Cockroach protease allergen induces allergic airway inflammation via epithelial cell activation

Sagar L. Kale¹, Komal Agrawal¹, Shailendra Nath Gaur² & Naveen Arora¹

Protease allergens are known to enhance allergic inflammation but their exact role in initiation of allergic reactions at mucosal surfaces still remains elusive. This study was aimed at deciphering the role of serine protease activity of Per a 10, a major cockroach allergen in initiation of allergic inflammation at mucosal surfaces. We demonstrate that Per a 10 increases epithelial permeability by disruption of tight junction proteins, ZO-1 and occludin, and enhances the migration of Monocyte derived dendritic cell precursors towards epithelial layer as exhibited by trans-well studies. Per a 10 exposure also leads to secretion of IL-33, TSLP and intracellular Ca²⁺ dependent increase in ATP levels. Further, in vivo experiments revealed that Per a 10 administration in mice elevated allergic inflammatory parameters along with high levels of IL-33, TSLP, IL-1α and uric acid in the mice lungs. We next demonstrated that Per a 10 cleaves CD23 (low affinity IgE receptor) from the surface of PBMCs and purified B cells and CD25 (IL-2 receptor) from the surface of PBMCs and purified T cells in an activity dependent manner, which might favour Th2 responses. In conclusion, protease activity of Per a 10 plays a significant role in initiation of allergic airway inflammation at the mucosal surfaces.

The prevalence of allergic diseases is increasing worldwide from past few decades in developing as well as developed countries. Allergic inflammatory responses are initiated on allergen uptake by APCs, predominantly DCs and subsequent polarisation of the T cell towards Th2. Though studies have tried to elucidate the protein characteristics that confer them with the ability to induce Th2 responses, it is still unclear why some proteins are allergenic while others are not. Recent studies have demonstrated the role of biological properties like proteolytic activity (HDM, cockroaches, fungal extracts) and phospholipase activity (bee venom) of some allergens in development of Th2 immune responses. Several allergens from a variety of sources viz., HDM, molds and cockroaches have proteolytic activity. These proteases have been shown to skew the immune response towards Th2 by the virtue of their proteolytic activity.

Dendritic cells, the professional antigen presenting cells of the immune system are the sentinels of immunity and tolerance. Dendritic cells act as a bridge between innate immune sensing and initiation of adaptive immune responses. Previously, Per a 10, a major serine protease allergen from American cockroaches (Periplaneta americana) has been shown to bias dendritic cell mediated polarisation of naïve T cells towards Th2. But the effect of Per a 10 exposure on the migration of DC precursors in the airway epithelium remains to be elucidated. Further, protease allergens are known to activate airway epithelial cells to secrete a wide array of chemokines and cytokines. Emerging studies have determined the role of epithelial cytokines, chemokines and mediators in orchestrating the local inflammatory responses and have directly implicated epithelial cells in Th2 responses. Per a 10, has also been shown to activate airway epithelial cells to secrete IL-6, IL-8 and GMCSF in an activity dependent manner via PAR-2 receptors. Airway epithelium acts as a first line of defence for the inhaled particles and forms a physical as well as a chemical barrier that shields immune cells from the external environment. For sensitization, allergens need to cross the epithelial barrier to interact with subepithelial dendritic cells so that they are processed and presented. This epithelial barrier is maintained by tight junctions that allow cell to cell contacts.

¹Allergy and Immunology Section, CSIR-Institute of Genomics and Integrative Biology, Delhi University Campus, Mall road, Delhi-110007, India. ²Department of Respiratory Medicine, Vallabhbhai Patel Chest Institute, University of Delhi, Delhi, India. Correspondence and requests for materials should be addressed to N.A. (email: naveen@igib.res.in)
and defective barrier plays a role in pathogenesis of allergic diseases like asthma and dermatitis. Proteases like Der p 1, Pen c 13 etc. disrupt airway epithelial barrier and contribute to sensitization and allergic responses. Mucosal epithelial cells not only form a barrier for inhaled environment but are able to modulate innate and adaptive immune responses.

Though proteolytic activity of Per a 10 causes allergic airway inflammation, its role in creating a Th2 microenvironment remains to be deciphered. In this study, we investigated the role of serine protease activity of Per a 10 in initiating and promoting allergic responses at mucosal surfaces.

**Results**

**Per a 10 increases epithelial permeability in a time and activity dependent manner.** Per a 10 was isolated from cockroach crude extract and it resolved as a single band at 28kDa on SDS-PAGE. The endotoxin levels in Per a 10 were determined by LAL assay and were below the detection limit (0.01 ng/mg of protein) of the kit. Native Per a 10 was proteolytically active but both recombinant Per a 10 (rPer a 10) and heat inactivated Per a 10 (ΔPer a 10) were proteolytically inactive. We investigated the effect of Per a 10 exposure on trans-epithelial permeability of Calu-3 cells, a human derived airway epithelial cell line cultured at air liquid interface, by using fluorescently labelled dextran. On exposure with native Per a 10 the transepithelial permeability increased in a time dependent manner but significant increase in the permeability was observed only after 8 h of exposure (Fig. 1a). Next, to investigate the role of serine protease activity of Per a 10 on epithelial permeability, Calu-3 cells grown at air liquid interface were exposed either to proteolytically active or inactive Per a 10. Both, rPer a 10 and ΔPer a 10 did not cause any significant change in the epithelial permeability as compared to the PBS exposed cells (Fig. 1b), demonstrating the role of protease activity in increased epithelial permeability.

Tight junctions are apical cell-cell contacts in polarised cells like epithelia and form a physical barrier. The significant increase in permeability of Calu-3 cells caused by Per a 10 exposure may be due to cleavage of inter junctional complexes. So, we next sought to determine the effect of Per a 10 exposure on tight junctional proteins ZO-1 and Occludin of Calu-3 cells by immunofluorescence. As shown in the Fig. 1(c) ZO-1 and Occludin staining...
in the control PBS exposed Calu-3 cells was continuous at the whereas in Per a 10 treated cells it was discontinuous and disrupted with reduced fluorescence (Fig. 1c) depicting cleavage of these proteins by Per a 10.

**Per a 10 induces migration of dendritic cell precursors.** CCL-2 levels are increased in bronchoalveolar lavage of asthmatic patients. Bronchial epithelial cells exposed to active Per a 10 showed increased secretion of CCL-2 in the supernatant as compared to inactive Per a 10 stimulated cells (Fig. 2a). PBMCs are the source for recruitment of DC precursors during allergen challenge; we isolated CD14+ monocytes from PBMCs, obtained from cockroach hypersensitive patients and cultured them in presence of rhIL-4 and rhGMCSF to generate DC precursors. We then checked the role of Per a 10 activated bronchial epithelial cells in migration of DC precursors by co-culture experiments. Per a 10 stimulated BEAS-2B cells caused migration of DCs from upper chamber to lower chamber of trans-wells as shown in Fig. 2b as compared to unstimulated or inactive Per a 10 stimulated epithelial cells.

**Per a 10 cleaves CD23, a low affinity IgE receptor.** CD23 is a regulator of IgE network. Binding of membrane bound CD23 (mCD23) on B cells to IgE mediates a negative feedback loop for IgE synthesis whereas sCD23 mediates IgE synthesis promoting pathway19. We tried to elucidate the effect of protease activity of Per a 10 on membrane bound CD23. PBMCs were isolated and incubated with different concentrations of Per a 10 and analysed for surface expression of CD23 by flow cytometry. Percent of cells expressing CD23 decreased with increasing concentration of Per a 10 (Fig. 3a). There is approximately half fold reduction in cells expressing CD23 on treatment with native active Per a 10 as compared to PBS and inactive Per a 10 exposed cells (Fig. 3b). Cells exposed to rPer a 10 and ΔPer a 10 showed no change in percent cells expressing CD23, as compared to PBS treated cells, suggesting role of proteolytic activity in reduced CD23 expression. Further, when purified B cells were exposed to Per a 10, percent cells expressing CD23 was significantly lower than ΔPer a 10 and rPer a 10 exposed cells (Fig. 3c).

**Per a 10 cleaves the α subunit of IL-2 receptor (CD25).** IL-2R is crucial for the propagation of Th1 cells and its cleavage can bias the immune response towards Th220. As Per a 10 cleaves CD23, we next checked for the cleavage of CD25 on PBMCs from healthy individuals that were exposed to native active Per a 10 or inactive ΔPer a 10 or rPer a 10 for 4 hrs. The surface expression of CD25 was analysed with flow cytometry. After exposure to Per a 10 the percentage of cells expressing CD25 reduced as compared to PBS stimulated cells in a dose dependent manner (Fig. 3d). There was no change in the percentage of cells expressing CD25 when exposed to inactive Per a 10 as compared to PBS stimulated cells (Fig. 3d). Similarly, purified CD25+ CD4+ T cells exposed to native Per a 10 showed significant decrease in CD25 expression as compared to ΔPer a 10 and rPer a 10 treated cells (Fig. 3f).

**Per a 10 activity induces secretion of TSLP and IL-33 from bronchial epithelial cell line BEAS-2B.** Previously, we have shown that Per a 10 activates airway epithelial cells to secrete proinflammatory cytokines12. The effect of Per a 10 activity on induction of IL-33 and TSLP from bronchial epithelial cell line BEAS-2B was assessed in this study. Serum starved BEAS-2B cells were exposed to 10 μg of active or inactive Per a 10 and the supernatant was analyzed for the secretion of IL-33 and TSLP by ELISA. As shown in the Fig. 3, active Per a 10 induces significant increase in secretion of IL-33 (Fig. 4a) and TSLP (Fig. 4b) from BEAS-2B cells. There was no significant change in the levels of cytokines in the supernatants of inactive Per a 10 (ΔPer a 10 and rPer a 10) stimulated BEAS-2B cells (Fig. 4a,b).
Figure 3. Per a 10 cleaves CD23, a low affinity IgE receptor and CD25, an IL-2 receptor subunit. PBMCs from normal healthy donors were cultured for 72 hours in the presence of PHA (5 μg/ml) in RPMI-1640 supplemented with 10% FBS. Cells were incubated with different stimuli for 4 hours, stained with FITC labelled anti CD23 or anti CD25 antibody and analysed by flow cytometry. B cells and CD4+ T cells were isolated from PBMCs using MACS, incubated with 10 μg of either active or inactive Per a 10 for 4 hrs and were stained with anti CD23 or CD25 antibody. (a) CD23 expression on PBMCs exposed with native active Per a 10 at varying concentrations, (b) CD23 expression on PBMCs exposed with 10 μg of Per a 10, ΔPer a 10 or rPer a 10. (c) Expression of CD23 on purified B cells exposed to 10 μg of Per a 10, ΔPer a 10 or rPer a 10, (d) CD25 expression on PBMCs exposed with Per a 10 at different concentrations, (e) CD25 expression on PBMCs exposed with 10 μg of Per a 10, ΔPer a 10 or rPer a 10, (f) Expression of CD25 on purified CD4+ T cells. Dot plots are representative of one of the three independent experiments. Data presented as mean ± SEM of three independent experiments. *P < 0.05 as compared to PBS exposed cells.
Per a 10 elevates ATP levels in BEAS-2B cells. Extracellular ATP, a danger alarmin has been previously shown to trigger the release of IL-33 and initiate Th2 type responses. As Per a 10 elevates the secretion of IL-33 from exposed BEAS-2B cells, we checked if Per a 10 also increases ATP release from BEAS-2B cells. Per a 10 exposure increased ATP levels in the BEAS-2B cell supernatant in a time dependent manner (Fig. 4d). Next, to assess whether intracellular Ca\(^{2+}\) mobilisation is involved in ATP release, BEAS-2B cells pre-exposed to BAPTA-AM (an intracellular calcium chelator) were stimulated with Per a 10 and ATP release with time was monitored. As shown in Fig. 4d chelation of intracellular Ca\(^{2+}\) prevented the release of ATP in Per a 10 exposed BEAS-2B cells indicating its upstream role in ATP release. Also, it has been demonstrated that IL-1\(\alpha\) triggers the release of IL-33. IL-1\(\alpha\) levels as shown in Fig. 4c were significantly elevated in Per a 10 stimulated BEAS-2B cells as compared to inactive Per a 10 immunised mice demonstrating the role of IL-1\(\alpha\) in Per a 10 induced epithelial activation.

Intranasal administration of Per a 10 induces allergic airway inflammation in mice. To determine the role of protease activity of Per a 10 on allergic airway inflammation, Balb/c mice were intranasally administered either with active or inactive Per a 10 (Fig. 5a) and allergic inflammatory parameters were analysed. Total cell count (Fig. 5b) and eosinophil peroxidase activity (EPO) (Fig. 5c) were higher in the BALF of Per a 10 immunised mice as compared to PBS administered mice and inactive Per a 10 (\(\Delta\)Per a 10 and rPer a 10) immunised mice. There was no significant change in the total cell count and EPO activity between PBS administered and inactive Per a 10 administered mice (Fig. 5b,c). Further, haematoxylin and eosin staining of the lung sections revealed increment in cellular infiltration in mice administered with active Per a 10 as compared to mice administered either with inactive Per a 10 or PBS (Fig. 5d). Active Per a 10 administered mice also showed elevated levels of IL-4 (Fig. 5e) in BALF and IgE (Fig. 5f) in serum in comparison to the mice exposed to PBS or inactive Per a 10.

**Figure 4. Per a 10 activates bronchial epithelial cells.** Serum starved BEAS-2B cells were stimulated either with 10\(\mu\)g of Per a 10, \(\Delta\)Per a 10 or rPer a 10 for 24 hours and (a) IL-33 (b) TSLP and (c) IL-1\(\alpha\) levels in the supernatant were assessed by ELISA. (d) ATP levels evaluated using luciferase/luciferin bioluminescence ATP determination kit (Invitrogen/Life technologies) and a luminometer, in supernatant of BEAS-2B cells grown in 96 well plate and stimulated with Per a 10 or PBS or cells incubated with BAPTA-AM and Per a 10. \(#P < 0.05\) as compared to Per a 10 stimulated cells.
Active Per a 10 elevates IL-33 and TSLP levels in the mice lungs. As Per a 10 induced secretion of IL-33 and TSLP from BEAS-2B cells, we checked whether Per a 10 exposure leads to any changes in the levels of these cytokines in mice model of allergic airway inflammation. Quantitative PCR analysis of IL-33 and TSLP mRNA expression in mice sensitized with active Per a 10 and inactive Per a 10 ($\Delta$Per a 10 and rPer a 10) revealed that the expression of IL-33 (Fig. 6a) and TSLP (Fig. 6b) is increased in active Per a 10 sensitized mice as compared to inactive Per a 10 sensitized mice.

Next we sought to determine the effect of Per a 10 sensitization on IL-33 and TSLP levels in the lung homogenate and BALF of mice sensitized with either active or inactive Per a 10. Levels of both the cytokines were elevated in the lungs of active Per a 10 sensitized mice as compared to PBS sensitized mice (Fig. 6c,d). There was no significant difference in the levels of both these cytokines in inactive Per a 10 sensitized mice as compared to PBS sensitized control group (Fig. 6c,d).

Per a 10 induces IL-1$\alpha$ secretion and uric acid release upon administration in the lungs. In corroboration with our in vitro results active Per a 10 exposure led to higher levels of IL-1$\alpha$ in the BALF as compared to inactive Per a 10 exposed mice (Fig. 6e). Uric acid is a danger associated molecule and precedes IL-33 secretion. The levels of uric acid in BALF of active Per a 10 sensitized mice were higher than in the inactive Per a 10 sensitized mice (Fig. 6f). Further, there was no difference in the levels of uric acid in inactive Per a 10 sensitized mice as compared to PBS sensitized control group (Fig. 6f). This indicates that respiratory mucosa on encountering protease allergen like Per a 10 releases uric acid in the airway lumen that can further lead to increased levels of IL-33 and TSLP, thus initiating allergen specific Th2 responses.

Discussion
Prevalence of allergic airway diseases is on the rise in past few decades, yet the mechanisms that lead to Th2 immune responses are not fully understood. Exogenous proteases from house dust mites, cockroaches, fungi and pollen or endogenous proteases viz. neutrophil elastase, mast cell tryptase etc. are implicated in initiating and exacerbating allergic responses$^{22–25}$. Airway epithelial cells act as a first line of defence for inhaled pathogens and aeroallergens$^{30}$. Lung epithelial cells form a tight, virtually impermeable barrier for macromolecules and regulates paracellular transport of inhaled material by the formation of tight junctions at the apical surfaces$^{18}$. In order to activate the immune cascade, allergens need to disrupt the tight epithelial barrier and gain access to the immune components of the host. Our study demonstrates that Per a 10 increases trans-epithelial permeability of Calu-3 cells by disruption of tight junctions which is in accordance with previous studies carried out with Der p 1, a cysteine protease and pollen proteases$^{32,37}$. Also, impaired epithelium is evident in the lungs of allergic asthmatics$^{34}$, suggesting possible damage by the inhaled allergens. Previously, Per a 10 has been shown to provide
adjuvant effect to other bystander allergens in the same environment. Per a 10, by compromising the epithelial barrier may facilitate the entry of other allergens across epithelial barrier and thus can act as an adjuvant to other bystander allergens in the same environment as reported earlier.

Allergens favour Th2 polarisation of naive T cells, but the mechanism of this polarisation is still not clearly understood. Our results demonstrate that Per a 10 cleaves CD25 in an activity and dose dependent manner. CD25 is a 55 kDa subunit of IL-2 receptor. IL-2 plays an important role in T cell proliferation and IL-2 receptor is pivotal for propagation of Th1 cells. Thus CD25 cleavage by Per a 10 can hamper Th1 development, favouring Th2 propagation. Previously, Der p 1 along with endogenous proteases like elastases and MMP-9 have been shown to cleave CD25. Th2 cells secrete myriad of Th2 cytokines, of which IL-4 induces IgE class switching in B cells, also patients with atopic diseases show elevated IgE levels. CD23 a low affinity IgE receptor (FcεRII) exists in membrane bound (mCD23) and soluble form (sCD23) and is a key regulator of IgE synthesis. Both mCD23 and sCD23 differentially regulate IgE. Free IgE binds to membrane bound CD23 and down regulates its own production through negative feedback loop. sCD23 binds to IgE, restricting its binding to membrane bound CD23 thus interrupting this feedback mechanism thereby promoting further IgE production. Endogenous proteases like disintegrin/metalloproteinase family members along with exogenous protease allergens (Der p 1) have been implicated in CD23 cleavage. Per a 10 cleaves CD23 and can disrupt the IgE regulation resulting in its increased production. This also explains elevated systemic and local IgE levels in mice sensitized and challenged with Per a 10 as observed in the present as well as other studies. Soluble CD23 levels are elevated in a variety of diseases including rheumatoid arthritis (joints and synovial fluids), Sjögren’s syndrome (plasma and saliva) and in systemic lupus erythematosus (systemic) patients. CD23 is a potential target for therapeutic intervention as it is involved in immune and IgE regulation that makes CD23 cleavage as an active area of research.

Further, our results demonstrate that bronchial epithelial cell activation by Per a 10 induces migration of dendritic cell precursors towards the epithelial layer. Dendritic cells are the sentinels of immune system and play a critical role in initiation and maintenance of allergic Th2 responses. Immature dendritic cells form a dense network below the epithelial cells in the lungs and sense the environment for inhaled particles in conjunction with epithelial cells. There exists an extensive crosstalk between epithelial cells and dendritic cells, which can influence the functioning of dendritic cells. Reports suggest that there is an increase in number of dendritic cells in the lungs of allergic asthmatic patients as compared to normal healthy individuals which increases further

Figure 6. Per a 10 elevates IL-33, TSLP, IL-1α and uric acid levels in mice lungs. Mice were sensitized intranasally with either active or inactive Per a 10 for three days a week for two weeks and euthanized on 15th day. BALF was collected: total RNA was isolated from the left lobe of the lung and expression of TSLP and IL-33 mRNA was assessed by quantitative realtime PCR. mRNA levels of (a) IL-33 and (b) TSLP in the lungs; (c) IL-33 in homogenate (d) TSLP (e) IL-1α and (f) Uric acid levels in the BALF of PBS, active Per a 10 and inactive Per a 10 sensitized mice. Data presented as mean ± SEM of 6 mice per group and are representative of one of the two independent experiments performed. *P < 0.05 as compared to active Per a 10 sensitized mice.
upon allergen challenge and that epithelial cells play a role in dendritic cell precursor migration to the lung epithelial surfaces.86

To assess the role of protease activity and epithelial cytokines in development of Th2 responses, we generated an intranasal Per a 10 sensitized and challenged mice model of allergic airway inflammation. Mice sensitized and challenged with proteolytically active Per a 10 developed allergic airway inflammation symptoms. Epithelial activation is associated with allergen sensitization and is a characteristic of allergic asthma and rhinitis.56 Emerging reports have implicated epithelial cytokines IL-33 and TSLP in allergic reactions.46–48 Our results indicate that Per a 10 activates airway epithelial cells to elevate IL-33 and TSLP levels. We next corroborated our results obtained in mice model of allergic airway inflammation in an in vitro model by exposing BEAS-2B cells with active and inactive Per a 10. BEAS-2B cells stimulated with active Per a 10 induced secretion of IL-33 and TSLP. IL-33 is a member of IL-1 family of cytokines and acts via its receptor ST2 that is expressed by a variety of cells like eosinophils, mast cells, basophils and natural killer cells.49 It is a constitutively expressed cytokine, usually found sequestered in the nucleus50 and is secreted upon epithelial cell activation. Proteases from Alternaria alternata and HDM are reported to induce release of IL-3351,52. Activation of IL-33/ST2 pathway triggers the release of proinflammatory cytokines, chemokines and mediators, induces systemic Th2 responses and contributes to allergic induced airway inflammation and hyper responsiveness53,54. TSLP is another epithelial cytokine implicated in pathogenesis of allergic diseases that directs DCs towards Th2 responses and links epithelial cell activation to dendritic cell mediated immune regulation55. TSLP expression is increased in bronchial mucosa of severe asthmatics.56 TSLP induces OX40L expression on DCs, which is required for polarisation of naïve CD4+ T cells towards TH2.57

Previously, Per a 10 has been shown to induce OX40L expression on DCs suggesting an important role of TSLP in the development of Th2 responses. DC-OX40L axis in initiating and maintaining Th2 responses during allergic inflammation. The protease activity of Per a 10, uric acid and ATP have been implicated in promoting allergic Th2 responses.1 We found elevated levels of uric acid and IL-1α in the BALF of active Per a 10 administered mice as compared to inactive Per a 10 and PBS administered mice. IL-1α levels were also elevated in the supernatant of active Per a 10 exposed BEAS-2B as compared to media and inactive Per a 10 stimulated cells. IL-1α plays an important role in Th2 sensitization to HDM. IL-1α was shown to act upstream of cytokine cascade that leads to activation of epithelial and dendritic cells in response to HDM sensitization.4 Uric acid, a by-product of metabolism, is an alarmin that is secreted by stressed cells.8 Intraperitoneal administration of OVA along with uric acid crystals led to Th2 immune responses like increased eosinophilia, goblet hyperplasia, along with induction of OVA specific IgE, IgG1 and IgG2a.9 Uric acid plays a critical role in peanut sensitization as peanut sensitized mice showed altered purine metabolism with increased serum uric acid levels which were also found to be elevated in peanut allergic children.50 Cysteine proteases like bromelin and papain have been shown to induce release of uric acid in the airway lumen.51 Further, administration of uric acid in the naive mice has been shown to induce IL-33 release and Th2 responses.52 Exposure of bronchial epithelial cells to Per a 10 leads to increased secretion of ATP. ATP is another danger signal that is elevated in BALF of asthmatics and mice models of allergic asthma.53 Previously, we have demonstrated the pivotal role protease activity of Per a 10 plays in the activation of airway epithelial cells and subsequent secretion of pro-inflammatory cytokines.13 Inactive Per a 10 did not show any change in the levels of TSLP, IL-33 and IL-1α, which were elevated after Per a 10 stimulation. The secretion of these cytokines seems to be dependent on the protease activity of Per a 10, the exact mechanism of which remains to be deciphered. It has been reported that the Alternaria alternata exposure leads to elevated ATP levels which in turn results in IL-33 release and is dependent on protease activity.54 The Alternaria alternata induced IL-33 secretion is mediated by ATP via activation of purinergic receptors. Also, the activation of PAR-2 receptors and Calcium ion mobilization were shown to be key features associated with Per a 10 induced epithelial cell activation.55 Jairaman et al., evaluated the underlying mechanism of Ca2+ mobilization after stimulation with cockroach and house dust mite extract and demonstrated that cockroach extract mediates Ca2+ mobilization via PAR-2 receptors and is independent of Ca2+ mobilization induced by purinergic receptor stimulation (P2Y) by UTP.63 In the present study ATP released after Per a 10 stimulation was dependent on intracellular Ca2+ mobilization but the exact pathway responsible for ATP secretion after protease allergen stimulation is still unknown. Based on the results of our study and relevant literature it is now known that there exists a complex interplay between PAR-2 receptors, Ca2+ mobilization, ATP secretion and purinergic receptors in the regulation of various cytokines leading to Th2 responses. A detailed mechanistic study taking into consideration the above mentioned factors will be helpful in unravelling the signaling mechanisms employed by protease allergens in activating airway epithelial cells.

In conclusion, our results demonstrate that Per a 10 enhances epithelial permeability, activates airway epithelial cells, enhances IgE production and induces Th2 responses by the virtue of its proteolytic activity. Our study indicates that apart from genetic predisposition, intrinsic biochemical activity of certain allergens (protease activity) plays a critical role in initiating and promoting allergic airway responses at mucusal surfaces.

Methods

Isolation of Per a 10. Per a 10 was isolated from whole body cockroach extract by affinity chromatography as described elsewhere.4 Per a 10 was cloned using gene specific primers, expressed in E.coli and purified using Ni-NTA affinity chromatography.58 Per a 10 was boiled at 90 °C for 90 minutes to inactivate its protease activity.

Permeability Assay. Calu-3 cells were cultured at air liquid interface in EMEM supplemented with 10% FBS, 100 U/ml penicillin and 100 ug/ml streptomycin at 37 °C in 5% CO2 to analyze the changes in transepithelial permeability.59 Briefly, the cells were cultured in a 24 well transwell plate till formation of monolayer and incubated overnight in serum free EMEM. Cells were stimulated with 10 μg of Per a 10, ΔPer a 10 and rPer a 10. RTIC
labeled dextran 70 S (50 μl, 100 μM) was added at the apical surface of transwell plates. For changes in transepithelial permeability 50 μl of media was collected from the basolateral compartment at different time points and its absorbance at excitation/emission of 530/590 nm was measured. Change in permeability was calculated as fold change with respect to PBS stimulated cells.

**Immunofluorescence staining of junctional proteins.** Calu-3 cells grown on coverslips till confluency were incubated in serum free media for 8 hours and exposed to either Per a 10 or PBS for 16 hours. The coverslips were removed and cells fixed in cold methanol for 10 mins, blocked with 3% (w/v) BSA in PBS. The cells were permeabilized with 0.3% triton X 100 for 10 mins at 4°C and blocked with BSA (10% w/v in PBS) for 1 hour at RT. Cells were stained with mouse monoclonal anti-occludin (Life Technologies, Carlsbad, CA, USA) or rabbit polyclonal anti ZO-1 antibody (Life Technologies, Carlsbad, CA, USA) diluted (1:200) in 3% BSA (w/v in PBS) for 1 hour at RT washed with PBS 3 times and incubated for 2 hrs with Alexaflour 488 labelled secondary antibody (Life Technologies, Carlsbad, CA, USA). These cover slips were mounted on slides and analyzed using fluorescent microscopy (Leica DMI 6000).

**Monocyte isolation, generation of MDDCs and co-culture experiments.** Monocytes were isolated from the PBMCs of cockroach hypersensitive patients, and MDDCs generated as described previously. The monocytes cultured for five days in these cytokines were characterised for the expression of CD11c and was used in migration studies.

For transwell migration assays BEAS-2B cells were cultured in the lower chamber of 24 well transwell plate till 70–80% confluency. The cells were starved in epithelial cell basal media overnight exposed to 10 μg of active Per a 10 and the MDDC precursors (1 × 10^6) were added to the upper chamber and incubated for 24 hours in serum free RPMI. After dissociating the BEAS-2B cells with a cell dissociating medium the cells were stained with PE labelled anti-human CD11c antibody or an isotype control and acquired using BD FACS calibur and analyzed by Cell Quest Pro.

**B cell and CD4+ T cell isolation.** B cells and CD4+ T cells were purified from PBMCs by using MACS B-CLL isolation kit and human CD4+ isolation kit II (Miltenyi Biotec, Gladbach, Germany) following manufacturers instruction.

**Stimulation of cells and staining for flow cytometry.** 1 × 10^6 Cells (PBMCs, B cells or CD4+ T cells) were incubated with Per a 10, rPer a 10 and Δ Per a 10 for 4 hours. Cells were washed with FACS buffer. The cells were stained with FITC labelled anti-CD23 (Life Technologies, Carlsbad, CA, USA) and co-cultured with 10 μg of active Per a 10 and bioluminescence was measured as per the manufacturer’s instructions. The concentration of ATP was calculated from a standard graph prepared using known ATP concentrations.

**Measurement of Per a 10 specific IgE in the sera of mice.** Relative levels of Per a 10 specific IgE were measured in the serum samples of mice by indirect ELISA. Briefly, microtiter plates were coated with 100 ng of Per a 10/well in 100 μl of 0.1 M carbonate buffer (pH 9.6) and incubated overnight at 4°C. After washing with PBS the plates were blocked with 3% defatted milk for 3 h at 37°C. Sera samples were diluted 1:10 in PBS and 100 μl of the diluted sera was added in triplicates. The plates were incubated at 4°C overnight. After washing with PBST (0.05% Tween-20 in PBS) to remove the unbound antibodies the plate was incubated with biotinylated anti-mouse IgE (2 μg/ml) at 25°C for 90 min followed by streptavidin-peroxidase (1:1000) for 30 min. The plate was washed and was developed using o-phenylenediamine and absorbance was read at 492 nm.

**Cell culture and stimulation with Per a 10.** BEAS-2B cells were cultured in BEGM (Invitrogen) till confluency at 37°C and 5% CO2. Cells were then incubated in BEBM (without growth factors) overnight and exposed to 10 μg of active (Per a 10) or inactive (Δ Per a 10 and rPer a 10) Per a 10. Supernatant was collected and measured in the serum samples of mice by indirect ELISA. Briefly, microtitre plates were coated with 100 ng of active (Per a 10) or inactive (Δ Per a 10 and rPer a 10) Per a 10. Sera samples were diluted 1:10 in PBS the plates were blocked with 3% defatted milk for 3 h at 37°C. Sera samples were diluted 1:10 in PBS and 100 μl of the diluted sera was added in triplicates. The plates were incubated at 4°C overnight. After washing with PBST (0.05% Tween-20 in PBS) to remove the unbound antibodies the plate was incubated with biotinylated anti-mouse IgE (2 μg/ml) at 25°C for 90 min followed by streptavidin-peroxidase (1:1000) for 30 min. The plate was washed and was developed using o-phenylenediamine and absorbance was read at 492 nm.

**ATP assays.** ATP release was assayed by using a luciferase/luciferin bioluminescence ATP determination kit (Invitrogen/Life technologies) and a luminometer. Cells were grown in white 96 well plates till confluency. The cells were exposed to 2 μg Per a 10 and bioluminescence was measured in real time as per the manufacturer’s instructions. The concentration of ATP was calculated from a standard graph prepared using known ATP concentrations.
Quantitative PCR. Total RNA was isolated from lung tissue by homogenizing in 2 ml of Trizol (Life technologies) and quantified using nanodrop. cDNA preparation and quantitative PCR was carried out as previously described24. The primers used were as follows: IL-33: forward 5′GCTCGCGTCTGGTGACACATT-3′ and reverse 5′CTCTCGATTTGCTCGAAGCTT-3′; TSLP: forward 5′GCGATGGGGCTAACTTACA-3′ and reverse 5′CTCTCGATTTGCTCGAAGCTT-3′; β-actin: forward 5′-GCGTTCGATGGCCAGGCTCTT-3′ and reverse 5′-GCGTACACTTCCATGATGGAATTGAG-3′.

Uric acid determination. The concentration of uric acid in the BALF of mice was determined by Amplex red uric acid/uricase kit. Briefly, 50 μl of samples were pipetted in to a 96 well plate and a 50 μl of working solution of Amplex Red reagent containing 0.4 U/ml HRP and 0.4 U/ml uricase was added to them. A standard curve ranging from 0–100 μM was prepared from the stock of 5 mM uric acid in 1X reaction buffer. The plate was incubated at 37 °C in dark for 30 minutes and absorbance was measured at 560 nm using a Benchmark plus microplate reader.

ELISA. The concentrations of cytokines hIL-33, hIL-1α, hCCL-2 (R&D systems), hTSLP, mIL-33, mTSLP, mIL-4 and mIL-1α (eBiosciences) in the cell supernatant, Lung homogenate and in the BALF were estimated using paired antibodies following manufacturer’s instructions.

Statistical analysis. Statistical analysis of results was done by using GraphPad Prism software (GraphPad Software, San Diego, CA, USA). The statistically significant difference was determined using one way ANOVA and p value < 0.05 was considered as statistically significant.

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