Pulmonary capillary haemangiomatosis: a distinct entity?

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Pulmonary capillary haemangiomatosis (PCH) is a rare condition that is clinically inseparable from pulmonary veno-occlusive disease. PCH can develop as a consequence of genetic mutations or pulmonary venous obstruction and remodelling. http://bit.ly/35TLAY4

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ABSTRACT Pulmonary capillary haemangiomatosis (PCH) is a rare and incompletely understood histopathological finding characterised by abnormal capillary proliferation within the alveolar interstitium, which has long been noted to share many overlapping features with pulmonary veno-occlusive disease (PVOD). But are PCH and PVOD distinct entities that occur in isolation, or are they closely intertwined manifestations along a spectrum of the same disease? The classic clinical features of both PCH and PVOD include signs and symptoms related to pulmonary hypertension, hypoxaemia, markedly impaired diffusion capacity of the lung and abnormal chest imaging with ground glass opacities, septal lines and lymphadenopathy. In recent years, increasing evidence suggests that the clinical presentation, histopathological features, genetic substrate and pathobiological mechanisms of PCH and PVOD are overlapping and usually indistinguishable. The discovery of biallelic mutations in the eukaryotic translation initiation factor 2 α kinase 4 (EIF2AK4) gene in heritable PCH and PVOD greatly advanced our understanding of the overlapping nature of these conditions. Furthermore, recognition of PCH and PVOD-like changes in other pulmonary vascular diseases and in conditions that cause chronic pulmonary venous hyper-perfusion or hypertension suggests that PCH/PVOD may develop as a reactive process to various insults or injuries to the pulmonary vasculature, rather than being primary angiogenic disorders.

Introduction
Pulmonary hypertension is defined as an elevation in the mean pulmonary arterial pressure (mPAP>20 mmHg), which can be caused by many conditions affecting the lungs, heart and pulmonary
vasculature [1]. Pulmonary hypertension is classified into five groups according to similar pathology, pathophysiology and treatment characteristics (table 1). Group 1 pulmonary hypertension includes related diseases of the pulmonary arteries, capillaries and veins with a pre-capillary haemodynamic profile. In 1977, WAGENVOORT et al. [2] described a patient presenting with diffuse lung infiltrates and respiratory failure who, at autopsy, had “aggressive” angiomatous growth of capillary-like channels in the pulmonary interlobular septae, a condition henceforth referred to as pulmonary capillary haemangiomatosis (PCH). Patients with PCH share many similarities to those with pulmonary veno-occlusive disease (PVOD) and with pulmonary arterial hypertension (PAH). Pathologically, PCH is characterised by abnormal capillary proliferation in the alveolar septae with frequent focal alveolar haemorrhage and PVOD is characterised by fibrotic narrowing and occlusion of the pulmonary veins [3].

Whether PCH is a distinct entity from PVOD has been debated over the years; however, the earliest descriptions of both diseases suggested overlapping features [4]. One of the first cases of PVOD reported by STOVIN and MITCHINSON [5] in 1965 noted the presence “capillary angiomatous anastamotic vessels” in the interlobular septae. Similarly, in the initial description of PCH, WAGENVOORT et al. [2] noted the extensive involvement of the pulmonary veins and venules “could well cause it to be confused with pulmonary veno-occlusive disease”. However, other histological observations of PCH without PVOD, and the presence of PCH-like lesions in patients without the occlusive changes in pulmonary veins suggested these could be unique conditions.

Some 26 years after the first description of PCH, the 2003 World Symposium on Pulmonary Hypertension in Venice, Italy, recognised that PCH and PVOD were clinically similar, potentially overlapping conditions [6, 7]. The discovery of pathogenic mutations in the eukaryotic translation initiation factor 2 α kinase 4 (EIF2AK4) gene in PCH and PVOD [8, 9] and recent advances in pulmonary microvascular pathobiology inform us that venous and capillary abnormalities are present across a spectrum of pulmonary vascular diseases, including PAH and chronic thromboembolic pulmonary hypertension (CTEPH) [10]. As such,

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**TABLE 1 Updated clinical classification of pulmonary hypertension (PH)**

| Group | Subgroup |
|-------|----------|
| 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 Haematological disorders | 5.2 Systemic and metabolic disorders | 5.3 Others | 5.4 Complex congenital heart disease |

PAH: pulmonary arterial hypertension; PVOD: pulmonary veno-occlusive disease; PCH: pulmonary capillary haemangiomatosis; LVEF: left ventricular ejection fraction. Reproduced from [1].

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the most recent 2018 6th World Symposium on Pulmonary Hypertension now classifies PVOD and PCH under Group 1.6: “PAH with overt features of venous/capillaries involvement” (table 1) [1]. Thus, PCH must be discussed within the context of PVOD and PAH. The objectives of this review are: 1) to summarise the histological features of PCH and PVOD; 2) to summarise potential mechanisms involved in PCH and PVOD pathogenesis across the spectrum of pulmonary vascular diseases; and 3) highlight the overlapping clinical features of PCH/PVOD and the potential methods of distinguishing them from PAH.

**Histological features in PCH and PVOD**

The hallmark histological feature of PCH is abnormal alveolar capillary proliferation, defined as $\geq 2$ rows of capillaries within the interstitium of the alveolar walls (figure 1) [7, 11, 12]. Atelectasis and congested capillaries may also mimic this appearance; however, reticulin and CD34 staining helps distinguish the proliferative nature [11, 12]. Foci of capillary haemangiomatosis may appear nodular at low power. These lesions have been reported to be widespread throughout the parenchyma with capillary infiltration extending into airway epithelium, bronchi, and vessel walls of the pulmonary veins and arteries [2, 7, 11, 13]. Scanning electron microscopy studies have shown a distortion of small alveolar capillaries with extensive and abnormally large capillaries budding off and overgrowing smaller ones [14]. The typical pathologic changes of PVOD are intimal fibrosis and luminal narrowing, arterialisation and/or obliteration of the septal veins and venules (figure 2) [3, 4, 15]. Patchy foci with capillary multiplication within the alveolar walls are very frequent. These PCH-like changes are usually associated with muscularised microvessels, rather resembling arterioles $>70$ µm in diameter. Fresh haemorrhage and haemosiderin-laden macrophages in the alveolar spaces are usually present in PCH/PVOD [14–16]. Cytological atypia is not present in the capillary endothelial cells and mitoses are not frequently observed despite the excessive proliferation [11–13]. Intimal fibrosis and medial hypertrophy are widespread and always present in the pulmonary arteries; however, plexiform lesions and thrombotic lesions are not typically present in PCH or PVOD [7, 11–13]. Rare cases of PCH with vascular thrombosis and infarction have been reported [13] and eccentric intimal fibrosis within the arteries may resemble thrombotic lesions. A mild interstitial lymphocytic infiltrate and interstitial fibrosis is often observed in both PCH and PVOD [12]. The thoracic

**FIGURE 1** a, b) Capillary haemangiomatosis foci are readily identified within lung parenchyma (star). They are characterised by the increase of the cell density, close to remodelled vessels. c) Alveolar septa are thickened with dilated and/or multiplication of capillaries (arrows). Increased cellularity is characterised by affluence of inflammatory cells within the alveolar septa. Endoalveolar collections of hemosiderin-laden macrophages are also visible (asterisk). Scale bars: a) 500 µm; b) 200 µm; c) 200 µm; d) 100 µm.
hilar and mediastinal lymph nodes are often enlarged due to lymphatic congestion from the extensive lymphatic vessel involvement in the interlobular septae, transformation of sinuses by capillary-like vascular channels and fibrosis, follicular hyperplasia and intrasinusal haemorrhage [17].

Mechanisms of disease in PCH and PVOD: two entities or a common final pathway?

Arguments for splitting PCH and PVOD are mostly based on histological observations that pulmonary venous obstruction is not observed in all cases of PCH and some cases of PVOD do not demonstrate the marked capillary proliferation typical of PCH. Infantile and paediatric cases of PCH without the intimal fibrosis and occlusive pulmonary venous changes typically observed in PVOD have also been described [18, 19]. Nevertheless, these are likely to be variable manifestations along a spectrum of the same process. In a large autopsy series of 35 cases, LANTUÉJOUL et al. [12] observed that 80% of cases classified as PCH had pulmonary venous obstruction and 73% of cases classified as PVOD had significant capillary proliferation, illustrating that there is considerable histopathological overlap. The presence of less pronounced capillary proliferation may have gone unrecognised in some cases of PVOD lacking electron microscopy to analyse vascular casts [12, 14]. Furthermore, PCH lesions may be multifocal and patchy through the lung [21], so lack of systematic tissue sampling could overlook PCH-like foci in cases initially diagnosed as PVOD.

Identification of EIF2AK4 genetic mutations as the cause of familial PCH and PVOD strongly support the notion that these are histopathological manifestations of the same disease [8, 9]. The first familial cases of PCH were described by LANGLEBEN et al. [22] in 1988, in whom an autosomal recessive pattern of inheritance was suspected. In this series of three French-Canadian siblings, histological analysis of the lungs at autopsy showed widespread capillary proliferation, including penetration into the alveolar spaces and venules; however, medial thickening and intimal fibrosis of the veins was noted [22]. In 2014, EYRIES et al. [9] used whole-exome sequencing to identify biallelic mutations in the EIF2AK4 gene on chromosome 15 in 13 families with histologically confirmed diagnoses of PVOD in at least one family member, most of whom had PCH-like changes. In the same year, BEST et al. [8] identified two loss-of-function mutations in EIF2AK4 in two brothers affected with PCH, with confirmation of asymptomatic carrier status in each of their parents. Novel mutations in EIF2AK4 gene were also identified in two additional patients with PCH without a family history. Familial PCH/PVOD due to biallelic EIF2AK4 mutations demonstrates an autosomal recessive inheritance pattern [8, 9, 23, 24]. In biallelic EIF2AK4 mutation carriers, penetrance is likely to be nearly complete by the age of 50, as supported by the careful analysis of several families [9] as well as from long-term follow-up of biallelic EIF2AK4 mutations carriers [25]. The relentless natural history of PCH/PVOD was illustrated in a patient with EIF2AK4 mutations and two affected sisters. She demonstrated progressive disease and disability over less than 10 years from mild exercise limitation, moderately reduced diffusing capacity for carbon monoxide (D\textsubscript{L,CO}), hypoxaemia, normal resting haemodynamics and mild pulmonary vascular disease at lung biopsy to a complete PCH/PVOD phenotype characterised by severe dyspnoea at rest, marked D\textsubscript{L,CO} reduction, severe hypoxaemia, end-stage pulmonary hypertension with characteristic PCH/PVOD histology on lung explants at the time of lung transplantation [25].

EIF2AK4 mutations are found in nearly all PCH/PVOD patients with a family history, but are also identified in 8.6–25% of sporadic cases of PCH/PVOD (figure 3) [9, 26, 27]. EIF2AK4 encodes a serine–threonine kinase known as general control nonderepressible 2 (GCN2), that phosphorylates the α-subunit.

FIGURE 2 Pulmonary veno-occlusive disease. a) Small pulmonary veins are markedly remodelled, showing fibro-oedematous increase of the intima, leading to significant narrowing of the vascular lumen. b) Alveolar septa appear thickened with increase of the cellularity and congestion. Scale bars: 200 µm.
of eukaryotic translation initiation factor (eIF2α) under amino acid deprivation, leading to preferential synthesis of stress proteins [9, 28, 29]. GCN2 alters gene expression and promotes immune tolerance through inhibition of an inflammatory response to cell apoptosis [9, 30]. Reduced GCN2 expression results in dysfunctional angiogenesis and remodelling in PCH/PVOD via mechanisms that are incompletely understood [28]. Biallelic mutations in EIF2AK4 lead to markedly decreased expression of GCN2, however PCH/PVOD non-carriers of EIF2AK4 also show significant reductions in GCN2 expression, as do patients with PAH without major venous remodelling [28]. Rats exposed to monocrotaline, mitomycin C and cyclophosphamide, which have been linked to PCH/PVOD pathogenesis [31, 32], also demonstrate markedly reduced or absent GCN2 expression [28]. Decreased GCN2 expression across a range of pulmonary vascular diseases, irrespective of EIF2AK4 mutation status, not only implicates an important role of GCN2 in the pathobiology of PAH and PCH/PVOD but suggests that these conditions belong to a clinical and anatomical spectrum. Recent data from our group indicate that GCN2 loss-of-function negatively regulates bone morphogenetic protein (BMP)-dependent Smad 1/5/9 signalling in pulmonary endothelial cells, underscoring possible biological similarities between heritable PAH and PCH/PVOD [33]. We showed, in vitro, that GCN2 loss-of-function negatively regulates BMP-dependent Smad 1/5/9 signalling in human primary pulmonary microvascular endothelial cell cultures (hPMEC). This molecular relationship was confirmed, in vivo, in the lungs of a newly created transgenic rat model knock out for Eif2ak4 (Δ152Ex1/Δ152Ex1). We found a four-fold decrease in Smad 1/5/9 phosphorylation in knock-out rats. Since those rats do not have spontaneous pulmonary hypertension, the decrease in Smad 1/5/9 phosphorylation is not the mere consequence of high pulmonary artery pressures. It appears in this “pure” background (same genetic background, same environment) that GCN2 deficiency robustly decreases Smad 1/5/9 phosphorylation. This regulation may be mediated through GCN2-dependent repression of chordin, a natural extracellular antagonist of BMP signalling [34]. Moreover, GCN2 inhibition induced a dramatic increase in hPMEC proliferation, which is highly relevant of PCH genesis. Interestingly, BMP9 treatment was able to block this exuberant proliferation, suggesting it may be considered as potential therapeutic option for PCH/PVOD.

Among a large cohort of patients with confirmed or suspected PCH/PVOD, carriers of bi-allelic mutations in EIF2AK4 were younger at diagnosis compared to non-carriers (median age 26 years versus 60 years) and had a 1:1 sex ratio compared with a male predominance in EIF2AK4 mutation non-carriers [35]. In addition to being older, PCH/PVOD patients lacking mutations in EIF2AK4 frequently had occupational exposure to organic solvents, such as trichloroethylene (42%) or chemotherapeutic agents, such as mitomycin C or cyclophosphamide (10%), which are risk factors for PCH/PVOD [35, 36]. Thus, clinical phenotypes of PCH/PVOD in mutation carriers and non-carriers differ in important ways, highlighting how variable predispositions and pulmonary vascular injury can converge to result in a pathological pattern of PCH/PVOD. Interestingly, the extent of pulmonary arterial and venous remodelling and the distribution of PCH lesions also differ by EIF2AK4 mutations status [28]. Carriers of EIF2AK4 mutations demonstrate more patchy PCH-like foci whereas non-carriers demonstrate more diffuse PCH-like changes (figure 4).

Other yet-unknown genetic mechanisms may be present in some PCH/PVOD cases. A series with three siblings affected by PCH in infancy [37] and another family with PCH and no identified EIF2AK4 mutation in the study by BEST et al. [8] had autosomal dominant transmission. A 16q deletion in the
region for the FOXF1 gene, which has previously been associated with alveolar capillary dysplasia and misalignment of pulmonary veins in children, is the only other mutation linked to PCH, described in a neonate diffuse capillary proliferation without venous obstruction or occlusive changes [38]. Heterozygous mutations in the Bone Morphogenetic Protein Receptor 2 gene (BMPR2) are found in the majority of families with heritable PAH [39–42]. Mutations in BMPR2 are also present in approximately 25% of PAH patients without a family history [43]; however, pathogenic BMPR2 mutations were not identified among 60 patients with PCH/PVOD [26]. A del44C mutation in BMPR2 was reported in a patient with PVOD without any capillary abnormalities on histology [44]; however, EIF2AK4 mutation status was not known since this case was published in 2003, before the discovery of the role of EIF2AK4 in PCH/PVOD.

Platelet-derived growth factor (PDGF)-B and PDGF receptor-β (PDGFR-β) have also been implicated in the pathogenesis of PCH lesions with increased expression of PDGF-B and PDGFR-β in perivascular cells, type II pneumocytes and endothelial cells within PCH lesions [45, 46]. PDGFR-β is also over-expressed in the pulmonary artery smooth muscle cells and endothelial cells of patients with idiopathic PAH [47], as well as in the arteries and veins of patients with systemic sclerosis-associated PAH and in PVOD [48]. Once again, the common importance of PDGF across the spectrum of PAH to PCH/PVOD suggests considerable overlap in the underlying mechanisms of disease. The use of imatinib, an inhibitor of PDGFR-β, has been associated with clinical improvements in several cases of PCH/PVOD [49–53], which may be due at least in part to its effects on pulmonary venous tone and post-capillary resistance [54]. However, no prospective controlled trials of imatinib in PCH/PVOD have been performed and a significant publication bias may exist with regards to imatinib’s effectiveness in isolated cases or case series. Imatinib also resulted in modest clinical improvements in PAH; however, it was ultimately not approved for clinical use due to deleterious side effects [55–57]. However, given the lack of alternative treatment options, further well-designed clinical studies of imatinib in PCH/PVOD would be interesting.
The occurrence of PCH/PVOD lesions in patients with systemic sclerosis [58–62], systemic lupus erythematosus [63], Takayasu arteritis [64] and post-lung transplantation [65, 66], suggest that PCH and PVOD may be the end result of other poorly understood inflammatory or immune-mediated mechanisms of vascular injury and remodelling.

Is PCH a primary or reactive process?

Although isolated PCH may occur in rare cases, an emerging hypothesis is that proliferative capillary lesions develop in reaction to chronic post-capillary obstruction. Recently, using microvascular corrosion casting, scanning electron microscopy and micro-computed tomography, NEUBERT et al. [67] suggested that capillary haemangiomatosis develops as a consequence of pulmonary venous narrowing, post-capillary pressure overload and shear stress, leading to capillary sprouting and intussusceptive neoangiogenesis in the capillaries (figure 5). Further supporting this hypothesis, PCH-like lesions are well described in other conditions that incite secondary pulmonary venous remodelling and venous obstruction due to chronic pulmonary venous hypertension. For example patients with left heart failure with reduced or preserved ejection fraction develop pulmonary venous intimal thickening and remodelling that resembles PVOD [68], while others have found PCH-like lesions in patients with left-sided valvular heart disease, hypertrophic cardiomyopathy or dilated cardiomyopathy with severe pulmonary venous congestion [69–72]. Interestingly, unilateral PCH-like foci also occur in the left lungs of patients with congenital heart defects that result in decreased right lung perfusion and hyperperfusion or congestion of the left lung. One case described PCH in the left lung of a patient with PH and surgically repaired Scimitar syndrome (corrected anomalous right pulmonary vein drainage to left atrium, closure of ventricular septal defect and ligation of patent ductus arteriosus) [73]. The post-operative catheterisation showed complete obstruction of the right pulmonary venous return with absent antegrade perfusion of the right lung, resulting in chronic hyper-perfusion of the left lung and hypoplasia of the right lung. There were extensive changes typical of PVOD and PCH in the left lung with only slight pulmonary arterial hypertrophy but no pulmonary venous or capillary changes in the right lung. Another 4-year-old with a congenital right pulmonary artery stenosis demonstrated unilateral PCH in the left lung on biopsy, presumably from

FIGURE 5 Structure and architecture of pulmonary veno-occlusive disease (PVOD) and pulmonary capillary haemangiomatosis (PCH). Prominent and back-to-back proliferation of capillaries in an area with PCH a) by scanning electron microscope and b) in a complementary haematoxylin–eosin stain. c, d) Scanning electron microscope following microvascular corrosion casting of affected lung tissue reveals capillary neo-formation by sprouting [green arrow in c] and intussusceptive vascular pillars [red arrow in c] next to non-remodelled pulmonary capillaries (red frames in d). Scale bars: a) 200 µm; b) 100 µm; c) 100 µm; d) 20 µm. Reproduced from [67] with permission.
congestion or hyper-perfusion [74]. In total, these cases support the hypothesis that PCH-like foci develop as a reactive process to chronically elevated pulmonary venous and capillary pressure, whether the primary disturbance is in the pulmonary veins or further “downstream” (figure 6).

Classic histological features of PCH and PVOD occurring in seemingly different pulmonary vascular diseases also argue in favour of a reactive mechanism, in which the systemic bronchial circulation likely plays an important role. For example, PCH-like foci, venous intimal fibrosis and arterialisation of pre-septal venules are histopathological features of the microvascular disease in CTEPH patients who have persistent pulmonary hypertension after pulmonary endarterectomy or inoperable distal disease, and in experimental CTEPH models [75]. The PCH-like lesions in CTEPH develop in close topographical association with segmental or subsegmental pulmonary arteries that had arterial thromboembolic occlusion. These capillary and venous changes in CTEPH are associated with anastomoses and shunting between the bronchial circulation and the pulmonary venous circulation [75]. Similarly, in PAH patients, systemic bronchial circulation anastomoses are seen with pulmonary arteries, capillaries and veins. The extent of bronchial artery hypertrophy and increased bronchial microvessel density are highly related to the degree of muscular hypertrophy and intimal fibrosis in pulmonary veins [76]. Furthermore, substantial PCH-like changes were demonstrated in 86% of patients with sickle-cell disease along with bronchial hypertrophy and venous thickening in the majority patients [77]. Therefore, exposure of the pulmonary venous circulation to higher bronchial circulation pressure and flow likely precipitate venous remodelling and secondary PCH-like changes [67].

The inseparable clinical presentation of PCH and PVOD

Patients with PCH present with nonspecific symptoms and signs and, in fact, it is not possible to distinguish PCH from PVOD based on clinical features. This is not surprising given the considerable histopathological and mechanistic overlap discussed above. Conversely, in clinical practice, differentiating PCH/PVOD from PAH is a more clinically relevant question. However, while the clinical distinction is important, in reality, discriminating PCH/PVOD from PAH as separate pathological entities can be difficult, since there can be considerable involvement of the pulmonary veins and capillaries in patients with PAH due to connective tissue disease or even in BMPR2 mutation carriers [62, 76].

Most patients with PCH/PVOD report symptoms related to high pulmonary arterial pressure and right ventricular dysfunction, such as dyspnoea, oedema and fatigue. In a pooled analysis of 64 cases of PCH, haemoptysis occurred in approximately one-third of patients [78], which was higher than reported in a large case series of PVOD patients [79]. Haemoptysis also occurs in PAH patients, particularly in BMPR2 carriers, which is due to the degree of bronchial vessel remodelling [76]. Spontaneous pneumothorax has been reported in patients without clinically evident pulmonary hypertension but with PCH-like foci on

FIGURE 6 Pathogenesis of vascular remodelling in pulmonary veno-occlusive disease (PVOD). 1) Marked fibrosis of the intima and hypertrophy of the media lead to occlusion of the pulmonary postcapillary vasculature. 2) Venous occlusions cause increasing blood pressure in the pulmonary capillaries (*). Pressure overload induces excessive neoangiogenesis by sprouting and intussusceptive pillar formation likely driven by increased flow and shear stress, which results in the formation of pulmonary capillary haemangiomatosis (PCH). 3) Congestion of the pulmonary capillary and postcapillary vasculature result in pulmonary hypertension associated with subsequent sclerotic remodelling of pulmonary arteries and arterioles. Reproduced from [67] with permission.
A history of occupational exposures to solvents like trichlorethylene or to chemotherapeutic agents is much more frequent in patients with PCH/PVOD than PAH [35, 36]. Hypoxaemia is more marked in PCH/PVOD than in PAH and the diffusion capacity for carbon monoxide ($D_{L\text{CO}}$) is markedly reduced, to a greater degree than in PAH [79]. However, the degree of hypoxaemia and $D_{L\text{CO}}$ reduction does not distinguish between PCH and PVOD [83]. Cardiopulmonary exercise testing may be useful as exercise capacity is lower and dyspnoea intensity during exercise is more severe in PCH/PVOD patients compared with matched PAH patients due to higher ventilatory demand, higher minute ventilation/exhaled CO$_2$ and physiological dead space (dead space/tidal volume), more severe gas exchange impairment and earlier onset lactic acidosis [84].

The features of PCH on chest computed tomography are also indistinguishable from PVOD and include: 1) hilar and mediastinal lymphadenopathy; 2) poorly circumscribed centrilobular ground glass nodular opacities; and 3) smooth thickening of the interlobular septae (figure 7) [79, 85, 86]. Having two or three of these computed tomography features has a sensitivity of 75% and specificity of 84.6% for PCH/PVOD.

FIGURE 7 High-resolution computed tomography scan of the chest in patients with pulmonary capillary haemangiomatosis and/or pulmonary veno-occlusive disease. a) Ground-glass opacities with centrilobular pattern, poorly defined nodular opacities, and septal lines in an adult carrier of eukaryotic translation initiation factor 2 $\alpha$ kinase 4 ($EIF2AK4$) bi-allelic mutations. b) Mediastinal lymph node enlargement in an adult carrier of $EIF2AK4$ bi-allelic mutations. c) A paediatric case carrying $EIF2AK4$ bi-allelic mutations. d) A paediatric case not carrying an $EIF2AK4$ bi-allelic mutation. e and f) The chest of an $EIF2AK4$ mutation carrier before initiation of specific therapy for pulmonary arterial hypertension showed mild abnormalities with septal lines and ground-glass opacities [e]; high resolution tomography of the chest was performed 2 months after initiation of endothelin receptor antagonists for rapid worsening dyspnoea and showed a substantial increase of radiological abnormalities evocative of pulmonary oedema [f]. Reproduced from [35] with permission.
These classic computed tomography features correspond to the lymph node congestion, capillary infiltration/proliferation, and venous remodelling in the interalveolar septae and focal alveolar haemorrhage observed on histology [79]. A small study of three patients with predominant PCH on histology and three patients with predominant PVOD suggested that ground glass nodule diameter may be larger in patients with PCH compared to patients with predominant PVOD changes [83]. However, the clinical relevance of this is unclear. Pleural effusions may also be present in PCH/PVOD but are usually small.

Bronchoscopy can be useful in certain situations to distinguish patients with suspected PCH/PVOD from PAH. As PCH and PVOD frequently have occult alveolar haemorrhage, identification of haemosiderin-laden macrophages in bronchoalveolar lavage (BAL) fluid [87, 88] or even in expectorated sputum analysis [89] is suggestive of PCH/PVOD over a diagnosis PAH. Practically, bronchoscopy is seldom needed and may be poorly tolerated due to the frequent presence of severe hypoxaemia. Open lung or thoracoscopic lung biopsies or transbronchial biopsies are not recommended to establish the diagnosis of PCH/PVOD over PAH because it remains a high-risk procedure in these frail patients [87, 90]. In most cases, the typical radiographic features, very low $D_{LCO}$ and disproportionate hypoxaemia are adequate to diagnose PCH/PVOD in a patient with precapillary pulmonary hypertension.

Genetic testing can be helpful to distinguish PCH/PVOD from PAH, particularly when there is a supportive family history, as EIF2AK4 mutations are uncommon in heritable PAH and BMPR2 mutations uncommon in PCH/PVOD [26, 91, 92]. The family tree and transmission pattern can be helpful in suggesting the specific pathology, as PCH/PVOD due to biallelic EIF2AK4 mutations are transmitted in an autosomal recessive pattern and there is often consanguinity, whereas heritable PAH due to BMPR2 mutations are transmitted via an autosomal dominant pattern with incomplete penetrance. Only rarely are EIF2AK4 mutations found in patients with suspected idiopathic PAH and no other suggestive features of PCH/PVOD on imaging or histology [91]. In one study, patients with clinically suspected idiopathic PAH but with biallelic EIF2AK4 mutations tended to be younger with lower $D_{LCO}$ even when classic computed tomography features were not present [91]. Among patients clinically diagnosed with PAH who were <50 years old and with a $D_{LCO} < 50\%$ predicted, biallelic EIF2AK4 mutations were present in 53%. Furthermore, patients with biallelic EIF2AK4 mutations did not respond to vasodilator therapies, indicating such patients had PCH/PVOD rather than PAH [91]. Since EIF2AK4 mutations are found in about 20% of sporadic cases of PCH/PVOD, a family history is not a prerequisite to genetic testing [9]. However, since the majority of PCH/PVOD cases do not carry EIF2AK4 mutations [26, 35], the sensitivity of genetic testing is likely to be low, especially in older patients. Therefore, negative genetic testing does not exclude PCH/PVOD but identification of biallelic EIF2AK4 mutations is highly supportive of the diagnosis.

The differentiation of PCH/PVOD from PAH is of utmost therapeutic importance, since the prognosis is poor in PCH/PVOD and the use of PAH-targeted therapies in PCH/PVOD is usually ineffective and can result in pulmonary oedema [3, 35, 93–95]. The pathophysiology of pulmonary oedema in these patients is thought to be related to proximal arterial vasodilation and increased pulmonary blood flow from the effect of PAH medical therapies, which results in increased capillary hydrostatic pressure in the face of down-stream obstructed pulmonary veins. A systematic review of 64 cases of PCH/PVOD treated with various types and doses of PAH therapies described improvements in 6-min walk distance and pulmonary vascular resistance in some cases, but pulmonary oedema was reported in 30 patients [95]. In a study by Montani et al. [35], 90% of the 94 patients with PCH/PVOD who had received PAH medical therapies but only three out of 47 patients with a follow-up assessment had achieved satisfactory clinical responses, despite statistically significant improvements in 6-min walk distance, cardiac index and pulmonary vascular resistance. Pulmonary oedema occurred in 23% of EIF2AK4 mutation carriers and 21% of non-carriers in that study. Patients with underlying connective tissue disease and suspected PAH are also prone to developing pulmonary oedema with prostacyclin therapy [96], which may reflect their known propensity to develop pulmonary venous remodelling [62]. The development of severe, life-threatening or even fatal pulmonary oedema from PAH therapies is well described and cannot be predicted by the presence or absence of clinical response in haemodynamics or 6-min walk distance. As such, these medications are not recommended in patients with known or suspected PCH/PVOD and the only potential treatment option is lung transplantation.

**Conclusions**

PCH is a histological finding that rarely exists in isolation and which shares nearly universal overlap with PVOD. The common risk factors, identical genetic substrate and indistinguishable clinical presentations of PCH and PVOD necessitate their consideration as a single disease and may warrant a uniform terminology to reduce confusion and heterogeneity in future clinical studies. However, since PCH/PVOD features occur across the spectrum of pulmonary vascular diseases and in many other conditions, establishing a single, precise definition may prove difficult. The current grouping of PCH/PVOD under
"PAH with overt features of venous/capillary involvement" is logical and the use of PAH targeted therapies should be avoided or used with extreme caution in such patients. There is currently no effective treatment for PCH/PVOD other than lung transplantation, emphasising the dire need for new innovative therapies in this rare and devastating condition.

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