Idiopathic pulmonary arterial hypertension

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Despite improved understanding of the pathobiology of pulmonary arterial hypertension (PAH), it remains a severe and progressive disease, usually culminating in right heart failure, significant morbidity and early mortality. Over the last decade, some major advances have led to substantial improvements in the management of PAH. Much of this progress was pioneered by work in animal models. Although none of the current animal models of pulmonary hypertension (PH) completely recapitulate the human disease, they do provide insight into the cellular pathways contributing to its development and progression. There is hope that future work in model organisms will help to define its underlying cause(s), identify risk factors and lead to better treatment of the currently irreversible damage that results in the lungs of afflicted patients. However, the difficulty in defining the etiology of idiopathic PAH (IPAH, previously known as primary pulmonary hypertension) makes this subset of the disease particularly difficult to model. Although there are some valuable existing models that are relevant for IPAH research, the area would value from the development of new models that more closely mimic the clinical pathophysiology of IPAH.

Pathobiology of IPAH

In PAH, blood flow through the pulmonary blood vessels becomes restricted. The right side of the heart must compensate to force blood through the small arteries and arterioles of the lungs. If left untreated, PAH has a very poor prognosis. There is currently no method to identify individuals who are at risk for developing the disease. No single mechanism can completely account for the etiology of PAH. Increased pulmonary vascular resistance may develop from vasoconstriction, vascular remodeling or in situ thrombosis.

At least some of the sustained vasoconstriction and concentric arterial remodeling is thought to come from the decreased expression and function of potassium ion (K⁺) channels, especially voltage-gated K⁺ channels, in pulmonary arterial smooth muscle cells (PASMCs). Decreased K⁺ channel activity causes membrane depolarization that subsequently increases the cytosolic calcium ion ([Ca²⁺]cyt) concentration ([Ca²⁺]cyt) in PASMCs by opening voltage-dependent Ca²⁺ channels, causing vasoconstriction. The resulting increase in [Ca²⁺]cyt stimulates PASMC migration and proliferation, which leads to medial hypertrophy and concentric vascular remodeling. The reduced activity of K⁺ channels also slows down apoptotic volume decrease and inhibits the apoptosis of PASMCs, which further contributes to pulmonary vascular medial hypertrophy (Pozeg et al., 2003; Yuan et al., 1998). Thus, K⁺ channel dysfunction is thought to significantly contribute to the pathology of PAH.

The endothelium also has a role in the disease process. It is not clear whether dysfunction, or injury of the endothelium, predisposes individuals to disease or whether it results from the increased PAP that is associated with disease. In either case, endothelial damage reduces the production of anti-proliferative and vasodilator substances such as prostacyclin and nitric oxide. Increased release of pro-proliferative and vasoconstrictive agents such as throm-
boxane A_2 and endothelin-1 further elevates vascular tone (Christman et al., 1992; Giaid, 1998). Smooth muscle cells, endothelial cells and fibroblasts all proliferate excessively in PAH, contributing to occlusion of vessel lumens and detrimental vasculopathy (Masri et al., 2007; Stenmark et al., 2006; Thomas et al., 2009; Yu et al., 2008). Several inheritable or acquired mutations or polymorphisms of genes that regulate the proliferation, apoptosis and differentiation of pulmonary vascular cells predispose individuals to IPAH. Some of these genes include those that encode the transforming growth factor β (TGF-β) receptor BMPR2, the serotonin transporter and transient receptor potential canonical 6 (TRPC6) (Hamidi et al., 2008; Machado et al., 2006; Rabinovitch, 2001; Yu et al., 2009).

Existing models for PH
Chronic hypoxic model of PH
Hypoxia-induced PH is consistent and reproducible in rats, despite some variability with age and between species. A chronically hypoxic (CH) model demonstrates pulmonary vasoconstriction, medial hypertrophy and increased muscularization of the small arteries with elevated smooth muscle α-actin. Remodeling of the precapillary arterioles results from increased medial thickening, and from smooth muscle cell hyperplasia and hypertrophy that occurs soon after disease onset. Maintaining CH rats at an altitude, to induce hypobaric conditions, elevates their PAP, decreases their numbers of aveoli and increases their PASMC endothelin production. These animals remodel their pulmonary vasculature. These characteristics mimic the symptoms found in humans, but CH rats develop concomitant systemic hypertension, which is absent in human PAH, suggesting that there are some important differences in their underlying physiology.

Non-rat CH models are used sometimes. Mice develop CH-induced PH, but their disease is substantially different from that seen in humans. It is associated with minimal vascular remodeling, whereas in humans pulmonary vascular remodeling, which is characterized by intimal and medial thickening, is the major known cause for elevated pulmonary vascular resistance. Another inconsistency of the mouse model is that its adventitial thickening and fibrosis occurs in the proximal pulmonary arteries, whereas in humans it often occurs in distal pulmonary arteries. Neonatal calves develop severe PH with significantly elevated PAP when exposed to reduced oxygen (12.5% O_2), which is associated with substantial vascular remodeling. Their medial and adventitial thickening is extensive, and mesenchymal progenitor cells and mononuclear cells accumulate in the arterial wall (Das et al., 2001; Davies et al., 1991), similar to in humans. The general use of these models is, however, fairly limited.

CH-induced PH in animal models is reversed when the animals are exposed to normal oxygen concentrations, making it fundamentally different from IPAH in humans, which occurs in normoxic conditions and causes irreversible intimal fibrosis and plexiogenic lesions. A modification of the CH model involves administration of Sugen 5416, a vascular endothelial growth factor (VEGF) receptor inhibitor, to CH rats (Taraseviciene-Stewart et al., 2001). This model emulates the hyperproliferative endothelial cell etiology and irreversible PH. Under these conditions, several animals develop persistent PH and right heart failure leading to deaths that look more similar to the disease progression in humans. Likewise, rats with an endothelin receptor B (ET_B) deficiency develop a more severe form of PH. These rats may offer a more relevant model and can perhaps offer some insight into the unique mechanisms that contribute to PH in animals compared with IPAH in humans.

Table 1. PAH: the facts

| 1.1 | Idiopathic |
| 1.2 | Heritable |
| 1.2.1 | BMPR2 |
| 1.2.2 | Alk-1, endoglin |
| 1.2.3 | Unknown |
| 1.3 | Drug and toxin induced |
| 1.4 | Associated with |
| 1.4.1 | Connective tissue diseases |
| 1.4.2 | HIV infection |
| 1.4.3 | Portal hypertension |
| 1.4.4 | Congenital heart disease |
| 1.4.5 | Schistosomiasis |
| 1.4.6 | Chronic hemolytic anemia |
| 1.5 | Persistent pulmonary hypertension of the newborn (PPHN) |

PH is a hemodynamic and pathophysiological state
PH is characterized by an increase in mean PAP to >25 mmHg at rest and >30 mmHg upon exercise, determined by right heart catheterization

PH is characterized by the presence of precapillary PH and includes a number of forms sharing the same clinical picture and pathological changes

PH forms Group 1 of the Dana Point clinical classification of PH (Simonneau et al., 2009) and is subdivided into five divisions

Case study
M.P. is a 34-year-old woman who, 14 months ago, presented with progressive dyspnea on exertion. She was initially diagnosed with asthma and prescribed an albuterol inhaler, but her dyspnea on exertion progressed, and seven months after her initial presentation she developed syncope after rapidly climbing a flight of stairs. At that time, the examination was notable for elevated jugular venous pressure, clear lungs and a tricuspid regurgitation murmur. Trace pedal edema was also noted. An echocardiogram was obtained and revealed an elevated pulmonary arterial systolic pressure of 67 mmHg, as well as right ventricular hypertrophy. An extensive evaluation of the patient’s PH was negative, and included a cardiac catheterization documenting PAP of 72/44 mmHg and a cardiac index of 2.4 l/min/m². Given the negative evaluation for associated conditions, the patient was diagnosed with IPAH. She was treated with inhaled iloprost, oxygen, warfarin and diuretics for seven months. Although her functional capacity remains limited by dyspnea on exertion, her overall functional status and symptoms improved and her six-minute walk distance increased by 37 meters to 234 meters. A repeat cardiac catheterization after six months of therapy showed a mild decrease in PAP to 64/39 mmHg, and an increase in cardiac index to 2.6 l/min/m².
Monocrotaline-induced pulmonary vascular disease

The pneumotoxin monocrotaline (MCT) can be used to generate another animal model of PH. A single subcutaneous injection of MCT undergoes oxidation in the liver to form monocrotaline pyrrole. By an unknown mechanism, MCT rapidly induces severe pulmonary vascular disease that, over a period of one to two weeks, is followed by pulmonary vascular remodeling and elevated PAP. An infiltration of mononuclear inflammatory cells into the adventitia often precedes the medial hypertrophy. Reminiscent of PAH in humans, MCT-treated rats develop right ventricular hypertrophy with the right ventricular systolic pressure reaching up to 80 mmHg (Hessel et al., 2006).

MCT-induced PH varies among strains and species of animals. The favored model is the rat, which has a more consistent and predictable response than the mouse. Larger mammals, such as dogs, also respond to MCT and undergo vascular remodeling and neointimal formation (Gust and Schuster, 2001), but there is disparity in the susceptibility of individual animals, which is thought to be the result of differences in the pharmacokinetics and hepatic metabolism of MCT. However, there is also a suggestion that differences between Sprague-Dawley and Fisher 344 rats reflect a unique pulmonary vascular response (Pan et al., 1993).

The MCT model is not perfect for human PAH for other reasons as well. It is relatively easy to cure MCT-induced PH, and over 30 agents have shown therapeutic benefits that prevent or reverse MCT-induced PH. One of these molecules is the anorexigen dexfenfluramine that, paradoxically, is associated with the development of IPAH in humans. These confusing data are summarized in a recent review (Stenmark et al., 2009). In the MCT model, single-sided pneumonectomy is required to produce neointimal formation and obliteration of the smaller arteriole lumens that are experienced commonly by human patients (White et al., 2007). The comparative pathologies between MCT mice with pneumonectomy and the standard MCT model are shown in Fig. 1. This modified model is now frequently utilized for etiologic studies.

It is interesting to note that inactivation of bone morphogenetic protein receptor (BMPR) signaling in both the MCT and CH models of PH may be crucial for pathogenesis (Long et al., 2009). This supports the human studies implicating BMPR type II (BMPR2) mutations as an underlying cause in over 70% of familial PAH cases and approximately 20% of IPAH cases. The expression of two related receptors, BMPR2 and BMPR1a, is reduced in patients with idiopathic PAH, suggesting that BMP–TGF–β signaling pathways are disrupted in PAH.

Genetically modified models of pulmonary hypertension

The association between PAH and BMPR2 gene mutations was first observed in 2000 (Lane et al., 2000), and heterologous mice (BMPR2+/–) were soon developed to determine the role of BMPR2 in PAH (Beppu et al., 2004; Song et al., 2008). However, these mice exhibit normal PAP at rest and require the presence of an additional factor, such as interleukin (IL)-1β or serotonin (Long et al., 2006; Song et al., 2005), to elevate their PAP. Homozygous knockout of BMPR2 (BMPR2–/–) is embryonically lethal, which suggests that a conditional knockout may prove valuable in the future. Still, around 20% of patients harboring BMPR2 gene mutations exhibit PH, suggesting that additional genetic and/or environmental factors are necessary to develop PAH.

In a search to uncover the mechanisms of neointimal lesion formation in humans, researchers have generated a variety of models. Blocking the VEGF receptor in CH rats with Sugen 5416 causes severe PH associated with precapillary arterial lesions and plexiform lesions (Taraseviciene-Stewart et al., 2001). Another model uses overexpression of S100A4 (MTS1) to cause plexiform lesions (Müller et al., 2004; Song et al., 2008). However, these animals exhibit normal PAP at rest and require the presence of an additional factor, such as interleukin (IL)-1β or serotonin (Long et al., 2006; Song et al., 2005), to elevate their PAP. Homozygous knockout of BMPR2 (BMPR2–/–) is embryonically lethal, which suggests that a conditional knockout may prove valuable in the future. Still, around 20% of patients harboring BMPR2 gene mutations exhibit PH, suggesting that additional genetic and/or environmental factors are necessary to develop PAH.

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The initiation and progression of human PAH involves multiple cellular and molecular mechanisms. The aforementioned work indicates that it is important to stimulate the generation of models to examine other pathways that might influence PAH. It is hoped that some of these genetic technologies may produce more informative models of PAH.

Moving research forward with new models

Current models have identified treatments for the acute symptoms of PAH; however, the etiology and underlying mechanisms of the disease remain unknown. This highlights the need to improve upon the current animal models of PH. An extensive, although not exhaustive, list of the current models of PH and the pathological charac-
teristics of the disease that they represent are featured in Table 2. Identification of the unique disease characteristics in model organisms and human patients should offer some insight into the mechanisms that contribute to disease.

**What constitutes a good model of PAH?**

An ideal model for PAH would recapitulate all aspects of the disease as it occurs in humans, including its pathological and hemodynamic alterations as well as its development and progression. The ‘ideal’ model of PAH should recapitulate the following pathological and hemodynamic features of patients with IPAH: (1) excessive pulmonary vascular remodeling characterized by intimal and medial hypertrophy, plexiform lesions and neointimal proliferation, obliteration and muscularization of small pulmonary arteries; (2) sustained pulmonary vasoconstriction; (3) in situ thrombosis in small arteries; (4) increased stiffness or decreased compliance of large and medium arteries; (5) significantly increased pulmonary vascular resistance and mean PAP; and (6) increased afterload in the right ventricle. This ideal model should also be free of unrelated toxic side effects.

Although no currently available model can fulfill all of these criteria, animal models have assisted enormously in understanding the pathophysiology of the disease. Prostacyclin analogues (including iloprost, epoprostanol and treprostinil) were identified and tested in the MCT and CH rat model of PH. Data from therapy studies in model organisms mostly agree with the clinical data showing that vasodilatation and exercise capacity improve with treatment (Lai et al., 2008; Lang et al., 2006; Obata et al., 2008). Prostanoids remain a mainstay for the treatment of PAH and are currently in clinical trial as part of a combinatorial therapy with treatments such as sildenafil, a phosphodiesterase-5 (PDE-5) inhibitor. The discovery of sildenafil as a therapeutic strat-

### Table 2. Summary of the currently available models to study PAH

<table>
<thead>
<tr>
<th>Experimental model</th>
<th>Species</th>
<th>Pathological similarity to PAH</th>
<th>Therapeutic developments using model</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chronic hypoxia (CH)</td>
<td>Rat, cow, mouse, pig, sheep</td>
<td>Increased muscularization of precapillary arterioles; medial and adventitial thickening; accumulation of mesenchymal progenitor cells and mononuclear cells</td>
<td>Calcium channel blockers: nifedipine and amlodipine</td>
<td>Hirenallur et al., 2008</td>
</tr>
<tr>
<td>VEGF-R2 inhibition + chronic hypoxia</td>
<td>Rat</td>
<td>As for CH, plus plexiform lesions, hyperproliferative endothelial cell etiology and irreversible PH</td>
<td>Statins: simvastatin</td>
<td>Taraseviciene-Stewart et al., 2001; Taraseviciene-Stewart et al., 2006</td>
</tr>
<tr>
<td>Monocrotaline (MCT)</td>
<td>Rat, dog</td>
<td>Increased muscularization, medial thickening and vascular inflammation; neointimal formation observed in dogs</td>
<td>Prostacyclin analogues: iloprost, epoprostanol, ONO-1301MS and treprostinil; ET receptor blockers: bosentan and BQ-123; PDE-5 inhibitors: ildenafil</td>
<td>Gust and Schuster, 2001; Lai et al., 2008; Mouchaers et al., 2009; Obata et al., 2008; van Albada et al., 2006</td>
</tr>
<tr>
<td>Monocrotaline + pneumonectomy</td>
<td>Rat</td>
<td>As for MCT, plus disorganized cellular proliferation in plexiform lesions</td>
<td>–</td>
<td>White et al., 2007</td>
</tr>
<tr>
<td>BMPR2 knockout</td>
<td>Mouse</td>
<td>Genetic predisposition resulting in increased muscularization</td>
<td>–</td>
<td>Lane et al., 2000; Song et al., 2005; West et al., 2004</td>
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Notably, existing therapeutic strategies owe their discovery to observations made in animal models of the disease. Prostacyclin analogues (including iloprost, epoprostanol and treprostinil) were identified and tested in the MCT and CH rat model of PH.

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Clinical and basic research opportunities

- To find biomarkers that predict the susceptibility of an individual to disease
- To identify the mechanistic differences that underlie PH-like disease in current model organisms versus human PAH
- To use genetic information collected from human patients to design models that can define the etiology of PAH
- To develop and characterize models for the identification and screening of therapeutics to prevent disease onset or to reverse some of the lasting consequences of PAH.

determine whether a model of pathogenesis will also be useful for large-scale pharmacological studies to identify novel therapeutic approaches; and (3) what types of organisms/animals (and species) can provide relevant models to understand IPAH. Since the pathogenic mechanisms of IPAH involve all pulmonary vascular cell types, various cell functions and multiple signaling pathways, a new strategy for generating an ideal model needs to aim at the multiple targets that are affected in the human disease, including a variety of cell types and genes. An animal model with multiple cellular and molecular abnormalities in various cell types would be more appropriate for pathogenic and therapeutic studies on the human disease.

Although animal models will undoubtedly continue to improve, it is unlikely that any one model will be able to successfully recapitulate the disease in humans. Human studies will always be an essential and mandatory step to scrutinize the efficacy and safety of novel treatment strategies for PAH.

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COMPETING INTERESTS

The authors declare no competing financial interests.

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