ORIGINAL ARTICLE


The European Paediatric Pulmonary Vascular Disease Network, endorsed by ISHLT and DGPK

Michael Kaestner,¹ Dietmar Schranz,² Gregor Warnecke,³,⁴ Christian Apitz,¹ Georg Hansmann,⁵ Oliver Miera⁵

ABSTRACT

Acute pulmonary hypertension (PH) complicates the course of several cardiovascular, pulmonary and other systemic diseases in children. An acute rise of RV afterload, either as exacerbating chronic PH of different aetiologies (e.g., idiopathic pulmonary arterial hypertension (PAH), chronic lung or congenital heart disease), or pulmonary hypertensive crisis after corrective surgery for congenital heart disease, may lead to severe circulatory compromise. Only few clinical studies provide evidence on how to best treat children with acute severe PH and decompenated RV function, that is, acute RV failure. The specific treatment in the intensive care unit should be based on the underlying pathophysiology and not only be focused on so-called ‘specific’ or ‘tailored’ drug therapy to lower RV afterload. In addition therapeutic efforts should aim to optimise RV preload, and to achieve adequate myocardial perfusion, and cardiac output. Early recognition of patients at high risk and timely initiation of appropriate therapeutic measures may prevent the development of severe cardiac dysfunction and low cardiac output. In patients not responding adequately to pharmacotherapy, (1) novel surgical and interventional techniques, temporary mechanical circulatory support with extracorporeal membrane oxygenation, (2) pumpless lung assist devices (3) and/or lung or heart-lung transplantation should be timely considered. The invasive therapeutic measures can be applied in a bridge-to-recovery or bridge-to-lung transplant strategy. This consensus statement focuses on the management of acute severe PH in the paediatric intensive care unit and provides an according treatment algorithm for clinical practice.

INTRODUCTION

Despite recent advances in the specific treatment of pulmonary hypertension (PH), RV failure following or in the context of severe rise of pulmonary vascular resistance (PVR) is a challenging complication of PH and is associated with substantial morbidity and mortality.

PH leading to a decompensation of the cardiovascular system can be considered a syndrome with non-specific signs and symptoms presenting late in the disease process and may be strongly associated with, or directly caused by several, very heterogeneous underlying conditions.¹ Distinction between precapillary and postcapillary aetiologies (or establishment of a combination of the two) is important to initiate specific individual therapy.

Pathophysiology of acute PH and RV failure

Chronic PH causes adaptation and remodelling of the RV to increased loading conditions. Pulmonary hypertensive crisis (PHC) occurs when compensatory mechanisms fail, RV systolic function decompenates and LV preload acutely decreases resulting in abolished cardiac output and coronary perfusion.² ³

Acute elevation of afterload (pulmonary artery pressure (PAP)) is poorly tolerated by the unprepared RV. In healthy adult individuals, the RV cannot acutely generate a mean PAP >40 mm Hg.⁴ Adaptive changes of the RV microstructure and function do not work in the setting of acute rise of RV afterload. Thus, although myocardial contractility may initially rise with RV concentric hypertrophy and preserved systolic and diastolic function, excessive pressure overload results in maladaptive remodelling with RV dilatation and failure.⁵ Among many others, arrhythmia, myocardial ischaemia and/or pulmonary disease such as infections or pulmonary arterial embolism may trigger an acute rise of PAP and PVR (figure 1). With subsequent RV dilatation, the contractile sarcocereus apparatus is damaged, resulting in RV failure. The combination of right-to-left septal shift with subsequent LV compression, related to limited space within the pericardial sac, results in low LV output, systemic arterial hypotension and elevated LV and RV end-diastolic pressure. With or without RV hypertrophy, these factors decrease coronary perfusion leading to myocardial ischaemia that perpetuates RV failure.⁶ The resulting metabolic acidosis further increases PVR and PAB. Moreover, a sudden increase in PAP causes airway obstruction due to arterial dis- tension of the smaller intrapulmonary arteries and lung oedema. The resulting dead space ventilation and ventilation/perfusion mismatch causes hypoxia and respiratory acidosis, eventually resulting in even more elevated PAP and PVR (figure 1).² ³ ⁶ ⁷

In children with congenital heart disease (CHD) and systemic-to-pulmonary shunt, PH crisis may occur following corrective surgery. Factors that promote the development of postoperative PH are...
not yet completely understood, however, endothelial cell dys-
function was found to be a contributing factor preoperatively and postoperatively.8–10 Inflammatory response to cardiopul-
monary bypass, and ischaemia-reperfusion injury and its impact
on heart-lung function contribute to the rise in PVR.11 Pain,
awakening reactions in mechanically ventilated children, tracheal
secretions or tracheal suctioning may trigger acute severe eleva-
tion of PVR presenting as PHC and potentially leading to cardi-
ocirculatory collapse and death (figure 1).

In this consensus statement, we distinguish between and elab-
orate on three common scenarios: (1) acute PH crisis after
cardiac surgery for CHD, with different management strategies
subdivided for patients with univentricular and biventricular
physiology (2) acute deterioration in a child with previously
known chronic PH (eg, a child with chronic parenchymal lung
disease and acute viral chest infection and acute deterioration in
a child with new diagnosis of group 1 PH (pulmonary arterial
hypertension (PAH), eg, idiopathic PAH), and (3) secondary
PH.12 We finally focus on the rapid assessment and efficient
management of acute severe PH in the paediatric intensive care
unit (PICU), including concise recommendations and a treat-
ment algorithm for clinical practice.

METHODS
The recommendations given in table 2 relate to the grading
system currently suggested by the European Society of Cardiology (ESC) and the American Heart Association (AHA),
and was based on paediatric data only (class of recommenda-
tion, level of evidence). The grading and voting process within
the writing group is outlined in the executive summary13 of this
online supplement. Computerised searches of the PubMed/
MEDLINE bibliographical database were conducted between
1990 and June 2015. Clinical trials, guidelines and reviews
limited to paediatric data were searched using the terms
'pulmonary hypertension' and 'intensive care', 'heart failure', 'con-
genital heart', 'postoperative', 'surgery', 'ECMO', 'lung assist',
'ventricular assist', 'mechanical circulatory support', 'cardiopul-
monary bypass', and 'pulmonary hypertensive crisis'. The
primary focus of this manuscript is on group 1 PH according to
the World Symposium on Pulmonary Hypertension (WSPH)

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**Figure 1** Schematic drawing of PAH crisis and contributing factors. Pulmonary hypertensive crisis develops with an acute increase of PAP. This leads to an increase in RV pressure and volume causing a shift of the interventricular septum towards the left side and reducing LV volume. Filling pressures of ventricles rise, compensatory tachycardia and the drop in systemic blood pressure compromise coronary perfusion pressure and flow, leading to low cardiac output and metabolic acidosis. Furthermore, the increase in PAP causes decreased pulmonary blood flow and airway obstruction related to arterial distension of the smaller intrapulmonary arteries and lung oedema. Consequently dead space ventilation increases; together with a mismatch of pulmonary ventilation and perfusion this causes hypoxia and respiratory acidosis. PAH, pulmonary arterial hypertension; RVEDV, RV end-diastolic volume; RVEDP, RV end-diastolic pressure; LVEDP, LV end-diastolic pressure; V/Q, pulmonary ventilation and perfusion; PH, pulmonary arterial hypertension; PAP, pulmonary artery pressure; PVR, pulmonary vascular resistance; CHD, congenital heart disease; NO, nitric oxide; CPB, cardiopulmonary bypass; LR-Shunt, left-to-right shunt; PBF, pulmonary blood flow.
and transthoracic echocardiography are mandatory to guide assessment, the following questions have to be addressed:

- Is the PH associated with systemic hypotension and/or hypoxaemia, low AVDO₂, and thus most likely low cardiac output and tissue hypoxia?
- Are there any precipitating factors that might be responsible for elevated PVR and RV dysfunction (eg, infection, acidosis, arrhythmia, pericardial effusion)?
- Are there any other causes that could explain the symptoms of PHC and RV failure (eg, pneumothorax, pulmonary embolism)?

After the initial clinical assessment of vital signs, chest X-ray and transthoracic echocardiography are mandatory to guide therapy. Monitoring of haemodynamics and organ function (brain, liver, kidney, coagulation system, etc) is an essential part of the routine set-up and procedures in the PICU.1

**ECG**
Signs of right atrial dilatation, right axis deviation, right bundle branch block or RV hypertrophy are typically seen in patients with chronic PH.14 15 In patients who have undergone surgery for CHD, the ECG may still display changes that occur with the underlying heart defect, but sometimes it may be normal. Loss of sinus rhythm (eg, in atrial flutter or junctional tachycardia) is an important diagnosis as it aggravates RV failure.

**Chest X-ray**
Chest X-ray helps to differentiate between PH with and without ventilation-perfusion mismatch and may show involvement of one or both lungs. In addition, in patients with chronic or newly diagnosed PH, signs of enlargement of the

### Table 1: Recommendations on the therapy of acute pulmonary hypertension in the paediatric ICU

<table>
<thead>
<tr>
<th>Recommendations</th>
<th>COR</th>
<th>LOE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxygen should be given when transcutaneous oxygen saturation &lt;95% in children with PH and normal cardiac anatomy²</td>
<td>I</td>
<td>C</td>
</tr>
<tr>
<td>Intravenous prostanoids should be considered to treat children with severe PH³</td>
<td>Ila</td>
<td>B</td>
</tr>
<tr>
<td>iNO may be considered for treatment of postoperative PH in mechanically ventilated patients to improve oxygenation and reduce the risk of pulmonary hypertensive crisis24</td>
<td>Ila</td>
<td>B</td>
</tr>
<tr>
<td>Concomitant sildenafil should be administered to prevent rebound PH in patients who have signs of increased PAP on withdrawal of iNO, and require re-start of iNO despite preceding gradual weaning of iNO.</td>
<td>I</td>
<td>B</td>
</tr>
<tr>
<td>Intravenous sildenafil may be considered for treatment of PH in critically ill patients, especially in those with an unsatisfactory response to iNO31</td>
<td>Ila</td>
<td>B</td>
</tr>
<tr>
<td>Inhaled iloprost may be as effective as iNO in children with postoperative PH25</td>
<td>Ila</td>
<td>B</td>
</tr>
</tbody>
</table>

In children who develop signs of low cardiac output or profound pulmonary failure despite optimal medical therapy, extracorporeal life support may be considered57 58

### Table 2: Medications used for treatment of pulmonary hypertension in the intensive care unit

<table>
<thead>
<tr>
<th>Drug</th>
<th>Dose</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Epoprostenol intravenous</td>
<td>Start with 1–3 ng/kg/min, increase gradually to 60 (and more) ng/kg/min intravenous</td>
<td>Caution: systemic arterial hypotension. Need to change drug vial/delivery system every 12–24 h</td>
</tr>
<tr>
<td>Iloprost, intravenous</td>
<td>0.25 µg/kg inhal, max. 10 µg; 6x/day or 1–5 ng/kg/min intravenous</td>
<td>Caution: systemic arterial hypotension</td>
</tr>
<tr>
<td>iNO</td>
<td>2–40 ppm inhal</td>
<td></td>
</tr>
<tr>
<td>Sildenafil, intravenous</td>
<td>0.4 mg/kg bolus over 3hrs IV (optional), then 1.6–2.4 mg/kg/day continuous infusion</td>
<td>do not exceed 30mg/d. Higher sildenafil doses up to 7.2mg/kg/day IV have been used in newborn infants with PPHN associated with congenital diaphragmatic hernia</td>
</tr>
<tr>
<td>oral</td>
<td>8–20 kg: 3×10 mg oral</td>
<td>In children weighing less than 8 kg, dosage of 1 mg/kg/3ime a day (oral not approved)</td>
</tr>
<tr>
<td>Epinephrine</td>
<td>0.01–1 µg/kg/min intravenous</td>
<td>Positive inotropy. Increases myocardial oxygen consumption, tachycardia. Moderate effects on PVR and SVR</td>
</tr>
<tr>
<td>Norepinephrine</td>
<td>0.01–1 µg/kg/min intravenous</td>
<td>Increases SVR and PVR</td>
</tr>
<tr>
<td>Vasopressin</td>
<td>0.0003–0.002 IU/kg/min intravenous</td>
<td>Probably does not increase PVR (advantage vs norepinephrine)</td>
</tr>
<tr>
<td>Terlipressin</td>
<td>5–10 ng/kg/min intravenous</td>
<td>Probably does not increase PVR (advantage vs norepinephrine)</td>
</tr>
<tr>
<td>Dobutamine</td>
<td>5–20 µg/kg/min intravenous</td>
<td>Increases myocardial oxygen consumption, tachycardia. Probably does not increase PVR</td>
</tr>
<tr>
<td>Milrinone</td>
<td>0.375–1.0 µg/kg/min intravenous</td>
<td>Lowers PVR.</td>
</tr>
<tr>
<td>Levosimendan</td>
<td>0.1 µg/kg/min intravenous</td>
<td>Lowers PVR. Caution: systemic arterial hypotension</td>
</tr>
</tbody>
</table>

iNO, inhaled nitric oxide; PPHN, persistent pulmonary hypertension in the newborn; PVR, pulmonary vascular resistance; SVR, systemic vascular resistance.
right heart and central pulmonary arteries are typically present. In more severe cases, one may recognise peripheral rarefication of the pulmonary vasculature (increased radiotranslucency of the lung fields). However, right heart enlargement may be best seen in the lateral plane (radiopaque retrosternal space that is filled by the dilated RV). In patients with underlying CHD, changes on the chest X-ray can be related to the cardiac lesion itself.\(^{15,16}\)

Echocardiography

Echocardiography is the most important tool for the assessment of ventricular function and RV-LV interaction.\(^ {17}\) Doppler analysis of tricuspid valve and pulmonary valve regurgitation to estimate PAP and diastolic inflow characteristics of both ventricles may help in guiding therapy in the ICU. The estimated PAP and right ventricular systolic pressure (RVSP) need to be interpreted in the context of the degree of RV dysfunction. Low pressure gradients across the tricuspid and pulmonary valves may be due to RV failure; they should be seen in relation to other variables (eg, tricuspid annular plane systolic excursion [TAPSE], and in light of the presence or absence of pericardial and pleural effusions, or ascites. If pulmonary vein obstruction is excluded, in most cases, the enlarged left atrial dimension defines the postcapillary component of PH and may indicate the need for decompression of the left atrial pressure through atrial septum stenting. Especially after cardiac surgery, the patient’s anatomic and haemodynamic status must be determined and taken into consideration.

Invasive monitoring

Invasive monitoring with an arterial and a central venous line should be established in all patients with cardiopulmonary compromise in whom vasopressor or inotropic therapy may be necessary. Insertion of invasive lines should be done with sufficient local anaesthesia and adequate analgesedation. Additional general anaesthesia might be hazardous (see anaesthesia and ventilation). Invasive monitoring is also indicated in patients at risk for systemic hypotension secondary to PH targeted therapy, even if this does not include vasopressor/inotropic therapy. There is a broad controversy on the most useful way to haemodynamically monitor a child with PH, in particular, we cannot make a clear recommendation for or against invasive PAP monitoring/Swan-Ganz catheters, in the PICU. Measurement of filling pressures of both ventricles may be of value to guide fluid management and escalation of therapy in distinct scenarios.

**THERAPY OF ACUTE PH IN THE ICU**

**Basic measures in the ICU**

**Oxygen**

As a potent pulmonary vasodilator and a weak systemic vasoconstrictor, oxygen is indicated in children with ventilation-perfusion mismatch based on arterial oxygen saturations of less than 95% (figure 2). Sufficient supply prevents anaerobic metabolism in peripheral organs.\(^ {2}\) In children with systemic-to-pulmonary shunts, supplemental oxygen augments pulmonary overcirculation with the risk of worsening cardiac and pulmonary function. In cyanotic heart disease with pulmonary-to-systemic (right-to-left) shunt flow, a higher haemoglobin level and shunt flow guarantees adequate systemic oxygen delivery. In these patients, oxygen is indicated in concomitant parenchymal lung disease or in deep cyanosis; arterial oxygen saturations of 75–85% are generally accepted as sufficient.

**Alkalisation**

Alkalisation is effective for immediate treatment of PH crisis.\(^ {18,19}\) Acidosis elevates PVR and impairs the effect of inotropic and vasopressor drugs.\(^ {18}\) Therefore, acidosis, as measured by negative base excess, should be abolished. Alkalisation with sodium bicarbonate to achieve a pH of 7.44 resulted in significantly reduced PVR.\(^ {18}\) Of note, neurodevelopmental outcome might be negatively affected after prolonged hypocapnic alkalosis in newborns.\(^ {20}\)

**Sedation**

Anxiety and agitation increase PVR and oxygen consumption and should be avoided. Sedation of a critically ill child has to be done with caution: in spontaneously breathing patients, hyperventilation and apnoea have to be avoided. In ventilated patients, loss of sufficient LV preload together with substantial decrease of systemic vascular resistance (SVR) can lead to circulatory arrest due to loss of coronary perfusion pressure.

**Anaesthesia and ventilation**

Anaesthesia, intubation and insertion of invasive lines are among the most crucial steps in the management of a child with imminent deterioration. Mechanical ventilation is indicated in severe PH with profound cyanosis, respiratory or metabolic acidosis not responding to initial therapy, respiratory failure or in cardiocirculatory arrest. In patients responding to medical therapy mechanical ventilation should be avoided. Anaesthesia should be performed by the most experienced person available. Induction is usually started with a rapidly acting sedative and muscle relaxant and may be followed by an opioid. Data regarding ketamine are ambiguous because its effect on PVR depends on comedication.\(^ {21}\) Induction of anaesthesia for intubation may cause a pronounced fall in SVR, and circulatory collapse. To overcome the fall in SVR vasopressor support may become necessary. Nursing care and respiratory therapy of ventilated patients requires awareness of cardiopulmonary interactions. Maneuuvres triggering pulmonary hypertensive crises such as insufficient sedation, rise of pCO\(_2\) or suctioning should be avoided. Moreover, positive pressure ventilation impairs cardiac filling and output\(^ {22}\) especially in the failing RV. Normoventilation (pCO\(_2\)-levels 35–40 mm Hg) and long expiratory times are recommended. Hyperventilation reduces cardiac output, increases SVR\(^ {19}\) and induces lung injury. In patients with failing RV or in univentricular circulation, the pulmonary perfusion pressure (flow) in relation to mean airway pressure has to be monitored to guarantee sufficient pulmonary flow.

**Fluid management**

RV function in neonates and patients with significant PH is preload dependent. Volume loss is poorly tolerated and in an acute PH crisis, volume challenge may be useful, but haemodynamic monitoring is mandatory. On the other hand, chronic RV failure is associated with fluid overload and systemic venous congestion. Nevertheless, rapid fluid removal with diuretic therapy or haemofiltration has to be used with caution because RV unloading can induce low cardiac output.

‘Specific’ pharmacotherapy to decrease RV afterload

**Prostanoids and nitric oxide**

PH targeted therapy improves pulmonary blood flow and decreases RV afterload. Intravenous prostanoids (epoprostenol, iloprost, treprostinil) should be considered in the critically ill patient, if a severe ventilation-perfusion mismatch has been
Since prostanoids lower SVR, concomitant systemic vasopressor therapy may become necessary.

Inhalative therapy with prostanoids or inhaled nitric oxide (iNO) has less impact on SVR and should be considered early, especially if the systemic blood pressure is low. In addition, inhaled or aerosolised application does not worsen the ventilation-perfusion mismatch in the same way as the intravenous route may do. Inhaled iloprost has been proposed as an alternative to iNO with comparable effects on PVR, but well designed prospective clinical trials are lacking. Acute haemodynamic effects of other inhaled drugs, such as nitroglycerine or milrinone, have been described in small case series, but their usefulness is not well established.

PDE5-inhibitors (sildenafil)
Oral sildenafil is reasonable to facilitate weaning from iNO. However, in the intensive care setting oral medications have the

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disadvantage of unpredictable absorption. Prophylactic use of sildenafil before surgical correction of CHD has been proven to be useful. The administration of intravenous sildenafil has been described but its effectiveness has not been clearly determined. Systemic arterial hypotension and impairment of oxygenation have been described as adverse events.

**Pharmacotherapy to increase myocardial perfusion and/or counteract right-to-left interventricular septal shift (LV compression and low LV output)**

**Inotropes**

In the case of severe RV failure in PH, inotropic support may be necessary. Milrinone and levosimendan can be useful as positive inotropic drugs since they have little or no effect on heart rate but lower PVR. Single centre studies have shown positive effects of levosimendan on PVR in children after cardiac surgery. Levosimendan’s major effect is thought to be improvement of myocardial contractility. Additionally, there is a vasodilator effect due to inhibition of PDE3 activity. Dobutamine and epinephrine improve RV contractility but may induce tachycardia which impairs diastolic filling and coronary perfusion subsequently lowering cardiac output.

**Vaspressors**

Vasopressor therapy with norepinephrine, vasopressin or terlipressin may be indicated to induce reshifting of the interventricular septum from left to right, to improve tissue perfusion in patients with systemic hypotension, and to treat low SVR caused by PH targeted therapy. Elevation of systemic blood pressure may become necessary to maintain coronary perfusion pressure and to reduce leftward septal shift. Vasopressin and terlipressin have been shown in small case series to lower PVR while increasing SVR. Patients on vasopressor therapy should be monitored closely for adequacy of cardiac output.

**Non-pharmacological therapy**

Surgical or interventional intra-atrial communication

On the basis of clinical experience in adult patients, atrial balloon septostomy should be avoided in patients with acute cardiac decompensation and in end stage RV failure (central venous pressure >20 mm Hg). However, in selected patients, the creation of an intra-atrial communication, decompressing the RA and RV, may be life-saving.

**Potts shunt**

As an alternative to lung transplantation, creation of an aortopulmonary shunt (Potts shunt) in children with suprasystemic idiopathic PH has been described in a small case series. Although these first long-term results are promising, no recommendation can be made because of the limited experience at this stage.

**Extracorporeal membrane oxygenation**

Depending on RV function venovenous (VV) or venoarterial (VA) extracorporeal membrane oxygenation (ECMO) may be considered as a bridge to recovery or bridge to transplantation. In patients with large atrial communications, circulatory support with VV-ECMO may be feasible even in the failing RV by providing oxygenated blood shunting right-to-left through the defect. The indication for mechanical support with ECMO depends on aetiology of RV or lung failure. Its short-term use in post cardiac surgery PH is generally accepted. However, longer support times had been described with new therapeutic strategies (e.g. awake-ECMO) making it feasible as bridge to transplantation in selected patients with other aetiologies. Timing of mechanical support in children is less well established compared with adults. The Interagency Registry for Mechanically Assisted Circulatory Support has defined clinical profiles of patients failing on optimal therapy for heart failure. Levels 1 and 2 are generally accepted indications for mechanical support in acute heart failure. In children after cardiac surgery, implantation of ECMO with metabolic acidosis or during cardiopulmonary resuscitation (CPR) is a risk factor for increased mortality and brain injury.

PH crisis after cardiac surgery may cause circulatory collapse requiring CPR. If spontaneous return of circulation cannot be achieved by conventional CPR measures, VA-ECMO implanted during CPR is an option (reversibility of disease). Overall survival in VA-ECMO implanted during CPR is 38%, with better outcome in the absence of severe metabolic acidosis before support; 11% of survivors suffered from cerebral seizures and 6% had evidence of brain injury in a CT scan.

In patients with idiopathic PH who deteriorate rapidly due to progressive RV failure, eligibility for transplantation has to be evaluated. If lung transplantation is deemed feasible, ECMO should be considered as bridging treatment. Acute decompensation in chronically ill children due to reversible disease (eg, in chronic lung disease and pneumonia) or in patients in whom targeted therapy for PH is suboptimal may require a bridge to recovery strategy. Although described in the literature, implantation of ECMO in children with idiopathic PH is controversial due to its irreversible nature and in consideration of the shortage of donor organs. If cannulated peripherally, patients can be awake and extubated, which allows longer support times and reduces the risks of ventilator associated complications. In severe RV compromise, VA-ECMO is needed to guarantee adequate cardiac output. Alternatively, a pumpless paracorporeal lung assist that is connected to the pulmonary artery and left atrium after sternotomy may be indicated in patients with hypoxaemia but sustained RV and LV functions.

This device decompresses the RV sufficiently, provides oxygenation and removes carbon dioxide. With normalisation of cardiac output after lung transplantation the chronically unloaded LV may develop substantially elevated filling pressures leading to pulmonary oedema and lung failure. Successful, scheduled postoperative VA-ECMO as a bridging strategy to LV reverse remodelling and recovery has been described in adults and children.

**Ventricular assist device**

Ventricular assist device treatment has not been proven to be effective in children with RV failure due to PH with preserved LV function. Pulsatile devices may lead to pulmonary haemorrhage due to high pulse pressures; while afterload dependent continuous flow devices may not provide sufficient pump flow in patients with high PVR. Therefore ventricular assist device implantation in RV failure due to PH with normal LV filling pressures is not recommended.

**Lung transplantation**

Bilateral lung transplantation should be considered in children with inadequate clinical response on maximal combination therapy who remain in functional class III or IV. Due to long waiting times and influence on outcome, transplantation should be considered before cardiopulmonary decompression has occurred. Mortality of children who are mechanically ventilated before transplantation is significantly increased (HR 2.6, CI...
1.72 to 4.07). The median survival rate after lung transplantation in children is between 5.6 years and 6.1 years.49

Therapeutic strategies in different clinical scenarios
CPR of children with acute PH crisis and cardiopulmonary arrest
Resuscitating a patient with severely elevated PVR, acute heart failure and cardiopulmonary arrest is particularly difficult (figures 2 and 3). Resuscitation is initiated and conducted following published guidelines.50 51 From a pathophysiological perspective, it is important to proceed with tailored, imminent treatment as mentioned above. Special attention should be given to coronary perfusion (pressure, flow), especially in the setting of severe RV hypertrophy, elevated RV filling pressures and/or tachycardia. RV hypertrophy, decreased aortic to RV end diastolic pressure gradient, and shortening of diastole contribute to RV myocardial ischaemia, RV dilation and subsequent LV compression, causing a rise in LV filling pressures and—together with low cardiac output and coronary perfusion

Figure 3  BAS, balloon atrioseptostomy; Clinical algorithm for the management of the scenarios for acute PH in the ICU. CHD, congenital heart disease; E-CPR, extracorporeal membrane oxygenation with cardiopulmonary resuscitation; ICU, intensive care unit; IPAH, idiopathic pulmonary arterial hypertension; PDE5, phosphodiesterase 5; PH, pulmonary hypertension; PHC, pulmonary hypertensive crisis; SaO2, arterial oxygen saturation; VA-ECMO, venoarterial extracorporeal membrane oxygenation; VAD, ventricular assist device.

Therapy of acute postoperative pulmonary hypertensive crisis (PHC)

Acute PHC is a life-threatening emergency and has to be treated aggressively since the risk of cardiocirculatory collapse necessitating cardiopulmonary resuscitation is high. Providing normal ventilation and normalising pCO₂ (if necessary with a manual ventilation bag), boluses of norepinephrine, sedative drugs and muscle relaxants restore circulation in the majority of cases. Medication lowering RV afterload, augmenting RV preload and contractility and maintaining adequate coronary perfusion pressure (as described above) preserve LV function and oxygen delivery in patients at risk. Adequate postoperative analgesia and sedation, minimal handling and minimising tracheal suctioning help to prevent acute PHC. Preoperative administration of PDE5 inhibitors such as sildenafil has been tried, but due to the paucity of clinical data, no corresponding recommendation can be made. VA-ECMO should be considered early on when pharmacological and ventilator strategies fail.

Patients with univentricular heart and partial (‘Glenn’) or total cavopulmonary anastomosis/ ‘Fontan-like’ circulation (ie, no subpulmonary ventricle).

Pulmonary vessels of newborns are highly reactive to stimuli which elevate PVR. In children with univentricular heart after Glenn operation, an increase of PVR may provoke deep cyanosis, stasis of blood in the shunt and shunt thrombosis. Management of suspected shunt thrombosis must be aggressive. No clinical studies comparing different treatment strategies of acute shunt thrombosis are available. Management generally includes: optimisation of ventilation, sedation, muscle relaxation if necessary, iNO, bolus of heparin and prompt treatment of arterial hypotension (bolus of volume if filling pressures are low; norepinephrine to increase afterload, epinephrine if systolic function is impaired). If immediate recovery of arterial oxygen saturations fails to occur, opening of the chest to massage the shunt or implantation of VA-ECMO may become necessary.

In patients with univentricular physiology and total cavopulmonary connection, a low PVR is a prerequisite for adequate ventricular filling and output. iNO, milrinone and ventilation with low mean airway pressure or better spontaneous breathing all contribute to improved outcomes in the early postoperative course. Negative pressure ventilation has been described in the failing Fontan circulation. In children who are mechanically ventilated long term, mild permissive hypercapnia may allow lower ventilator settings without elevation of PVR as long as there is no respiratory acidosis (figure 3).

Therapy of PH in a child with acute deterioration due to new diagnosis of PH

The principles of treatment are comparable to those described above (see Therapy of PH in a child with acute deterioration due to previously known chronic PH). The clinical condition of a critically ill child initially often precludes a complete diagnostic workup. However, diagnoses and conditions which change the therapeutic regimen such as acute pulmonary embolism must not be missed.

Conclusions

Acute PH is a serious complication in children at risk, including those after surgery for CHD, and has major impact on clinical outcome in patients with and without chronic PH. Treatment in the ICU should be based on the underlying pathophysiology but ultimately needs to focus on the basic goals of lowering RV afterload and augmenting RV preload and contractility. Furthermore, maintaining adequate coronary perfusion pressure and flow will help to preserve myocardial oxygen and energy supply, and thus RV and LV systolic function and oxygen delivery. Early recognition of patients at particular risk, and timely establishment of efficient therapeutic actions may prevent the development of severe cardiac dysfunction, low cardiac output and death.

Author affiliations

1Department of Paediatric Cardiology, University Children’s Hospital Ulm, Ulm, Germany
2Paediatric Heart Centre, University Hospital of Giessen and Marburg, Giessen, Germany
3Department of Cardiothoracic, Transplantation and Vascular Surgery, Hannover Medical School, Hannover, Germany
4German Centre for Lung Research, BREATHE, Hannover, Germany
5Department of Paediatric Cardiology and Critical Care, Hannover Medical School, Hannover, Germany
6Department of Congenital Heart Disease and Paediatric Cardiology, Deutsches Herzzentrum Berlin, Berlin, Germany

Correction notice

In table 2 the Sildenafil intravenous dosage has been updated from 0.4mg to 0.4mg/kg since this paper was first published online.
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