The multifaceted roles of the matrikine Pro-Gly-Pro in pulmonary health and disease

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ABSTRACT Matrikines are bioactive fragments of the extracellular matrix (ECM) that are fundamental in regulating a diverse array of physiological processes. The tripeptide Proline-Glycine-Proline (PGP) is a collagen-derived matrikine that has classically been described as a neutrophil chemoattractant. In this article, we describe our current understanding of the pathways that generate, degrade and modify PGP to dictate its bioavailability and stability. Additionally, we discuss our expanding appreciation of the capacity of PGP to regulate diverse cell types and biological processes, independent of its activity on neutrophils, including a putative role in wound repair. We argue that PGP functions as a primitive and conserved damage-associated molecular pattern, which is generated during infection or injury and subsequently acts to shape ensuing inflammatory and repair processes. As a fragment of the ECM that accumulates at the epicentre of the action, PGP is perfectly positioned to focus neutrophils to the exact site required and direct a localised repair response. However, it is essential that PGP is efficiently degraded, as if this matrikine is allowed to persist then pathology can ensue. Accordingly, we discuss how this pathway is subverted in chronic lung diseases giving rise to persistent inflammation and pathological tissue remodelling.

Introduction
The extracellular matrix (ECM) is a non-cellular component of all tissues and organs, which serves as a scaffold for constituent cells. The individual components and three-dimensional structure of the ECM imparts signals to resident cells to shape fundamental processes such tissue development, maintenance of homeostasis, inflammation and wound repair. The ECM is a dynamic entity that undergoes continuous degradation and resynthesis. This can result in the proteolytic liberation of bioactive fragemns termed “matrikines” that function to further modulate cell behaviour. Classically, we consider the complex interplay of mediators such as cytokines, chemokines, growth factors and eicosanoids in regulating fundamental physiological processes but overlook the significance of matrikines. However, matrikines, derived from ECM macromolecules such as collagen, elastin, laminin and hyaluronan, have been implicated in the regulation of diverse biological processes [1]. Localised perturbations within a tissue will inevitably result in alterations to the ECM and frequently the release of matrikines. Consequently, matrikines are perfectly positioned to sense microenvironmental fluctuations and dictate an appropriate

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response to the exact location and at the precise time when required to meet the immediate needs of the tissue. However, it is mandatory that ECM turnover be tightly regulated as an aberrant ECM and dysregulated matrikine production are implicated in the pathology of many chronic diseases. This article will focus on our current understanding of the collagen-derived matrikine Proline-Glycine-Proline (PGP). We detail pathways that define its bioavailability, its central role in regulating basic biological processes, and how dysregulation of the pathway is implicated in disease states.

**PGP: a multifaceted matrikine**

Collagen represents in excess of 90% of the total protein mass of the ECM in mammals, and proteolytic processing of native collagen can yield the matrikine PGP. The prevalence of the PGP sequence within collagen molecules (28 PGP sequences per type I collagen fibril, 43 per type III collagen, 25 per type IV collagen and 44 per type V collagen) results in a potentially abundant bioactive signalling moiety that remains cryptic within the triple helical structure of collagen until liberated by proteolytic processing. Once released, the small and hydrophilic nature of PGP means that it can readily diffuse through the dense ECM. Furthermore, its unusual structure owing to the cycling back of the proline side chains onto the backbone amino group results in a matrikine that is resistant to generic protease degradation.

Peister et al. [2] first identified N-terminal acetylated PGP (AcPGP) and N-terminal methylated PGP in an alkali eye injury model in rabbits, demonstrating their capacity to drive neutrophil recruitment and ensuing corneal ulceration [3–5]. Whilst conventional glutamic acid-leucine-arginine+ (ELR+) CXC neutrophil chemokines are functional at nanomolar levels, AcPGP was subsequently demonstrated to operate at a micromolar level [6]. The unacetylated peptide (PGP) also evokes neutrophil chemotaxis but is 4–7-fold less potent [7]. Weatherington et al. [6] convincingly demonstrated that PGP functioned as a neutrophil chemoattractant by mimicking key sequences found in classical neutrophil chemokines and signalling through CXCR1/2. Accordingly, intratracheal instillation of AcPGP dose dependently elicited neutrophilic inflammation in the airways of mice that was abolished in cxcr2−/− animals [6, 8]. Subsequent studies have highlighted downstream signalling events following CXCR1/2 engagement by AcPGP [9, 10], and demonstrated the capacity of AcPGP to drive neutrophil superoxide production [6] and matrix metalloproteinase (MMP)-9 [9] and CXCL8 release [10]. More recently, Afonso et al. [11] demonstrated that collagen engagement of discodin domain receptor 2 on the surface of neutrophils promoted the release of PGP-generating enzymes at the leading edge of the neutrophil and the generation of a localised PGP gradient. This acted as a compass to focus neutrophil migration through a three-dimensional matrix [11], rationalising how this pathway may operate physiologically.

Increasingly, it is apparent that PGP can have profound effects on other cells independent of its classical activity as a neutrophil chemoattractant. Hahn et al. [12] demonstrated that PGP and AcPGP could function at nanomolar concentrations to promote paracellular permeability in endothelial cells both in vitro and in vivo, in a process that was CXCR2 dependent. Subsequently, the same group demonstrated that AcPGP could also drive the release of pro-inflammatory mediator endothelin-1 from aortic endothelial cells both via ligation of CXCR2. Additionally, recent studies have suggested a role for PGP in facilitating wound repair. AcPGP was demonstrated to drive CXCR2-dependent migration, proliferation and tube-forming activity in endothelial progenitor cells at nanomolar concentrations. Subsequently, topical application of AcPGP promoted cutaneous neovascularisation and wound healing [13]. Furthermore, our group have recently demonstrated a profound capacity of PGP/AcPGP to promote proliferation, radial spreading and prominent lamellipod formation in human lung bronchial epithelial cells [14], cardinal features of epithelial wound repair responses.

**Pathways defining PGP bioavailability**

Generation of PGP from collagen is a multistep enzymatic process that requires the concerted action of specific MMPs and prolylendopeptidase (figure 1). The initial cleavage of native collagen by MMP-1, MMP-8 or MMP-9 yields fragments that are a suitable substrate size for prolylendopeptidase, which subsequently acts to liberate PGP [7, 15]. MMPs can be derived from a variety of cellular sources [16], whilst prolylendopeptidase has been reported to be expressed by neutrophils [15], airway macrophages [17] and epithelial cells [17, 18]. Since neutrophils are an abundant source of the proteases that generate PGP it is anticipated that this pathway can drive a self-sustaining cycle of neutrophilic inflammation [9].

We rationalised that pathways must be in place to ensure that PGP-mediated inflammation is resolved. Accordingly, we identified an anti-inflammatory pathway whereby PGP is degraded during episodes of acute neutrophilic inflammation by a novel extracellular aminopeptidase activity of the enzyme leukotriene A₄ hydrolase (LTA₄H) [19]. LTA₄H classically functions intracellularly within myeloid cells as an epoxide hydrolase to convert leukotriene A₄ to pro-inflammatory lipid mediator LTB₄ [20, 21]. LTB₄ drives the recruitment and activation of an array of cells including neutrophils and is implicated in protection against
invading microorganisms but also in the pathology of a multitude of chronic diseases [22]. Thus LTA₄H represents a highly unusual enzyme with directly opposing pro- and anti-inflammatory activities that dictate the amplitude and persistence of neutrophilic inflammation (figure 2) [23]. Whilst the epoxide hydrolase activity of LTA₄H operates intracellularly primarily within myeloid cells, the cellular source of the extracellular enzyme is less clear.

Whilst PGP is readily broken down by LTA₄H, AcPGP is completely protected from degradation [19] and thus the acetylation process is seemingly a key event. Acetylation of peptides is classically the function of N-acetyl transferases. However, the small size and N-terminal proline of PGP seemingly preclude it from being a favourable substrate for such enzymes [24, 25]. Subsequently, we have demonstrated that PGP can be chemically acetylated through the action of reactive aldehydes to yield a species (AcPGP) that is resistant to LTA₄H-mediated degradation. The AcPGP can, however, be degraded by the action of angiotensin converting enzyme.

PGP in health
In response to infection/injury, we believe that PGP essentially functions as a damage-associated molecular pattern, operating to fine-tune neutrophil recruitment to the specific site required (figure 3). Subsequently, as neutrophils navigate through the ECM they generate “breadcrumbs” of PGP to guide other proximal neutrophils. Indeed, it is feasible that whilst neutrophil-derived proteases function to breach the collagen basement membrane during extravasation that they are concomitantly generating PGP to attract
neutrophils to a focal point of exit from the vasculature. More recent studies pertaining to endothelial progenitor cells [13] and bronchial epithelial cells [14], suggests that PGP also possesses a previously unheralded function in wound healing and may repair demarcated airway epithelium and promote neo-angiogenesis. Thus, in essence, a fragment of the ECM liberated during inflammation and injury could remarkably be functioning to direct a localised repair response at the specific site of the inflammatory reaction (figure 3).

A constant dynamic exists between generation and degradation of PGP/AcPGP. In acute self-resolving settings it is absolutely the norm for PGP/AcPGP to be efficiently degraded and not persist [19, 23, 33]. High levels of ACE are present in the plasma [34], and we have demonstrated that airway ACE levels are elevated during episodes of inflammation as consequence of increased vascular permeability and an influx of plasma enzyme [29]. We have also observed very high levels of extracellular LTA4H circulating in plasma, and thus airway levels may again be dictated, in part, by vascular permeability. It is noteworthy, therefore, that neutrophils and PGP itself impart changes in endothelial cells to promote vascular permeability, which would in turn enable an influx of extracellular LTA4H and ACE to degrade PGP/AcPGP. Thus PGP/AcPGP can potentiate their own degradation to facilitate the efficient resolution of inflammation and restoration of homeostasis.

**PGP in chronic lung disease**

A protease–anti-protease imbalance is a hallmark of many chronic lung diseases, resulting in disruption of tissue architecture and excessive matrikine production that can perpetuate inflammation and remodelling.
There is accumulating evidence that PGP persistence can drive pathologies observed in chronic lung diseases (figure 4). We have demonstrated that a failure to degrade PGP in a murine model of acute self-resolving airway inflammation resulted in substantially augmented neutrophils and macrophages, a general protease imbalance and ECM degradation [33]; hallmark features of many chronic lung diseases. Furthermore, whilst PGP may function to promote bronchial epithelial repair, we have seen that a failure to degrade PGP in a mouse model of allergic airways disease results in pathological epithelial remodelling, mucus hypersecretion and exacerbated airway hyperresponsiveness [14]. Given the capacity of PGP to promote endothelial permeability [12] and neovascularisation [13], we believe that persistence of this matrikine could also contribute to oedema and pathological angiogenesis observed in some chronic lung diseases.

Chronic obstructive pulmonary disease (COPD) describes a progressive, irreversible and predominantly smoking-induced small airway and/or emphysematous disease associated with airflow limitation. Chronic exposure of mice to cigarette smoke causes PGP/AcPGP accumulation, neutrophilic inflammation and emphysema. Reducing/neutralising PGP/AcPGP in this cigarette smoke model ameliorates inflammation and pathology [17, 19, 35, 36], whilst promoting PGP degradation reduces inflammation and emphysema [37]. Furthermore, it is noteworthy that chronic administration of AcPGP alone into the airways of mice is sufficient to elicit an emphysematous phenotype [6, 8]. Elevated PGP/AcPGP have been reported in sputum, bronchoalveolar lavage fluid and serum of COPD patients, often correlating with neutrophil surrogate marker myeloperoxidase (MPO) [6, 15, 35, 38, 39]. This is true for all Global Initiative for Chronic Obstructive Pulmonary Disease stages of disease and in patients that have stopped smoking [35]. Preliminary studies suggest that sputum PGP levels are highest around the time of COPD exacerbation and decline with successful treatment [38], potentially supportive of PGP/AcPGP being utilised as a biomarker for disease severity. Intriguingly, a recent study has also highlighted that AcPGP accumulation can drive tumour cell dissemination to lung parenchyma [40], potentially rationalising the association between inflammatory processes in smokers and an increased risk of lung cancer.

Rationalising the accumulation of PGP/AcPGP in COPD patients, we have demonstrated that LTA4H-mediated PGP degradation is perturbed by cigarette smoke. Cigarette smoke contains large amounts of LTA4H, which may contribute to the persistence of PGP/AcPGP in COPD patients.
amounts of reactive aldehydes that chemically acetylate PGP, enhancing its chemotactic activity and protecting it from degradation by LTA4H [19, 23, 26]. Additionally, cigarette smoke selectively abrogates the peptidase activity of LTA4H [19, 35, 41], again seemingly driven by intrinsic reactive aldehydes. Reactive aldehyde scavengers such as carbocisteine are able to prevent cigarette smoke-induced PGP acetylation and LTA4H inhibition. Thus, it is pertinent that these compounds have demonstrated clinical benefit in COPD patients and reduced inflammation in animal models of emphysema [42–44]. ACE levels and activity in COPD patients remain largely unexplored though there are suggestions that it too may be reduced [29].

Cystic fibrosis (CF) is a lethal genetic disorder caused by defective function of the CF transmembrane conductance regulator, which predisposes patients to infection, chronic airway inflammation (with a prominent neutrophilia) and airway remodelling. PGP/AcPGP is elevated in CF patients, correlating with PGP-generating enzymes and neutrophil surrogate MPO [7]. Importantly, PGP levels were shown to spike with disease exacerbation, decline with inpatient therapy and inversely correlate with forced expiratory volume in 1 s. Whilst lung transplantation is the therapeutic modality frequently used for end-stage lung disease, recipients often have a poor prognosis due to the development of bronchiolitis obliterans.
syndrome (BOS), a condition characterised by a neutrophil influx and ECM remodelling. BOS patients have elevated levels of PGP in their bronchoalveolar lavage fluid, again showing a striking correlation with levels of MMP-9, prolylendopeptidase and MPO [45]. Importantly, HARDISON et al. [45] demonstrated that the capacity of BOS bronchoalveolar lavage fluid to elicit neutrophil chemotaxis ex vivo was predominantly attributable to the intrinsic PGP component, which was relatively more important as a driver of the neutrophil chemotaxis than the CXCL8 present. Asthma is a heterogeneous disease with variable inflammatory, remodelling and clinical phenotypes. Severe asthma phenotypes have been described that exhibit pronounced neutrophilic inflammation [46], and pathological epithelial remodelling that contributes to progressive decline in lung function [47]; hallmark features of PGP persistence. Intriguingly, we have recently demonstrated that AcPGP accumulates specifically in the sputum of severe asthmatics [14]. The rationale for why PGP/AcPGP is able to accumulate in CF, BOS and severe asthma patients is the subject of ongoing investigation.

An over-exuberant or persistent PGP response is clearly implicated in the pathologies of certain chronic lung diseases, but a recent study paradoxically suggests that too little PGP can yield a distinct pathology. Idiopathic pulmonary fibrosis (IPF) and COPD are anticipated to share similar aetiologies, with cigarette smoke the major risk factor for the development of both. However, IPF displays quite distinct clinical and pathological features manifesting as an interstitial fibrosis and honeycombing [48]. Whilst PGP/AcPGP is prevalent in COPD patients, it is undetectable in patients with IPF [29]. Furthermore, therapeutic administration of AcPGP/PGP in a murine belomycin model of fibrosis completely abrogated pulmonary fibrosis [29]. Accordingly, we demonstrated the pathogenic function attributed to ACE in fibrosis to be a consequence of overzealous AcPGP degradation [29]. Thus PGP/AcPGP can seemingly have very divergent roles: pathogenic in their capacity to drive neutrophilic inflammation and ECM degradation in the context of COPD, but protective in their capacity to limit fibrosis in IPF. Accordingly, we believe that disparate perturbations of the PGP–LTA₄H/AcPGP–ACE axis contribute to these distinct pathologies.

Conclusions

We anticipate that PGP essentially functions as a primitive and conserved damage-associated molecular pattern, which is generated during infection or injury and acts to shape ensuing inflammatory and repair processes. As a fragment of the ECM that accumulates at the epicentre of the action, PGP is perfectly positioned to focus neutrophils to exactly where they are required, as well as initiate proximal repair responses. However, it is critical the PGP is efficiently degraded as its persistence can drive pathological features of chronic lung diseases. We have previously discussed strategies that could be utilised to target this matrikine therapeutically to potentially ameliorate chronic lung disease pathologies; be it neutralisation of the peptide itself, inhibition of PGP-generating enzymes or receptor blockade [49]. However, our increasing realisation of the potential role of PGP in wound repair processes and the surprising recent developments that it is protective against pulmonary fibrosis, highlights the necessity to maintain homeostatic levels and questions the feasibility of targeting this matrikine. Furthermore, we should now consider implications and opportunities for LTA₄H and ACE inhibitors, which have been developed to reduce LTB₄ and angiotensin II, respectively, if they inadvertently cause PGP and AcPGP accumulation [31, 50].

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