Bronchopulmonary dysplasia-associated pulmonary hypertension

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Abstract: Chronic respiratory sequelae are a major health problem in surviving premature infants. Despite immense improvement in the care of premature infants, the rate of bronchopulmonary dysplasia (BPD) and pulmonary morbidities have remained the same or even slightly increased over the past decades. BPD results from the impairment of lung development following premature birth, and the resultant dysregulated pulmonary vasculature leads to chronic pulmonary hypertension (PH). It is seen more commonly in the setting of moderate to severe BPD. A high index of suspicion is necessary to diagnose PH in BPD. Diagnosis is most commonly done by echocardiogram, but cardiac catheterization may be required to delineate the cardiac structure, cardio-pulmonary vasculature and myocardial function. Once the diagnosis is made, a multidisciplinary approach for management of these patients is suggested, aiming to optimize management of BPD and use of various pulmonary vasodilators. The various therapeutic agents available help reduce pulmonary vascular resistance by acting on the nitric oxide (NO), prostacyclin and/or endothelin pathways. Serial echocardiograms and measurement of biomarkers are used to evaluate the response to therapy. Currently there is not much evidence to support optimal management approaches, but the recent guidelines published by the Pediatric Pulmonary Hypertension Network (PPHNet) provide practical recommendations for screening and management of these infants.

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Introduction

The term ‘bronchopulmonary dysplasia’ (BPD) was first used by Northway et al. to describe a chronic form of neonatal lung injury associated with barotrauma in preterm infants (1). Although there has been significant advancement in this field over the last couple of decades including sophisticated, newer and gentler methods of mechanical ventilation (MV), surfactant replacement therapy etc., the prevalence of BPD remains quite high. Classical or “old BPD”, present in the pre-surfactant era, was characterized by diffuse proliferative lung disease and postnatal lung injury. Jobe coined the term “new BPD” in 1999, to refer to the chronic lung disease of preterm infants at that time (2). This “new BPD” demonstrated much less alveolar septal fibrosis and airway damage when compared to its old counterpart, and was characterized with alveolar simplification and dysmorphic microvasculature (3). This dysmorphic growth and function of the pulmonary vasculature can cause pulmonary hypertension (PH) in infants with BPD. Bonikos et al. (4) first described cardiac atrial or ventricular stress, including cor pulmonale, in infants with BPD and since then it has been increasingly recognized in this population. A recent meta-analysis reported an accumulative estimated prevalence of PH in BPD to be 20%. The prevalence was estimated to
be as high as 40% in severe BPD (5). BPD and BPD-associated PH (BPD-PH) have been correlated with devastating short- and long-term consequences for infants born prematurely. Retrospective studies demonstrate a 2-year morbidity rate of 26–47% in patients with BPD-PH (6,7). The pathophysiology of PH in BPD is not well understood at this time and there is a lack of standardized criteria for identifying preterm infants that are most at risk for developing BPD-PH. In this article we will review the pathophysiology, clinical presentation, diagnosis, suggested management and outcomes of infants with PH in the context of BPD.

**Pathophysiology of BPD-PH**

The pathogenesis of PH in BPD is affected by various maternal, genetic and environmental factors. Premature birth exposes the developing lung to an environment that can alter the normal developmental process and result in impaired lung development (8-10). Increased risk of BPD and PH has been shown in premature infants exposed to pre-eclampsia (11,12) and in infants with intrauterine growth restriction (13,14) due to disruption in vascular signaling pathways. Abnormal growth of the pulmonary circulation in BPD is characterized by decreased vascular branching, an altered pattern of vascular distribution within the lung interstitium, and persistent intrapulmonary venous anastomoses (15). Disruption in vascular growth results in decreased vessel density throughout the pulmonary microcapillary network (16). This leads to decreased cross sectional area for pulmonary blood flow and increased pulmonary vascular resistance, which in turn alters pulmonary vasoreactivity and causes vascular remodeling. This dysregulated pulmonary vasculature has overall less area for gaseous exchange. In patients with BPD who have airway and parenchymal injury and inflammation, this results in hypoxic vasoconstriction and impaired pulmonary blood flow, especially under stress (17). Chronic hypoxia also contributes to pulmonary arterial vasoreactivity, leading to vascular remodeling with intimal hyperplasia and increased muscularization of small pulmonary arteries. Endothelial injury due to environmental stressors induces smooth muscle cell proliferation in the media of the small pulmonary arteries, precocious maturation of immature mesenchymal cells into mature smooth muscle cells and incorporates fibroblasts into the vessel wall. Such pulmonary vascular remodeling, along with failure of the lung vasculature to catch up with the normal term lung, may contribute to the pulmonary vessel disease associated with BPD (18).

While the lung pathology of infants with PH and BPD has many common elements, it is important to note that not all infants with BPD develop PH; it is more commonly seen in infants with severe BPD (6). The index of suspicion for diagnosing PH is also higher in this subgroup and they are screened more frequently for PH. Infants with other associated conditions like large patent ductus arteriosus, pulmonary vein stenosis (PVS) or aorto-pulmonary collaterals are more at risk for developing severe BPD and PH (7). Infants with lower baseline oxygen saturations or frequent hypoxic episodes are also at higher risk for developing PH due to recurrent episodes of hypoxia and increased vasoreactivity. Maternal chorioamnionitis and postnatal lung infections also have a negative influence on lung development and may lead to abnormal lung vasculature growth (19). Lastly, BPD has a strong genetic predisposition (8), and so it can be postulated that development of PH in BPD also has some genetic influence (20,21). A proposed schema of the interacting factors involved in the pathogenesis of BPD-PH is illustrated in Figure 1.

**Clinical features**

Infants with BPD who develop PH may be asymptomatic or may present with increased need for respiratory support, failure to wean from respiratory support, frequent hypoxic events and/or impaired somatic growth due to increased metabolic demands. Many institutions screen infants with BPD for PH. Asymptomatic infants may be diagnosed as a result of such screening echocardiograms or incidentally while looking for other pathology. Symptomatic infants may have sudden deterioration in their clinical status and have a pulmonary hypertensive crisis. Such episodes may be idiopathic in origin or may be precipitated by infection (systemic sepsis, respiratory or urinary tract infections), aspiration events or changes in lung pathology (atelectasis, mucous plugging, etc.). On the other hand, some infants may have gradual deterioration in their clinical status and have a more insidious onset. They may initially present with a gradual increase in oxygen requirement and/or ventilator support and may progress to develop right heart dysfunction (19). It is important to have a high index of suspicion for PH in these infants so as to be able to timely diagnose them and adequately manage this disease.
Screening and diagnosis of BPD-PH

The Pediatric Task Force at the 6th World Symposium on PH suggested a recent update to the definition of pediatric PH, where they now suggest including patients with mean pulmonary artery pressure >20 mmHg and pulmonary vascular resistance ≥3 Wood units (WU) to identify pre-capillary PH (22). Most neonatal intensive care units use a clinical algorithm for the identification and evaluation of infants at risk for BPD-PH. However, adoption of a standardized clinical algorithm is a challenge given the lack of agreed-upon definition of PH in this setting and inadequacy of echocardiography to detect PH, which is the most commonly used screening tool. In 2015, the American Heart Association (AHA) and American Thoracic Society (ATS) published recommendations for the care of patients with pediatric PH. These guidelines provided the initial framework for screening and care of infants with BPD-PH (23). In 2017, the Pediatric Pulmonary Hypertension Network (PPHNet), a multi-disciplinary panel of PH experts published more detailed guidelines that provide practical clinical recommendations for the evaluation, diagnosis and management of infants with BPD-PH (24).

These guidelines suggest obtaining an echocardiogram for premature infants with severe hypoxic respiratory failure with minimal parenchymal disease and those that still need MV at 7 days or more in order to identify infants with acute PH that might benefit from therapy. The guidelines also suggest getting an echocardiographic evaluation for infants with diagnosis of mild or moderate BPD (at 36 weeks postmenstrual age) and those with sustained need for significant respiratory support at any age. They suggest monitoring infants with mild BPD clinically, with echocardiography to be obtained only if they clinically deteriorate (24). At our institution, we screen all patients with the diagnosis of BPD for PH, irrespective of the

Figure 1 Pathogenesis of pulmonary hypertension in BPD. Dysregulated vascular and alveolar development in BPD leads to morphological changes in the lung causing pulmonary hypertension. ROS, reactive oxygen species; BPD, bronchopulmonary dysplasia.
severity of their disease. Figure 2 illustrates the protocol for screening infants with BPD for PH used at St. Christopher's Hospital for Children, Philadelphia, PA, USA. The most commonly used modalities in the screening and diagnosis of PH are discussed below.

**Echocardiogram**

This is the most commonly used screening tool for PH in infants with BPD. However, the echocardiogram uses somewhat subjective criteria for diagnosis, and it does not have great inter-rater reliability. The PPHNet guidelines suggest that a complete echocardiogram for PH screening in preterm infants should include, at a minimum: (I) a complete anatomic evaluation, to identify and characterize the physiologic contribution of structural abnormalities, shunts and pulmonary veins; (II) assessment of right and left ventricular size, hypertrophy; systolic and diastolic function; (III) systolic and diastolic interventricular septal position; (IV) tricuspid regurgitation jet velocity (TRJV) when present; and (V) documentation of the systemic blood pressure at the time of the study (24). Although TRJV is the most frequently used measurement for evaluating PH, many infants do not have a detectable TRJ at the time of the study and may also be affected by the lung inflation (25). In a study done to evaluate clinical utility of echocardiograms in the diagnosis of PH in infants with chronic lung disease, Mourani et al. found that when compared with cardiac
catheterization measurements, echocardiographic estimates of systolic pulmonary artery pressure correctly diagnosed the presence or absence of PH in 79% of the studies but determined the severity of PH (severe PH was defined as pulmonary/systemic pressure ratio of ≥0.67) correctly in only 47% of those studies. In the absence of a measurable TRJV, qualitative echocardiographic findings, including right atrial enlargement, right ventricular hypertrophy, right ventricular dilation, pulmonary artery dilation, and septal flattening, either alone or in combination, have relatively poor predictive value (26).

A growing body of evidence also suggests that acquired PVS can result in obstructed pulmonary blood flow and contribute to increased pulmonary pressures. A recent study of 213 infants with severe BPD demonstrated the presence of PVS in 5% of these infants and these infants were noted to have worse outcomes with higher mortality (27). Although echocardiography may be able to identify most of these cases, cardiac catheterization or detailed imaging may be required for better evaluation of the vessel involvement.

**Biomarkers**

Currently there is no single reliable biomarker that can diagnose, predict severity and monitor response to therapy. Serum brain natriuretic peptide (BNP) and N-terminal-pro-brain natriuretic peptide (NT-proBNP), which are released by the myocardium in response to stretch, are the most commonly used biomarkers for monitoring changes in disease status in infants with known PH, but are not specific for right ventricular stress or PH. The PPHNet guidelines suggest using serial BNP or NT-proBNP levels to monitor disease progression/regression, response to therapy, in conjunction with echocardiography rather than in isolation (24). Various other biomarkers are currently being explored in the setting of BPD that can be used to identify the infants most at risk for developing severe disease and/or PH. Identification of such novel biomarkers will potentially greatly facilitate the diagnosis and management of these infants (29).

**Management**

The evaluation and management of infants with BPD-PH is an area of immense debate and the practices vary greatly between individuals and institutions. The current evidence suggests that there are significant comorbidities present in infants with this diagnosis. Hence, they should undergo a comprehensive evaluation of the parenchymal lung and large airway disease prior to initiation for therapy for PH. This should include evaluation for hypoxemia, gastroesophageal reflux, aspiration, vocal cord abnormalities, subglottic stenosis, laryngo-bronchomalacia, pulmonary stenosis, cardiac dysfunction and other cardiac anomalies. Oxygen therapy should be optimized with a goal to keep target saturations between 92–95% to prevent the adverse effects of hypoxia without causing additional lung injury by hyperoxia (24).

The suggested threshold for initiation of vasodilators includes TRJV >3 m/second, estimated right ventricle/left ventricle systolic pressure >0.5, and septal flattening in absence of a significant left to right shunt. For cardiac catheterization significant measurements would include a ratio of pulmonary artery to systemic pressure ≥0.5, indexed pulmonary venous resistance ≥3 WU, or pulmonary/systemic resistance (Rp/Rs) ≥0.5, and a normal wedge or left ventricle end diastolic pressure without evidence of significant PVS. The various agents used in the treatment...
of PH in BPD are discussed below.

**Management of acute hypertensive crisis and use of nitric oxide (NO)**

Acute exacerbations of PH occur in infants with BPD, e.g., due to viral infections or mechanical airway obstruction. These infants can have a labile respiratory status and have profound hypoxemia with hypotension in response to routine daily care interventions. An initial evaluation should be performed with a chest X ray to rule out atelectasis, pneumonia and/or other parenchymal lung pathologies. Ventilator support must be optimized, and treatment of the underlying cause should be done—including antibiotics for infection if warranted, treatment of gastroesophageal reflux and steroids for inflammation, if needed (24). Once optimized, a dose of 10–20 ppm of inhaled NO (iNO) should be initiated for the acute PH crisis. Once the infant is stabilized and improving, iNO should be weaned initially aggressively to 5 ppm and then gradually by 1 ppm at selected intervals based on the infant’s tolerance. NO produced by endothelial cells stimulate soluble guanylate cyclase to increase cyclic guanosine monophosphate (cGMP), which causes relaxation of the vascular smooth muscle cell. iNO acts as a potent pulmonary vasodilator by acting on the same pathway (30). cGMP is an intracellular mediator that controls vascular contractility via NO activity. Increased concentration of PDE-5 isoenzyme causes degradation of cGMP through hydrolysis. Sildenafil acts by inhibiting the PDE-5 and increasing levels of cGMP, thereby augmenting vasodilation. In 2005 the Food and Drug Administration (FDA) approved the use of sildenafil for PH in adults and it has been widely used off-label for PH in neonates due to its ease of administration compared to other PH medications. Despite widespread use, there is minimal data supporting its efficacy in treating PH in neonates and infants (32,33). In 2012 the FDA issued a warning against the use of sildenafil in children between 1–17 years of age based on the data from the STARTS-2 trial that showed a high mortality in older children taking higher dose of sildenafil (>3 mg/kg/day) (34). In a retrospective cohort study of 269 pediatric patients with PH (50% of which had BPD-PH), use of low dose sildenafil (2 mg/kg/day or less) was well tolerated and was found to be safe with an acceptable side effect profile. Infant with BPD-PH were most likely to show improvement in PH, allowing for discontinuation of sildenafil in 45% of patients in that category (35). Tadalafil is a longer acting PDE5 inhibitor that has been approved for management of adult PH, but its use and efficacy in pediatric PH is still not known.

**Pharmacotherapy**

PH present in BPD consists of 2 components—a fixed component, due to decreased cross sectional area of pulmonary vasculature in the setting of arrested lung development and remodeling of pulmonary arteries; and a responsive component presumably due to pulmonary vascular smooth muscle, which responds to vasodilators. The mainstay of management has been optimization of BPD management to promote lung development and use of pulmonary vasodilators. The therapeutic agents used are either vasodilators that act on the NO pathway or prostacyclin pathway, or inhibitors of vasoconstrictors that modulate the endothelin pathway (30). These drugs have been summarized in Table 1.

**Drugs acting on the NO pathway**

**Sildenafil**

cGMP is an intracellular mediator that controls vascular contractility via NO activity. Increased concentration of PDE-5 isoenzyme causes degradation of cGMP through hydrolysis. Sildenafil acts by inhibiting the PDE-5 and increasing levels of cGMP, thereby augmenting vasodilation. In 2005 the Food and Drug Administration (FDA) approved the use of sildenafil for PH in adults and it has been widely used off-label for PH in neonates due to its ease of administration compared to other PH medications. Despite widespread use, there is minimal data supporting its efficacy in treating PH in neonates and infants (32,33). In 2012 the FDA issued a warning against the use of sildenafil in children between 1–17 years of age based on the data from the STARTS-2 trial that showed a high mortality in older children taking higher dose of sildenafil (>3 mg/kg/day) (34). In a retrospective cohort study of 269 pediatric patients with PH (50% of which had BPD-PH), use of low dose sildenafil (2 mg/kg/day or less) was well tolerated and was found to be safe with an acceptable side effect profile. Infant with BPD-PH were most likely to show improvement in PH, allowing for discontinuation of sildenafil in 45% of patients in that category (35). Tadalafil is a longer acting PDE5 inhibitor that has been approved for management of adult PH, but its use and efficacy in pediatric PH is still not known.

**Milrinone**

Milrinone is a PDE-3 inhibitor that increases cyclic adenosine monophosphate (cAMP) by inhibiting its breakdown in the arterial smooth muscle cells and myocardium. By increasing cAMP in cardiac muscle and vascular cells, it causes relaxation of vascular smooth muscle, enhances myocardial contractility (inotropy) and improves myocardial relaxation (lusitropy) (36). Hence it improves cardiac function both directly and by reducing afterload. It can also be used as an adjuvant to iNO to promote pulmonary vasodilation (37). Since it can only be used as a continuous infusion, its use is limited for hospitalized infants with acute exacerbations.
Drugs acting of the endothelin pathway

Bosentan
Endothelin-1 (ET-1) is a potent vasoconstrictor that is produced in response to hypoxia. It promotes smooth muscle proliferation, endothelial cell dysfunction, inflammation and fibrosis (37). ET-1 acts via two G-protein coupled receptors—ET_A that occurs on vascular smooth muscle cells, and ET_B that is present on endothelial and vascular smooth muscle cells. ET-1 is strongly implicated in the pathogenesis of PH in various clinical and preclinical studies (38,39). Bosentan is an antagonist of both ET_A and ET_B receptors (with a slightly higher affinity for ET_A) and is used extensively in adults with PH (39). In 2017, bosentan was approved for use by the FDA for PH in children 3 years of age or older and its experience in pediatric patients has been favorable with improved outcomes (40). There is limited data available for its use in patients with BPD-PH; however, its use is increasingly been seen in this population, especially as maintenance therapy for PH (41).

Other endothelin receptor antagonists include ambrisentan, which is a selective antagonist of ET_A and macitentan, which antagonizes both ET_A and ET_B receptors. However, there is no published data regarding their use in infants with BPD-PH.

Drugs acting of the prostacyclin pathway

Epoprostenol
Epoprostenol (Flolan) is an arachidonic acid metabolite and is produced endogenously by the vascular endothelium. It mediates vasodilation by activating adenylyl cyclase and increasing cAMP in the pulmonary arterial smooth muscle cell (25). Several different PGI_2 analogs have been used in adults and children with PH and are discussed below.

**Table 1 Drugs used in treatment of BPD-PH**

<table>
<thead>
<tr>
<th>Name of drug</th>
<th>Formulations available</th>
<th>Dose</th>
<th>Adverse effects</th>
</tr>
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<tbody>
<tr>
<td>Sildenafil (phosphodiesterase-5 inhibitor)</td>
<td>Oral: tablets and suspension; intravenous</td>
<td>PO: 1 mg/kg/dose q 6–8 h; start with low dose (0.3–0.5 mg/kg/dose) and gradually increase to 1 mg/kg/dose q 8 h as tolerated; intravenous: 0.25–0.5 mg/kg/dose q 6–8 h</td>
<td>Systemic hypotension, irritability, bronchospasm, rarely priapism</td>
</tr>
<tr>
<td>Milrinone (phosphodiesterase-3 inhibitor)</td>
<td>Intravenous continuous infusion</td>
<td>0.15–0.5 mcg/kg/min</td>
<td>Systemic hypotension, arrhythmia, risk of decreased myocardial perfusion</td>
</tr>
<tr>
<td>Bosentan (endothelin receptor antagonist)</td>
<td>Oral dispersible tablets</td>
<td>0.5–1 mg/kg BID, may increase up to 2 mg/kg BID in 2–4 weeks</td>
<td>Liver dysfunction (monitor liver function tests monthly), VQ mismatch, hypotension, anemia, known teratogen (precautions for pregnant caregiver)</td>
</tr>
<tr>
<td>Epoprostenol (Flolan)</td>
<td>Intravenous</td>
<td>Start at 1–2 ng/kg/min, titrate up slowly every 4–6 h to 20 ng/kg/min</td>
<td>Hypotension, VQ mismatch, GI disturbances, tachyphylaxis and risk of rebound PH</td>
</tr>
<tr>
<td>Iloprost</td>
<td>Inhaled</td>
<td>2.5–5 mcg every 2–4 h. Can be given as continuous inhalation during mechanical ventilation</td>
<td>Bronchospasm, hypotension, pulmonary hemorrhage, ventilator tube blocking due to crystallization, GI disturbances</td>
</tr>
<tr>
<td>Treprostinil</td>
<td>Intravenous; subcutaneous</td>
<td>Start at 2 ng/kg/min and titrate every 4–6 h up to 20 ng/kg/min</td>
<td>Similar to epoprostenol. Subcutaneous form: local site pain</td>
</tr>
</tbody>
</table>

BPD-PH, bronchopulmonary associated-pulmonary hypertension; PO, per os; VQ, ventilation-perfusion; GI, gastrointestinal.
Iloprost

Iloprost has a half-life of 20–30 minutes and is a more stable compound and so can be delivered by continuous inhalation to limit the systemic side effects (45). There is limited data from case series and case reports that suggest improved oxygenation in infants with BPD-PH with use of iloprost either alone or in combination with sildenafil (46,47). It has also shown promise in improving oxygenation in the setting of PPHN (48,49).

Treprostinil

It is a stable PGI₂ analogue that can be delivered by intravenous or subcutaneous route. Continuous subcutaneous delivery can be used to transition children who were previously stable on continuous IV epoprostenol (50). Limited experience with the use of this drug in the setting of BPD-PH shows improvement in right ventricular function and PH with serial echocardiograms and decreased respiratory support within weeks of therapy initiation without significant side effects (51,52).

New and emerging drugs—prostacyclin receptor agonists

Selexipag

It is an oral, long acting, highly selective prostacyclin receptor agonist that causes vasodilation and inhibits smooth muscle cell proliferation. It has been used in adults as oral triple combination therapy (with ET receptor antagonist and PDE inhibitor) in adults with severe PH. However, data on its use in pediatric population is limited, with the first off-label use reported in 2017 (53,54). Since then there have been limited data describing its use in children ranging in age from 1.5 to 17 years with idiopathic pulmonary arterial hypertension or PH with congenital heart disease (55). Its use has not been reported in the setting of BPD-PH.

Beraprost

It was the first oral prostacyclin receptor agonist. However, it was not approved by FDA in the United States due to its adverse effects but is approved for use in Japan and South Korea in patients with PH (56).

A suggested sequence of these of these drugs is to initiate with iNO at 10–20 ppm. Obtain an echocardiogram and monitor infant for change in clinical status. If clinical status remains unchanged or worsens, sildenafil should be added. Sildenafil can also be used for chronic maintenance therapy after iNO has been weaned off. Treatment with prostacyclin derivatives or ET-1 antagonists should be considered in patients unresponsive to the above treatment. Echocardiogram should be obtained when changing the treatment modality to evaluate the response to therapy. Cardiac catheterization should be considered if first- and second-line treatments are ineffective and patient is stable. Serial BNP or NT-proBNP levels may be obtained weekly or twice weekly in unstable patients to assess response to therapy (24,30).

Outcomes of BPD-PH

The mortality rate in infants with PH is high, especially in infants with severe PH. In a retrospective study evaluating outcomes of 36 patients with BPD and severe/moderate PH, follow-up at 35 months showed improvement in PH in 15 patients after treatment; persistent severe PH in 4 patients despite treatment and closure of shunts; spontaneous improvement in 4 patients who were not on treatment; while 8 patients (26%) had died (7). In another study, evaluating 42 premature infants with BPD and PH, estimated survival rates were 64%±8% at 6 months and 53%±11% at 2 years after diagnosis of pulmonary artery hypertension. Severe PH and small for gestational age were associated with worse survival rates in this study. Among 26 survivors, improvement in PH was observed in 24 patients (92%) (6). Presence of PVS has also been shown to increase mortality in infants with BPD-PH. Among infants with severe BPD-PH, survival was lower among infants with PVS than those without PVS [5/9 (56%) vs. 26/30 (87%); P=0.01] (27). Lung transplantation is a therapeutic option for infants and children with severe PH unresponsive to therapy. However it is reserved for patients who do not have other comorbidities. As such, children with severe BPD-PH are not good candidates for this treatment since many of them have severe comorbidities associated with prematurity.

Conclusions

BPD-PH is a complex disease with immense variability in its clinical phenotype. Although we have come a long way in understanding its pathophysiology, clinical evidence is still needed for the use of novel biomarkers in the diagnosis and management and to test the efficacy and safety of various therapeutic agents. Routine screening of infants with BPD is important for the early detection of BPD-PH. Using a multidisciplinary approach to optimize the management of infants with BPD will also help to prevent the short- and
long-term morbidities associated with BPD-PH.

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None.

Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

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