

This is a repository copy of *Current and future treatments of pulmonary arterial hypertension*.

White Rose Research Online URL for this paper: https://eprints.whiterose.ac.uk/159665/

Version: Published Version

Article:

Sommer, N., Ghofrani, A., Pak, O. et al. (6 more authors) (2021) Current and future treatments of pulmonary arterial hypertension. British Journal of Pharmacology, 178 (1). pp. 6-30. ISSN 0007-1188

https://doi.org/10.1111/bph.15016

Reuse

This article is distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs (CC BY-NC-ND) licence. This licence only allows you to download this work and share it with others as long as you credit the authors, but you can't change the article in any way or use it commercially. More information and the full terms of the licence here: https://creativecommons.org/licenses/

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



eprints@whiterose.ac.uk https://eprints.whiterose.ac.uk/

REVIEW ARTICLE THEMED ISSUE



Current and future treatments of pulmonary arterial hypertension

David G. Kielv⁶

Natascha Sommer¹ | Hossein A. Ghofrani^{1,7} | Oleg Pak¹ | Sebastien Bonnet² | Steve Provencher² | Olivier Sitbon³ | Stephan Rosenkranz⁴ | Marius M. Hoeper⁵ |

¹Cardiopulmonary Institute (CPI), University of Giessen and Marburg Lung Center (UGMLC), Member of the German Center for Lung Research (DZL), Justus-Liebig-University, Giessen, Germany

²Groupe de recherche en hypertension pulmonaire Centre de recherche de IUCPQ, Universite Laval Quebec, Quebec City, Ouebec, Canada

³Université Paris-Sud, Faculté de Médecine, Université Paris-Saclay, Le Kremlin-Bicêtre, France, AP-HP. Service de Pneumologie. Hôpital Bicêtre, Le Kremlin-Bicêtre, France, Inserm UMR S 999, Hôpital Marie-Lannelongue, Le Plessis-Robinson, France

⁴Klinik III für Innere Medizin, Cologne Cardiovascular Research Center (CCRC). Heart Center at the University of Cologne, Cologne, Germany

⁵Department of Respiratory Medicine, Hannover Medical School, Member of the German Center for Lung Research (DZL), Hanover, Germany

⁶Sheffield Pulmonary Vascular Disease Unit, Royal Hallamshire Hospital and Department of Infection Immunity and Cardiovascular Disease, University of Sheffield, Sheffield, UK

⁷Department of Medicine, Imperial College London, London, UK

Therapeutic options for pulmonary arterial hypertension (PAH) have increased over the last decades. The advent of pharmacological therapies targeting the prostacyclin, endothelin, and NO pathways has significantly improved outcomes. However, for the vast majority of patients, PAH remains a life-limiting illness with no prospect of cure. PAH is characterised by pulmonary vascular remodelling. Current research focusses on targeting the underlying pathways of aberrant proliferation, migration, and apoptosis. Despite success in preclinical models, using a plethora of novel approaches targeting cellular GPCRs, ion channels, metabolism, epigenetics, growth factor receptors, transcription factors, and inflammation, successful transfer to human disease with positive outcomes in clinical trials is limited. This review provides an overview of novel targets addressed by clinical trials and gives an outlook on novel preclinical perspectives in PAH.

This is an open access article under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs License, which permits use and distribution in any medium, provided the original work is properly cited, the use is non-commercial and no modifications or adaptations are made. © 2020 The Authors. British Journal of Pharmacology published by John Wiley & Sons Ltd on behalf of British Pharmacological Society

1

Abbreviations: 6MWD, 6-min walk distance; ALK1, activin receptor-like kinase 1; ANP, atrial natriuretic peptide; ASK1, apoptosis signal-regulating kinase 1, MAP3K5; Bcl-2, B-cell lymphoma 2; BMP, bone morphogenetic protein; BMPR2, bone morphogenetic protein receptor 2; BNP, brain natriuretic peptide; BRD4, bromodomain-containing protein 4; CO, cardiac output; CTD-PAH, connective tissue disease-associated pulmonary arterial hypertension; CTEPH, chronic thromboembolic pulmonary hypertension; DCA, dichloroacetate; DHEA, dehydroepiandrosterone; DMT, DNA methyltransferase; E2, oestradiol; ERAs, endothelin receptor antagonists; FDA, U.S. Food and Drug Administration; FHIT, fragile histidine triad; FKBP12, FK506-binding protein; FOX, forkhead box protein; HDACs, histone deacetylases; HIF, hypoxia-inducible factor; HMGB1, high mobility group box-1; IPAH, idiopathic pulmonary arterial hypertension; KCNK3, potassium channel subfamily K member 3 gene; mPAP, mean pulmonary arterial pressure; mTOR, mechanistic target of rapamycin; NAD+, nicotinamide adenine dinucleotide; NFAT, nuclear factor of activated T-cells; Nrf2, nuclear factor erythroid 2-related factor 2; p21, cyclin-dependent kinase inhibitor 1; p27, cyclin-dependent kinase inhibitor 1B; PA, pulmonary artery; PAEC, pulmonary arterial endothelial cells; PAH, pulmonary arterial hypertension; PASMCs, pulmonary arterial smooth muscle cells; PAWP, pulmonary arterial wedge pressure; PH, pulmonary hypertension; PVR, pulmonary vascular resistance: RCT, randomised controlled trial: ROCK, Rho-associated protein kinase: RTK, recentor TK; RUNX2, runt-related transcription factor 2; RV, right ventricle; SIRT, Sirtuin; SMURF-1, Smad ubiquitin regulatory factor 1; STAT, signal transducer and activator of transcription; TGF \$\beta R, TGF\$ receptor; TPH1, tryptophan hydroxylase 1; UCP2, uncoupling protein 2; VCAM1, vascular cell adhesion molecule 1; VE/VCO2, minute ventilation/carbon dioxide production; VIP, vasoactive intestinal peptide; VO2max, maximal oxygen uptake.



Correspondence

Natascha Sommer, Cardiopulmonary Institute (CPI), University of Giessen and Marburg Lung Center (UGMLC), Member of the German Center for Lung Research (DZL), Justus-Liebig-University, Giessen, Germany. Email: natascha.sommer@innere.med.unigiessen.de

Funding information Justus-Liebig-University

1 | DEFINITION AND CLASSIFICATION OF PULMONARY HYPERTENSION

Pulmonary hypertension (PH) was defined until recently by an elevated mean pulmonary arterial pressure (mPAP) of ≥25 mmHg (Galie et al., 2015). However, as the mPAP in healthy subjects was determined to be 14.0 ± 3.3 mmHg (Kovacs, Berghold, Scheidl, & Olschewski, 2009), the upper limit of normal, defined as two SDs above the mean value, mPAP should not exceed 20 mmHg (Simonneau et al., 2019). In the past, the relevance of the arbitrary definition of PH with a mPAP ≥25 mmHg has been questioned. Values >20 mmHg have clinical impact and affect outcome in populations at risk for PH (e.g., patients with systemic sclerosis; Simonneau et al., 2019; Valerio, Schreiber, Handler, Denton, & Coghlan, 2013). Thus, new cut-off values to define PH were suggested at the 6th World Symposium on PH in Nice 2018 defining PH as a mPAP >20-mmHg concomitant with a pulmonary vascular resistance (PVR) ≥ 3 Wood Units (WU) for pre-capillary forms of PH to account for the effect of cardiac output (CO) and pulmonary arterial wedge pressure (PAWP) on mPAP [PVR = (mPAP-PAWP)/CO] (Simonneau et al., 2019). It is currently unknown whether patients with mPAP between 21 and 24 mmHg and PVR ≥3 Wood Units would benefit from vasoactive treatment.

PH is currently classified into five separate groups with distinct pathophysiological characteristics (Galie et al., 2015). The clinical classification underwent minor revision at the 6th World Symposium (see Table 1; Simonneau et al., 2019). It ranges from rare forms such as pulmonary arterial hypertension (PAH, Group 1) and PH due to pulmonary artery obstructions (Group 4, primarily chronic thromboembolic PH [CTEPH]), to more common usually mild elevations of pressure seen in significant cardiac (PH due to left heart disease, Group 2) and respiratory disease (PH due to lung diseases and/or hypoxia, Group 3) and PH with unclear and/or multifactorial mechanisms (Group 5). The present review article will focus exclusively on PAH.

PAH has multiple causes but symptoms are similar in all forms of PAH, although their evolution and pattern vary depending on the aetiology. The most common symptoms are progressive breathlessness, fatigue, syncope, and clinical signs of heart failure. The diagnosis is usually first suggested by echocardiography. Confirmation of PAH requires right heart catheterisation and a systematic approach to investigation (Galie et al., 2015) including extensive imaging (Kiely et al., 2019) and integration with clinical features, to enable accurate classification which defines treatment. It is anticipated that advances in genotyping, improved imaging, and the application of artificial intelligence approaches to analysing data sets will aid refinement of disease classification and aid the study of treatment interventions. An overview of important landmarks in PAH research is given in Figure 1.

2 | OVERVIEW OF PATHOGENETIC MECHANISMS IN PAH

Pathological alterations have been recently reviewed in detail (Humbert et al., 2019). The initial concept that PAH is largely caused by mechanisms of vasoconstriction has been expanded over the last decades to a more complex picture in which multiple genetic, epigenetic, and environmental mechanisms lead to pulmonary vascular remodelling (Humbert et al., 2019). In some regards, PAH may even be considered as a pseudo-malignant disease with similar features to cancer (apoptotic resistance, altered metabolism, and overexpression of growth factor receptors; Boucherat, Vitry, et al., 2017). It is now accepted that curative therapeutic approaches must address not only vasoconstriction but also vascular remodelling, by inhibiting proliferative and activating anti-proliferative mechanisms (reverse remodelling; El Kasmi et al., 2014).

Pulmonary vascular remodelling underlying PAH is characterised by medial hypertrophy/hyperplasia, intimal and adventitial fibrosis, (in situ) thrombotic lesions, and plexiform lesions, as well as perivascular infiltration of inflammatory cells (B- and T-lymphocytes, mast cells, dendritic cells, macrophages, etc.; Humbert et al., 2019). It affects mainly distal muscular-type pulmonary arterial vessels and small pre-capillary arterioles (with diameters of 70–500 μ m and 20–70 μ m, respectively, in humans), but also to a varying degree postcapillary veins and bronchial arteries. The mechanisms for the latter are incompletely understood but may be connected by bronchial arterio-venous shunting (Humbert et al., 2019).

There is evidence demonstrating that all cell types of the vascular wall (fibroblasts, pulmonary arterial endothelial cells [PAEC], pulmonary arterial smooth muscle cells [PASMC], myofibroblasts, and pericytes) contribute to pulmonary vascular remodelling (Humbert et al., 2019). Several triggering factors combined with genetic/epigenetic susceptibility can initiate a phenotypical change of PAEC and PASMC characterised by apoptosis resistance, increased proliferation, and migration (Humbert et al., 2019). Damage of PAEC may play a particular role in the initiation of this process. Despite overabundance and/or overactivation of several pro-angiogenic

TABLE 1 Clinical classification of PH and haemodynamic definitions (World Symposium on PH, Nice 2018)

Classification	Haemodynamics ^a	Therapy
Group 1: PAH 1.1 Idiopathic PAH 1.2 Heritable PAH 1.2.1 BMPR2 mutations 1.2.1 other mutations (ALK1, endoglin [with or without hereditary haemorrhagic telangiectasia]) 1.3 Drug- and toxin-induced PAH 1.4 PAH associated (APAH) 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 involvement (pulmonary veno-occlusive disease/pulmonary capillary haemangiomatosis) 1.7 Persistent PH of the newborn syndrome	Pre-capillary mPAP > 20 mmHg PAWP ≤ 15 mmHg PVR ≥ 3 WU	Specific pulmonary vasoactive drugs Supportive therapy APAH: therapy of underlying disease
 Group 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 	Isolated post-capillary mPAP > 20 mmHg PAWP > 15 mmHg Combined pre-/post-capillary mPAP > 20 mmHg PAWP > 15 mmHg PVR ≥ 3 WU	Therapy of underlying disease
 Group 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 	Pre-capillary mPAP > 20 mmHg PAWP ≤ 15 mmHg PVR ≥ 3 WU Severe PH: mPAP ≥ 35 mmHg	Therapy of underlying disease Oxygen supplementation
 Group 4: PH due to pulmonary artery obstructions 4.1 Chronic thromboembolic PH 4.2 Other pulmonary artery obstructions 4.2.1 Sarcoma (high or intermediate grade) or angiosarcoma 4.2.2 Other malignant tumours 4.2.3 Non-malignant tumours 4.2.4 Arteritis without connective tissue disease 4.2.5 Congenital pulmonary artery stenoses 4.2.6 Parasites: Hydatidosis 	Pre-capillary mPAP > 20 mmHg PAWP ≤ 15 mmHg PVR ≥ 3 WU	Anticoagulation ^b Pulmonary endarterectomy ^b Balloon pulmonary angioplasty ^b Specific pulmonary vasoactive drugs ^b
 Group 5: PH with unclear and/or multifactorial mechanisms 5.1 Haematological disorders: chronic haemolytic anaemia, myeloproliferative disorders 5.2 Systemic and metabolic disorders: pulmonary Langerhans cell 	Pre-capillary mPAP > 20 mmHg PAWP ≤ 15 mmHg PVR ≥ 3 WU Isolated post-capillary mPAP > 20 mmHg	Therapy of underlying disease

BRITISH PHARMACOLOGICAL

BJF



TABLE 1 (Continued)

Classification	Haemodynamics ^a	Therapy
histiocytosis, Gaucher disease,	PAWP > 15 mmHg	
glycogen storage disease,	PVR < 3 WU	
neurofibromatosis, sarcoidosis		
5.3 Others: chronic renal failure with or	Combined pre-/post-capillary	
without haemodialysis, fibrosing	mPAP > 20 mmHg	
mediastinitis	PAWP > 15 mmHg	
5.4 Complex congenital heart disease	PVR ≥ 3 WU	

Note. Grey shading means main post-capillary form of PH.

Abbreviations: ALK, activin receptor-like kinase; APAH, associated pulmonary arterial hypertension; BMPR, bone morphogenic protein receptor; HIV, human immunodeficiency virus; LVEF, left ventricular ejection fraction; PAH, pulmonary arterial hypertension.

^aAccording to the 6th World Symposium on PH (Simonneau et al., 2019).

^bTreatment of chronic thromboembolic PH requires a multifaceted approach which may include multiple interventions.



FIGURE 1 History of PH and development of therapeutic drugs. FDA, U.S. Food and Drug Administration; WHO, World Health Organisation

pathways, angiogenesis and pericyte-associated tube formation of endothelial cells is disturbed leading to progressive obliteration of pre-capillary arteries ("vascular pruning" or "dead-tree" picture; Humbert et al., 2019; Yuan et al., 2015). Furthermore, fibroblasts become hyperproliferative and are involved in remodelling of the extracellular matrix promoting the alterations of PASMC and PAEC, as well as increasing stiffness of the large elastic main, lobar, and segmental pulmonary arteries (Humbert et al., 2019).

PAH is a multifactorial and heterogeneous disease in which multiple and different pathogenetic alterations have been observed in similar phenotypes (Figure 2): (1) Environmental trigger factors include air pollution (Sofianopoulou et al., 2019), as well as (particularly for PH Group 3) hypoxia and smoke exposure, shear stress (PH Group 1.4.4) and infections (PH Groups 1.4.2 and 1.4.5). (2) Different mutations leading to genetic susceptibility (bone morphogenetic protein receptor 2 [BMPR2], activin receptor-like kinase 1 [ALK1], voltage-

gated potassium channel 1.5 [Kv1.5], and potassium channel subfamily K member 3 [KCNK3]) have been identified, and early life epigenetic imprinting may also play a role. (3) PAH is promoted or triggered by systemic and circulating factors such as hormones and metabolites, as well as a pro-coagulatory and inflammatory disposition, particularly in PH Group 1.4.1. (4) Endothelial dysfunction leads to an imbalanced release of endothelial factors (endothelin, Tx, NO, and prostacyclin) and is currently targeted by PAH therapy. Further endothelial mechanisms include reduced anticoagulatory endothelial properties, increased expression of adhesion molecules (E-selectin, intercellular adhesion molecule 1, and vascular cell adhesion molecules), and endothelial release of different chemokines, cytokines, and growth factors (Huertas et al., 2018). (5) Cellular mechanisms in PAEC, PASMC, and fibroblasts include altered expression/function of ion channels and growth factor receptors, activation or deactivation of transcription factors (e.g., nuclear factor of



FIGURE 2 Patho-mechanisms underlying PAH. For details, please refer to text. DMT, DNA methyltransferase; ET, endothelin; Fb, fibroblasts; IC, immune cell; IL1R1, IL-1 receptor type 1; PA, pulmonary artery; PV, pulmonary veins; SiMFis, singular millimetric fibrovascular lesions

activated T-cells [NFAT], hypoxia-inducible factor [HIF] 1, signal transducer and activator of transcription [STAT] 3, and forkhead box protein [FOX] O1), and dysregulated cellular metabolism. (6) Finally, repair mechanisms counteracting remodelling are disturbed, including DNA and endothelial repair mechanisms (Boucherat, Vitry, et al., 2017; Humbert et al., 2019).

The multifactorial nature of PAH may be one of the reasons why finding a cure for PAH is challenging. For example, despite the importance of BMPR2 in the pathobiology of PAH, disease penetrance in carriers is only ~20% (higher in females than males), indicating that other factors are required for development of the phenotype. There are currently studies ongoing in large populations with idiopathic PAH (IPAH) and heritable PAH to identify further mutations or polymorphisms that could represent a "second hit." Several candidates have been identified, including variants in the prostacyclin synthase gene and mutations in the potassium voltage-gated channel sub-family A member 5 gene (Ghataorhe et al., 2017).

3 **CURRENT TREATMENT OPTIONS** IN PAH

Treatment of PAH currently includes supportive therapy such as diuretics, supervised rehabilitation, birth control advice, and oxygen supplementation, if needed (Ulrich et al., 2019). Pharmacological therapies targeting the pulmonary vasculature are currently only approved for patients with PAH (Group 1) and selected patients with CTEPH (Group 4). In other forms of PH, clinical trials using therapies approved for PAH have yet to demonstrate benefit, and current treatment strategies aim to optimise therapy of the underlying disease (see Table 1).

5

In PAH, current pharmacological therapies target three pathways regulating endothelial factors with vasoconstrictive/vasodilatory and proliferative/mitogenic properties: (1) NO-cGMP signalling is targeted by PDE5 inhibitors (tadalafil and sildenafil) and a soluble GC stimulator (riociguat); (2) endothelin receptor antagonists (ERAs) target both ET_A and ET_B receptors non-selectively (bosentan) or ET_A receptors selectively (ambrisentan and macitentan. This selectivity may be advantageous because stimulation of ETA receptors causes vasoconstriction whereas stimulation of ET_B receptors causes vasodilation); and (3) prostacyclin signalling is increased by either parenteral prostacyclin analogues (epoprostenol, treprostinil, and iloprost) or an orally available IP receptor agonist (selexipag; Galie et al., 2015). Milestones in the development of PAH therapies are shown in Figure 1, and pivotal studies are summarised in Table 2. In addition, a small group of patients with idiopathic, heritable, or drug-induced PAH who respond to a vasodilator challenge with a significant drop in pulmonary artery pressures and who have a sustained response to high-dose calcium channel blockade (around 5-10% of patients with IPAH) have excellent long-term survival (Rich, Kaufmann, & Levy, 1992; Sitbon

TABLE 2 Clinical trials in PAH

Trial name and identification number	Type of trial	Primary endpoint	Reference or trial duration/start date and status
A. Landmark clinical trials in PAH po	ositive for primary endpoint and resultir	ng in approval of therapy	
Continuous infusion of i.v. epoprostenol	RCT: 81 patients with IPAH	Change in 6MWD and PVR	Barst R.J., N Engl J Med 1996; 334: 296–301
Continuous s.c. infusion of treprostinil	RCT: 470 patients with PAH	Change from baseline in 6MWD at 12 weeks	Simonneau G. et al., Am J Respir Crit Care Med 2002; 165: 800- 804
Aerosolised iloprost (AIR)	RCT: 203 treatment-naïve patients with mixed types of PH	An increase of at least 10% in 6MWD and 1 NYHA FC at 12 weeks	Olschewski H. et al., N Engl J Med 2002; 347: 322-329
Bosentan (BREATHE-1)	RCT: 213 treatment-naïve patients with either IPAH or CTD-PAH	Change from baseline in 6MWD at 16 weeks	Rubin L.J. et al. N Engl J Med 2002; 346: 896–903
Sildenafil (SUPER)	RCT: 278 treatment-naïve patients with PAH	Change from baseline in 6MWD at Week 12	Galiè N. et al., N Engl J Med 2005; 353: 2148–2157
Ambrisentan (ARIES-1 and ARIES-2)	RCT: 202 (ARIES-1) and 192 (ARIES-2) treatment-naïve patients with PAH	Change from baseline in 6MWD at Week 12	Galiè N. et al., <i>Circulation</i> 2008; 117: 3010-3019
Tadalafil (PHIRST-1)	RCT: 405 treatment-naïve or pretreated patients with PAH	Change from baseline in 6MWD at Week 16. Endpoint not met in pretreated patients	Galiè N. et al., <i>Circulation</i> 2009; 119: 2894–2903
Inhaled treprostinil (TRIUMPH I)	RCT: 235 pretreated patients with PAH	Change from baseline in 6MWD at 12 weeks	McLaughlin V. et al., J Am Coll Cardiol 2010; 55: 1915-1922
Macitentan (SERAPHIN)	RCT: 742 treatment-naïve or pretreated patients with PAH	Time from initiation of treatment to first occurrence of a composite morbidity or mortality endpoint	Pulido T. et al., N Engl J Med 2013; 369: 809–818
Riociguat (PATENT-1/2)	RCT: 443 treatment-naïve or pretreated patients with PAH	Change from baseline in 6MWD at Week 12	Ghofrani H.A. et al. N Engl J Med 2013; 369: 330–340
Selexipag (GRIPHON)	RCT: 1156 treatment naïve or pretreated patients with PAH	Time from initiation of treatment to first occurrence of a composite morbidity or mortality endpoint	Sitbon O. et al N Engl J Med 2015; 373: 2522–2533
Clinical trials of up-front combination	on therapy in PAH positive for primary e	endpoint	
Ambrisentan and tadalafil in combination versus montherapy (AMBITION)	RCT: 500 treatment naïve patients with PAH	Time from initiation of treatment to first occurrence of a composite morbidity or mortality endpoint	Galie N. et al., N Engl J Med 2015; 373: 834-844
B. Trials in PAH not positive for primary endpoint, terminated early or other results			
Terguride: 5-HT receptor antagonist	Phase 2 trial: 78 patients with PAH	Change from baseline in PVR at Week 16	Ghofrani et al., 2012 (abstract) Endpoint not reached
Nilotinib: TK inhibitor (AMN107) NCT01179737	Phase 2 trial: 23 patients with PAH	Change from baseline in PVR at 6 months	2010–2014 Terminated due to serious adverse event
Aspirin and simvastatin: COX and HMG-CoA-reductase inhibitor	Phase 2 trial: 92 patients with PAH	Change from baseline in 6MWD at 6 months	Kawut S.M. et al., <i>Circulation</i> 2011; 123 (25): 2985–2993 Endpoint not reached
Atorvastatin: HMG-CoA- reductase inhibitor (APATH)	Phase 2 trial: 220 patients with PAH and CTEPH	Change from baseline in 6MWD at 24 weeks	Zeng W.J. et al., 2012; 40 (1): 67– 74 Endpoint not reached
Inhaled aviptadil: VIP	Phase 2 trial: 56 patients with PAH	Change in PVR acute or after 3 months	Published as abstract: Galiè et al., Am J Respir Crit Care Med 2010; 181: A2516

(Continues)

TABLE 2 (Continued)



Trial name and identification number	Type of trial	Primary endpoint	Reference or trial duration/start date and status
			Said S.I., Am J Respir Crit Care Med 2012 Apr 1; 185 (7): 786; author reply 786 Endpoint not reached
Imatinib: TK inhibitor (QTI571; IMPRES)	Phase 3 trial: 202 patients with PAH	Change from baseline in 6MWD at Week 24	Hoeper M.M. et al., <i>Circulation</i> 2013; 127 (10): 1128–1138. Endpoint reached but safety concerns
Selonsertib: ASK1 inhibitor (ARROW)	Phase 2 trial: 150 patients with PAH	Change from baseline in PVR at Week 24	Published as abstract: Boucherat O. et al., Am J Respir Crit Care Med 197 (3): 284–286, 2018 Endpoint not reached
FK506: calcineurin inhibitor (tacrolimus)	Phase 2 trial: 23 patients with PAH	Frequency of adverse events during 16 weeks	Spiekerkoetter E. et al., <i>Eur Respir J.</i> 2017; 50 (3) Low-level FK506 is well tolerated and increases BMPR2 in subsets of patients with PAH
Pioglitazone: dipeptidyl peptidase 4 inhibitor NCT00825266	Phase 2 trial: 2 patients with PAH	Insulin resistance profile change at Week 16	2009–2017 Terminated (difficulty in finding eligible patients)
Ubenimex: aminopeptidase inhibitor (LIBERTY1/2) NCT02664558/ NCT02736149	Phase 2 trial: 61/51 patients with PAH	Change from baseline in PVR at Week 24/frequency of adverse events	2016–2018 Terminated due to failure of efficacy
Racecadotril: neprilysin inhibitor	Phase 2 trial: 21 patients with PAH	Maximum change in circulating ANP concentration after 14 days	Hobbs A.J. et al. Br J Pharmacol 2019 May; 176 (9): 1251–1267 Increase in ANP
Anakinra: IL-1 inhibitor	Phase 1 trial: 7 patients with PAH	VO ₂ max, VE/VCO ₂ slope at 14 days	Trankle C.R. et al., Am J Respir Crit Care Med 2019; 199 (3): 381– 384 Endpoint not reached
Ambrisentan plus spironolactone: ERA plus aldosterone antagonist NCT02253394	Phase 4 trial: 30 patients with PAH	Cardiopulmonary fitness at 200 days	2014–2019 Terminated (low enrolment)
Fulvestrant: oestrogen antagonist NCT02911844	Phase 2 trial: 5 patients with PAH	Change from baseline in oestradiol, tricuspid annular plane systolic excursion, 6MWD, and N- terminal pro-brain natriuretic peptide at 9 weeks	Kawut S.M. et al., Ann Am Thorac Soc. 16 (11): 1456–1459, 2019 No significant changes of endpoints
Tocilizumab: anti-IL-6 antibody (TRANSFORM-UK) NCT02676947	Phase 2 trial: 29 patients with PAH	Adverse events; change from baseline in PVR at 6 months	2016–2018 Completed, results posted online Endpoint not reached
C. Current ongoing clinical trials on PAH			
Clinical trials of up-front combination therapy in PAH using licensed therapies			
Macitentan and tadalafil and selexipag versus macitentan and tadalafil in combination (TRITON) NCT02558231	RCT: 238 treatment naïve patients with PAH	Change from baseline in PVR at Week 26	2016 Recruiting
Hormonal modulators			
Anastrozole: aromatase inhibitor (PHANTOM) NCT03229499	Phase 2 trial: 84 patients with PAH	Change from baseline in 6MWD at 6 months	2017 Recruiting
	Phase 2 trial: 24 patients with PAH		2018 Recruiting
			(Continues)



TABLE 2 (Continued)

Trial name and identification number	Type of trial	Primary endpoint	Reference or trial duration/start date and status
Tamoxifen: oestrogen receptor inhibitor (T3PAH) NCT03528902		Change from baseline in tricuspid annular plane systolic excursion at Week 24	
DHEA (EDIPHY) NCT03648385	Phase 2 trial: 24 patients with PH	Change in RV longitudinal strain between DHEA and placebo at Weeks 18 and 40	2018 Recruiting
Spironolactone: aldosterone antagonist NCT01712620	Phase 2 trial: 70 patients with PAH	Change from baseline in 6MWD at 6 months	2012 Recruiting
rhACE2: GSK2586881 NCT03177603	Phase 2 trial: 24 patients with PAH	Change from baseline in PVR up to 4 hr	2017 Recruiting
KAR5585: tryptophan hydroxylase 1 inhibitor NCT02746237	Phase 1 trial: 120 healthy individuals	Adverse events up to 4 days	2016–2016 Completed, results pending
Escitalopram: SSRI NCT00190333	Phase 3 trial: 30 patients with PH	Change from baseline in 6MWD at Week 16	2005–2008 Completed, results pending
Fluoxetine: SSRI NCT03638908	Phase 2 trial: 8 patients with WHO group 1 PH	Change from baseline in PVR at Week 24	2018 Active, not recruiting
PB1046: VIP analogue NCT03315507	Phase 1 trial: 10 patients with WHO group 1 PH	Adverse events 28 days after last dose, laboratory parameters, change in PVR	2017 Active, not recruiting
GPCR pathways			
Apelin (EXAP) NCT01590108	Phase 1 trial: 12 healthy individuals and patients with IPAH	Cardiopulmonary performance at 6 months	2012–2019 Completed, results pending
Mitochondrial and metabolic adapta	ations		
Ranolazine: sodium channel inhibitor, partial FAO inhibitor NCT01839110	Phase 4 trial: 22 patients with PAH	Change in RVEF by MRI at 6 months	2013–2019 Completed, results pending
Ranolazine NCT02829034	Phase 4 trial: 22 patients with PAH	Change from baseline in RVEF by MRI at Week 26	2016-2019 Completed, results pending
Trimetazidine: FAO inhibitor NCT02102672	Phase 2 trial: 25 patients with PAH	Changes in RV function assessed by 3D echo at 3 months	2014 Recruitment state unknown
Metformin: biguanide NCT03617458	Phase 2 trial: 160 patients with PAH	Change from baseline in 6MWD at Week 12	2018 Recruiting
Ferinject or CosmoFer: iron infusion NCT01447628	Phase 2 trial: 40 patients with PAH	Change from baseline in PVR and endurance at Week 12	2011–2019 Completed, results pending
Epigenetic alterations and interaction with metabolic pathway			
Olaparib: PARP inhibitor (OPTION) NCT03782818	Phase 1 trial: 20 patients with PAH	Adverse events at Week 24	2019 Not yet recruiting
Apabetalone: BRD4 inhibitor (APPRoAcH-p) NCT03655704	Phase 1 trial: 10 patients with PAH	Change from baseline in PVR at Week 16	2019 Not yet recruiting
Oxidative stress related pathways			
Bardoxolone methyl: IkB kinase and NF-kB inhibitor, Nrf2 activator (LARIAT) NCT02036970	Phase 2 trial: 166 patients with PH	Change from baseline in 6MWD at Week 16	2014–2019 Completed, results pending
Bardoxolone methyl (CATALYST) NCT02657356	Phase 3 trial: 200 patients with CTD-PAH	Change from baseline in 6MWD at Week 24	2016 Recruiting
Bardoxolone methyl (RANGER) NCT03068130	Phase 3 trial: 414 patients with PH	Long-term safety up to 5 years	2017 Recruiting
CXA-10: nitrated fatty acid compound (PRIMEx) NCT03449524	Phase 2 trial: 96 patients with PAH	Change in RVEF and PVR at 6 months	2018 Recruiting

TABLE 2 (Continued)

Trial name and identification number	Type of trial	Primary endpoint	Reference or trial duration/start date and status
Inflammatory mediators			
Rituximab: anti-CD20 antibody NCT01086540	Phase 2 trial: 58 patients with systemic sclerosis associated- PAH	Change in PVR at Week 24	2010 Active, not recruiting
Elafin: elastase-specific protease inhibitor NCT03522935	Phase 1 trial: 30 healthy individuals	Adverse events at 28 days	2018 Recruiting
Growth factor receptors			
Sotatercept: activin receptor type 2A fusion protein acting as a ligand trap (SPECTRA) NCT03738150	Phase 2 trial: 25 patients with PAH	Change from baseline in VO ₂ max at Week 24	2018 Recruiting
Sotatercept (PULSAR) NCT03496207	Phase 2 trial: 100 patients PAH	Change from baseline in PVR at Week 24	2018 Recruiting
Transcriptional factors			
ABI-009: mTOR inhibitor NCT02587325	Phase 1 trial: 25 patients with PAH	Adverse events at Week 16	2015 Recruiting
Stem cell therapy			
Allogeneic cardiosphere-derived stem cells NCT03145298	Phase 1 trial: 26 patients with PAH	Primary safety endpoints within 72 hr of infusion	2017 Recruiting
eNOS-enhanced EPCs NCT03001414	Phase 2 trial: 45 patients with PAH	Change from baseline in 6MWD at Month 6	2016 Recruiting

Abbreviations: 6MWD, 6-min walk distance; ANP, atrial natriuretic peptide; ASK1, apoptosis signal-regulating kinase 1; BRD4, bromodomain-containing protein 4; CTD-PAH, connective tissue disease-associated pulmonary arterial hypertension; DHEA, dehydro-epiandrosterone; eNOS, endothelial NOS; EPCs, endothelial progenitor cells; FAO, fatty acid oxidation; HMG-CoA, 3-hydroxy-3-methyl-glutaryl-CoA; mTOR, mechanistic target of rapamycin; Nrf2, nuclear factor erythroid 2-related factor 2; NYHA FC, New York Heart Association functional classification; RCT, randomised controlled trial; rhACE2, recombinant human ACE2; RVEF, right ventricular ejection fraction; SSRI, selective serotonin reuptake inhibitor; VE/VCO₂, minute ventilation/carbon diox-ide production; VIP, vasoactive intestinal peptide; VO₂max, maximal oxygen uptake.

et al., 2005). This serves to illustrate the heterogeneous nature of IPAH and the importance of developing a personalised approach to drug therapy.

Recent improvements in outcomes for patients with PAH reflect a move to the use of combination drug therapy targeting multiple pathways. An overview of clinical trials of combination therapy in PAH was recently provided (Humbert & Ghofrani, 2016). Current treatment algorithms are based on interval multiparameter risk assessment with the aim of achieving a low-risk status (Galie et al., 2019). Lung transplantation remains an important therapeutic option for patients deteriorating despite maximal medical therapy. Despite treatment advances and an increase in survival by more than twofold during the last two decades, PAH remains a fatal disease, and the identification of new targets and development of new therapies is required.

4 | NOVEL TREATMENT OPTIONS TARGETING SPECIFIC PATHWAYS IN PAH

Preclinical and clinical research on novel drugs currently focusses on PAH where typical characteristics of pulmonary vascular remodelling exist. This review describes novel concepts that have been established in preclinical models and have been transferred to clinical trials (Figure 3; ongoing clinical trials are summarised in Table 2). Furthermore, some very new studies on preclinical approaches are described. Although preclinical models only partly reflect features of PAH, they have been successfully used to test treatment options in PAH. The most common preclinical models are the chronic hypoxic model of the mouse and rat leading to rather mild PH without characteristic plexiform lesions, the combined application of hypoxia and the VEGF inhibitor Sugen 5416 (Sugen/hypoxia) in rats which is characterised by more severe PH including plexiform lesions, and the monocrotaline model of rats which is largely driven by inflammatory stimuli and causes severe PH (Stenmark, Meyrick, Galie, Mooi, & McMurtry, 2009). Several other models with knockdown of relevant pathways have been described, but none of the models reproduces all the histopathological patterns of PAH.

4.1 | Circulating hormones

4.1.1 | Sex hormones

Systemic factors, mainly hormones or nutrients, may either trigger or promote the development of PAH. The best known hormone-

Η ΔΓΟΙ Ο ΓΓΔΙ



FIGURE 3 Novel treatment options in PAH. Substances shown in the Figure are currently tested in clinical trials. For details, please refer to text. EPCs, endothelial progenitor cells; ERs, oestrogen receptors; LTA₄H, LTA₄ hydrolase; SERT, 5-HT (serotonin) transporter; VPAC, vasoactive intestinal peptide receptor

related phenomenon in this regard is the "oestrogen paradox": Women have a higher incidence of IPAH and heritable PAH than men, but female patients with PAH have better outcomes than male patients. The role of different oestradiol (E2) metabolites and receptors is not completely elucidated. Some of the metabolites have proliferative effects (e.g., 16α -hydroxyoestrone), while others also have anti-proliferative and anti-inflammatory properties (e.g., 2-hydroxyoestradiol and 2-methoxyoestradiol). High E2 levels were associated with PAH in male patients and reduced exercise capacity in female patients, but E2 may also have protective effects on the right ventricle (RV; recently reviewed in Tofovic & Jackson, 2019). Thus, different levels of E2 and its metabolites at different disease stages could explain sex-dependent differences. Moreover, genes of the Y-chromosome are up-regulated in PH and have protective anti-remodelling properties in hypoxic mice (Umar et al., 2018). Decreasing circulating E2 by inhibiting conversion of androgens to E2 using the aromatase inhibitor anastrozole and inhibition of the E2 receptor with tamoxifen inhibited PH in different animal models of PH, partly in a sex-dependent way (Tofovic & Jackson, 2019). By contrast, supplementation with E2 increased exercise capacity in male and female rats with severe PH, probably by exerting protective effects on the RV (Lahm et al., 2016). Most importantly, anastrozole showed good safety and tolerability in male and female patients with PAH in a small "proof-of-principle" trial and significantly reduced E2 levels and improved 6-min walk distance (6MWD) but not RV function (Kawut et al., 2017). These pilot data supported initiation of the randomised, controlled PH and Anastrozole Trial (PHANTOM; ClinicalTrials.gov identifier: NCT03229499). Furthermore, the effect of oestrogen inhibition with tamoxifen will be investigated in patients with PAH in a small randomised controlled trial (RCT; Tamoxifen Therapy to Treat PAH [T3PAH]; ClinicalTrials.gov identifier: NCT03528902). In contrast to anastrozole, tamoxifen can be used in pre-menopausal female patients without inducing menopause. An open-label trial of the oestrogen antagonist fulvestrant recently (ClinicalTrials.gov identifier: NCT02911844) was completed. However, due to the low number of participants, no final conclusions on the effects of fulvestrant on PAH can be drawn. Nevertheless, it was generally well tolerated (Kawut et al., 2019).

4.1.2 | Dehydroepiandrosterone

Dehydroepiandrosterone (DHEA) is a steroid hormone that serves as a precursor for both oestrogen and testosterone synthesis. DHEA prevented and reversed PH and RV dysfunction in different animal models of PH (Lahm, Tuder, & Petrache, 2014). Accordingly, higher levels of E2 and lower levels of DHEA were associated with increased risk of PAH in men (Ventetuolo et al., 2016) and increased risk and severity of PAH in post-menopausal women (Baird et al., 2018). DHEA treatment significantly improved 6MWD,

11

BRITISH PHARMACOLOGICAL

pulmonary haemodynamics, and diffusion capacity of patients with PH associated with chronic obstructive pulmonary disease, without worsening gas exchange (de La Roque et al., 2012). Currently, DHEA is being tested in a crossover trial in a small number of patients with PAH (Effects of DHEA in PH [EDIPHY]; ClinicalTrials. gov identifier: NCT03648385). The primary outcome is change in RV longitudinal strain, as determined by cardiac MRI.

4.1.3 | Renin-angiotensin-aldosterone system

The concept of transferring therapeutic approaches from the systemic to the pulmonary circulation has refocussed attention on the role of the neurohumoral and renin-angiotensin-aldosterone systems (Maron & Leopold, 2015). Antagonists of the receptor for aldosterone, a steroid hormone that binds the mineralocorticoid receptor in the heart and pulmonary vasculature, have been used for fluid management in PAH. In animal studies, aldosterone antagonists attenuated or partially reversed PH (Preston et al., 2013). However, despite the potential of aldosterone antagonists to exert beneficial effects on the RV and pulmonary vasculature beyond their diuretic properties, no RCT has vet evaluated their efficacy in PAH. The "Combination Ambrisentan Plus Spironolactone in Pulmonary Arterial Hypertension Study" (CAPS-PAH)-a crossover study investigating whether addition of spironolactone to ambrisentan affects exercise capacity in patients with PAH-was terminated because of low enrolment (ClinicalTrials.gov identifier: NCT02253394). Another RCT will examine the effect of spironolactone on 6MWD and clinical worsening in patients with PAH (ClinicalTrials.gov identifier: NCT01712620).

Another novel approach addresses the peptidase ACE2 which converts the peptide hormones angiotensin I and angiotensin II to their vasodilator derivates (angiotensin-(1-9) and angiotensin-(1-7), respectively) with a preference for angiotensin II degradation. However, ACE2 may also degrade other vasodilatory factors such as apelin (see below; Kazemi-Bajestani, Patel, Wang, & Oudit, 2012). Decreased ACE2 levels and ACE2 autoantibodies have been reported in PAH (Tan, Liao, Zhou, Mei, & Wong, 2018). An openlabel pilot study on acute haemodynamic responses after a single infusion of recombinant human ACE2 in patients with PAH showed improved CO without a significant change in mPAP or systemic pressures, and reduced markers of oxidant and inflammatory stress (Hemnes et al., 2018). A dose-escalation study in PAH is currently recruiting patients (ClinicalTrials.gov identifier: NCT03177603).

4.1.4 | Atrial natriuretic peptide

Atrial natriuretic peptide (ANP) and brain natriuretic peptide (BNP) are released from atrial cardiomyocytes or the ventricles respectively. Natriuretic peptides increase renal sodium excretion, cause vasodilation via cGMP release, and decrease heart fibrosis. Natriuretic peptides are degraded by the metalloprotease neprilysin. Recently, a combination of the angiotensin receptor blocker valsartan and the neprilysin inhibitor sacubitril was approved for treatment of left heart failure (McMurray et al., 2014). The combination treatment was used because neprilysin degrades not only natriuretic peptides but also vasoconstrictive factors such as endothelin and angiotensin II. Studies of neprilysin in animal models of PH showed conflicting results. While neprilysin-deficient mice exhibited an exaggerated response to chronic hypoxia (Dempsey et al., 2009), pharmacological inhibition of neprilvsin alone or in combination with PDE5 inhibition attenuated PH in chronic hypoxic rats without affecting systemic circulation (Baliga et al., 2008; Thompson, Sheedy, & Morice, 1994). Recently, the efficacy and safety of the neprilysin inhibitor racecadotril was tested in an RCT in PAH. Acute administration increased plasma ANP and cGMP levels and slightly decreased PVR. Similar to the ACE2 trial, mPAP did not change significantly, indicating that the decrease in PVR was mostly driven by an increase in CO. Systemic BP was not decreased significantly, and plasma endothelin levels did not change following exposure to racecadotril (Hobbs et al., 2019). Further studies are necessary to investigate the effect of these vasodilators on PAH.

4.1.5 | Neurohormonal regulation

Neurohormonal regulation plays an important role in pulmonary vascular tone and RV function (Maron & Leopold, 2015). In fact, β-blockers (β-adrenoceptor antagonists) are one of the main treatment strategies in left ventricular systolic dysfunction and may thus have a place in PAH therapy (beyond their indication for supraventricular tachycardia which is a frequent co-morbidity in PAH). Animal studies suggest that β -blockers may have beneficial effects on RV dysfunction and maladaptive remodelling. It has also been suggested that some β -blockers have favourable effects on the pulmonary vasculature in PAH (reviewed in Ameri et al., 2016). Although β -blocker use in PAH was long regarded as non-beneficial, careful administration in stable PAH without decompensated heart failure was recently shown to have acceptable safety. Several small studies tested the effect of β -blockers on RV function and functional capacity with different outcomes. Thus, larger and longer studies are required to establish which patients (if any) might benefit from specific β -blockers in PAH. Currently, the use of β -blockers is not recommended in patients with PAH unless required for comorbidities (Perros et al., 2017). More recently, the impact of sympathetic denervation of the pulmonary arteries has been examined; Chinese investigators demonstrated a drop in pulmonary artery pressure using catheter radiofrequency ablation (Chen et al., 2013). Although serious concerns were raised regarding the study design and patient characteristics (Galie & Manes, 2013), a clinical trial is ongoing to examine intravascular ultrasound-directed pulmonary artery denervation in patients on dual oral combination therapy (Treatment of PH 1 Study [TROPHY1]; ClinicalTrials.gov identifier: NCT02835950/NCT02516722).

4.1.6 | 5-HT

5-HT (serotonin) has been implicated in the development of PAH since anorexigens, which increase the availability of 5-HT by inducing its release from platelets and inhibiting its reuptake and degradation by MAO, were noted to increase the risk of PAH (Abenhaim et al., 1996). Inhibition of the 5-HT_{2A} and 5-HT_{2B} receptors can inhibit development of PH in mouse models (Delaney et al., 2018; West et al., 2016). Nevertheless, the 5-HT_{2A} and 5-HT_{2B} receptor inhibitor terguride showed no clinical benefit in a Phase 2 study in PAH (Ghofrani et al., 2012), possibly because the 5-HT_{1B} receptor is the most highly expressed 5-HT receptor in the pulmonary arteries in PAH and mediates PASMC proliferation in humans (Lythgoe et al., 2016). However, pre-specified subgroup analysis indicated an improvement of PVR in patients on PAH background therapy with ERAs (Ghofrani et al., 2012). Thus, further studies are planned with a selective inhibitor of tryptophan hydroxylase 1 (TPH1) which is the rate-limiting enzyme in 5-HT biosynthesis. In preclinical PH models. the TPH1 inhibitor significantly reduced PH and showed an additive effect when applied together with ambrisentan, but not tadalafil (Aiello et al., 2017). A clinical Phase 1 study with the TPH1 inhibitor KAR5585 (ClinicalTrials.gov identifier: NCT02746237) showed a good safety profile and a decrease in circulating 5-HT (Paralkar et al., 2017). Furthermore, a trial with the 5-HT uptake inhibitor escitalopram was completed in 2008, but no results were published (ClinicalTrials.gov identifier: NCT00190333). Another trial with the selective 5-HT uptake inhibitor fluoxetine is planned (ClinicalTrials.gov identifier: NCT03638908), although a recent analysis using the "Registry to Evaluate Early and Long-term PAH Disease Management" (REVEAL) showed that incident selective 5-HT uptake inhibitor use was associated with increased mortality and a greater risk of clinical worsening, albeit without adjustment for all confounders (Sadoughi et al., 2013).

4.1.7 | Other hormones

Several other hormones have effects on the cardiopulmonary system. One of the long-standing hormones investigated in the field of PAH is vasoactive intestinal peptide (VIP), a peptide hormone that stimulates contractility in the heart and causes dilation of smooth muscles of different organs, including blood vessels. Despite failure of a previous clinical trial with inhaled administration of VIP (Said, 2012), a future study with a s.c. injected, sustained-release analogue of VIP (PB1046) is planned (ClinicalTrials.gov identifier: NCT03315507), because there were serious concerns that the inhaled route of administration may have affected the outcome negatively (Lythgoe et al., 2016). The best examples of endocrine dysfunction in PAH are thyroid disorders, which are common in PAH and associated with poor prognosis (Simonneau et al., 2019).

In summary, there are multiple approaches to balance the hormonal state in PAH. However, despite promising preliminary results, potential beneficial (and non-beneficial) effects of hormones independent of their action on the pulmonary vasculature/RV need to be carefully investigated.

4.2 | GPCR pathways

4.2.1 | Rho-associated protein kinase

Rho-associated protein kinase (ROCK) belongs to the family of serine-threonine kinases and can be activated via the GTPase RhoA by several cellular receptors, including GPCRs which are stimulated by various vasoactive substances such as angiotensin II or 5-HT. ROCK is involved in many cellular functions, for example, smooth muscle cell contraction, cell migration, and stress fibre formation, and has been implicated in the pathogenesis of PAH (Antoniu, 2012). ROCK inhibitors have shown promising results in animal studies (Antoniu, 2012). However, in a clinical trial, the 6MWD was not improved by the ROCK inhibitor **fasudil** (Fukumoto et al., 2013). Novel ROCK inhibitors are under development (Vaidya et al., 2017).

4.2.2 | Apelin

Apelin is an endogenous vasodilatory and inotropic peptide acting via the G protein-coupled **apelin receptor**. Apelin is down-regulated in human PAH and can inhibit PH in animal models. Use of apelin for PAH might be challenging due to its short $t_{1/2}$ and systemic vasodilatory effect (Kazemi-Bajestani et al., 2012; Kim, 2014). The effect of apelin infusion on cardiopulmonary performance in healthy volunteers and patients with IPAH was recently investigated in a small clinical trial (ClinicalTrials.gov identifier: NCT01590108). Results are still pending.

4.2.3 | Novel preclinical targets

Future approaches may include activation of NO release and inhibition of RhoA/ROCK signalling by activation of the G protein-coupled adenosine A2A receptor (Alencar, Montes, Barreiro, Sudo, & Zapata-Sudo, 2017). Salidroside, an active ingredient isolated from Rhodiola rosea, inhibited chronic hypoxia-induced PH and pulmonary arterial remodelling by increasing A_{2A} receptor expression and enhancing A_{2A} receptor-related mitochondria-dependent apoptosis (Huang et al., 2015). Another potential target is the Wnt/planar cell polarity pathway which involves the Frizzled family of GPCRs. Activation of the Wnt/planar cell polarity pathway was recently shown to be required for the establishment of human pulmonary endotheliumpericyte interactions. Loss of Wnt/planar cell polarity signalling could reduce the viability of newly formed vessels in PAH and thus contribute to vascular pruning. Interestingly, mice lacking endothelial expression of Wnt5a showed a similar response to chronic hypoxia as wildtype mice but failed to recover after re-exposure to normoxia (Yuan et al., 2019).

4.3 | Ion channels

Ion channels are crucial for cellular calcium homeostasis and thus regulate vasoconstriction and calcium-dependent transcription factors, but they also regulate cytosolic concentrations of other ions (e.g., potassium) which contribute to cellular survival. Dysregulated cell-type-specific ion channels have been extensively described in PAH and include potassium channels, different types of transient receptor potential channels, calcium sensor proteins, and chloride channels (Boucherat et al., 2015: Lambert et al., 2018). Moreover, the identification of novel heterozygous loss-of-function mutations in the KCNK3 gene in PAH has drawn attention to channelopathies as underlying mechanisms (Ma et al., 2013). in vivo pharmacological activation of KCNK3 had beneficial effects in monocrotaline-induced PH, demonstrating therapeutic relevance (Antigny et al., 2016). Very recent studies also suggest a role for glutaminergic NMDA receptors (Dumas et al., 2018). Despite these promising preclinical approaches, clinical trials addressing ion channels are currently lacking in PAH.

4.4 | Mitochondrial and metabolic adaptation

4.4.1 | Cellular metabolic alterations—The Warburg effect

The Warburg effect has been described in different animal models of PH (Bonnet et al., 2006; McMurtry et al., 2004), and several studies in humans have underlined the relevance of mitochondrial dysfunction and metabolic alterations in PAH (Farha et al., 2016; Rhodes et al., 2017; Xu et al., 2007). As the Warburg effect was originally described in cancer cells, its discovery in PH essentially contributed to the concept of the "cancer-hypothesis" in PAH (Archer et al., 2008). The Warburg effect is characterised by a switch of cellular metabolism from glucose oxidation to glycolysis despite the presence of oxygen and provides metabolic intermediates for macromolecule synthesis. Metabolic intermediates from the tricarboxylic acid cycle (e.g., acetyl-CoA or oxidised nicotinamide adenine dinucleotide [NAD+]) can support epigenetic alterations thereby promoting proliferation (see below).

The picture of mitochondrial alterations in PH is becoming increasingly detailed, including mitochondrial hyperpolarization, dysbalanced mitochondrial fission and fusion, altered mitochondrial calcium handling due to decreased expression of the mitochondrial calcium uniporter and disturbed interaction with the endoplasmic reticulum, increased glutaminolysis and accumulation of mitochondrial heat shock protein 90 (Boucherat et al., 2018; reviewed in detail in Chan & Rubin, 2017, and Culley & Chan, 2018). Most importantly, a causative role for mitochondrial alterations in the development of PH has been supported by the fact that targeting mitochondrial alterations, for example, using dichloroacetate (DCA), which promotes glucose oxidation, or the mitochondrial fission inhibitor Mdivi or by inhibiting fatty acid oxidation, inhibited PH in animal models (Chan & Rubin, 2017; Culley & Chan, 2018).

Encouraged by the promising preclinical results, an open-label clinical trial was performed to test the effect of DCA in patients with IPAH (Michelakis et al., 2017). After 4 months of DCA treatment, the average mPAP and PVR decreased and functional capacity increased, but some of the patients did not respond to treatment. Four of the 20 patients withdrew from the study due to neuropathic side effects (all of them from the highest dose group). Detailed analysis showed that lack of treatment response was associated with loss-of-function polymorphisms of specific genes that affect mitochondrial metabolism (sirtuin 3 [SIRT3] and uncoupling protein 2 [UCP2]). Interestingly, mice deficient in UCP2 exhibited PH under baseline conditions (Dromparis et al., 2013; Pak et al., 2013). SIRT3 deficiency has also been connected to spontaneous development of PH (Paulin et al., 2014), but animal studies have not been completely consistent (Waypa et al., 2013). The role of SIRT3 is described in more detail in Section 4.6. Although the DCA trial only showed limited treatment effects, it was an important step to show the feasibility of mitochondrial targeted therapy in PH and to demonstrate the practical use of individualised therapy. Further large-scale studies are needed to investigate the efficacy of DCA in specific patients at different doses. However, DCA may not be optimal for PAH treatment, because of its rapid clearance, highly variable exposure (with individually varying inactivation by GSH transferase zeta 1) and small therapeutic window (Rowlands, 2016), as well as the dose-limiting risk of neuropathy in adults. Further research is necessary to identify novel mitochondrial targets or optimised alternatives to DCA as a promising approach for treatment of PAH.

The concept of mitochondrial dysfunction in PAH has also been applied to the RV, with promising preclinical results (see Chan & Rubin, 2017). However, one must consider that altering mitochondrial function may have opposing effects on the RV and pulmonary vasculature. For example, although UCP2 deficiency is associated with spontaneous PH and RV dysfunction under baseline conditions (Esfandiary et al., 2019; Pak et al., 2013), UCP2-deficient mice show better RV adaptation to increased afterload (induced by pulmonary arterial banding) than wild-type mice (Esfandiary et al., 2019). More details on mitochondrial dysfunction and anti-apoptotic mechanisms were recently summarised by Boucherat, Vitry, et al. (2017).

4.4.2 | Systemic metabolic alterations

PAH has been associated with glucose intolerance and insulin resistance. Glucose intolerance was found to be a predictor of mortality in PAH (Belly et al., 2012). A more detailed study recently showed similar glucose intolerance in patients with PAH and controls when matched for the metabolic syndrome, but profound alterations in lipid metabolism and lipid-related insulin resistance (Hemnes et al., 2019). It remains unclear if these alterations contribute to the pathogenesis of PH or are consequences of PH. Nevertheless, optimisation of blood glucose level may have beneficial effects in PAH, although this has not yet entered the BJP BRITISH PHARMACOLOGICAL

guidelines for PAH treatment beyond optimal standard of care for diabetes. Several clinical trials now address this question in PAH and PH due to heart failure with preserved ejection fraction. A Phase 2 clinical trial is investigating the impact of 12 weeks of metformin treatment on 6MWD and further functional parameters, RV and left ventricular performance, and several metabolic and hormonal parameters (ClinicalTrials.gov identifier: NCT03617458). Inhibitors of dipeptidyl peptidase 4 (gliptins) which increase glucagon-like peptide 1 (GLP-1) and insulin release may also be of interest; sitagliptin alleviated PH induced by monocrotaline, bleomycin, or hypoxia in rats (Xu et al., 2018). The GLP-1 receptor agonist liraglutide both prevented and reversed monocrotalineinduced PH, RV hypertrophy, and pulmonary vascular wall remodelling (Lee et al., 2016). The concept of using statins to improve PAH by addressing systemic lipid metabolism has been unsuccessful (Kawut et al., 2011; Zeng et al., 2012).

Another concept identified in the guidelines for PAH treatment is related to iron metabolism. Although the underlying mechanisms are not completely understood, a recent open-label study showed that i.v. iron supplementation in patients with PAH increased exercise endurance capacity, although 6MWD was not changed significantly (Ruiter et al., 2015). This study included only patients without anaemia, and blood Hb content did not change significantly after iron infusion. In this regard, several groups have shown that iron deficiency in the absence of anaemia is common and associated with reduced survival in PAH (Rhodes et al., 2011). Another pilot study demonstrated an increase in 6MWD in patients with PAH and iron deficiency receiving i.v. iron supplementation compared with matched patients without iron deficiency who did not receive iron supplementation. However, Hb levels were also increased in the treated patients (Viethen et al., 2014). A double-blind study of iron infusion in iron-deficient patients with PAH was recently completed, and results are pending (ClinicalTrials.gov identifier: NCT01447628).

4.5 | Epigenetic alterations and interaction with metabolic pathways

The role of epigenetics in PAH is a fast-growing field of research. The major epigenetic phenomena include DNA methylation, histone modifications, and modulations of non-coding RNAs (reviewed in Chelladurai et al., 2019; Pullamsetti, Perros, Chelladurai, Yuan, & Stenmark, 2016). Despite encouraging preclinical studies, few of them have reached the clinical stage in PAH yet, mostly because their modulation is often associated with unwanted effects and strategies to therapeutically deliver non-coding RNA mimics or inhibitors to the lungs remain in their infancy (Meloche, Paulin, Provencher, & Bonnet, 2015).

Recently, clinical interest in histone deacetylase (HDAC) inhibition in PAH has been rekindled by the discovery that the cytosolic HDAC6 is involved in both pulmonary arterial remodelling and RV failure (Boucherat, Chabot, et al., 2017). This isoenzyme represents an important pharmacological target for selective inhibition that may reduce the toxicity related to the off-target effects of pan-HDAC inhibitors previously described in PAH (Bogaard et al., 2011). Other HDACs known as Sirtuins are also implicated in PAH. The Sirtuins are NAD+-dependent HDACs regulating important metabolic pathways involved in many biological processes such as cell survival, proliferation, apoptosis, DNA repair, and cell metabolism all of which are critical to PAH development. Consistent with these findings, mice lacking SIRT3, a mitochondrial deacetylase, have increased acetylation and inhibition of many mitochondrial enzymes and complexes, suppressing mitochondrial oxidative metabolism. These mice spontaneously develop PH; a loss-of-function SIRT3 polymorphism is associated with PAH development in humans (Paulin et al., 2014). The importance of this metabolism-epigenetics axis has been further highlighted by the recent clinical trial results using DCA, a pyruvate dehydrogenase kinase inhibitor known to promote glucose oxidation (see above). The DCA trial showed that functional variants of SIRT3 and UCP2 largely influenced the clinical and haemodynamic response to DCA (Michelakis et al., 2017). Although not fully understood, the downregulation of both SIRT1 and SIRT3 in PAH might also result from activation of PARP-1 (Meloche et al., 2014), which could cause depletion of NAD+ (SIRT substrate) levels, which inhibits SIRT1 activity (Meloche et al., 2014). Inhibition of PARP1 in conjunction with standard combination therapy (ERA + PDE5 inhibitor) in an experimental PH model showed greater efficacy than standard combination therapy alone (Meloche et al., 2014), and thus, the U.S. Food and Drug Administration-approved PARP1 inhibitor olaparib is under clinical investigation in the Olaparib for PAH study (OPTION: ClinicalTrials. gov identifier: NCT03782818).

The epigenetic/metabolism/DNA damage response axis described above in PAH is very similar to that described in cancer. This cancer theory of PAH (Boucherat, Vitry, et al., 2017) is further reinforced by the implication of the newly described epigenetic reader, bromodomain-containing protein 4 (BRD4), in PAH. Similar to cancer cells in which BRD4 has been shown to promote several oncogenes implicated in PAH pathogenesis, including c-Myc, B-cell lymphoma 2 (Bcl-2), cyclin-dependent kinase inhibitor 1 (p21), cyclin-dependent kinase inhibitor 1B (p27), Runt-related transcription factor 2 (RUNX2), and FOXM1 (Belkina & Denis, 2012), we recently documented that BRD4 is significantly overexpressed in human PAH, accounting for the up-regulation of the oncogenic NFAT, Bcl-2, Survivin, and p21 and triggering the proliferation/apoptosis imbalance in PAH-PASMCs (Meloche et al., 2015; Meloche et al., 2017). BRD4 was similarly up-regulated in PH rat models in which its inhibition improved pulmonary haemodynamics, RV function, and distal pulmonary arterial remodelling. Although these effects were attributed to the modulation of NFAT, Bcl-2, Survivin, and p21, other mechanisms cannot be excluded. The mechanisms accounting for BRD4 inhibitor efficacy in PAH are the subject of numerous published and ongoing preclinical studies. In addition to its effects on the cancer-like phenotype of PASMCs, the inhibition autoimmune-mediated/inflammatory vascular of iniuries. RUNX2-mediated pro-calcification processes, and its influence on metabolism and DNA damage are suggested mechanisms of

14

therapeutic intervention by BRD4 inhibitors in PAH. Recently, a clinically available BRD4 inhibitor reversed PH in two independent animal studies potentially through FOXM1 (Van der Feen et al., 2019). Altogether, these findings support a therapeutic role for BRD4 inhibitors in PAH, which is currently being explored in the Apabetalone for PAH Pilot Study (APPRoAcH-p; ClinicalTrials. gov identifier: NCT03655704).

4.6 | Oxidative stress-related pathways

Oxidative stress has been implicated in the development of PAH, although animal studies in this regard are inconclusive, depending on the model used (Fulton et al., 2017). Increased mitochondrial ROS release might be more relevant for RV remodelling than PH, at least in the hypoxic mouse model (Pak et al., 2018). Increased levels of markers of oxidatively modified proteins (e.g., nitrotyrosine), fatty acids (e.g., malondialdehyde), and DNA have been detected in lung sections, plasma, or urine of patients with PAH (Bowers et al., 2004; Cracowski et al., 2001; Irodova, Lankin, Konovalova, Kochetov, & Chazova, 2002) but the clinical relevance remains unclear. A small open-label clinical trial with the antioxidant ubiquinol showed a reduction of indirect signs of oxidative stress and right atrial pressure after 3 months of treatment. but no alterations of BNP/6MWD could be detected (Sharp et al., 2014). Non-specific inhibition of ROS may result in disappointing outcomes, because low levels of ROS are necessary to maintain physiological cellular functions.

The pleiotropic effects of ROS on metabolism and inflammation in PH were recently explored. Inhibition of apoptosis signalregulating kinase 1 (ASK1; MAP3K5) a serine/threonine kinase that is activated by oxidative stress, promoting inflammation, ROS production, proliferation, fibrosis, apoptosis, mitochondrial damage, and, under certain contexts, insulin resistance) reduced pathological remodelling and halted progression of PH in rodent models (Budas et al., 2018). Despite these encouraging results, a Phase 2 study (ClinicalTrials.gov identifier: NCT02234141) evaluating the ASK1 inhibitor selonsertib did not achieve its primary endpoint in PAH (Boucherat, Provencher, & Bonnet, 2018). Conversely, a Phase 2 clinical trial evaluating bardoxolone, an activator of the nuclear factor erythroid 2-related factor 2 (Nrf2), a transcription factor that regulates antioxidant proteins and suppresses activation of the proinflammatory factor NF-kB, showed significant improvements in exercise capacity in PAH (LARIAT; ClinicalTrials.gov identifier: NCT02036970), albeit mainly in connective tissue disease (CTD)associated PAH (Oudiz et al., 2017). Thus, a follow-up trial in patients with CTD-PAH was initiated (CATALYST; ClinicalTrials.gov identifier: NCT02657356), and a Phase 3 clinical trial evaluating long-term safety of bardoxolone in patients with PAH is ongoing (RANGER; ClinicalTrials.gov identifier: NCT03068130). Results of a Phase 2 clinical trial evaluating CXA-10, another Nrf2 activator and NF-kB suppressor, are also pending (PRIMEx; ClinicalTrials.gov identifier: NCT03449524).

P _____BRITISH _____15

4.7 | Inflammatory mediators

4.7.1 | Immune modulators in PAH

Immune modulation is an established concept in PAH, but classic antiinflammatory drugs such as corticosteroids or acetylsalicylic acid have shown beneficial effects only in specific forms of PAH (Sanchez, Sitbon, Jais, Simonneau, & Humbert, 2006) or no beneficial effects (Kawut et al., 2011). However, interest in the field has been revived by an improved understanding of immune regulation in PAH and the notion that perivascular inflammatory infiltrates (macrophages, B-cells, T-cells, and dendritic cells) often precede structural pulmonary vascular remodelling (Tamosiuniene et al., 2011). Histopathological studies have demonstrated the presence of complement system components, autoantibodies, and inflammatory cells (neutrophils) in the vessel lumen, which can bind to the endothelium and may infiltrate the medial muscular layer. The inflammatory infiltrate in the neointimal laver is composed of T- and B-lymphocytes, with macrophages, mast cells, and dendritic cells present in the adventitial layer. Lymphoid follicles, characterised by T-cells, B-cells, and plasmacytoid dendritic cells, are found in the periadventitial space (see Rabinovitch, Guignabert, Humbert, & Nicolls, 2014). Although current knowledge of the immune system is far from complete, increasing evidence suggests that excessive local secretion of inflammatory mediators by pulmonary vascular cells (e.g., IL-1^β, IL-6, LTB₄, macrophage migration inhibitory factor, leptin, and $TNF-\alpha$) as well as dysregulated immune responses (innate [through macrophages and monocytes] and adaptive [impaired T-regulatory cell function and T-helper 17 cell immune polarisation]) are major drivers of pulmonary remodelling in PAH with or without autoimmune diseases (see Kuebler, Bonnet, & Tabuchi, 2018; Rabinovitch et al., 2014).

Several immune modulatory approaches have been successfully tested in animal models, including an IL-1 receptor antagonist, IL-6 antibodies, mycophenolate, dexamethasone, cyclosporine. tacrolimus, LTB₄ pathway inhibitors, and TNF-related apoptosisinducing ligand (Rabinovitch et al., 2014), but none have yet made the transition to treatment of human PAH beyond CTD-PAH. However, several clinical trials addressing immune modulation are ongoing, including a clinical trial of the anti-CD20 monoclonal antibody rituximab (which targets B-lymphocytes) in PAH associated with systemic sclerosis (ClinicalTrials.gov identifier: NCT01086540). Rituximab is often used to treat conditions for which there is a clonal source of autoantibodies. Interestingly, plasmablasts in IPAH displayed clonal expansions similar to those observed in autoimmune disease (Blum et al., 2018). Anakinra is a recombinant IL-1 receptor antagonist used to treat rheumatoid arthritis. The safety of anakinra in PAH was recently evaluated in an open-label study. After 14 days of treatment, high-sensitivity C-reactive protein and symptom burden were significantly reduced (Trankle et al., 2019). The recently completed trial of the IL-6 inhibitor tocilizumab in PAH (TRANSFORM-UK; ClinicalTrials.gov identifier: NCT02676947) showed no significant change in PVR from baseline at 6 months in the 19 patients who completed the study protocol (published online at https://www. clinicaltrialsregister.eu/ctr-search/trial/2015-002799-26/results). A Phase 2 open-label extension study with **ubenimex** was recently terminated after the original study failed to demonstrate efficacy (ClinicalTrials.gov identifier: NCT02736149). A different approach uses the endogenous protein elafin which inhibits the neutrophilderived serine proteases elastase and proteinase-3. Elafin reduced or reversed PH in rats exposed to hypoxia or Sugen/hypoxia respectively (Nickel et al., 2015; Zaidi, You, Ciura, Husain, & Rabinovitch, 2002). Elafin has now progressed to a clinical trial in healthy volunteers (ClinicalTrials.gov identifier: NCT03522935).

4.7.2 | Novel preclinical approaches

Recently, it was reported that fucoidan, a polysaccharidic ligand of the adhesion molecule P-selectin, exhibits anti-proliferative properties and can attenuate hypoxia-induced PH in mice. P-selectin is expressed on endothelial cells and is involved in recruitment of leukocytes and platelets to areas of inflammation and vascular injury respectively (Novoyatleva et al., 2019). Moreover, the damageassociated high mobility group box-1 (HMGB1)-which is secreted from immune cells, activates macrophages as well as lymphocytes, and triggers autoimmunity by binding to the toll-like receptor 4 (TLR4)-was shown to be involved in development of PH. Expression of both HMGB1 and TLR4 was elevated in the lungs of patients with PAH. An inhibitor of binding between HMGB1 and TLR4 reduced PH severity and mortality in animal models, even when given to animals with established disease (Goldenberg et al., 2019). Thus, further novel pathways involved in inflammation may be exploited in future for treatment of PH.

4.8 | Growth factor receptors

Receptor TKs (RTKs) and the TGF β superfamily of growth factor receptors are both critically involved in PAH.

4.8.1 | RTKs

RTKs are cell surface receptors for growth factors, cytokines and hormones. RTKs which have been implicated in PAH include PDGF, EGF, FGF, VEGF, and nerve growth factor receptors. Upon activation by growth factors, RTKs dimerise and become autophosphorylated. Binding of factors such as Src and PLCγ leads to activation of several downstream kinases including PI3K/PKB (Akt), MAPKs, PKC, JAK, STAT, and cyclin-dependent kinases (Lemmon & Schlessinger, 2010). Several RTK inhibitors are used to treat different kinds of cancer, including **imatinib**, **dasatinib**, and **nilotinib** (inhibiting the PDGF **receptor**), the multikinase inhibitors **sorafenib** and **sunitinib** (inhibiting the PDGF and VEGF receptors), and nilotinib (inhibiting the BCR-ABL TK; Lemmon & Schlessinger, 2010). The PDGF receptor inhibitor imatinib was the first RTK used in clinical PAH and the first drug used in PAH to directly target vascular remodelling. However, despite reducing PVR and increasing mean placebo-corrected 6MWD (by 32 m) in a clinical Phase 3 trial (Imatinib in PAH, a Randomised Efficacy Study [IMPRES]), imatinib was not approved for PAH owing to an unfavourable risk-benefit ratio; the imatinib-treated group did not improve time to clinical worsening and showed subdural haematoma in eight patients receiving both imatinib and anticoagulation with vitamin K antagonists (Hoeper et al., 2013). However, treatment with imatinib is currently the only approach beyond the classical, approved and mainly vasodilator therapy that was shown to offer considerable benefit at least in specific patients (Ghofrani et al., 2010; Ghofrani, Seeger, & Grimminger, 2005; Hoeper et al., 2013). Thus, interest in further trials with imatinib and defining groups of patients with an optimal risk-benefit ratio remains high. Moreover, the efficacy of imatinib opened the door to the quest for alternative RTKs with more favourable profiles. Unfortunately, none of the tested RTKs met expectations. A clinical trial with nilotinib was terminated due to severe adverse events (ClinicalTrials.gov identifier: NCT01179737). Exposure to dasatinib was even associated with the development of PAH (Guignabert et al., 2016; Montani et al., 2012). Nintedanib was unsuccessful in animal models and when used to treat single patients with PAH on a compassionate basis (Richter et al., 2018). Sorafenib was tested in an open-label study and showed some efficacy (improvement of BNP, uric acid and cardiothoracic ratio of more than 10% from the original value continued for 1 month) in two out of nine patients. However, CO also decreased in most patients treated with sorafenib, raising serious concerns about further trials of sorafenib in PAH (Kimura et al., 2017; Weatherald, Humbert, Guignabert, & Montani, 2017). A similar decrease of CO was observed in a previous open-label 1b trial (Gomberg-Maitland et al., 2010).

4.8.2 | TGF β superfamily of growth factors

An imbalance in TGF β /bone morphogenetic protein (BMP) signalling has long been known as an important pathogenetic mechanism in PAH, since mutations in *BMPR2*, a TGF β receptor (TGF β R) subtype, were identified in heritable PAH in 2000 (Lane et al., 2000). Recently, heterozygous mutations of *BMPR2* were identified in 15.3% of patients from a U.K. PAH cohort (76% in familial PAH, 12% in sporadic cases, and 8% in anorexigen-exposed PAH cases; Graf et al., 2018). Decreased **BMPR2** signalling has also been detected in patients with PAH without *BMPR2* mutations. However, attempts to address TGF β pathways therapeutically have only recently been transferred to the clinic.

TGF β /BMP signalling is transduced by a combination of different type I (ALK1 to ALK7) and type II (e.g., TGF β R2, BMPR2, and **activin** 2A and 2B receptors) receptors as well as co-receptors (betaglycan and endoglin). These multicomponent receptors activate canonical downstream signalling via either Smad 2/3 (e.g., through TGF β R2 or activin receptors together with ALK1, ALK2, ALK3, or ALK6) or Smad 1/5/8 (e.g., through BMPR2 or activin receptors together with ALK4, ALK5, or ALK7), or non-canonical signalling via different kinases

BRITISH PHARMACOLOGICAL

(e.g., LIM domain kinase 1). Different ligands (e.g., TGF β , BMPs, and activin) can activate the receptors with different affinity and induce partially opposing effects with regard to development of PAH. While activation of Smad 2/3 signalling (e.g., by TGF β) results in gene expression promoting PAH, activation of Smad 1/5/8 signalling (e.g., by BMPs) inhibits alterations associated with PAH such as vascular smooth muscle growth (Rol, Kurakula, Happe, Bogaard, & Goumans, 2018). Thus, inhibition or activation of the respective receptor pathway may represent a promising strategy to treat PAH, although balancing the response might be challenging due to overlapping ligand and receptor functions.

The most straightforward therapeutic approach may be activation of BMPR2-Smad 1/5/8 signalling using BMPR2 receptor agonists (e.g., specific ligands) or inhibitors of degradation of BMPR2, such as chloroquine (Thompson & Lawrie, 2017). With regard to BMPR2 activation, tacrolimus (FK506) was recently found to promote BMPR2-Smad1/5/8 signalling by interacting with its pharmacological target, 12-kDa FK506-binding protein (FKBP12), thereby removing FKBP12 from all three BMP type 1 receptors (ALK1, ALK2, and ALK3), including those preferred by BMPR2 (ALK1 and ALK3; Sitbon et al., 2019). Tacrolimus prevented the development of PAH in mice with endothelial deletion of BMPR2 and reversed established PAH in two rat models (Spiekerkoetter et al., 2013). Compassionate treatment with tacrolimus showed promising results in single patients with IPAH (Spiekerkoetter et al., 2015) but did not attenuate the progression of PAH in a patient with an ALK1 mutation (Sommer et al., 2019). A Phase 2 clinical trial of tacrolimus in patients with PAH showed favourable safety, but efficacy signals were low (Spiekerkoetter et al., 2017).

Another strategy to address TGFβ signalling involves inhibition of the Smad 2/3 signalling pathway using sotatercept, an activin receptor type 2A fusion protein that acts as a ligand trap. In a preclinical study, sotatercept reversed pulmonary vascular remodelling and restored RV function (Joshi, Liu, Pearsall, Li & Kumar, 2019). A Phase 2 open-label study (SPECTRA; ClinicalTrials.gov identifier: NCT03738150) and a Phase 2 RCT (PULSAR; ClinicalTrials.gov identifier: NCT03496207) are currently recruiting patients with PAH. Primary outcomes will be peak oxygen uptake and PVR respectively.

4.8.3 | Novel preclinical approaches

Smad ubiquitin regulatory factor 1 (SMURF-1) is involved in the degradation of BMPR2 and different Smads and is up-regulated in PAH. Nebulised administration of microRNA-140-5p, shown to target and repress SMURF-1, attenuated PAH in rat models (Rothman et al., 2016). Use of chloroquine to inhibit lysosomal degradation of BMPR2 also restored BMPR2 signalling in vitro and prevented development of monocrotaline-induced PH in rats (Long et al., 2013).

Application of recombinant **BMP9** reversed PH in several animal models, including a mouse model with a human loss-of-function mutation in *Bmpr2* (Long et al., 2015). By contrast, BMP9 deficiency or inhibition prevented development of PH in mice exposed to chronic

hypoxia (Tu et al., 2019). It was recently argued that because of the complexity of the BMP system, inhibition of endogenous BMP9 and application of exogenous BMP9 may not be comparable and that inhibition of endogenous BMP9 may inhibit muscularisation but also induce an endothelial pathology similar to that observed in hereditary haemorrhagic telangiectasia (Ormiston, Godoy, Chaudhary, & Stewart, 2019).

Recently, fragile histidine triad (FHIT) was discovered as a novel BMPR2 modifier by a high-throughput screen. FHIT works as a tumour suppressor, and its expression is reduced in patients with PAH. Up-regulation of FHIT by **enzastaurin** reversed PH in the Sugen/hypoxia rat model (Dannewitz Prosseda et al., 2019).

4.9 | Transcription factors

4.9.1 | PPAR-γ

PPAR- γ belongs to a group of nuclear receptor proteins and functions as a transcription factor. When activated, PPAR-y improves insulin sensitivity and has anti-inflammatory properties. Both preclinical and human studies have demonstrated the importance of PPAR-γ in PAH. Activation of PPAR-y signalling by rosiglitazone and pioglitazone (PPAR-y agonists used to treat Type 2 diabetes mellitus) inhibited development of PH in animal models (reviewed in Prins et al., 2019). However, despite the promising preclinical results, a clinical trial with pioglitazone (ClinicalTrials.gov identifier: NCT00825266) was terminated early. Furthermore, rosiglitazone has been withdrawn from the market in certain countries due to suspected cardiovascular side effects, and pioglitazone is suspected to cause bladder cancer. Moreover, the doses in preclinical PH studies were significantly higher than the maximal doses used in patients (Prins et al., 2019). Nevertheless, the quest for novel therapeutics exploiting PPAR-y signalling continues. An inhaled therapy combining sildenafil and rosiglitazone was successfully applied in rats with PH in a recently published study (Rashid, Nozik-Grayck, McMurtry, Stenmark, & Ahsan, 2019).

4.9.2 | HIF and mechanistic target of rapamycin signalling

Hypoxia causes pulmonary vasoconstriction, and hypoxic signalling is also suspected to play a role in PAH. This concept is supported by the finding of decreased expression of prolyl hydroxylase domaincontaining protein 2 (PHD2), which inhibits HIF, in PAEC from plexiform lesions, and by the fact that PHD2-deficient mice develop severe, HIF-2 α -dependent PH (Dai, Li, Wharton, Zhu, & Zhao, 2016). Accordingly, the HIF-2 α translation inhibitor C76 reduced PH in three different rodent models (Dai et al., 2018). At least for hypoxia-induced PH, HIF-1 α also seems to be important (Ball et al., 2014).

One strategy to address HIF signalling is via inhibition of mechanistic target of rapamycin (mTOR) which increases HIF expression. mTOR can act as a serine/threonine kinase or TK and is

activated (for example) by growth factor receptors, but it also integrates signals from high nutrient availability to regulate multiple cellular functions, such as proliferation, motility, survival, and cell cycle progression (e.g., in T-cells). Thus, mTOR inhibitors such as **rapamycin** are used as immunosuppressants, and clinical trials of rapamycin in (for example) pancreatic cancers are currently ongoing. Inhibition of mTOR reduced PH in several animal models (Goncharov et al., 2014; Paddenberg et al., 2007). Moreover, PAH was recently improved in a patient treated with rapamycin for pancreatic cancer (Wessler, Steingart, Schwartz, Harvey, & Schaffer, 2010). Currently, a Phase 1 open-label clinical trial of ABI-009, an albumin-bound version of rapamycin, is recruiting patients with severe PAH (ClinicalTrials.gov identifier: NCT02587325).

4.9.3 | Novel preclinical targets

Another transcription factor with high potential as a therapeutic target is FOXO1, which integrates several anti-proliferative and pro-apoptotic signalling pathways in PASMC. Total FOXO1 levels were decreased in animal models of PH and in human PAH, and activation of FOXO1 reversed PH in animal models (Savai et al., 2014). Targeting the FOXO1 pathway with paclitaxel-which is currently used as a chemotherapeutic agent for different types of cancer-could be a viable strategy for the treatment of PAH. Many other transcription factors have been implicated in the development of PH, including Notch receptor 3, STAT3, large tumour suppressor 1 (central component of the Hippo pathway), myocyte enhancer factor 2, tumour protein p53, kruppel-like factor 4. CCAAT-enhancer binding proteins. RUNX2. activator protein 1, NF-κB, β-catenin, the Twist family basic helix-loop-helix transcription factor 1, and SLUG. Moreover, transcription factor coactivators have also been implicated in the development of PH (see Humbert et al., 2019; Pullamsetti et al., 2016).

5 | WHERE TO GO IN FUTURE?

There is consensus that PAH therapy must go beyond targeting primarily vasodilation. Although a wide range of novel targets addressing vascular and RV remodelling have been identified by preclinical studies, the challenge will be prioritising targets and designing clinical trials for successful target validation. The success rate of clinical trial programmes (measured as the proportion of drugs entering clinical development that reach the marketplace) has decreased from 21% in the early 1990s to 16%; about 50% of drugs entering clinical development fail because of lack of efficacy (25% fail because of safety concerns; Lythgoe et al., 2016). In order to prioritise targets, one has to consider the set-up of the preclinical study with regard to limitations of different animal models, species-specific pathways, and drug dose requirements.

How can we overcome the gap between animal studies and clinical trials? Considering the high success rate in animal models,

one should ask if the animal models oversimplify the patho-mechanisms. The predominant stimulus in the chronic hypoxic animal model is HIF-1 stabilisation, while the monocrotaline model is largely driven by inflammation. By contrast, human PAH is triggered by more than one stimulus acting over a long time period. Multifactorial pathogenesis and individually varying pathways may hamper disease targeting in individual patients (Lythgoe et al., 2016; Provencher et al., 2018). Thus, the use of several animal models (possibly in a multicenter approach) and animal models induced by more than one hit (e.g., Sugen/hypoxia or BMPR2 knockout plus hypoxia) might be a good step forward for preclinical studies of potential PAH therapies, although animal models reflecting end-stage disease are probably still lacking. The use of mostly very young and often male mice for preclinical studies may also be a limiting factor. Moreover, potential new therapies should be tested on a background of existing PAH therapy, as most patients in clinical trials will be pretreated with approved therapy. Finally, one should consider that many pathways (particularly immunological pathways, Mestas & Hughes, 2004, but also metabolic pathways and distribution of specific vascular receptors and enzymes) differ between mice and humans. Thus, transfer to the human system at any step of research is mandatory and should include more sophisticated models, such as organ culture, cultured lung slices, or induced pluripotent stem cells to derive endothelial cells. Nevertheless, animal models may offer optimal conditions to identify specific pathways that trigger the disease in a particular patient population.

Identifying the patient population which may profit most from a specific treatment ("enrichment") and applying individualised medicine will allow successful targeting of very specific pathways (Sitbon et al., 2019). Great efforts have been made in the past years using genomics, epigenomics, and metabolomics to identify sub-clusters within patient groups (Pulmonary Vascular Disease Phenomics [PVDOMICS]; ClinicalTrials.gov identifier: NCT02980887). Thus, it will be more important for future clinical trials to utilise biomarkers to focus on specific patient subgroups (although this will further limit the number of patients eligible for enrolment). Moreover, detailed patient phenotyping can help to find pathways in common with animal studies and thus prioritise targets. Finally, specific effects of treatment on the pulmonary vasculature and RV have to be taken into account in future.

5.1 | Nomenclature of targets and ligands

Key protein targets and ligands in this article are hyperlinked to corresponding entries in http://www.guidetopharmacology.org, the common portal for data from the IUPHAR/BPS Guide to PHARMACOLOGY (Southan et al., 2016), and are permanently archived in the Concise Guide to PHARMACOLOGY 2019/20 (Alexander, Christopoulos et al., 2019; Alexander, Cidlowski et al., 2019; Alexander, Fabbro et al., 2019a, 2019b; Alexander, Kelly et al., 2019a, 2019b).

ACKNOWLEDGEMENTS

Claire Mulligan (Beacon Medical Communications Ltd, Brighton, UK) edited the final draft, funded by the Justus-Liebig-University, Giessen, Germany.

CONFLICT OF INTEREST

N.S. has received personal fees from Actelion Pharmaceuticals and Bayer. S.P. is clinician-scientist of the Fonds de Recherche en Santé du Québec and has received speaker and consultant fees from Actelion Pharmaceuticals and research grants from Actelion Pharmaceuticals. Astra-Zeneca (in-kind), Boehringer-Ingelheim, and Resverlogix. S.B. holds a Canadian research chair; CIHR and heart and stroke foundation grants. He received a grant from Resverlogix and Astra Zeneca provided in kind contribution (provided the drug) for a clinical trial. M.H. has received fees for lectures and consultations from Actelion Pharmaceuticals, Bayer, MSD, and Pfizer. D.K. has received personal fees, research funding, and funding to attend educational meetings from Actelion Pharmaceuticals and GSK and personal fees and funding to attend meetings from MSD and Bayer.

REFERENCES

- Abenhaim, L., Moride, Y., Brenot, F., Rich, S., Benichou, J., Kurz, X., ... Bégaud, B. (1996). Appetite-suppressant drugs and the risk of primary pulmonary hypertension. International Primary Pulmonary Hypertension Study Group. The New England Journal of Medicine, 335, 609–616. https://doi.org/10.1056/NEJM199608293350901
- Aiello, R. J., Bourassa, P. A., Zhang, Q., Dubins, J., Goldberg, D. R., de Lombaert, S., ... Paralkar, V. (2017). Tryptophan hydroxylase 1 inhibition impacts pulmonary vascular remodeling in two rat models of pulmonary hypertension. *The Journal of Pharmacology and Experimental Therapeutics*, 360, 267–279. https://doi.org/10.1124/jpet.116. 237933
- Alencar, A. K. N., Montes, G. C., Barreiro, E. J., Sudo, R. T., & Zapata-Sudo, G. (2017). Adenosine receptors as drug targets for treatment of pulmonary arterial hypertension. *Frontiers in Pharmacology*, *8*, 858. https://doi.org/10.3389/fphar.2017.00858
- Alexander, S. P. H., Christopoulos, A., Davenport, A. P., Kelly, E., Mathie, A., Peters, J. A., ... CGTP Collaborators (2019). THE CONCISE GUIDE TO PHARMACOLOGY 2019/20: G protein-coupled receptors. *British Journal of Pharmacology*, 176, S21–S141. https://doi.org/10. 1111/bph.14748
- Alexander, S. P. H., Cidlowski, J. A., Kelly, E., Mathie, A., Peters, J. A., Veale, E. L., ... CGTP Collaborators (2019). THE CONCISE GUIDE TO PHARMACOLOGY 2019/20: Nuclear hormone receptors. *British Journal of Pharmacology*, 176, S229–S246. https://doi.org/10.1111/bph. 14750
- Alexander, S. P. H., Fabbro, D., Kelly, E., Mathie, A., Peters, J. A., Veale, E. L., ... CGTP Collaborators (2019a). THE CONCISE GUIDE TO PHARMACOLOGY 2019/20: Catalytic receptors. *British Journal* of Pharmacology, 176, S247–S296. https://doi.org/10.1111/bph. 14751
- Alexander, S. P. H., Fabbro, D., Kelly, E., Mathie, A., Peters, J. A., Veale, E. L., ... CGTP Collaborators (2019b). THE CONCISE GUIDE TO PHARMACOLOGY 2019/20: Enzymes. *British Journal of Pharmacol*ogy, 176, S297–S396. https://doi.org/10.1111/bph.14752
- Alexander, S. P. H., Kelly, E., Mathie, A., Peters, J. A., Veale, E. L., Faccenda, E., ... CGTP Collaborators (2019a). THE CONCISE GUIDE TO PHARMACOLOGY 2019/20: Introduction and Other Protein Targets. British Journal of Pharmacology, 176, S1–S20. https://doi.org/10. 1111/bph.14747

- Alexander, S. P. H., Kelly, E., Mathie, A., Peters, J. A., Veale, E. L., Armstrong, J. F., ... CGTP Collaborators (2019b). THE CONCISE GUIDE TO PHARMACOLOGY 2019/20: Transporters. *British Journal* of *Pharmacology*, 176, S397–S493. https://doi.org/10.1111/bph. 14753
- Ameri, P., Bertero, E., Meliota, G., Cheli, M., Canepa, M., Brunelli, C., & Balbi, M. (2016). Neurohormonal activation and pharmacological inhibition in pulmonary arterial hypertension and related right ventricular failure. *Heart Failure Reviews*, 21, 539–547. https://doi.org/10.1007/ s10741-016-9566-3
- Antigny, F., Hautefort, A., Meloche, J., Belacel-Ouari, M., Manoury, B., Rucker-Martin, C., ... Perros, F. (2016). Potassium channel subfamily K member 3 (KCNK3) contributes to the development of pulmonary arterial hypertension. *Circulation*, 133, 1371–1385. https://doi.org/10. 1161/CIRCULATIONAHA.115.020951
- Antoniu, S. A. (2012). Targeting RhoA/ROCK pathway in pulmonary arterial hypertension. Expert Opinion on Therapeutic Targets, 16, 355–363. https://doi.org/10.1517/14728222.2012.671811
- Archer, S. L., Gomberg-Maitland, M., Maitland, M. L., Rich, S., Garcia, J. G., & Weir, E. K. (2008). Mitochondrial metabolism, redox signaling, and fusion: a mitochondria-ROS-HIF-1α-Kv1.5 O₂-sensing pathway at the intersection of pulmonary hypertension and cancer. *American Journal of Physiology. Heart and Circulatory Physiology*, 294, H570-H578. https://doi.org/10.1152/ajpheart.01324.2007
- Baird, G. L., Archer-Chicko, C., Barr, R. G., Bluemke, D. A., Foderaro, A. E., Fritz, J. S., ... Ventetuolo, C. E. (2018). Lower DHEA-S levels predict disease and worse outcomes in post-menopausal women with idiopathic, connective tissue disease- and congenital heart diseaseassociated pulmonary arterial hypertension. *The European Respiratory Journal*, 51, 1800467. https://doi.org/10.1183/13993003.00467-2018
- Baliga, R. S., Zhao, L., Madhani, M., Lopez-Torondel, B., Visintin, C., Selwood, D., ... Hobbs, A. J. (2008). Synergy between natriuretic peptides and phosphodiesterase 5 inhibitors ameliorates pulmonary arterial hypertension. *American Journal of Respiratory and Critical Care Medicine*, 178, 861–869. https://doi.org/10.1164/rccm.200801-121OC
- Ball, M. K., Waypa, G. B., Mungai, P. T., Nielsen, J. M., Czech, L., Dudley, V. J., ... Schumacker, P. T. (2014). Regulation of hypoxiainduced pulmonary hypertension by vascular smooth muscle hypoxiainducible factor-1α. *American Journal of Respiratory and Critical Care Medicine*, 189, 314–324. https://doi.org/10.1164/rccm.201302-0302OC
- Belkina, A. C., & Denis, G. V. (2012). BET domain co-regulators in obesity, inflammation and cancer. *Nature Reviews Cancer*, 12, 465–477. https://doi.org/10.1038/nrc3256
- Belly, M. J., Tiede, H., Morty, R. E., Schulz, R., Voswinckel, R., Tanislav, C., ... Reichenberger, F. (2012). HbA1c in pulmonary arterial hypertension: a marker of prognostic relevance? *The Journal of Heart and Lung Transplantation*, 31, 1109–1114. https://doi.org/10.1016/j.healun.2012. 08.014
- Blum, L. K., Cao, R. R. L., Sweatt, A. J., Bill, M., Lahey, L. J., Hsi, A. C., ... Robinson, W. H. (2018). Circulating plasmablasts are elevated and produce pathogenic anti-endothelial cell autoantibodies in idiopathic pulmonary arterial hypertension. *European Journal of Immunology*, 48, 874–884. https://doi.org/10.1002/eji.201747460
- Bogaard, H. J., Mizuno, S., Hussaini, A. A., Toldo, S., Abbate, A., Kraskauskas, D., ... Voelkel, N. F. (2011). Suppression of histone deacetylases worsens right ventricular dysfunction after pulmonary artery banding in rats. *American Journal of Respiratory and Critical Care Medicine*, 183, 1402–1410. https://doi.org/10.1164/rccm.201007-1106OC
- Bonnet, S., Michelakis, E. D., Porter, C. J., Andrade-Navarro, M. A., Thebaud, B., Bonnet, S., ... Weir, E. K. (2006). An abnormal mitochondrial-hypoxia inducible factor-1α-Kv channel pathway

5Η Μάροι οιςτραί disrupts oxygen sensing and triggers pulmonary arterial hypertension in fawn hooded rats: similarities to human pulmonary arterial hypertension. *Circulation*, 113, 2630–2641. https://doi.org/10.1161/ CIRCULATIONAHA.105.609008

- Boucherat, O., Chabot, S., Antigny, F., Perros, F., Provencher, S., & Bonnet, S. (2015). Potassium channels in pulmonary arterial hypertension. *The European Respiratory Journal*, 46, 1167–1177. https://doi. org/10.1183/13993003.00798-2015
- Boucherat, O., Chabot, S., Paulin, R., Trinh, I., Bourgeois, A., Potus, F., ... Bonnet, S. (2017). HDAC6: A novel histone deacetylase implicated in pulmonary arterial hypertension. *Scientific Reports*, 7, 4546. https:// doi.org/10.1038/s41598-017-04874-4
- Boucherat, O., Peterlini, T., Bourgeois, A., Nadeau, V., Breuils-Bonnet, S., Boilet-Molez, S., ... Bonnet, S. (2018). Mitochondrial HSP90 accumulation promotes vascular remodeling in pulmonary arterial hypertension. *American Journal of Respiratory and Critical Care Medicine*, 198, 90–103. https://doi.org/10.1164/rccm.201708-1751OC
- Boucherat, O., Provencher, S., & Bonnet, S. (2018). Therapeutic value of ASK1 inhibition in pulmonary arterial hypertension. *American Journal* of Respiratory and Critical Care Medicine, 197, 284–286. https://doi. org/10.1164/rccm.201708-1767ED
- Boucherat, O., Vitry, G., Trinh, I., Paulin, R., Provencher, S., & Bonnet, S. (2017). The cancer theory of pulmonary arterial hypertension. *Pulm Circ*, 7, 285–299. https://doi.org/10.1177/2045893217701438
- Bowers, R., Cool, C., Murphy, R. C., Tuder, R. M., Hopken, M. W., Flores, S. C., & Voelkel, N. F. (2004). Oxidative stress in severe pulmonary hypertension. American Journal of Respiratory and Critical Care Medicine, 169, 764–769. https://doi.org/10.1164/rccm.200301-147OC
- Budas, G. R., Boehm, M., Kojonazarov, B., Viswanathan, G., Tian, X., Veeroju, S., ... Schermuly, R. T. (2018). ASK1 inhibition halts disease progression in preclinical models of pulmonary arterial hypertension. *American Journal of Respiratory and Critical Care Medicine*, 197, 373–385. https://doi.org/10.1164/rccm.201703-0502OC
- Chan, S. Y., & Rubin, L. J. (2017). Metabolic dysfunction in pulmonary hypertension: From basic science to clinical practice. *European Respira*tory Review, 26, 170094. https://doi.org/10.1183/16000617.0094-2017
- Chelladurai, P., Boucherat, O., Stenmark, K., Kracht, M., Seeger, W., Bauer, U. M., ... Pullamsetti, S. S. (2020). Targeting histone acetylation in pulmonary hypertension and right ventricular hypertrophy. *British Journal of Pharmacology*, 1–18. https://doi.org/10.1111/bph.14932
- Chen, S. L., Zhang, F. F., Xu, J., Xie, D. J., Zhou, L., Nguyen, T., & Stone, G. W. (2013). Pulmonary artery denervation to treat pulmonary arterial hypertension: The single-center, prospective, first-in-man PADN-1 study (first-in-man pulmonary artery denervation for treatment of pulmonary artery hypertension). *Journal of the American College of Cardiology*, *62*, 1092–1100. https://doi.org/10.1016/j.jacc. 2013.05.075
- Cracowski, J. L., Cracowski, C., Bessard, G., Pepin, J. L., Bessard, J., Schwebel, C., ... Pison, C. (2001). Increased lipid peroxidation in patients with pulmonary hypertension. *American Journal of Respiratory* and Critical Care Medicine, 164, 1038–1042. https://doi.org/10.1164/ ajrccm.164.6.2104033
- Culley, M. K., & Chan, S. Y. (2018). Mitochondrial metabolism in pulmonary hypertension: beyond mountains there are mountains. *The Journal of Clinical Investigation*, 128, 3704–3715. https://doi.org/10.1172/ JCI120847
- Dai, Z., Li, M., Wharton, J., Zhu, M. M., & Zhao, Y. Y. (2016). Prolyl-4 hydroxylase 2 (PHD2) deficiency in endothelial cells and hematopoietic cells induces obliterative vascular remodeling and severe pulmonary arterial hypertension in mice and humans through hypoxiainducible factor-2α. *Circulation*, 133, 2447–2458. https://doi.org/10. 1161/CIRCULATIONAHA.116.021494

- Dai, Z., Zhu, M. M., Peng, Y., Machireddy, N., Evans, C. E., Machado, R., ... Zhao, Y. Y. (2018). Therapeutic targeting of vascular remodeling and right heart failure in pulmonary arterial hypertension with a HIF-2α inhibitor. *American Journal of Respiratory and Critical Care Medicine*, 198, 1423–1434. https://doi.org/10.1164/rccm.201710-2079OC
- Dannewitz Prosseda, S., Tian, X., Kuramoto, K., Boehm, M., Sudheendra, D., Miyagawa, K., ... Spiekerkoetter, E. (2019). FHIT, a novel modifier gene in pulmonary arterial hypertension. *American Journal of Respiratory and Critical Care Medicine*, 199, 83–98. https://doi. org/10.1164/rccm.201712-2553OC
- de La Roque, E. D., Savineau, J. P., Metivier, A. C., Billes, M. A., Kraemer, J. P., Doutreleau, S., ... Baulieu, É. É. (2012). Dehydroepiandrosterone (DHEA) improves pulmonary hypertension in chronic obstructive pulmonary disease (COPD): A pilot study. Annales d'Endocrinologie, 73, 20–25. https://doi.org/10.1016/j.ando.2011. 12.005
- Delaney, C., Sherlock, L., Fisher, S., Maltzahn, J., Wright, C., & Nozik-Grayck, E. (2018). Serotonin 2A receptor inhibition protects against the development of pulmonary hypertension and pulmonary vascular remodeling in neonatal mice. *American Journal of Physiology. Lung Cellular and Molecular Physiology*, 314, L871–L881. https://doi.org/10. 1152/ajplung.00215.2017
- Dempsey, E. C., Wick, M. J., Karoor, V., Barr, E. J., Tallman, D. W., Wehling, C. A., ... Miller, Y. E. (2009). Neprilysin null mice develop exaggerated pulmonary vascular remodeling in response to chronic hypoxia. *The American Journal of Pathology*, 174, 782–796. https://doi. org/10.2353/ajpath.2009.080345
- Dromparis, P., Paulin, R., Sutendra, G., Qi, A. C., Bonnet, S., & Michelakis, E. D. (2013). Uncoupling protein 2 deficiency mimics the effects of hypoxia and endoplasmic reticulum stress on mitochondria and triggers pseudohypoxic pulmonary vascular remodeling and pulmonary hypertension. *Circulation Research*, 113, 126–136. https://doi. org/10.1161/CIRCRESAHA.112.300699
- Dumas, S. J., Bru-Mercier, G., Courboulin, A., Quatredeniers, M., Rucker-Martin, C., Antigny, F., ... Vocelle, M. (2018). NMDA-type glutamate receptor activation promotes vascular remodeling and pulmonary arterial hypertension. *Circulation*, 137, 2371–2389. https://doi.org/10. 1161/CIRCULATIONAHA.117.029930
- El Kasmi, K. C., Pugliese, S. C., Riddle, S. R., Poth, J. M., Anderson, A. L., Frid, M. G., ... Stenmark, K. R. (2014). Adventitial fibroblasts induce a distinct proinflammatory/profibrotic macrophage phenotype in pulmonary hypertension. *Journal of Immunology*, 193, 597–609. https://doi. org/10.4049/jimmunol.1303048
- Esfandiary, A., Kutsche, H. S., Schreckenberg, R., Weber, M., Pak, O., Kojonazarov, B., ... Schlüter, K. D. (2019). Protection against pressure overload-induced right heart failure by uncoupling protein 2 silencing. *Cardiovascular Research*, 115, 1217–1227. https://doi.org/10.1093/ cvr/cvz049
- Farha, S., Hu, B., Comhair, S., Zein, J., Dweik, R., Erzurum, S. C., & Aldred, M. A. (2016). Mitochondrial haplogroups and risk of pulmonary arterial hypertension. *PLoS ONE*, 11, e0156042. https://doi.org/10. 1371/journal.pone.0156042
- Fukumoto, Y., Yamada, N., Matsubara, H., Mizoguchi, M., Uchino, K., Yao, A., ... Shimokawa, H. (2013). Double-blind, placebo-controlled clinical trial with a rho-kinase inhibitor in pulmonary arterial hypertension. *Circulation Journal*, 77, 2619–2625. https://doi.org/10.1253/ circj.CJ-13-0443
- Fulton, D. J. R., Li, X., Bordan, Z., Haigh, S., Bentley, A., Chen, F., & Barman, S. (2017). Reactive oxygen and nitrogen species in the development of pulmonary hypertension. *Antioxidants (Basel)*, *6*, 54. https:// doi.org/10.3390/antiox6030054
- Galie, N., Channick, R. N., Frantz, R. P., Grunig, E., Jing, Z. C., Moiseeva, O., ... McLaughlin, V. V. (2019). Risk stratification and medical therapy of pulmonary arterial hypertension. *The European Respiratory Journal*, 53, 1801889. https://doi.org/10.1183/13993003.01889-2018

- Galie, N., Humbert, M., Vachiery, J. L., Gibbs, S., Lang, I., Torbicki, A., ... Hoeper, M. (2015). 2015 ESC/ERS guidelines for the diagnosis and treatment of pulmonary hypertension: The joint task force for the diagnosis and treatment of pulmonary hypertension of the European Society of Cardiology (ESC) and the European Respiratory Society (ERS): Endorsed by: Association for European Paediatric and Congenital Cardiology (AEPC), International Society for Heart and Lung Transplantation (ISHLT). *The European Respiratory Journal*, 46, 903–975. https://doi.org/10.1183/13993003.01032-2015
- Galie, N., & Manes, A. (2013). New treatment strategies for pulmonary arterial hypertension: Hopes or hypes? *Journal of the American College* of Cardiology, 62, 1101–1102. https://doi.org/10.1016/j.jacc.2013. 06.032
- Ghataorhe, P., Rhodes, C. J., Harbaum, L., Attard, M., Wharton, J., & Wilkins, M. R. (2017). Pulmonary arterial hypertension—Progress in understanding the disease and prioritizing strategies for drug development. *Journal of Internal Medicine*, 282, 129–141. https://doi.org/10. 1111/joim.12623
- Ghofrani, H. A., Morrell, N. W., Hoeper, M. M., Olschewski, H., Peacock, A. J., Barst, R. J., ... Pascoe, S. (2010). Imatinib in pulmonary arterial hypertension patients with inadequate response to established therapy. American Journal of Respiratory and Critical Care Medicine, 182, 1171–1177. https://doi.org/10.1164/rccm.201001-01230C
- Ghofrani, H. A., Seeger, W., & Grimminger, F. (2005). Imatinib for the treatment of pulmonary arterial hypertension. *The New England Journal of Medicine*, 353, 1412–1413. https://doi.org/10.1056/NEJMc051946
- Ghofrani, H. A., Al-Hiti, H., Vonk-Noordegraaf, A., Behr, J., Neurohr, C., Jansa, P., ... Rosenkranz, S. (2012). Proof-of-concept study to investigate the efficacy, hemodynamics and tolerability of terguride vs. placebo in subjects with pulmonary arterial hypertension: results of a double blind, randomised, prospective phase Ila study. *American Journal of Respiratory and Critical Care Medicine* 2012, 185, A2496. (Link: https://www.atsjournals.org/doi/pdf/10.1164/ajrccmconference.2012.185.1_MeetingAbstracts.A2496)
- Goldenberg, N. M., Hu, Y., Hu, X., Volchuk, A., Zhao, Y. D., Kucherenko, M. M., ... Kuebler, W. M. (2019). Therapeutic targeting of high-mobility group box-1 in pulmonary arterial hypertension. American Journal of Respiratory and Critical Care Medicine, 199, 1566–1569. https://doi.org/10.1164/rccm.201808-1597LE
- Gomberg-Maitland, M., Maitland, M. L., Barst, R. J., Sugeng, L., Coslet, S., Perrino, T. J., ... Ratain, M. J. (2010). A dosing/cross-development study of the multikinase inhibitor sorafenib in patients with pulmonary arterial hypertension. *Clinical Pharmacology and Therapeutics*, 87, 303–310. https://doi.org/10.1038/clpt.2009.217
- Goncharov, D. A., Kudryashova, T. V., Ziai, H., Ihida-Stansbury, K., DeLisser, H., Krymskaya, V. P., ... Goncharova, E. A. (2014). Mammalian target of rapamycin complex 2 (mTORC2) coordinates pulmonary artery smooth muscle cell metabolism, proliferation, and survival in pulmonary arterial hypertension. *Circulation*, 129, 864–874. https:// doi.org/10.1161/CIRCULATIONAHA.113.004581
- Graf, S., Haimel, M., Bleda, M., Hadinnapola, C., Southgate, L., Li, W., ... Morrell, N. W. (2018). Identification of rare sequence variation underlying heritable pulmonary arterial hypertension. *Nature Communications*, 9, 1416. https://doi.org/10.1038/s41467-018-03672-4
- Guignabert, C., Phan, C., Seferian, A., Huertas, A., Tu, L., Thuillet, R., ... Humbert, M. (2016). Dasatinib induces lung vascular toxicity and predisposes to pulmonary hypertension. *The Journal of Clinical Investigation*, 126, 3207–3218. https://doi.org/10.1172/JCI86249
- Hemnes, A. R., Luther, J. M., Rhodes, C. J., Burgess, J. P., Carlson, J., Fan, R., ... Brittain, E. L. (2019). Human PAH Is Characterized by a Pattern of Lipid-Related Insulin Resistance. JCI Insight, 4. https://doi.org/ 10.1172/jci.insight.123611
- Hemnes, A. R., Rathinasabapathy, A., Austin, E. A., Brittain, E. L., Carrier, E. J., Chen, X., ... West, J. (2018). A potential therapeutic role for angiotensin-converting enzyme 2 in human pulmonary arterial

hypertension. The European Respiratory Journal, 51, 1702638. https://doi.org/10.1183/13993003.02638-2017

- Hobbs, A. J., Moyes, A. J., Baliga, R. S., Ghedia, D., Ochiel, R., Sylvestre, Y., ... MacAllister, R. J. (2019). Neprilysin inhibition for pulmonary arterial hypertension: a randomized, double-blind, placebo-controlled, proofof-concept trial. *British Journal of Pharmacology*, 176, 1251–1267. https://doi.org/10.1111/bph.14621
- Hoeper, M. M., Barst, R. J., Bourge, R. C., Feldman, J., Frost, A. E., Galie, N., ... Ghofrani, H.-A. (2013). Imatinib mesylate as add-on therapy for pulmonary arterial hypertension: Results of the randomized IMPRES study. *Circulation*, 127, 1128–1138. https://doi.org/10.1161/ CIRCULATIONAHA.112.000765
- Huang, X., Zou, L., Yu, X., Chen, M., Guo, R., Cai, H., ... Wang, L. (2015). Salidroside attenuates chronic hypoxia-induced pulmonary hypertension via adenosine A2a receptor related mitochondriadependent apoptosis pathway. *Journal of Molecular and Cellular Cardiology*, 82, 153–166. https://doi.org/10.1016/j.yjmcc.2015. 03.005
- Huertas, A., Guignabert, C., Barbera, J. A., Bartsch, P., Bhattacharya, J., Bhattacharya, S., ... Wilkins, M. R. (2018). Pulmonary vascular endothelium: the orchestra conductor in respiratory diseases: Highlights from basic research to therapy. *The European Respiratory Journal*, *51*, 1700745. https://doi.org/10.1183/13993003.00745-2017
- Humbert, M., & Ghofrani, H. A. (2016). The molecular targets of approved treatments for pulmonary arterial hypertension. *Thorax*, 71, 73–83. https://doi.org/10.1136/thoraxjnl-2015-207170
- Humbert, M., Guignabert, C., Bonnet, S., Dorfmuller, P., Klinger, J. R., Nicolls, M. R., ... Rabinovitch, M. (2019). Pathology and pathobiology of pulmonary hypertension: state of the art and research perspectives. *The European Respiratory Journal*, *53*, 1801887. https://doi.org/10. 1183/13993003.01887-2018
- Irodova, N. L., Lankin, V. Z., Konovalova, G. K., Kochetov, A. G., & Chazova, I. E. (2002). Oxidative stress in patients with primary pulmonary hypertension. *Bulletin of Experimental Biology and Medicine*, 133, 580–582. https://doi.org/10.1023/a:1020238026534
- Joshi, S. R., Liu, J., Pearsall, R. S., Li, G., & Kumar, R. (2019). ACTRIIA-Fc (Sotatercept) reverses pulmonary vascular remodeling to attenuate pulmonary arterial hypertension (PAH) by rebalancing TGF-b/BMP signaling in a preclinical model. *American Journal of Respiratory and Critical Care Medicine* 2019, 199, A4395. Link: https://www.atsjournals.org/ doi/abs/10.1164/ajrccm-conference.2019.199.1_MeetingAbstracts. A4395
- Kawut, S. M., Archer-Chicko, C. L., DeMichele, A., Fritz, J. S., Klinger, J. R., Ky, B., ... Propert, K. J. (2017). Anastrozole in pulmonary arterial hypertension. A randomized, double-blind, placebo-controlled trial. *American Journal of Respiratory and Critical Care Medicine*, 195, 360–368. https://doi.org/10.1164/rccm.201605-1024OC
- Kawut, S. M., Bagiella, E., Lederer, D. J., Shimbo, D., Horn, E. M., Roberts, K. E., ... ASA-STAT Study Group (2011). Randomized clinical trial of aspirin and simvastatin for pulmonary arterial hypertension: ASA-STAT. *Circulation*, 123, 2985–2993. https://doi.org/10.1161/ CIRCULATIONAHA.110.015693
- Kawut, S. M., Pinder, D., Al-Naamani, N., McCormick, A., Palevsky, H. I., Fritz, J., ... DeMichele, A. (2019). Fulvestrant for the treatment of pulmonary arterial hypertension. *Annals of the American Thoracic Society*, 16, 1456–1459. https://doi.org/10.1513/AnnalsATS.201904-328RL
- Kazemi-Bajestani, S. M., Patel, V. B., Wang, W., & Oudit, G. Y. (2012). Targeting the ACE2 and apelin pathways are novel therapies for heart failure: Opportunities and challenges. *Cardiology Research and Practice*, 2012, 823193.
- Kiely, D. G., Levin, D., Hassoun, P., Ivy, D. D., Jone, P. N., Bwika, J., ... Moledina, S. (2019). EXPRESS: Statement on imaging and pulmonary hypertension from the Pulmonary Vascular Research Institute (PVRI). *Pulm Circ*, 9, 204589401984199. https://doi.org/10.1177/ 2045894019841990

1 ACUI URICA CULUGICA

- Kim, J. (2014). Apelin-APJ signaling: A potential therapeutic target for pulmonary arterial hypertension. Molecules and Cells, 37, 196-201. https://doi.org/10.14348/molcells.2014.2308
- Kimura, G., Kataoka, M., Inami, T., Fukuda, K., Yoshino, H., & Satoh, T. (2017). Sorafenib as a potential strategy for refractory pulmonary arterial hypertension. Pulmonary Pharmacology & Therapeutics, 44, 46-49. https://doi.org/10.1016/j.pupt.2017.03.009
- Kovacs, G., Berghold, A., Scheidl, S., & Olschewski, H. (2009). Pulmonary arterial pressure during rest and exercise in healthy subjects: A systematic review. The European Respiratory Journal, 34, 888-894. https://doi.org/10.1183/09031936.00145608
- Kuebler, W. M., Bonnet, S., & Tabuchi, A. (2018). Inflammation and autoimmunity in pulmonary hypertension: Is there a role for endothelial adhesion molecules? (2017 Grover Conference Series). Pulm Circ, 8, 204589321875759. https://doi.org/10.1177/2045893218757596
- Lahm, T., Frump, A. L., Albrecht, M. E., Fisher, A. J., Cook, T. G., Jones, T. J., ... Brown, M. B. (2016). 17β-Estradiol mediates superior adaptation of right ventricular function to acute strenuous exercise in female rats with severe pulmonary hypertension. American Journal of Physiology. Lung Cellular and Molecular Physiology, 311, L375-L388. https://doi. org/10.1152/ajplung.00132.2016
- Lahm, T., Tuder, R. M., & Petrache, I. (2014). Progress in solving the sex hormone paradox in pulmonary hypertension. American Journal of Physiology. Lung Cellular and Molecular Physiology, 307, L7-L26. https://doi.org/10.1152/ajplung.00337.2013
- Lambert, M., Capuano, V., Olschewski, A., Sabourin, J., Nagaraj, C., Girerd, B., ... Antigny, F. (2018). Ion channels in pulmonary hypertension: A therapeutic interest? International Journal of Molecular Sciences, 19. https://doi.org/10.3390/ijms19103162
- Lane, K. B., Machado, R. D., Pauciulo, M. W., Thomson, J. R., Phillips, J. A. 3rd, Loyd, J. E., ... Trembath, R. C. (2000). Heterozygous germline mutations in BMPR2, encoding a TGF- β receptor, cause familial primary pulmonary hypertension. Nature Genetics, 26, 81-84. https://doi. org/10.1038/79226
- Lee, M. Y., Tsai, K. B., Hsu, J. H., Shin, S. J., Wu, J. R., & Yeh, J. L. (2016). Liraglutide prevents and reverses monocrotaline-induced pulmonary arterial hypertension by suppressing ET-1 and enhancing eNOS/sGC/-PKG pathways. Scientific Reports, 6, 31788. https://doi.org/10.1038/ srep31788
- Lemmon, M. A., & Schlessinger, J. (2010). Cell signaling by receptor tyrosine kinases. Cell, 141, 1117-1134. https://doi.org/10.1016/j.cell. 2010.06.011
- Long, L., Ormiston, M. L., Yang, X., Southwood, M., Graf, S., Machado, R. D., ... Morrell, N. W. (2015). Selective enhancement of endothelial BMPR-II with BMP9 reverses pulmonary arterial hypertension. Nature Medicine, 21, 777-785. https://doi.org/10.1038/nm. 3877
- Long, L., Yang, X., Southwood, M., Lu, J., Marciniak, S. J., Dunmore, B. J., & Morrell, N. W. (2013). Chloroquine prevents progression of experimental pulmonary hypertension via inhibition of autophagy and lysosomal bone morphogenetic protein type II receptor degradation. Circulation Research, 112, 1159-1170. https://doi.org/10.1161/ CIRCRESAHA.111.300483
- Lythgoe, M. P., Rhodes, C. J., Ghataorhe, P., Attard, M., Wharton, J., & Wilkins, M. R. (2016). Why drugs fail in clinical trials in pulmonary arterial hypertension, and strategies to succeed in the future. Pharmacology & Therapeutics, 164, 195-203. https://doi.org/10.1016/j. pharmthera.2016.04.012
- Ma, L., Roman-Campos, D., Austin, E. D., Eyries, M., Sampson, K. S., Soubrier, F., ... Chung, W. K. (2013). A novel channelopathy in pulmonary arterial hypertension. The New England Journal of Medicine, 369, 351-361. https://doi.org/10.1056/NEJMoa1211097
- Maron, B. A., & Leopold, J. A. (2015). Emerging concepts in the molecular basis of pulmonary arterial hypertension: Part II: Neurohormonal signaling contributes to the pulmonary vascular and right ventricular

pathophenotype of pulmonary arterial hypertension. Circulation, 131, 2079-2091. https://doi.org/10.1161/CIRCULATIONAHA.114. 006980

- McMurray, J. J., Packer, M., Desai, A. S., Gong, J., Lefkowitz, M. P., Rizkala, A. R., ... PARADIGM-HF Investigators and Committees (2014). Angiotensin-neprilysin inhibition versus enalapril in heart failure. The New England Journal of Medicine, 371, 993-1004. https://doi.org/10. 1056/NEJMoa1409077
- McMurtry, M. S., Bonnet, S., Wu, X., Dyck, J. R., Haromy, A., Hashimoto, K., & Michelakis, E. D. (2004). Dichloroacetate prevents and reverses pulmonary hypertension by inducing pulmonary artery smooth muscle cell apoptosis. Circulation Research, 95, 830-840. https://doi.org/10.1161/01.RES.0000145360.16770.9f
- Meloche, J., Lampron, M. C., Nadeau, V., Maltais, M., Potus, F., Lambert, C., ... Bonnet, S. (2017). Implication of inflammation and epigenetic readers in coronary artery remodeling in patients with pulmonary arterial hypertension. Arteriosclerosis, Thrombosis, and Vascular Biology, 37, 1513-1523. https://doi.org/10.1161/ATVBAHA.117. 309156
- Meloche, J., Paulin, R., Provencher, S., & Bonnet, S. (2015). Therapeutic potential of microRNA modulation in pulmonary arterial hypertension. Current Vascular Pharmacology, 13, 331-340. https://doi.org/10. 2174/15701611113119990010
- Meloche, J., Pflieger, A., Vaillancourt, M., Paulin, R., Potus, F., Zervopoulos, S., ... Bonnet, S. (2014). Role for DNA damage signaling in pulmonary arterial hypertension. Circulation, 129, 786-797. https:// doi.org/10.1161/CIRCULATIONAHA.113.006167
- Meloche, J., Potus, F., Vaillancourt, M., Bourgeois, A., Johnson, I., Deschamps, L., ... Bonnet, S. (2015). Bromodomain-containing protein 4: The epigenetic origin of pulmonary arterial hypertension. Circulation Research, 117, 525-535. https://doi.org/10.1161/CIRCRESAHA.115. 307004
- Mestas, J., & Hughes, C. C. (2004). Of mice and not men: differences between mouse and human immunology. Journal of Immunology, 172, 2731-2738. https://doi.org/10.4049/jimmunol.172.5.2731
- Michelakis, E. D., Gurtu, V., Webster, L., Barnes, G., Watson, G., Howard, L., ... Wilkins, M. R. (2017). Inhibition of pyruvate dehydrogenase kinase improves pulmonary arterial hypertension in genetically susceptible patients. Science Translational Medicine, 9, eaao4583. https://doi.org/10.1126/scitranslmed.aao4583
- Montani, D., Bergot, E., Gunther, S., Savale, L., Bergeron, A., Bourdin, A., ... Humbert, M. (2012). Pulmonary arterial hypertension in patients treated by dasatinib. Circulation, 125, 2128-2137. https://doi.org/10. 1161/CIRCULATIONAHA.111.079921
- Nickel, N. P., Spiekerkoetter, E., Gu, M., Li, C. G., Li, H., Kaschwich, M., ... Rabinovitch, M. (2015). Elafin reverses pulmonary hypertension via caveolin-1-dependent bone morphogenetic protein signaling. American Journal of Respiratory and Critical Care Medicine, 191, 1273-1286. https://doi.org/10.1164/rccm.201412-2291OC
- Novoyatleva, T., Kojonazarov, B., Owczarek, A., Veeroju, S., Rai, N., Henneke, I., ... Schermuly, R. T. (2019). Evidence for the fucoidan/Pselectin axis as a therapeutic target in hypoxia-induced pulmonary hypertension. American Journal of Respiratory and Critical Care Medi-1407-1420. https://doi.org/10.1164/rccm.201806cine. 199. 1170OC
- Ormiston, M. L., Godoy, R. S., Chaudhary, K. R., & Stewart, D. J. (2019). The Janus faces of bone morphogenetic protein 9 in pulmonary arterial hypertension. Circulation Research, 124, 822-824. https://doi.org/ 10.1161/CIRCRESAHA.119.314753
- Oudiz, R. J., Meyer, C., Chin, M., Feldman, J., Goldsberry, A., McConnell, J. W., ... White, J. (2017). Results of interim analysis of the efficacy and safety of bardoxolone methyl in patients with pulmonary arterial hypertension associated with connective tissue disease (CTD) (the LARIAT study). American Journal of Respiratory and Critical Care Medicine 2017, 195, A6896. Link: https://www.atsjournals.org/doi/

abs/10.1164/ajrccm-conference.2017.195.1_MeetingAbstracts. A6896

- Paddenberg, R., Stieger, P., von Lilien, A. L., Faulhammer, P., Goldenberg, A., Tillmanns, H. H., ... Braun-Dullaeus, R. C. (2007). Rapamycin attenuates hypoxia-induced pulmonary vascular remodeling and right ventricular hypertrophy in mice. *Respiratory Research*, *8*, 15. https://doi.org/10.1186/1465-9921-8-15
- Pak, O., Scheibe, S., Esfandiary, A., Gierhardt, M., Sydykov, A., Logan, A., ... Sommer, N. (2018). Impact of the mitochondria-targeted antioxidant MitoQ on hypoxia-induced pulmonary hypertension. *The European Respiratory Journal*, *51*, 1701024. https://doi.org/10.1183/13993003. 01024-2017
- Pak, O., Sommer, N., Hoeres, T., Bakr, A., Waisbrod, S., Sydykov, A., ... Weissmann, N. (2013). Mitochondrial hyperpolarization in pulmonary vascular remodeling. Mitochondrial uncoupling protein deficiency as disease model. American Journal of Respiratory Cell and Molecular Biology, 49, 358–367. https://doi.org/10.1165/rcmb.2012-0361OC
- Paralkar, V., Pearson, P., Mason, J. W., Li, S.-X., Curry, S., Aiello, R., & Feig, P. U. (2017). KAR5585, a first-in-class oral tryptophan hydroxylase 1 (TPH1) inhibitor as a novel candidate for the treatment of pulmonary arterial hypertension. *American Journal of Respiratory and Critical Care Medicine 2017*, 195, A4193. Link: https://www. atsjournals.org/doi/abs/10.1164/ajrccm-conference.2017.195.1_ MeetingAbstracts.A4193
- Paulin, R., Dromparis, P., Sutendra, G., Gurtu, V., Zervopoulos, S., Bowers, L., ... Michelakis, E. D. (2014). Sirtuin 3 deficiency is associated with inhibited mitochondrial function and pulmonary arterial hypertension in rodents and humans. *Cell Metabolism*, 20, 827–839. https://doi.org/10.1016/j.cmet.2014.08.011
- Perros, F., de Man, F. S., Bogaard, H. J., Antigny, F., Simonneau, G., Bonnet, S., ... Humbert, M. (2017). Use of β-blockers in pulmonary hypertension. *Circulation. Heart Failure*, 10, e003703. https://doi.org/ 10.1161/CIRCHEARTFAILURE.116.003703
- Preston, I. R., Sagliani, K. D., Warburton, R. R., Hill, N. S., Fanburg, B. L., & Jaffe, I. Z. (2013). Mineralocorticoid receptor antagonism attenuates experimental pulmonary hypertension. *American Journal of Physiology*. *Lung Cellular and Molecular Physiology*, 304, L678–L688. https://doi. org/10.1152/ajplung.00300.2012
- Prins, K. W., Thenappan, T., Weir, E. K., Kalra, R., Pritzker, M., & Archer, S. L. (2019). Repurposing medications for treatment of pulmonary arterial hypertension: What's old is new again. *Journal of the American Heart Association*, 8, e011343.
- Provencher, S., Archer, S. L., Ramirez, F. D., Hibbert, B., Paulin, R., Boucherat, O., ... Bonnet, S. (2018). Standards and methodological rigor in pulmonary arterial hypertension preclinical and translational research. *Circulation Research*, 122, 1021–1032. https://doi.org/10. 1161/CIRCRESAHA.117.312579
- Pullamsetti, S. S., Perros, F., Chelladurai, P., Yuan, J., & Stenmark, K. (2016). Transcription factors, transcriptional coregulators, and epigenetic modulation in the control of pulmonary vascular cell phenotype: Therapeutic implications for pulmonary hypertension (2015 Grover Conference series). *Pulm Circ*, *6*, 448–464. https://doi.org/10.1086/ 688908
- Rabinovitch, M., Guignabert, C., Humbert, M., & Nicolls, M. R. (2014). Inflammation and immunity in the pathogenesis of pulmonary arterial hypertension. *Circulation Research*, 115, 165–175. https://doi.org/10. 1161/CIRCRESAHA.113.301141
- Rashid, J., Nozik-Grayck, E., McMurtry, I. F., Stenmark, K. R., & Ahsan, F. (2019). Inhaled combination of sildenafil and rosiglitazone improves pulmonary hemodynamics, cardiac function, and arterial remodeling. *American Journal of Physiology. Lung Cellular and Molecular Physiology*, 316, L119–L130. https://doi.org/10.1152/ajplung.00381.2018
- Rhodes, C. J., Ghataorhe, P., Wharton, J., Rue-Albrecht, K. C., Hadinnapola, C., Watson, G., ... Wilkins, M. R. (2017). Plasma metabolomics implicates modified transfer RNAs and altered bioenergetics

in the outcomes of pulmonary arterial hypertension. *Circulation*, 135, 460–475. https://doi.org/10.1161/CIRCULATIONAHA.116.024602

- Rhodes, C. J., Howard, L. S., Busbridge, M., Ashby, D., Kondili, E., Gibbs, J. S., ... Wilkins, M. R. (2011). Iron deficiency and raised hepcidin in idiopathic pulmonary arterial hypertension: Clinical prevalence, outcomes, and mechanistic insights. *Journal of the American College of Cardiology*, 58, 300–309. https://doi.org/10.1016/j.jacc.2011. 02.057
- Rich, S., Kaufmann, E., & Levy, P. S. (1992). The effect of high doses of calcium-channel blockers on survival in primary pulmonary hypertension. *The New England Journal of Medicine*, 327, 76–81. https://doi. org/10.1056/NEJM199207093270203
- Richter, M. J., Ewert, J., Grimminger, F., Ghofrani, H. A., Kojonazarov, B., Petrovic, A., ... Gall, H. (2018). Nintedanib in severe pulmonary arterial hypertension. *American Journal of Respiratory and Critical Care Medicine*, 198, 808–810. https://doi.org/10.1164/rccm.201801-0195LE
- Rol, N., Kurakula, K. B., Happe, C., Bogaard, H. J., & Goumans, M. J. (2018). TGF-β and BMPR2 Signaling in PAH: Two black sheep in one family. *International Journal of Molecular Sciences*, 19, 2585. https://doi.org/ 10.3390/ijms19092585
- Rothman, A. M., Arnold, N. D., Pickworth, J. A., Iremonger, J., Ciuclan, L., Allen, R. M., ... Francis, S. E. (2016). MicroRNA-140-5p and SMURF1 regulate pulmonary arterial hypertension. *The Journal of Clinical Investi*gation, 126, 2495–2508. https://doi.org/10.1172/JCI83361
- Rowlands, D. J. (2016). Mitochondria dysfunction: A novel therapeutic target in pathological lung remodeling or bystander? *Pharmacology & Therapeutics*, 166, 96–105. https://doi.org/10.1016/j.pharmthera. 2016.06.019
- Ruiter, G., Manders, E., Happe, C. M., Schalij, I., Groepenhoff, H., Howard, L. S., ... de Man, F. S. (2015). Intravenous iron therapy in patients with idiopathic pulmonary arterial hypertension and iron deficiency. *Pulm Circ*, 5, 466–472. https://doi.org/10.1086/682217
- Sadoughi, A., Roberts, K. E., Preston, I. R., Lai, G. P., McCollister, D. H., Farber, H. W., & Hill, N. S. (2013). Use of selective serotonin reuptake inhibitors and outcomes in pulmonary arterial hypertension. *Chest*, 144, 531–541. https://doi.org/10.1378/chest.12-2081
- Said, S. I. (2012). Vasoactive intestinal peptide in pulmonary arterial hypertension. American Journal of Respiratory and Critical Care Medicine, 185, 786author reply 786. https://doi.org/10.1164/ajrccm.185.7.786
- Sanchez, O., Sitbon, O., Jais, X., Simonneau, G., & Humbert, M. (2006). Immunosuppressive therapy in connective tissue diseases-associated pulmonary arterial hypertension. *Chest*, 130, 182–189. https://doi. org/10.1378/chest.130.1.182
- Savai, R., Al-Tamari, H. M., Sedding, D., Kojonazarov, B., Muecke, C., Teske, R., ... Schermuly, R. T. (2014). Pro-proliferative and inflammatory signaling converge on FoxO1 transcription factor in pulmonary hypertension. *Nature Medicine*, 20, 1289–1300. https://doi.org/10. 1038/nm.3695
- Sharp, J., Farha, S., Park, M. M., Comhair, S. A., Lundgrin, E. L., Tang, W. H., ... Erzurum, S. C. (2014). Coenzyme Q supplementation in pulmonary arterial hypertension. *Redox Biology*, 2, 884–891. https://doi.org/10. 1016/j.redox.2014.06.010
- Simonneau, G., Montani, D., Celermajer, D. S., Denton, C. P., Gatzoulis, M. A., Krowka, M., ... Souza, R. (2019). Haemodynamic definitions and updated clinical classification of pulmonary hypertension. *The European Respiratory Journal*, *53*, 1801913. https://doi.org/10. 1183/13993003.01913-2018
- Sitbon, O., Gomberg-Maitland, M., Granton, J., Lewis, M. I., Mathai, S. C., Rainisio, M., ... Rubin, L. J. (2019). Clinical trial design and new therapies for pulmonary arterial hypertension. *The European Respiratory Journal*, 53, 1801908. https://doi.org/10.1183/13993003.01908-2018
- Sitbon, O., Humbert, M., Jais, X., Ioos, V., Hamid, A. M., Provencher, S., ... Simonneau, G. (2005). Long-term response to calcium channel blockers in idiopathic pulmonary arterial hypertension. *Circulation*,

24

111, 3105-3111. https://doi.org/10.1161/CIRCULATIONAHA.104. 488486

- Sofianopoulou, E., Kaptoge, S., Graf, S., Hadinnapola, C., Treacy, C. M., Church, C., ... Morrell, N. W. (2019). Traffic exposures, air pollution and outcomes in pulmonary arterial hypertension: A UK cohort study analysis. *The European Respiratory Journal*, *53*, 1801429. https://doi. org/10.1183/13993003.01429-2018
- Sommer, N., Droege, F., Gamen, K. E., Geisthoff, U., Gall, H., Tello, K., ... Pullamsetti, S. (2019). Treatment with low-dose tacrolimus inhibits bleeding complications in a patient with hereditary hemorrhagic telangiectasia and pulmonary arterial hypertension. *Pulm Circ, 9*, 204589401880540. https://doi.org/10.1177/2045894018805406
- Southan, C., Sharman, J. L., Benson, H. E., Faccenda, E., Pawson, A. J., Alexander, S. P., ... NC-IUPHAR (2016). The IUPHAR/BPS Guide to PHARMACOLOGY in 2016: Towards curated quantitative interactions between 1300 protein targets and 6000 ligands. *Nucleic Acids Research*, 44, D1054–D1068. https://doi.org/10.1093/nar/gkv1037
- Spiekerkoetter, E., Sung, Y. K., Sudheendra, D., Bill, M., Aldred, M. A., van de Veerdonk, M. C., ... Zamanian, R. T. (2015). Low-dose FK506 (Tacrolimus) in end-stage pulmonary arterial hypertension. American Journal of Respiratory and Critical Care Medicine, 192, 254–257. https://doi.org/10.1164/rccm.201411-2061LE
- Spiekerkoetter, E., Sung, Y. K., Sudheendra, D., Scott, V., Del Rosario, P., Bill, M., ... Zamanian, R. T. (2017). Randomised placebo-controlled safety and tolerability trial of FK506 (tacrolimus) for pulmonary arterial hypertension. *The European Respiratory Journal*, 50, 1602449. https:// doi.org/10.1183/13993003.02449-2016
- Spiekerkoetter, E., Tian, X., Cai, J., Hopper, R. K., Sudheendra, D., Li, C. G., ... Rabinovitch, M. (2013). FK506 activates BMPR2, rescues endothelial dysfunction, and reverses pulmonary hypertension. *The Journal of Clinical Investigation*, 123, 3600–3613. https://doi.org/10.1172/ JCI65592
- Stenmark, K. R., Meyrick, B., Galie, N., Mooi, W. J., & McMurtry, I. F. (2009). Animal models of pulmonary arterial hypertension: The hope for etiological discovery and pharmacological cure. *American Journal of Physiology. Lung Cellular and Molecular Physiology*, 297, L1013–L1032. https://doi.org/10.1152/ajplung.00217.2009
- Tamosiuniene, R., Tian, W., Dhillon, G., Wang, L., Sung, Y. K., Gera, L., ... Nicolls, M. R. (2011). Regulatory T cells limit vascular endothelial injury and prevent pulmonary hypertension. *Circulation Research*, 109, 867–879. https://doi.org/10.1161/CIRCRESAHA.110.236927
- Tan, W. S. D., Liao, W., Zhou, S., Mei, D., & Wong, W. F. (2018). Targeting the renin-angiotensin system as novel therapeutic strategy for pulmonary diseases. *Current Opinion in Pharmacology*, 40, 9–17. https://doi. org/10.1016/j.coph.2017.12.002
- Thompson, A. A. R., & Lawrie, A. (2017). Targeting vascular remodeling to treat pulmonary arterial hypertension. *Trends in Molecular Medicine*, 23, 31–45. https://doi.org/10.1016/j.molmed.2016.11.005
- Thompson, J. S., Sheedy, W., & Morice, A. H. (1994). Neutral endopeptidase (NEP) inhibition in rats with established pulmonary hypertension secondary to chronic hypoxia. *British Journal of Pharmacology*, 113, 1121–1126. https://doi.org/10.1111/j.1476-5381.1994.tb17112.x
- Tofovic, S. P., & Jackson, E. K. (2019). Estradiol metabolism: Crossroads in pulmonary arterial hypertension. *International Journal of Molecular Sci*ences, 21, 116. https://doi.org/10.3390/ijms21010116
- Trankle, C. R., Canada, J. M., Kadariya, D., Markley, R., De Chazal, H. M., Pinson, J., ... Grinnan, D. (2019). IL-1 blockade reduces inflammation in pulmonary arterial hypertension and right ventricular failure: A singlearm, open-label, phase IB/II pilot study. *American Journal of Respiratory* and Critical Care Medicine, 199, 381–384. https://doi.org/10.1164/ rccm.201809-1631LE
- Tu, L., Desroches-Castan, A., Mallet, C., Guyon, L., Cumont, A., Phan, C., ... Guignabert, C. (2019). Selective BMP-9 inhibition partially protects against experimental pulmonary hypertension. *Circulation Research*, 124, 846–855. https://doi.org/10.1161/CIRCRESAHA.118.313356

- Ulrich, S., Saxer, S., Hasler, E. D., Schwarz, E. I., Schneider, S. R., Furian, M., ... Bloch, K. E. (2019). Effect of domiciliary oxygen therapy on exercise capacity and quality of life in patients with pulmonary arterial or chronic thromboembolic pulmonary hypertension: A randomised, placebo-controlled trial. *The European Respiratory Journal*, 54, 1900276. https://doi.org/10.1183/13993003.002762019
- Umar, S., Cunningham, C. M., Itoh, Y., Moazeni, S., Vaillancourt, M., Sarji, S., ... Eghbali, M. (2018). The Y chromosome plays a protective role in experimental hypoxic pulmonary hypertension. *American Journal of Respiratory and Critical Care Medicine*, 197, 952–955. https:// doi.org/10.1164/rccm.201707-1345LE
- Vaidya, B., Pangallo, M., Ruffenach, G., Cunningham, C. M., Perron, J. C., Kolluru, S., ... Gupta, V. (2017). Advances in treatment of pulmonary arterial hypertension: Patent review. *Expert Opinion on Therapeutic Patents*, 27, 907–918. https://doi.org/10.1080/13543776.2017. 1313232
- Valerio, C. J., Schreiber, B. E., Handler, C. E., Denton, C. P., & Coghlan, J. G. (2013). Borderline mean pulmonary artery pressure in patients with systemic sclerosis: Transpulmonary gradient predicts risk of developing pulmonary hypertension. Arthritis and Rheumatism, 65, 1074–1084. https://doi.org/10.1002/art.37838
- Van der Feen, D. E., Kurakula, K., Tremblay, E., Boucherat, O., Bossers, G. P. L., Szulcek, R., ... Goumans, M.-J. (2019). Multicenter preclinical validation of BET inhibition for the treatment of pulmonary arterial hypertension. *American Journal of Respiratory and Critical Care Medicine*, 200, 910–920. https://doi.org/10.1164/rccm.201812-2275OC
- Ventetuolo, C. E., Baird, G. L., Barr, R. G., Bluemke, D. A., Fritz, J. S., Hill, N. S., ... Kawut, S. M. (2016). Higher estradiol and lower dehydroepiandrosterone-sulfate levels are associated with pulmonary arterial hypertension in men. *American Journal of Respiratory and Critical Care Medicine*, 193, 1168–1175. https://doi.org/10.1164/rccm. 201509-1785OC
- Viethen, T., Gerhardt, F., Dumitrescu, D., Knoop-Busch, S., ten Freyhaus, H., Rudolph, T. K., ... Rosenkranz, S. (2014). Ferric carboxymaltose improves exercise capacity and quality of life in patients with pulmonary arterial hypertension and iron deficiency: A pilot study. *International Journal of Cardiology*, 175, 233–239. https://doi. org/10.1016/j.ijcard.2014.04.233
- Waypa, G. B., Osborne, S. W., Marks, J. D., Berkelhamer, S. K., Kondapalli, J., & Schumacker, P. T. (2013). Sirtuin 3 deficiency does not augment hypoxia-induced pulmonary hypertension. *American Journal of Respiratory Cell and Molecular Biology*, 49, 885–891. https://doi. org/10.1165/rcmb.2013-01910C
- Weatherald, J., Humbert, M., Guignabert, C., & Montani, D. (2017). Response to the article "orafenib as a potential strategy for refractory pulmonary arterial hypertension". *Pulmonary Pharmacology & Therapeutics*, 45, 11–12. https://doi.org/10.1016/j.pupt.2017.03.017
- Wessler, J. D., Steingart, R. M., Schwartz, G. K., Harvey, B. G., & Schaffer, W. (2010). Dramatic improvement in pulmonary hypertension with rapamycin. *Chest*, 138, 991–993. https://doi.org/10.1378/ chest.09-2435
- West, J. D., Carrier, E. J., Bloodworth, N. C., Schroer, A. K., Chen, P., Ryzhova, L. M., ... Merryman, W. D. (2016). Serotonin 2B receptor antagonism prevents heritable pulmonary arterial hypertension. *PLoS ONE*, 11, e0148657. https://doi.org/10.1371/journal.pone. 0148657
- Xu, J., Wang, J., He, M., Han, H., Xie, W., Wang, H., & Kong, H. (2018). Dipeptidyl peptidase IV (DPP-4) inhibition alleviates pulmonary arterial remodeling in experimental pulmonary hypertension. *Laboratory Investigation*, *98*, 1333–1346. https://doi.org/10.1038/s41374-018-0080-1
- Xu, W., Koeck, T., Lara, A. R., Neumann, D., DiFilippo, F. P., Koo, M., ... Erzurum, S. C. (2007). Alterations of cellular bioenergetics in pulmonary artery endothelial cells. *Proceedings of the National Academy of*

Sciences of the United States of America, 104, 1342–1347. https://doi. org/10.1073/pnas.0605080104

- Yuan, K., Orcholski, M. E., Panaroni, C., Shuffle, E. M., Huang, N. F., Jiang, X., ... de Jesus Perez, V. A. (2015). Activation of the Wnt/planar cell polarity pathway is required for pericyte recruitment during pulmonary angiogenesis. *The American Journal of Pathology*, 185, 69–84. https://doi.org/10.1016/j.ajpath.2014.09.013
- Yuan, K., Shamskhou, E. A., Orcholski, M. E., Nathan, A., Reddy, S., Honda, H., ... de Jesus Perez, V. A. (2019). Loss of endotheliumderived Wnt5a is associated with reduced pericyte recruitment and small vessel loss in pulmonary arterial hypertension. *Circulation*, 139, 1710–1724. https://doi.org/10.1161/CIRCULATIONAHA.118. 037642
- Zaidi, S. H., You, X. M., Ciura, S., Husain, M., & Rabinovitch, M. (2002). Overexpression of the serine elastase inhibitor elafin protects transgenic mice from hypoxic pulmonary hypertension. *Circulation*, 105, 516–521. https://doi.org/10.1161/hc0402.102866

Zeng, W. J., Xiong, C. M., Zhao, L., Shan, G. L., Liu, Z. H., Xue, F., ... Atorvastatin in pulmonary arterial hypertension (APATH) Study Group (2012). Atorvastatin in pulmonary arterial hypertension (APATH) study. *The European Respiratory Journal*, 40, 67–74. https://doi.org/10. 1183/09031936.00149011

How to cite this article: Sommer N, Ghofrani HA, Pak O, et al. Current and future treatments of pulmonary arterial hypertension. *Br J Pharmacol.* 2020;1–25. <u>https://doi.org/10.</u> 1111/bph.15016