Smouldering fire or conflagration? An illustrated update on the concept of inflammation in pulmonary arterial hypertension

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Abstract

Pulmonary arterial hypertension (PAH) is a rare condition that is characterised by a progressive increase of pulmonary vascular resistances that leads to right ventricular failure and death, if untreated. The underlying narrowing of the pulmonary vasculature relies on several independent and interdependent biological pathways, such as genetic predisposition and epigenetic changes, imbalance of vasodilating and vasoconstrictive mediators, as well as dysimmunity and inflammation that will trigger endothelial dysfunction, smooth muscle cell proliferation, fibroblast activation and collagen deposition. Progressive constriction of the pulmonary vasculature, in turn, initiates and sustains hypertrophic and maladaptive myocardial remodelling of the right ventricle.

In this illustrated review, we will focus on the role of inflammation and dysimmunity in PAH which is generally accepted today, although existing PAH-specific medical therapies still lack targeted immune-modulating approaches.

Introduction

Pulmonary arterial hypertension (PAH) is a rare condition with an inevitable critical evolution and fatal outcome, if untreated. The disease is characterised by a progressive increase of pulmonary vascular resistance that eventually leads to right ventricular failure [1]. Diagnosis mostly relies on right heart catheterisation and exclusion of other aetiologies, such as thromboembolism, through imaging or ventilation/perfusion scan, since the remodelling process of lung vessels that causes an increase of pressure within the pulmonary circuit involves small peripheral vessels and microvessels that cannot be directly visualised by non-invasive techniques [1].

The pathophysiology of PAH is complex and has many pillars, such as genetic predisposition and epigenetic changes, an imbalance of vasodilating and vasoconstrictive mediators, as well as dysimmunity and inflammation that eventually lead to endothelial dysfunction, smooth muscle cell proliferation and fibroblast activation/collagen deposition within the thickening vessel walls. This in turn leads to hypertrophic and maladaptive myocardial remodelling of the right ventricle (figure 1) [2].

In this illustrated review, we will focus on the role of inflammation and dysimmunity, a generally accepted branch of the condition’s pathophysiology, although existing PAH-specific medical therapies still lack targeted anti-inflammatory or immune-modulating approaches [3].

While the presence of perivascular inflammatory cells and auto-antibody generating lymphoid tertiary follicles in the vicinity of pulmonary arteries has been described in the past and the increase in...
pro-inflammatory cytokines and chemokines and auto-antibodies in the peripheral blood repeatedly reported, interpretations of the role, mechanistic rank and pathophysiologic relevance of these inflammatory and dysimmune hallmarks are still controversial [4]. This controversy is further fuelled by the fact that the most
prominent PAH-associated mutations are critical for immune-regulatory processes, in addition to their relevance in vascular homeostasis and function [5]. It should be kept in mind, however, that PAH is frequently associated with primary autoimmune and immune disorders, such as connective tissue disease (CTD)-associated or HIV-associated PAH, where pulmonary hypertension (PH) develops in a clearly pro-inflammatory context (figure 2). For these conditions at least, increases in pulmonary arterial pressures can be positively influenced by anti-inflammatory and/or immune-modulatory therapy, a strategy that has not yet made its way to PAH-specific therapies in general, despite recent pharmacological trials [6].

One fundamental question remains still unanswered: why do obstructive lesions of small pulmonary arteries/arterioles/venules and perivascular chronic inflammatory infiltrate concur? Where does the story begin? Occlusion and reactive inflammation, or inflammation and reactive occlusion? The latter hypothesis has been a much-used rationale for many scientific reports in the past 30 years. The former option appears less inspiring, but should not be discarded too quickly, since the pathophysiologic importance of disease evolution can come from self-sustaining, smouldering and long-standing secondary fires, not only from initial conflagrations (figure 3).

FIGURE 3 The two types of mechanism that underlie pulmonary arterial hypertension pathogenesis.

FIGURE 4 Structural alterations as a cause of inflammation. Disturbed blood flow and stiffness-induced inflammation. ECM: extra-cellular matrix.
The smouldering fire hypothesis

We first consider the smouldering fire hypothesis. Increased vessel wall stiffness through the deposition of extra-cellular matrix, from whatever aetiology, leads to a marked pulsatile flow (with higher pressure amplitudes) that will provoke a pro-inflammatory response within the directly exposed vascular compartment, the lumen-facing endothelial cell layer of small muscular-type arteries and arterioles that lack greater amounts of elastic (amplitude-soothing) fibres. In addition, increased shear stress (due again to higher pressure amplitudes) will activate NF-κB, a transcription factor that is essential for most immune/inflammatory and cell stress responses, and contribute to the pro-inflammatory state of endothelial cells during PAH (figure 4) [7, 8].

In addition, the mere hypoxic state that is associated with so many respiratory and cardiovascular diseases leads to inflammatory responses by itself: at altitude, for example, hypoxia increases the levels of circulating interleukin (IL)-6 and C-reactive protein, even in healthy humans [9]. Hypoxia induces inflammation within the pulmonary artery through local expression of pro-contractile and proliferative growth factors and pro-inflammatory mediators like vascular endothelial growth factor, brain-derived neurotrophic factor or thymic stromal lymphopoietin that are generated in resident cells of the vessel wall [10]. Moreover, hypoxia induces the auto-oxidation of haemoglobin within red blood cells, and thereby increases superoxide production: higher amounts of H$_2$O$_2$ are spilled into the pulmonary circulation, and oxidative stress at the level of microvascular endothelial cells triggers the activation of the NF-κB pathway and local recruitment of leukocytes (figure 5) [11].

However, there are other, less obvious players that might be involved in maintaining the low-level smouldering fire of inflammation within and around the pulmonary micro-vasculature. The intestinal microbiome is such a potential factor: enteric congestion due to right ventricular decompensation may lead to a dysfunctional intestinal barrier with eventual permeation of bacterial pro-inflammatory components like lipopolysaccharide into the systemic circulation [12]. These so-called pathogen-associated molecular patterns (PAMPs) induce inflammation and lead to cellular damage and stress in resident host cells; for example, in pulmonary endothelial cells that will, in turn, generate so-called damage-associated molecular patterns (DAMPs) that are part of the sterile inflammatory response. Both PAMPs and DAMPs bind to pattern-recognition receptors like Toll-like receptors or cytoplasmic NOD-like receptors and invoke strong

**FIGURE 5** Hypoxia-induced inflammation. BDNF: brain-derived neurotrophic factor; PAEC: pulmonary arterial endothelial cell; PASMC: pulmonary arterial smooth muscle cell; TSLP: thymic stromal lymphopoietin; VEGF: vascular endothelial growth factor.
local reactions with leukocytic influx, more inflammation and eventually reactive vascular remodelling [13] (figure 6). Besides pro-inflammatory properties, the commensal human microbiome could also directly influence the pulmonary vasculature [14]. Significant taxonomic and functional changes in microbial communities in the PAH cohort have been observed and some of the previously identified potential biomarkers of PAH could be derived from altered bacterial functions in PAH patients [15].

The prominent role of transcription factors with regard to inflammation and vascular remodelling has been reported and discussed in the past years. Numerous studies suggest that ETS-related gene (ERG) and Friend leukaemia integration 1 transcription factor are involved in endothelial function, proliferation, homeostasis, cellular interaction, angiogenesis, fibrosis and vascular inflammation through the gene clusters they control [16]. In experimental models with ERG-knockout, most mice die in utero, but a small fraction of survivors develops pulmonary veno-occlusive disease (PVOD), spontaneously. It has been shown that ERG is strongly diminished in human PVOD [17]. In addition, PVOD-like disease that can be induced in rats with mitomycin treatment is concomitant with loss of ERG expression, and this condition combines vascular occlusive remodelling with perivascular eosinophilic inflammation [18]. Other transcription factors that have been associated with the development of PAH-specific vascular lesions and inflammation are FoxO, hypoxia-inducible factors and twist basic helix–loop–helix transcription factor 1 (figure 7) [19].

The conflagration hypothesis
Let us have a look at what arguments could be in favour of a primary inflammatory process that initiates the pathophysiology of PAH; in other words, the conflagration hypothesis (figure 8). Immune self-tolerance is organised and controlled by a specialised subpopulation of T-lymphocytes that are known as regulatory T-cells (Tregs). Tregs attenuate the misled immune auto-reactivity that, to a certain extent, permanently occurs, even in the healthy [20]. When the Treg-dependent immune balance is functional, autoimmune disease is avoided [21]. It has been reported that the number and function of Tregs is substantially altered in PAH [22]. In experimental models, Treg deficiency in athymic SU5416- or chronic hypoxia-treated rats leads to a severe form of PH that is more accentuated in female animals. Treg reconstitution in those animals coherently protects against the disease [23, 24]. Moreover, it has been shown that Treg signalling through leptin, a pleotropic hormone involved in physiological immune

**FIGURE 6** Pattern recognition receptor (PRR)-induced inflammation in pulmonary arterial hypertension. DAMP: damage-associated molecular pattern; HMGB1: high mobility group box-1; LPS: lipopolysaccharide; PAMP: pathogen-associated molecular pattern; RAGE: receptor for advanced glycation end products; TLR4: toll-like receptor 4.
Functional Treg Recruitment in affected arteries

High activation and proliferation

Polarisation

Pathogenic targets under investigation

Reduced function and amount

CD4

Th17

IL-17

MHC

Autoimmune mediated reaction

Figure 8: Inflammation as a cause of structural alterations: autoimmunity in pulmonary arterial hypertension (PAH). CCL20: chemokine (C-C motif) ligand 20; CXCL13: chemokine ligand 13; FOXP: Forkhead box P; IL-17: interleukin-17; MHC: major histocompatibility complex; MoDC: monocyte-derived dendritic cell; NK: natural killer cell; NKT: natural killer T-cell; Th17: T-helper 17; tLT: tertiary lymphoid tissue.
modulation, is hampered in patients with idiopathic PAH (IPAH), as well as in patients with CTD-associated PAH and in non-PAH patients with CTD in a similar fashion, connecting the dots from PAH to mere autoimmune disease (figure 8) [25].

The link between innate and adaptive immunity is physiologically secured by dendritic cells (DCs) [26]. In the past, our group has identified and reported intense DC recruitment into the vessel wall of pulmonary arteries from patients with PAH and in PAH animal models [27–29]. We have shown that, in PAH, DCs trigger activation and proliferation of CD4+ T-cells, in concomitance with a T-helper 17 (Th17) immune primary analysis that includes data through week 24, the observed improvement in 6-min walking distance arteries from patients with PAH and in PAH animal models [27–29]. We have shown that, in PAH, DCs trigger activation and proliferation of CD4+ T-cells, in concomitance with a T-helper 17 (Th17) immune primary analysis that includes data through week 24, the observed improvement in 6-min walking distance

In a rat model of heritable PAH linked to BMPR2 mutation, pulmonary IL-6 overexpression discriminated –29]. We have shown that, in PAH, DCs

The conflagration concept of autoimmunity in PAH is further supported by the phenomenon of lymphoid neogenesis in lungs from patients with PAH: lymphocytic follicles which can be observed in the vicinity of diseased pulmonary arteries are part of so-called tertiary lymphoid tissues (tLTs) [32] that can emerge in many organs and tissues in response to persistent autoimmune-triggered inflammation and that go along with progressive organ dysfunction and degeneration, as it can be seen in RA [33]. Of note, one classic hallmark in RA histopathology of the lung is tertiary lymphocytic follicles in association with fibrotic non-specific interstitial pneumonia (NSIP)- or usual interstitial pneumonia (UIP)/idiopathic pulmonary fibrosis (IPF)-like changes of the lung parenchyma [34]. In contrast, NSIP and especially UIP virtually always present with pulmonary arterial and venous remodelling [35]. The constitution of tLT depends on chemokines, such as chemokine (C-C motif) ligand 20 and chemokine ligand 13 that have been shown to be overexpressed in explanted lungs of PAH patients [32]. IL-17, another important cytokine with regard to the constitution of tLTs and associated with human and experimental PH, drives the formation of tertiary lymphoid follicles in lung tissue as a pillar of local antimicrobial protection, yet also enhances local inflammatory responses and autoimmune disease [30, 36, 37]. The formation of tLTs could also be favoured by the functional and numeral loss of lymphocytic populations, such as Tregs and natural killer cells (natural killer cells and natural killer T-cells) [22, 25, 38, 39].

Importantly, it has been shown that tLTs produce auto-antibodies that may be directed against the endothelial cells of pulmonary vessels, and that this aggression induces apoptosis and, eventually, critical reactive hyperproliferation of surviving endothelial cells [40]. Although auto-antibodies against smooth muscle cells and fibroblasts have been reported in PAH [41], no major common auto-antigen has been clearly associated to the disease (figure 8).

**Targeting inflammation and dysregulated immunity in PAH**

From these points of view, inflammation and dysregulated immunity play a significant role in a spectrum of causes of PAH. From the perspective of identifying pathways that are targetable, IL-6 has emerged as a strong candidate. Lung-specific IL-6 overexpression transgenic mice develop PH with occlusive neointimal angioproliferative lesions that worsen with hypoxia and are composed of endothelial cells and T-lymphocytes [42]. IL-21, a Th17 cytokine, has been identified as a downstream target of IL-6 signalling in PAH. IL-6 blockade by the monoclonal anti-IL-6 receptor antibody, MR16-1, ameliorated hypoxia-induced PH and prevented the hypoxia-induced accumulation of Th17 cells and M2 macrophages in the lungs [43]. In a rat model of heritable PAH linked to BMPR2 mutation, pulmonary IL-6 overexpression discriminated rats that developed spontaneous PAH from rats that did not develop the condition [44]. In humans, plasma IL-6 provides incremental prognostic information in PAH, especially in patients with low brain natriuretic peptide levels [45] and explanted lungs from PAH patients display ectopic upregulation of membrane-bound IL-6 receptor on pulmonary artery smooth muscle cells [46]. However, therapeutic blockade of IL-6 signalling with tocilizumab demonstrated no significant effects on haemodynamics or exploratory secondary endpoints in heritable or IPAH. A potential improvement was noted in the small subgroup of patients with CTD-associated PAH. This raises the possibility that disease is driven by inflammation in only a subset of patients [47]. Other cytokines represent relevant therapeutic targets like tumour necrosis factor (TNF). Transgenic mice expressing chronically low levels of human TNF (TNF-Tg mice; line 3647) mimic human CTD-PAH, in which patient lungs demonstrate increased TNF signalling and significant similarities in genomic pathway dysregulation [48]. Treatment with TNF-α blocker etanercept (receptor Fc-based cytokine blocker) prevented and reversed monocrotaline-induced PAH in rats by reducing inflammatory cell infiltration [49]. Lastly, IL-4-/IL-13-deficient mice do not develop Schistosomiasis-associated PH, suggesting that Th2 cytokine production by CD4+ T-cells may be required to drive PH in this setting [50].

In a proof-of-concept, randomised, placebo-controlled study of B-cell depletion, rituximab treatment demonstrated an acceptable safety and tolerability profile with stable systemic sclerosis PAH [51]. In the primary analysis that includes data through week 24, the observed improvement in 6-min walking distance
favoured rituximab-treated patients, but this difference did not reach statistical significance. Interestingly, in a post hoc analysis, the use of machine learning allowed the identification of a panel of cytokines (low levels of rheumatoid factor, IL-2 and IL-17) that predicted a response to rituximab.

With regard to the genetic predisposition to PAH, an increasing number of genes has been linked to PAH pathophysiology. The most frequently encountered mutations concern the gene BMPR2 for hereditary PAH and an important fraction of IPAH, and the gene EIF2AK4 for PAH subgroup 1.6, a particular form of PAH with predominating veno-occlusive remodelling and capillary proliferation (PVOD) [52]. The dysfunction of transmembrane receptor bone morphogenetic protein receptor type 2 (BMPR2) diminishes the endothelial barrier function and increases the permeability towards circulating inflammatory mediators, such as cytokines or chemokines that build a chemoattractant gradient, along which different types of leukocytes can migrate into the vessel wall of pulmonary arteries and veins (figure 9) [53, 54].

The identification of loss-of-function mutations in KCNK3 (TASK-1) [55] and ABCC8 (SUR1) [56], or gain-of-function mutations in ABCC9 (SUR2) [57], as well as polymorphisms in KCNAl5 (Kv1.5) [58], which encode two potassium (K+) channels and two K+ channel regulatory subunits, has revived the interest in K+ channels in the pathobiology of PAH [59]. Indeed, potassium and calcium channels are involved in the regulation of pulmonary vascular tone [60]. The opening of plasma membrane K+ channels leads to hyperpolarisation of pulmonary artery smooth muscle cells resulting in the closure of voltage-sensitive Ca²⁺ channels and subsequent vasodilation, while acute contraction of pulmonary artery smooth muscle cells is activated in part by plasma membrane K⁺ channel inhibition-induced membrane depolarisation and subsequent Ca²⁺ entry through L-type Ca²⁺ channels. K⁺ channels are also strongly involved in the proliferation of leukocytes [61, 62]. The importance of K⁺ channels is especially critical in T-cell activation and cytokine release. Their opening leads to plasma membrane hyperpolarisation, which in turn increases the driving force for Ca²⁺ influx shared by Ca²⁺ release-activated Ca²⁺ channels [63].

Potassium two pore domain channel subfamily K member 3 (KCNK3) dysfunction contributes to the development of PAH and experimental PH [64–66]. Moreover, KCNK3-mutated rats are predisposed to develop PH [67]. Paradoxically, PH spontaneously develops only slowly over 12 months in these rats, and

![FIGURE 9 Heritable pulmonary arterial hypertension: predisposing mutations involved in immune regulation. BMPR2: bone morphogenetic protein receptor type 2; ER: endoplasmic reticulum; GCN2: general control nonrepressible 2; IL: interleukin; KCNK3: potassium two pore domain channel subfamily K member 3; PVOD: pulmonary veno-occlusive disease; TGF-β: transforming growth factor-β.]
while loss of KCNK3 is associated with impaired effector function in T-cells and protection against autoimmune disorders [68], KCNK3-mutated rats and mice and human PAH patients with KCNK3 mutation show increased cytokine induction and chronic immune cell activation (evidence of T-cell exhaustion) [64, 67, 69]. The question arises of how KCNK3 causes these defects. One possibility is that KCNK3 is most important not at the cell membrane, but at the membrane of the endoplasmic reticulum (ER), where it has been reported to regulate ER calcium homeostasis [70]. This explanation, recently investigated by West et al. [69], supposes that KCNK3 mutation primarily impacts ER calcium handling, resulting in ER stress and attendant metabolic shift, ending sequentially in cytokine/chemokine induction, cytokine/chemokine receptors activation and eventually in T-lymphocyte polarisation and recruitment (and other inflammatory cell types like macrophages) (figure 9).

General control nonderepressible 2 (GCN2), the protein product of EIF2AK4, is part of the integrated stress response and supports cell adaptation during restricted amino acid supply [71]. In myeloid cells, GCN2 is activated by catalytic degradation of tryptophan by indolamine-2,3-dioxygenase (IDO) [72]. The activation of the IDO pathway is induced by interferons during inflammation, coming from DCs, monocytes and macrophages. In addition, activation of the IDO pathway generates metabolites with immunosuppressive properties, such as the kynurenins [73]. In order to adapt to restricted amino acid conditions, GCN2 can activate autophagy [74]. However, the latter phenomenon opposes the inflammatory signals of macrophages during the immune response that are initiated by mitochondrial stress and the expression of cytokines such as IL-1β and IL-17 [75]. Hence, GCN2 appears to display a negative feedback control with regard to inflammation and its loss diminishes tolerance with regard to apoptotic cells and cellular debris and thereby shifts the balance towards autoimmunity, at least in animal models. GCN2-deficient CD8 T-cells have T-cell-intrinsic proliferative and trafficking defects not observed in CD4 T-cells [76]. Thus, GCN2 is also required for normal cytotoxic T-cell function, and this could explain the alterations of circulating cytotoxic cell subpopulations described in PVOD [38]. Its role in the development of human PVOD, however, remains less clearly defined (figure 9) [66].

Here are some additional thoughts on a possible inflammatory background in PAH development that underscore the difficulty of translating this concept into efficient therapeutic targets (figure 10): When analysing the explanted lungs of PAH patients who underwent year-long treatment with modern-era PAH specific therapy, we observe unresolved, stable-appearing collagen-rich obstructive vascular lesions, as well as a constant moderate chronic inflammatory infiltrate in the vessel’s perimeter [77].

![FIGURE 10 Controversies and consequences of pulmonary arterial hypertension therapy.](https://doi.org/10.1183/16000617.0161-2021)
surprising to some point, since modern PAH-specific therapy has been at least experimentally shown to combine vasodilating and antiproliferative properties with immune-modulatory qualities that may attenuate exuberant inflammation [4]. Loss of immunoregulation in PAH appears to favour an ongoing chronic inflammatory process, even in end-stage PAH at a lower level [30]. It would be of particular interest to study the possible mechanisms of resistance with regard to immune modulation in patients with PAH in order to establish more personalised and hence efficient immunotherapies.

It is important to state and to admit that blocking one specific pro-inflammatory cytokine will hardly change the patient’s fate or influence outcome, despite the observed and reported association of high overall cytokine levels and mortality [78]. In fact, during PAH, a vast inflammatory cytokine armada appears to be involved: therapeutically targeting only one single cytokine might be ineffective due to an important biologic redundancy of active immune mediators [79]. The dual roles of cytokines are another level of complexity. For instance, human-induced pluripotent stem cells with KCNK3 mutation and the lungs from KCNK3-mutated mice both expressed lower levels of IL-6 [69], which might be similar to the fact that IL6 maintains a stoichiometric balance in shifting signals from an active immune response to a suppressive state [80]. One should also note that IL-6 signalling prevents neointimal remodelling in Schistosomiasis-associated PH [81], possibly highlighting a dual role for IL-6. Of relevance, Sotatercept, a ligand trap with high selectivity for multiple members of the TGF-β superfamily, significantly improved pulmonary-vascular, cardiovascular and exercise-related outcomes in PAH patients [82]. It is believed to act as a reverse-remodelling agent designed to rebalance bone morphogenetic protein/activin signalling. However, it is well known that IL-6 promotes the differentiation of Th17 cells from naïve T-cells, presumably in cooperation with TGF-β in the lung [43]. It is possible that Sotatercept may work in other ways than those initially expected and, thus, targets some crucial cytokine/cytokine networks.

**Conclusion**

In conclusion, different pro-inflammatory and dysimmune triggers and pathways may – in sensible/vulnerable patients, preferably with a genetic predisposition [83] – ignite and fuel persistent pulmonary vascular inflammation and eventually irreversible obstructive remodelling of the vessel wall. The precise mechanisms through which autoimmunity triggers vascular dysfunction are yet to be clarified.

As we have seen, there are numerous pro-inflammatory pathways and pathomechanisms, including perturbated blood-flow, vessel wall stiffness, hypoxia, bacterial translocation with pattern recognition receptor-induced inflammation and the influence of various transcription factors. Finally, up to now, immune-modulating therapeutic approaches, in general, show only very limited efficacy in PAH patients with regard to haemodynamic and clinical parameters.

We have learned from the rituximab and the tocilizumab trials that there could also be a potential benefit of addressing cells over soluble factors and that future stratified medicine approaches based on underlying (autoimmune) aetiology and inflammatory biomarkers may identify responder populations to immunomodulation. Like in other fields of applied biomedical research, for instance in novel personalised cancer therapies, these observations should not dishearten researchers within the PH community, but rather be understood as a challenge to pioneer a promising therapeutic landscape, almost in the range of our fingertips (figure 11).
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