Exercise-Induced Pulmonary Hypertension: How to Define, Diagnose, and Treat At-Risk Patients

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ABSTRACT

Introduction: Pulmonary hypertension (PH) is a devastating disease that progresses rapidly, leading to high patient mortality. This condition is characterized by high blood pressure in the pulmonary vasculature and poor pulmonary perfusion, resulting in patient fatigue, dyspnea, and syncope—especially upon physical exertion. A sub-clinical form of PH is referred to as exercise induced pulmonary hypertension (EIPH), where patients display normal resting hemodynamic properties but abnormal pulmonary responses to exercise.

Discussion: Recent evidence suggests that early intervention and treatment of PH can improve patient outcomes. However, little clinical evidence exists to support effective treatments of EIPH. This lack is due in part to the removal of EIPH from official guidelines, such as the European Respiratory Society in 2008. EIPH was removed from clinical guidelines because of a lack of consensus on what constitutes EIPH and a lack of agreement on standardized testing procedures for diagnosing EIPH. Emerging evidence suggests that exercise testing following a standardized protocol of stress echocardiography or right heart catheterization may allow for better classification of EIPH. This review proposes that a mean pulmonary artery pressure/cardiac output slope > 3 mm Hg/L/min or mean pulmonary artery pressure > 30 mm Hg with a pulmonary vascular resistance > 3 Wood units is both a sensitive and specific enough threshold to diagnose EIPH and is sufficient to consider specific treatment options.

Conclusion: Providing evidence for a consensus on the definition of EIPH, along with a validated and standardized testing procedure, will hopefully foster the progression of research on EIPH and further the development of treatments to improve patient outcomes.

KEYWORDS
Pulmonary hypertension, Exercise-induced pulmonary hypertension, Cardiopulmonary exercise testing.
Pulmonary hypertension (PH) is a relatively rare yet devastating disease that is estimated to affect 1% of the global population under 65 and 10% over the age of 65. (1) PH is a progressive disease that results in decreasing exercise intolerance, dyspnea, fatigue, altered hemodynamic and cardiopulmonary function, and low long-term survival rates. (2, 3) If left untreated, the average length of survival for patients is approximately three years from the time of diagnosis. This delay is of concern because the average time from patient-described onset of PH symptoms to diagnosis of the disease is approximately 4 years. (4) In recent years, the prognosis for PH patients has been improved by the advent of new and improved therapies like pulmonary vasodilators and diuretics; however, a diagnosis of PH in Canada still increases the 1-year standardized mortality ratio by 7.2-fold. (5) Recent studies suggest that interventions conducted at earlier stages, or even pre-clinical stages of PH, may result in increased longevity and quality of life for patients. (6, 7) Therefore, a need exists for improved diagnostic methods to identify and treat patients with PH early in the development of the disease.

PH is characterized by abnormal pulmonary hemodynamics and high pulmonary blood pressure. This increased pulmonary blood pressure can be the result of several different underlying pathologies; however, the increase is typically caused by an obstruction to flow downstream of the right ventricle, either in the pulmonary arteries or the aortic valve. (8) Recently, the definition of PH was updated to be a resting mean pulmonary arterial pressure (mPAP) of 20 mmHg with a pulmonary artery systolic pressure (PASP) > 50 mmHg, or a pulmonary capillary wedge pressure (PCWP) < 20 mmHg during exercise. (8) However, in 2008, the official guidelines for pulmonary hypertension diagnosis and treatment removed EIPH as a subset of PH. (8) The old definition of EIPH was abandoned because of its failure to accurately and consistently differentiate healthy patients from those experiencing true EIPH. Failure to differentiate healthy individuals was especially true in patients with increased cardiac outputs (CO), as their mPAP values would often exceed the 30 mm Hg threshold established for EIPH. (19, 20) In one example, a study found that 26% of healthy patients displayed a mPAP > 30 mm Hg with exercise, especially as COs exceeded 10 L/min. (21) Another issue that resulted in the old definition being abandoned was disparity in the methodology and exertion metrics used to diagnose EIPH. (22) Some researchers proposed that stress echocardiography (SE) measurements during exercise were sufficient to diagnose EIPH, while others, including the Canadian Thoracic Society maintained that right heart catheterization (RHC) was necessary for diagnosis. (11, 23) Additionally, the formulas used to attain pulmonary pressure and vascular resistance varied between groups, as well as the actual exercise parameters used (ie, cycle ergometry vs treadmill tests). Therefore, consistency and accuracy in measurements was a major concern that ultimately forced the definition to be abandoned.

More recently, new thresholds for defining EIPH have been put forth; these new thresholds are demonstrated to be more sensitive and specific for defining EIPH. The
improvement of diagnostic techniques and methodology has led to more consistent measurements of hemodynamic properties. The definitions that were recently suggested for EIPH are a mPAP to CO slope of > 3 mm Hg/L/min (24, 25) or a mPAP >30 mm Hg with total peripheral resistance > 3 Wood units during exercise. (21) Since EIPH is viewed as a potentially more treatable stage of PH it is crucial to develop a standard diagnostic approach with reliable threshold parameters to accurately diagnose EIPH. (17) This review will briefly describe the etiology of PH and focus on creating a definition for EIPH, illustrating the gold standards for diagnosis, and discussing relevant interventions for patients.

2 | ETIOLOGY OF PH

PH is a progressive disease of abnormal pulmonary hemodynamics and increased pulmonary circulation pressures. (14) Currently there are 5 different clinical classifications of PH (Table 1) with different subclassifications based upon the mechanisms of onset. (9) The two most common causes of increased pulmonary pressure are increased PVR and increased left atrial pressure (LAP) (13, 26) In many cases, the initial onset of PH is idiopathic, however, the progression of EIPH typically stems from changes in the pulmonary vasculature that induce inflammation, vasoconstriction, cell proliferation, and hypertrophy. The result is exacerbated pulmonary damage inducing strain and dysfunction of the right ventricle (RV). (27)

It is important to try to understand the underlying etiology of the condition as it can help guide the treatment approach and the success of the therapy. For conditions with a known etiology, the treatment is often adapted to address the underlying cause of the PH. For example PH due to left heart disease is often treated with a mechanical intervention such as surgery to repair the defect (i.e. a valve replacement for a valvular disease). (8) For other etiologies where a mechanical intervention is not obvious or relevant, the endothelin (ET), nitric oxide (NO), and prostacyclin (PC) dependent pathways are currently being utilized as therapeutic targets in Canada. (23) ET activates two different receptor subtypes (ETA and ETB) present on vascular smooth muscle cells (vSMC) and endothelial cells (EC) (ETB only). Activation of ET receptors on vSMCs induces vasoconstriction, cellular proliferation, and hypertrophy, while activation on ECs results in vasodilation. (28) In PH it is believed there is a shift increasing ET expression on vSMCs that results in the deleterious phenotype associated with PH; therefore, ET antagonists such as Ambrisentan have been employed with relative success. (23, 29, 30) NO is a potent vasodilator that also inhibits platelet aggregation and thrombosis. Under normal conditions, ECs continuously produce NO using nitric oxide synthase (NOS), causing relaxation of vSMCs and inhibition of cellular proliferation. (31, 32) In PH, levels of NOS are greatly reduced resulting in an increased vasoconstriction and cellular hypertrophy. (29) Prostacyclin (PC) is produced in ECs through prostacyclin synthase and acts as a potent vasodilator. (27) PC binds to prostaglandin receptors and induces relaxation of vSMCs. In PH PC levels are diminished, contributing to the phenotype of vasoconstriction and cellular hypertrophy observed in PH. (33) Since all three of these pathways contribute to rising PVR and increased pulmonary pressures, they have been investigated extensively and make up the bulk of therapeutic targets for patients suffering from PH. (8, 23)

3 | MEASURING EIPH AND COMMON TOOLS USED FOR DIAGNOSIS

A common complaint of patients with undiagnosed EIPH or PH is exertional fatigue/dyspnea. Therefore, SE is a common starting diagnostic procedure followed by chest x-rays, electrocardiograms (EKG), echocardiograms (ECG), chest CT or MRI scans, and blood tests to identify the underlying cause and rule out conditions other than PH. (14) As exertional dyspnea/fatigue is the most frequent presenting symptom, it is often useful to attempt diagnosis of EIPH or PH using cardiopulmonary exercise testing (CPX). (22) However, some con-
cerns have been raised about making patients with compromised cardiopulmonary systems complete exercise testing. Despite concerns, a comprehensive study involving over 4000 patients who underwent CPX while presenting with high risk cardiac diseases found an adverse event rate of only 0.16%. (34) Looking at the 194 patients with PH in the study, there were 0 adverse events after CPX in PH patients providing reasonable proof of the safety of CPX in PH patients. CPX usually includes monitoring ventilatory-perfusion parameters along with right and left heart functioning either through non-invasive ECG or invasive RHC. (16, 22) From these tools, values of CO, PVR, PCWP (which estimates left atrial pressure), right ventricle regurgitation velocity, mPAP, PASP, ventricular elasticity and wall thickness, right atrial pressure, peak VO2, lactic acidosis, CO2 output, oxygen saturation, and minute ventilation (VE/VCO2) can be generated and used to assess the presence and extent of the progression of disease.

4 | CHALLENGES IN DIAGNOSING IN EIPH

Many of the parameters used to diagnose EIPH are estimated from derivations of the following formula for mPAP, assuming an ohmic linear relationship between the parameters. (22)

\[ mPAP = PVR \times CO + LAP \]

The assumption being that any changes in PVR or CO will result in linear increases of mPAP, and that increases in LAP through backflow will increase mPAP in a 1:1 ratio. However, this does not consider the dynamic responses of the pulmonary vasculature in responses to changes in pressure. In normal healthy individuals, pressure increases are quickly met with pulmonary dilatation to normalize flow. Therefore, past predictions of mPAP from this formula may be inaccurate. Herve and colleagues (21) demonstrated that this formula for PVR only provided a 48% sensitivity, unless the patient had a purely vascular etiology (implying limited distensibility of the vasculature leading to a more linear relation). A more comprehensive alternative formula has been proposed to take into account the distensibility of the venous system. (22, 35)

\[ mPAP = \frac{(1 + aLAP)^5 + (5aRo \times Q)^\frac{1}{5} - 1}{\alpha} \]

Where Ro is the resting total pulmonary resistance, as defined by the ratio of mPAP/CO, Q is pulmonary blood flow and \( a \) is the distensibility coefficient of the vasculature. The coefficient can be obtained by measuring the pulse pressure of a large artery, but is often assumed as 2% in most species. (22) This formula better represents the pressure-flow relation shown in a compliant vasculature and could potentially lead to more accurate mPAP approximations.

Another issue that arises from the way values are commonly derived from functional tests is the calculation of CO. It is often cited in literature that CO can be approximated equivocally from thermodilution or direct Fick measurements. (36) However, previous CO measurements had only been validated in resting patients and Hsu and colleagues found that using thermodilution actually greatly overestimated the prevalence of EIPH in patients, with a 20% increase in false diagnoses. (37) As CO increased in patients, the separation in predicted Fick and thermodilution COs increased, with thermodilution underpredicting CO relative to Fick measurements. This is extremely concerning, given the importance of accurate CO measurements in predicting other important parameters for diagnosing EIPH, such as total pulmonary resistance and PVR. Therefore, it is possible that the prevalence of EIPH has been overreported in the literature where studies used thermodilution.

Currently, there is a consensus on what constitutes normal resting values for many of the parameters described above, (i.e. mPAP <20mmHg). However, the definitions for what constitutes normal hemodynamic parameters during exercise remain elusive. Especially since some of the techniques used to diagnose PH at rest have not been well validated for measurements of
the same parameters during exercise.

4.1 | Difficulty in consistent measurements

A major difficulty in diagnosing EIPH is making consistent measurements on patients during exercise. Performing invasive RHC or SE on an actively moving patient can be a difficult procedure. In addition, normalizing the workload between patients to establish a normal PAP is challenging. (16) Often a cycle ergometer is used with patients in an upright position, with work values ranging from 10-30 W; however, other methods have been used such as treadmills or supine leg press. (22) This can lead to disparity in results, since patient stance/posture can have substantial effects on intrathoracic pressure and concomitantly on pulmonary pressure. (17, 38) SE is often used as a diagnostic tool as it is readily available in most clinics and is a non-invasive measurement, but SE has not been well validated for use during exercise. (18) One issue with using SE during exercise is that predictions of right atrial pressure often rely on measurements of the distension of the inferior vena cava. However, it is known that exercise lowers venous compliance and therefore can reduce the accuracy of observations made by SE. (18, 39) However, performing SE after the patient has finished exercise runs the risk of missing important data, as hemodynamic parameters quickly return to resting levels within five minutes post-exercise. (40) Another issue with SE, is that PCWP, which is a standard measurement in the diagnosis of PH, cannot be calculated without direct invasive measurements such as RHC. Yet invasive measurements are not without their own problems, as they can be difficult to obtain during exercise because of large swings in intrathoracic pressure that accompany heavy breathing during exercise. (16)

4.2 | Variation in measurements of healthy individuals

One of the main reasons EIPH was removed from the official guidelines in 2008, and not reinstated in the updated guidelines released in 2019, was the lack of consensus among experts on what constitutes normal pulmonary pressures during exercise. Initially EIPH was defined as a mPAP>30 mmHg, but multiple studies have found that healthy individuals consistently exceed mPAP values of 30 mmHg during strenuous exercise, especially in well-trained athletes. (19-21) The caveat being that the increased mPAP and PASP seen with these individuals were accompanied by an increased CO, which fits in line with the observation that healthy individuals do not exceed a mPAP/CO slope of 3 mmHg/L/min. (19, 41) As well, studies have shown that 6% of otherwise healthy individuals over 50 have a PASP>40 mmHg as well as 5% of people with a BMI >30. (42) More recent studies have shown that healthy (non-athletes) cannot exceed PASP values of 40-45 mmHg at CO <20L/min during exercise. (41, 43) However, these studies excluded athletes.

PASP is most often estimated from the following formula:

\[ \text{PASP} = 4V^2 + \text{RAP} \]

Where V is the maximum velocity of the tricuspid valve regurgitant jet and RAP is the right atrial pressure. Tricuspid valve regurgitation itself has been suggested as a defining criteria for EIPH as a regurgitant jet velocity of >3.0m/s, but in line with the findings that PASP can often exceed EIPH definitions, researchers found that athletes had an average peak tricuspid regurgitation velocity of 3.41m/s. (44) Therefore, the large variations of hemodynamic parameters in healthy individuals and trained athletes adds to the challenge of finding a single parameter to define EIPH.

4.3 | Predictive ability of Different Diagnostic Tools

The multifactorial nature of PH requires a large repertoire of diagnostic techniques to pinpoint the etiology of each patient’s PH. Several methods that are effective in the initial screening of patients have proven to
be ineffective at diagnosing EIPH when used in isolation. Electrocardiograms, although useful for providing direction for further investigation, were shown to be inadequate at diagnosing PH with a sensitivity (true positive rate) of only 55% and a specificity (true negative rate) of 70%. (45) Minute ventilation (VE/VCO2) at ventilatory threshold and peak alveolar-arterial difference in PO2 were not sensitive enough to distinguish EIPH from normal. (18) The lack of standardized procedures for SE has led to both over and underestimations of PAP. (3) When compared to RHC, SE displays variable correlation values between 0.57-0.93 with PASP measurements (11). A more recent study found that 35 of 100 patients identified to have EIPH according to SE, actually had normal hemodynamics via RHC results. (15) Therefore caution must be used when interpreting SE results.

The current gold standard for diagnosing EIPH or PH is RHC. The biggest downfall of RHC is lack of availability and the inconvenience for the patient in using an invasive measurement. For this reason and also for simplicity, SE is often the preferred first method for screening patients of suspected EIPH or PH. (22, 23) Despite the limitations of SE, it allows for quick, easy and relatively reliable estimates, when conducted properly, of PASP, PVR, and pericardial effusions, as well as overall morphology of the heart and valves allowing for diagnoses of RV size and function, RA size, diastolic dysfunctions, valvular functions, patent foramen ovale, and intrapulmonary shunts. (45) However, the importance of standardized techniques used during SE to ensure more consistent and better interpretation of results cannot be understated.

5 | PROPOSED STANDARD PROTOCOL AND DEFINITION OF EIPH

Creating a definition for EIPH is challenging as it exists in an intermediary state between normal values and overt PH. Normal hemodynamic values used for diagnosis of PH have been shown to vary substantially even among healthy control patients during exercise, making it difficult to isolate clear standards for the subset of patients displaying EIPH. The traditional definition of EIPH being a mPAP > 30 mmHg, PASP > 50, or PCWP < 20 mmHg during exercise, proved to be inefficient at adequately separating the distinct patient populations (healthy vs EIPH vs overt PH). Therefore, efforts have been made to revise and improve upon these thresholds to find parameters that are more sensitive and specific. It seems clear that a singular definition to define EIPH is not reasonable. Instead, a consensus between several parameters should be used for diagnosis. The best and most current thresholds for an EIPH diagnosis is: a mPAP > 30 mm Hg with CO < 10 L/min or a mPAP/CO slope > 3 mmHg/L/min; or a mPAP > 30 mmHg with a resistance greater than 3 Wood units (Table 2). These thresholds are shown to be 93% sensitive and 100% specific. (21, 24, 25) Several other parameters are useful for unlocking the etiology of EIPH, albeit insufficient for diagnosis. VE/VCO2 can be useful for separating stages of PH, as VE/VCO2 increases as PH worsens, but is unchanged in EIPH patients compared to normal. (46) Whereas PASP and PCWP can be used as indicators for the presence of heart disease (45) Barst and colleagues describe the ideal testing methodology in their review, from which we propose a standardized methodology to test for EIPH (Figure 1). (45)

6 | SHOULD WE TREAT EIPH

The lack of a current formally acknowledged definition for EIPH does not mean the disease does not merit treatment. However, at the time of this review, a limited number of studies have been conducted to investigate the efficacy of treating patients with EIPH. This phenomenon is despite the fact that numerous studies have shown that the earlier the intervention, the better the prognosis for patients with PH. (8, 47-49) PH therapy has been shown to slow and prevent further disease progression as well as improve hemodynamic properties of the patients. (6) Kovacs et al (47) reported greater decrease in PAP and PVR with intervention earlier in the stages of disease progression (lower WHO functional class). Ear-
1. Include patients with exercise intolerance whose resting mPAP < 20 mmHg.

2. Dynamic exercise in supine position on a stress echocardiography bed, if possible, to allow for constant measurement with SE while patient pedals on a cycle ergometer.

3. Use estimated exercise capacity to design number of work steps and work increments to reach maximum tolerable workload within 10-15 minutes.

4. Take baseline measurements while patient is pedaling at a workload of 0 watts.

5. Increase workload increments at constant rate, between 10-30 W.

6. Constantly measure mPAP using peak pulmonary regurgitation Doppler signal.*


8. Stop procedure once the thresholds for EIPH have been met (mPAP/CO slope > 3 mmHg/L/min or mPAP > 30 mmHg + TPR > 3 WU, or patient reaches maximal tolerable workload).

9. If uncertainty in the results of SE proceed to RHC and repeat steps 2-8.

**FIGURE 1** Proposed standardized protocol for exercise hemodynamic testing.

*For a comprehensive review on SE techniques, see reference (51).

lifer treatment has also been suggested to lead to less disease morbidity. (48, 49) A recent study demonstrated that patients with EIPH (defined as a mPAP/CO slope > 3 mmHg/L/min during exercise) had a 2-fold increase in the hazard ratio of a future cardiovascular event or death compared to patients with a normal mPAP/CO slope during exercise. (50) As EIPH is considered by many to be a distinct and early stage in the progression of PH, it stands to reason that early intervention will delay the progression into classically defined PH, extend the expected lifespan of patients, and greatly improve patients’ future quality of life. Not only does early diagnosis lead to a better prognosis but also to more robust treatment options for the patients. Late stages of the disease often require continuous IV administration of drugs, which greatly diminishes patient quality of life. Whereas for earlier stages, patients often receive oral therapies which are less intrusive and more convenient.

Currently, no drugs are specifically tailored for treatment of EIPH; however, some clues can be taken from a study that looked at scleroderma patients displaying normal pulmonary hemodynamics but abnormal exercise hemodynamics at rest. Saggar et al (48) showed that invasive measurements were easily able to distinguish different stages of abnormal hemodynamic progression, and the authors suggested that early treatment of patients with EIPH-like symptoms using a pulmonary vasodilator (ambrisentan) may yield promising results. Many patients whose PH is derived from pulmonary vascular disorders are prescribed pulmonary vasodilators, and the results from these studies may yield some additional insight into EIPH treatment. The more recent therapy of choice for treatment of early stages of PH are endothelin blockers (ETA and ETB). One such drug currently on the market is bosentan, which has been shown to improve exercise capacity (increased Six-Minute Walk Test levels), vascular resistance, and cardiac index. (6) Bosentan was also shown to decrease brain natriuretic peptide levels which along with improved vascular resistance are predictive factors for survival. The most promising characteristic of endothelin blockers so far is the lack of reported adverse side-effects. The most common reported side-effect has been increased liver amminotransferase levels and has yet to be correlated with adverse events. (6) Other drug targets include prostanoids and phosphodiesterase-5 inhibitors. However, these therapies are less preferred for treatment of earlier stages of EIPH because they either have more adverse side-effects or call for less-convenient administration methods, like IV injections. (27)

In Canada, there are currently 10 PH-specific treatments that are approved and have significantly improved patient quality of life and long-term survival. (23) Despite the promising results of these therapies in treating many conditions, they have proved less effective in treating PH caused by systolic and/or diastolic dysfunctions. (49) Typically, anti-clotting agents, such as warfarin, and diuretic agents are prescribed to patients with resting PH; however, whether these treatments would be beneficial for patients suffering from EIPH is uncertain. (27) Calcium channel blockers may also be pre-
scribed to PH patients, but these drugs’ safety and efficacy profiles may not warrant use in patients with EIPH. Finally, dual therapy has been shown to provide an additional advantage when compared to monotherapy, yet again, no evidence exists for efficacy of dual therapy in treating EIPH patients.

In general, very little data supports the efficacy of treating EIPH. Nevertheless, studies evaluating the effect of treating early stages of PH appear to be advantageous. With the fast progression of PH and high mortality rate associated with the disease, more studies evaluating the efficacy of EIPH treatment are necessary. As most cases of EIPH are believed to be the result of pulmonary vascular diseases or left heart disease (both of which increase PVR and pulmonary pressure), pulmonary vasodilators and ET antagonists would appear to be the most promising therapy for treating patients with EIPH. (22, 27) Consequently, further studies are needed to provide a conclusive argument for the efficacy of treating EIPH.

7 | CONCLUSION

EIPH has consistently been shown as a distinct entity that is intermediate between normal hemodynamics and overt pulmonary hypertension. However, the lack of consensus on defining criteria for EIPH and an absence of standardized treatment methodologies has led to the poor state of knowledge on treatment efficacy for this subset of patients. With evidence showing that treatment of the early stages of PH leads to increased longevity and better quality of life for patients, it is crucial to evaluate if these results are transferable to patients suffering from EIPH. This review proposes that a mPAP/CO slope > 3 mmHg/L/min or mPAP >30 mmHg + TPR > 3 Wood units is both a sensitive and specific enough threshold to diagnose EIPH and is sufficient to consider specific treatment options. Having a set EIPH definition for clinicians along with the standardized protocol provided here will allow research to move forward on treatments for EIPH, and hopefully lead to a better future prognosis for patients who develop pulmonary hypertension.

REFERENCES


1. PAH
   1.1 Idiopathic PAH
   1.2. Heritable PAH
   1.3. Drug- and toxin-induced PAH
   1.4. PAH associated with:
      1.4.1. Connective tissue disease
      1.4.2. HIV infection
      1.4.3. Portal hypertension
      1.4.4. Congenital heart disease
      1.4.5. Schistosomiasis
   1.5. PAH long-term responders to calcium channel blockers
   1.6. PAH with overt features of venous/capillaries (PVOD/PCH) involvement
   1.7. Persistent PH of the newborn syndrome
2. PH due to left heart disease
   2.1. PH due to heart failure with preserved LVEF
   2.2. PH due to heart failure with reduced LVEF
   2.3. Valvular heart disease
   2.4. Congenital/acquired cardiovascular conditions leading to post-capillary PH
3. PH due to lung diseases and/or hypoxia
   3.1. Obstructive lung disease
   3.2. Restrictive lung disease
   3.3. Other lung disease with mixed restrictive/obstructive pattern
   3.4. Hypoxia without lung disease
   3.5. Developmental lung disorders
4. PH due to pulmonary artery obstructions
   4.1. Chronic thromboembolic PH
   4.2. Other pulmonary artery obstructions
5. PH with unclear and/or multifactorial mechanisms
   5.1. Hematological disorders
   5.2. Systemic and metabolic disorders
   5.3. Others
   5.4. Complex congenital heart disease

LVEF, left ventricular ejection fraction; PAH, pulmonary arterial hypertension; PCH, pulmonary capillary hemangiomatosis; PH, pulmonary hypertension; PVOD, pulmonary veno-occlusive disease. Reproduced from Simonneau et al. under Creative Commons Attribution-NonCommercial 4.0 International (CC BY-NC 4.0). (9)

**TABLE 1** Updated clinical classification of pulmonary hypertension
### Proposed hemodynamic definitions of pulmonary hypertension and exercise induced pulmonary hypertension

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<thead>
<tr>
<th>Definition</th>
<th>Characteristics</th>
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<tr>
<td>Pulmonary hypertension</td>
<td>Resting mPAP 20 mmHg</td>
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<td></td>
<td>Resting PVR 3 Wood Units</td>
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<tr>
<td>Exercise Induced pulmonary hypertension</td>
<td>mPAP/CO slope &gt; 3 mmHg/L/min during exercise</td>
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<tr>
<td></td>
<td>mPAP &gt; 30 mmHg with a PVR &gt;3 Wood Units during exercise</td>
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