes to RV fibrosis and contractile dysfunction. Influx of inflammatory cells and expression of proinflammatory cytokines and adhesion molecules is observed rapidly (within 2 h) after onset of RVPO (96, 98, 250). For example, animal models of PE, an influx of neutrophils and M1 macrophages is seen in the RV outflow tract within 18 hours (95, 98, 250). On the other hand, 6 weeks after acute RVPO, secretory M2 macrophages predominate, likely contributing to RV repair processes (95). In chronic RVPO, abundant expression of proinflammatory cytokines is noted (223), although the degree of inflammation appears to be less in chronic RVF than in RVF from acute RVPO (96).

**Current knowledge gaps.** Whether inflammation is a cause or a consequence of RVF is unknown. Initiators of RV inflammation (e.g., ischemia, apoptosis, impaired efferocytosis, mitochondrial dysfunction, oxidative stress), and potential links between inflammation and fibrosis, apoptosis, and metabolic dysfunction need to be identified. Similarly, it is unknown whether a certain degree of inflammation is beneficial for repair (and if so, how much). In LV failure, distinct macrophage populations have been linked to adaptive and maladaptive responses (251); it is unclear whether such a dual macrophage function also plays a role in RVF. Detailed investigation in human RVs (especially from patients not in end-stage disease) are lacking. Mechanisms mediating regression of inflammation, and whether this translates into clinical improvement, are poorly understood. Last, a better understanding of effects of established and novel PAH therapies as well as specific antiinflammatory strategies (e.g., ketorolac [250]) on acute and chronic RV inflammation is needed.

**Recommendations/pathway for progress.** Further characterization of inflammatory processes in the RV using biopsy (222) and novel imaging...
approaches (225) targeting specific inflammatory mediators in various stages of RVF and in distinct PAH populations (e.g., idiopathic vs. scleroderma-associated) would provide a better understanding of the role of inflammation in RVF. Preclinical mechanistic loss- or gain-of-function studies focused on individual cell types, mediators, and signaling pathways would provide a better understanding of the inflammatory contribution to RVF. Studying isolated cardiomyocytes and other relevant cell types (e.g., cardiac ECs and fibroblasts) will provide novel mechanistic insights into inflammatory signaling, especially if the proinflammatory environment of the failing RV can be recapitulated in vitro. Identification of mediators of inflammation development and regression and correlation of findings with human phenotypes will provide a better understanding of mechanisms of RV inflammation. Human samples, stereology, biomarkers, progenitor cells, sex and age, and sequencing techniques should be studied as outlined in Table 6.

Assessment of Oxidative Stress

Rationale. Oxidative stress is a purported contributor to RV dysfunction, providing a strong rationale for studying the oxidant/antioxidant imbalance of the stressed myocardium in both acute and chronically failing RV tissue. Please see the online supplement for detailed discussion of this topic.

Assessment of Contractile Signaling and Calcium Handling

Rationale. The RV in PAH is characterized by hypertrophy and both systolic and diastolic dysfunction (182). Perturbations in calcium handling have been implicated in all of these processes. A better understanding of calcium signaling in the failing RV may lead to novel diagnostic and/or therapeutic interventions.

Current state of knowledge. The RV in PAH exhibits changes in both the passive and active portions of the contractile apparatus. Sarcomere stiffening was identified in isolated RV cardiomyocytes (182). Both increased collagen accumulation and intrinsic stiffening of RV cardiomyocyte sarcomeres contribute to RV myocardial stiffness. In mild RVF, increased myofibril stiffness predominately contributes to myocardial stiffening, whereas in severe RVF, both myofibril- and fibrosis (collagen)-mediated stiffness contribute to increased RV myocardial stiffness (252). Sarcomere stiffening and RV diastolic dysfunction is caused by titin hypophosphorylation (182). In addition, there is dysfunction of active contractile regulator mechanisms. Downregulation of SERCA2a (sarcoplasmic reticulum Ca^{2+}-ATPase 2a) appears to play a critical role in the pulmonary vasculature in PAH (253). Although less is known about calcium handling in RV myocytes, recent data from a pig pulmonary vein banding model demonstrated downregulation of SERCA2a and endoplasmic reticulum stress (88). In a recent study, intracellular calcium recycling appeared similar to normal LVs in patients with idiopathic PAH but was depressed in patients with scleroderma-PAH (167). A follow-up study using skinned cardiomyocytes obtained from RV biopsies revealed significant sarcomere dysfunction in patients with scleroderma-PAH (254). Apelin, a potent proangiogenic and procontractile peptide that exerts critical effects in the LV (255), may also be of importance in the RV (223, 256).

Current knowledge gaps. More insight is needed regarding mechanisms of calcium homeostasis in the RV myocyte. The occurrence, timing, and pathologic relevance of changes in SERCA2a during RVH in PAH is unknown. Although selective pulmonary SERCA2a gene transfer may offer benefit as a therapeutic intervention for the vascular bed in PAH (253), it is unknown if cardiac-targeted SERCA2a therapy, an approach that was (so far, unsuccessfully) taken to treat patients with LV failure (257), would be beneficial for the RV. The role of mitochondrial calcium handling has not been assessed in RVF; this may be important, as the influx of calcium via the mitochondrial calcium uniporter is a major regulator of both cytosolic calcium and the intramitochondrial calcium stores, which regulate key enzymes in oxidative metabolism (258). Effects of neurohormonal modulators (e.g., inhibitors of the renin-angiotensin/aldosterone system) and B-adrenergic receptor antagonists on RV contractile function require further study. In particular, the effect of the latter is of great interest, because currently available studies yielded conflicting results of these drugs on clinical outcomes and RV function (259–263). RV inotropic agents may improve short- and/or long-term outcomes in acute and/or chronic PH, but their role (and potential for adverse events) is incompletely explored.

Recommendations/pathway for progress. More studies of the mitochondrial calcium uniporter and sarcoplasmic reticulum calcium handling system in preclinical models and humans are required. Studies of active and passive mechanisms that control RV diastolic and systolic function likely will identify new insights. In addition to addressing changes in fibrosis (which determines passive stiffness), more study of calcium pumps, gap junctions, and ion channels (which determine active stiffness) in RV myocytes is required. More studies focusing on the regulation of SERCA2a in RVF are needed. Acute and chronic effects of

Table 7. Research Priorities to Be Addressed in the Next 5 Years

<table>
<thead>
<tr>
<th>Number</th>
<th>Priority</th>
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<tr>
<td>1.</td>
<td>Understand the mechanisms underlying adaptive vs. maladaptive RV adaptation to pulmonary vascular load, including role of fibrosis, angiogenesis, metabolic shifts</td>
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<tr>
<td>2.</td>
<td>Develop a series of plasma and/or imaging biomarkers (by echocardiography and CMR) that could be used to accurately evaluate RV function, perfusion, and RV-PA coupling and as endpoints in clinical trials</td>
</tr>
<tr>
<td>3.</td>
<td>Develop and validate or improve on current animal models of RV dysfunction in the setting of increased pulmonary vascular load</td>
</tr>
<tr>
<td>4.</td>
<td>Develop therapies aimed at targeting RV myocardial contractility (e.g., calcium-sensitizing agents targeting sarcoplasmic reticulum and/or sarcomere function)</td>
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Definition of abbreviations: CMR = cardiac magnetic resonance imaging; PA = pulmonary artery; RV = right ventricular. The list was generated by surveying working group members to identify their top five priorities. The most commonly mentioned topics and themes are included in the table.


205. Turek Z, Grandtner M, Kreuzer F. Cardiac hypertrophy, capillary and muscle fiber density, muscle fiber diameter, capillary radius and diffusion distance in the myocardium of growing rats adapted to a simulated altitude of 3500 m. F Fl u g e r s Arch 1972;335:19–28.


