Innate function of house dust mite allergens: robust enzymatic degradation of extracellular matrix at elevated pH

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Abstract

Background: Exposure to the house dust mite Dermatophagoides pteronyssinus (D.p.) increases the risk for developing allergic diseases in humans and their best friend, the dogs. Here, we explored whether this allergenic mite via its enzymes may impact the cutaneous extracellular matrix (ECM), which critically determines epithelial barrier integrity both structurally and functionally.

Methods: Two extracts obtained from either dust-purified or cultured D.p. bodies were used in the present study. To assess the potential impact of D.p. on protein components of the ECM, proteolytic activity of the D.p. extracts were determined by casein and gelatin gel zymography, and their N-acetyl-β-hexosaminidase activity determined colorimetrically. In addition, IgE-dependent and innate degranulation potential of D.p. was examined in canine MPT-1 mast cells and neurite outgrowth assay using rat pheochromocytoma PC-12 cells.

Results: In gel zymography, both extracts digested the substrates casein and gelatin in a dose-dependent manner, especially at alkaline pH, and effective in a wide range of temperatures (30 °C−42 °C). In particular, a 25-kDa band corresponding to Der p 1, the major D.p. allergen for humans, was found enzymatically active in both casein and gelatin gels regardless of the presence of metal ions and of alkaline conditions. Besides protease activity, N-acetyl-β-hexosaminidase activity was detected in both extracts, suggesting that D.p. affects the cutaneous ECM through deteriorating both proteins and glycosaminoglycans. While both D.p. extracts induced IgE-dependent mast cell degranulation, much less innate effects on mast- and neuronal cells were observed.

Conclusions: Our data highlight that D.p. is a robust source of several distinct enzymes with protease- and N-acetyl-β-hexosaminidase activities. In alkaline milieu they can degrade components of the ECM. Therefore, D.p. may contribute to epithelial barrier disruption especially when the skin surface pH is elevated.

Keywords: Dermatophagoides pteronyssinus, Allergens, Epithelial barrier, Extracellular matrix, Proteases, N-acetyl-β-hexosaminidase

Background

Dermatophagoides pteronyssinus (D.p.) represents one of the most frequently encountered house dust mite (HDM) species worldwide [1]. Exposure to D.p. allergens raises the risk of developing allergic diseases, such as asthma [2] or atopic dermatitis (AD) [3]. Especially, those allergens categorized as group 1 (Der p 1 in D.p.; Der f 1 in Dermatophagoides farinae, D.f.), potently induce allergic responses, and also break tight junctions between airway epithelial cells by cysteine protease activity, and thereby alter the permeability [4]. In accordance,
papain, a structural homolog of group 1 allergens, affects tight junction permeability of the *epidermis* [5], and activates mast cells (MCs) [6].

This seems important as epithelial barrier dysfunction, and even more -disruption, is accepted as the initial event before cutaneous sensitization occurs [7]. In line with the “outside-inside-outside” concept a defect in tissue-related genes, as the example of loss-of-function Filaggrin alleles in AD [8], was proposed to support hypersensitivity. In addition to the genetic predisposition, environmental factors play a crucial role and recent studies indicate that a rise in skin surface pH might be a major determinant for a barrier defect [9, 10]. The enzymatic activity of Der p 1 as outlined above is an example of an environmental factor; however, dogs, which share common immune principles [11] and milieu with humans [12], are less likely to react to group 1 allergens, in spite of the fact that HDMs are a major source of environmental allergens for dogs as well as for humans [13]. Accordingly, the mechanisms by which D.p. frequently cause hypersensitivity in humans and animals, seem more than the action of a single enzyme.

The cutaneous extracellular matrix (ECM) consists mainly of proteins associated with glycosaminoglycans (GAGs) and glycoproteins. The architecture and composition of these components determines the biophysical properties of the skin, such as stiffness, compliance and resilience [14]. The basement membrane, representing a thin but firm meshwork of collagen and laminin, underlies the basal layer of the epidermis and thereby shields the host from the outside environment [15]. The body’s own matrix metalloproteases (MMPs) are responsible for the tissue remodeling and ECM degradation. However, several invading pathogens produce ECM-degrading enzymes or may alter host-derived proteolytic activity [16].

In addition to structural defects, ECM breakdown leads to an alteration in behavior of multiple cell types in the skin. Collagen fragmentation, which is frequently associated with aged [17] or photo damaged skin [18], results in dysfunction of keratinocytes or fibroblasts. Cleavage products of hyaluronan (HA), known as a major GAG in ECM, serve as signals of tissue injury, which enhance inflammatory cytokine production and differentially modulate function of macrophages and dendritic cells through pattern recognition receptors [19]. Therefore, exposure to foreign enzymes with potential for ECM deterioration is a likely cause of epidermal barrier defects.

In this study, we aimed to determine in vitro whether and under which conditions, D.p. allergens have the potency to affect the cutaneous ECM and thereby initiate skin barrier disruption. Further, to verify whether such potency is compatible with allergenic activity, we examined the innate effect of D.p. allergens on MCs ex vivo, and the specific effect using canine serum obtained from dogs with canine atopic dermatitis (CAD) which is clinically and pathologically similar to the human disease [20].

**Methods**

**Extracts from mite bodies**

The whole mite bodies: the purified D.p. from dust samples 4966 (Allergon, Aengelholm, Sweden) for extract I; the cultured D.p. LG-8444 (Cosmo Bio LSL, Tokyo, Japan) for extract II, were homogenized in sterile phosphate-buffered saline with 0.1 mM PMSE, a serine protease inhibitor. This supernatant was collected by centrifugation. Proteins were quantified by the method of Bradford.

**Reagents**

Polyacrylamide gels were prepared using 30% acrylamide solution (AppliChem, Darmstadt, Germany). In zymography, gels were copolymerized casein (C3400) or gelatin (G2500) (both from Sigma-Aldrich, St. Luis, MO) as a protein substrate. In a colorimetric measurement of N-acetyl-β-hexosaminidase (β-HEX) activity, its chromogenic substrate 4-nitrophenyl N-acetyl-β-D-glucosaminide (NP-GlcNAc) (Sigma-Aldrich) was applied. For the induction of MC degranulation, dog IgE P115 was purchased from Bethyl Laboratories (Montgomery, TX), anti-dog anti-IgE antibody SM1498P from Acris Antibodies (Herford, Germany), and Ca-ionophore A23187 from Sigma-Aldrich. For Western blot analysis, horseradish peroxidase-conjugated goat anti-dog anti-IgE antibody NB7346 was purchased from Novus Biologicals (Littleton, CO). In neurite outgrowth bioassay, recombinant mouse β-nerve growth factor (NGF) was purchased from R&D Systems (Minneapolis, MN). Unless otherwise indicated, all chemicals used in this study were obtained from Sigma-Aldrich.

**Zymography**

D.p. extracts or recombinants were diluted with 125 mM Tris-HCl (pH 6.8) and the following 2 × sample buffer (20% glycerol, 4% sodium dodecyl sulfate: SDS, 125 mM Tris-HCl, 0.01% bromophenol blue, pH 6.8) with neither 2-mercaptoethanol nor boiling. Samples were separated at 4 °C in 10% SDS-polyacrylamide gel electrophoresis (SDS-PAGE) copolymerized with 0.1% casein or gelatin. Migrated enzymes were renatured from SDS by soaking in 2.5% Triton X-100 for 30 min twice, and then activated in 50 mM Tris-HCl, pH 8.0, for casein; 50 mM Tris-HCl, 5 mM CaCl₂, 1 mM ZnCl₂, pH 7.6, for gelatin, at 37 °C for 16 h. To chelate metal ions, after separation and after SDS removal, gelatin gels were incubated in the reaction buffer supplemented with 50 mM EDTA for Ca²⁺ or 10 mM 1,10-phenanthroline.
The membrane was blocked by 1% PAGE and transferred onto PVDF membrane (Millipore, Bedford, MA). The optimal pH and temperature were evaluated by changing the buffer conditions during digestion. Thermal susceptibility was determined by heating extracts (10 μg) prior to an assay. Gels were stained with 0.2% Coomassie Brilliant Blue (CBB) R-250 solution. Digestion bands were analyzed using the Java-based image processing program ImageJ (The National Institutes of Health, Bethesda, MD).

**Recombinant allergens**

The recombinant D.p. allergens used in our experiments are described in Additional file 1: Table S1. In brief, allergen proteins were expressed in *Escherichia coli* and purified to homogeneity. The purified proteins were analysed by SDS-PAGE with CBB or silver staining. For SDS-PAGE, recombinants were diluted with 2 × sample buffer containing 10% 2-mercaptoethanol, and then boiled at 95 °C for 7 min. Zymography was performed as mentioned above. A specification of the recombinants is shown in Additional file 1: Table S1.

**Measurement of β-HEX activity**

To determine β-HEX activity, various concentrations of D.p. extracts (0.1–50 μg/ml) or recombinants (25 μg/ml) were incubated at 37 °C for 1 h with 3 mM NP-GlcNAc in 100 mM citrate buffer, pH 4.5. The enzymatic reaction was stopped by 200 mM glycine, pH 10.7. The accumulation of degrading products was measured by the absorbance at 405 nm using the Infinite M200 PRO plate reader (TECAN, Maennedorf, Switzerland). Thermal lability was determined by heating extracts (50 μg/ml) prior to a colorimetric assay and the relative β-HEX activity was calculated as a ratio of the optical density (OD) value without heating. All experiments were performed in duplicate.

**Canine sera**

Sera were obtained from 26 canine patients diagnosed with atopic dermatitis (CAD according to ICADA guidelines [21]), in the Dermatology Department of the University of Veterinary Medicine Vienna (Vienna, Austria). In line with the routine diagnostic procedures, all atopic dogs underwent an intradermal skin test. The total IgE levels and the serological reactivity to D.p. were assessed in all specimens. Those that specifically reacted to D.p. allergens were used in this study (Additional file 1: Table S2).

**Western blot analysis**

Recombinant allergens were diluted with 2 × reducing sample buffer and then boiled at 95 °C for 7 min. Samples, 168.8 ng/lane for recombinant (r) Der p 15; 143.8 ng/lane for rDer p 18, were separated in 10% SDS-PAGE and transferred onto PVDF membrane (Millipore, Bedford, MA). The membrane was blocked by 1% bovine serum albumin/Tris-buffered saline, 0.05% Tween-20 and incubated overnight at 4 °C with diluted canine sera (the final IgE concentration: 30–100 ng/ml). Allergen-specific IgE bonds were detected with horse-radish peroxidase-labeled anti-IgE.

**MC degranulation assay**

MC degranulation assay was performed by measurement of β-HEX release in the canine MC line MPT-1, which harbors the high-affinity IgE receptor FcεRI, essentially as described [22], using anti-IgE (5 μg/ml) or A23187 (5–50 μM). Cells were cultured in α-MEM with 10% fetal bovine serum (FBS) (both from Life Technologies, Carlsbad, CA) and antibiotics. Cells were preloaded with serum IgE from canine patients allergic to D.p., at 37 °C for 2 h. The optimal IgE concentration had been determined by titrating each serum in pre-experiments to 5–10 μg/ml IgE. Then preloaded cells were challenged (37 °C, 1 h) with D.p. extracts (50 μg/ml). Following incubation, cell-free conditioned medium was examined for β-HEX content using NP-GlcNAc as mentioned earlier. To eliminate the direct effect of extracts on the substrate, the OD values obtained from those cells which were incubated at 4 °C with D.p. extracts (50 μg/ml), were subtracted. All experiments were performed in triplicate. The degranulation rate was expressed as present of total content by the following formula: degranulation rate (%) = (OD<sub>supernatant</sub>/(OD<sub>supernatant</sub> + OD<sub>lysate</sub>)) × 100.

**Neurite outgrowth bioassay**

Neurite outgrowth experiments were performed in the rat pheochromocytoma PC-12 (Japanese Collection of Research Bioresources Cell Bank, Osaka, Japan) using NGF (50 ng/ml). Cells were maintained in RPMI1640 (Life Technologies) with 5% FBS, 10% horse serum (Thermo Fisher Scientific, Waltham, MA) and antibiotics. In bioassays, following NGF priming for 7 days, cells were treated with D.p. extracts (100 μg/ml) for 48 h in a collagen-coated dish. In control experiments, cells were incubated with or without NGF. A percentage of neurite-bearing cells, which extended neurites longer than their cell body, was calculated from random 5 fields under microscopy.

**Statistics**

Statistical examinations were performed using SigmaPlot® 11 (Systat Software, San Jose, CA). All data represent the mean ± standard error of the mean (SEM). p value of <0.05 was considered statistically significance in difference (*, p < 0.05; **, p < 0.01; ***, p < 0.001).
Results
Protease activity in D.p. extracts
The proteolytic activity repertoire of D.p. was examined by gel zymography. This technology is based on non-reducing SDS-PAGE copolymerized substrate proteins (i.e., casein or gelatin). Following electrophoretic separation of the enzyme of interest, protease activity can be visualized by CBB staining as a digestion band at the molecular weight (MW) of the enzyme. To evaluate the potential impact of any contaminating commensal factors, 2 different D.p. extracts of distinct origin were analyzed: D.p. extract I from dust-purified mites, and extract II from cultured mites. In fact SDS-PAGE depicted slight differences in their protein composition (Additional file 1: Figure S1). Zymography showed dose-dependent protein digestion, albeit with distinct patterns of MW (Fig. 1a, b and Additional file 1: Figure S2a, b).

MMPs belong to a family of Ca^{2+}-dependent, Zn^{2+}-containing endopeptidases [23] whose homologs are found in genomes of both vertebrates and invertebrates [24]. Some of these enzymes potently lyse gelatin; to clarify their involvement in gelatinolysis, catalytic dependencies on the metal ions were determined by use of chelators: EDTA, for Ca^{2+}; Phen, for Zn^{2+}. In D.p. extract I, the 70- and 100-kDa bands were markedly inhibited by both agents (Fig. 1c). The digestion of the 40-kDa band was partially diminished by EDTA, but not affected by Phen (Fig. 1c). Similarly, gelatinolysis with high MW detected in extract II were effectively inhibited by the chelators (Additional file 1: Figure S2c). In contrast, the approximately 25-kDa band, which was detected in casein and gelatin gels using both extracts, was significantly suppressed by Phen, but not by EDTA (Fig. 1c and Additional file 1: Figure S2c).

Optimal conditions for D.p. protease activity
The optimal pH and temperature for the catalytic activity were determined with a change of buffer conditions. After electrophoretic separation and after SDS removal, gels were incubated 37 °C at pH 5.0 or pH 9.0. In both extracts, casein digestion hardly occurred at pH 5.0, while it was dramatically enhanced at pH 9.0 (Fig. 2a and Additional file 1: Figure S3a). Although gelatin digestion remained predominant at pH 5.0 towards low MW compounds of extract I, elevation in pH again intensified protease activity (Fig. 2b and Additional file 1: Figure S3b). These results, together with data in Fig. 1 and Additional file 1: Figure S2, indicate that most of these proteases are activated optimally at neutral to alkaline pH (i.e., pH 8.0, for casein; pH 7.6, for gelatin). On the other hand, protein digestion capacity of the D.p. extracts was only prevented by incubation at 4 °C, but operative over a wide range of temperatures from 30 °C to 42 °C (Fig. 3a).

Protease activity of recombinant allergens
The thermal susceptibility and tolerance of D.p. protease activity was evaluated by heating the extracts, prior to enzymatic analysis, at a series of temperatures from 40 °C to 60 °C for 30 min, which are common temperatures for washing clothes. Heating at 56 °C for 30 min had seemingly no effects on gelatin digestion by both extracts (Fig. 3b), neither 40 °C, 45 °C nor 50 °C (data not shown). Heating at 60 °C for 30 min only partially diminished gel digestion (Fig. 3b). However, when the temperature reached 100 °C, neither of extracts degraded gelatin gels at
a detectable level (Fig. 3c). Overall, there were no marked differences in heat susceptibility between the proteases.

Der p 1 is a 25-kDa allergen that is a homolog to cysteine proteases [25]. Its conformational epitopes are readily denatured at 75 °C for 1 h or 100 °C for 10 min [26]. It has been also reported that this allergen is irreversibly denatured in acidic pH (pH 2.0) but resistant to mild alkaline conditions (pH 10.0) [26]. Accordingly, Der p 1 is supposed to be responsible for the digestion band appearing at 25 kDa, and purified rDer p 1 was applied to verify its activity upon the substrates. SDS-PAGE showed a protein migrating at approximately 25 kDa under reducing conditions (Fig. 4a). Zymography showed robust degradation by rDer p 1 on both gels, albeit with seemingly different mobility, most likely due to dimerization of recombinants (Fig. 4b). However, the other allergens within the range of recombinants available to us for this study (Additional file 1: Table S1) had no effect on casein and gelatin gels (Additional file 1: Figure S4a, b).

β-HEX activity in D.p. extracts

The hydrolytic activity towards GAG components is of specific relevance for chitinase allergens, and was examined with focus on β-HEX (the Enzyme Commission number: EC 3.2.1.52), now including N-acetyl-β-glucosaminidase (EC 3.2.1.30). This enzyme is known as a lysosomal enzyme that hydrolyzes glycosidic linkages between N-acetyl-β-glucosamine (GlcNAc) and glucuronate. Hence, β-HEX activity in D.p. extracts was colorimetrically measured using its chromogenic substrate NP-GlcNAc. Both extracts induced a dose-dependent increase in the OD at 405 nm, indicating accumulation.
tested samples reacted via IgE to one or both allergens Der p 15 and Der p 18. Twenty four sera of 26 specific IgE, as revealed by testing on blotted canine major allergens from pet dogs suffering from CAD containing D.p.-specific IgE were determined by the effect of D.p. extracts on MCs. Sera basophils [29]. Thus, allergenic activity of D.p. was demonstrated not only by the increased OD values, but also by increased mediator release from cells preloaded with canine sera. Results were evaluated by one-way ANOVA with Holm-Sidack tests. Error bars, ± SEM (n = 3); n.s., no significance in difference; D.p.; extract I; D.p.II, extract II; Ctrl, buffer alone.

Chitinase (EC 3.2.1.14) hydrolyzes glycosidic bonds between GlicNAcs of chitin, known as the most abundant acid which can be used as an initiator in various biological processes. Interestingly, IgE reacts to group 15 (chitinases) and 18 (chitinase-like proteins) of mite allergens at high frequencies in dogs [27]. Therefore, any potential actions of D.p. on nerve elongation were determined by neurite outgrowth bioassay using PC-12 cells. While NGF in doses-dependently triggered by canine IgE-FcεRI cross-linkage, by A23187, or canine sera and extracts. Results were evaluated by one-way ANOVA with Holm-Sidack, and by Ca-ionophore A23187 (Fig. 6c). Hence, MPT-1 cells were adapted to a novel ex vivo bioassay using the canine sera. As shown in Fig. 6d, mediator release from cells preloaded with canine D.p.-IgE could be triggered by both D.p. extracts, but none of the extracts had any effect in the absence of specific IgE (data not shown); taking into account specific IgE levels and skin test positivity to D.p., 6 serum samples were selected for further analysis (Fig. 6a and Additional file 1: Table S2). Degranulation in canine MPT-1 cells could be dose-dependently triggered by canine IgE-FcɛRI cross-linking (Fig. 6b) comparable with that induced by human antibodies [22], and also by Ca-ionophore A23187 (Fig. 6c). Hence, MPT-1 cells were adapted to a novel ex vivo bioassay using the canine sera. As shown in Fig. 6d, mediator release from cells preloaded with canine D.p.-IgE could be triggered by both D.p. extracts, but none of the extracts had any effect in the absence of specific IgE (data not shown). Intractable pruritus in AD is resistant to conventional treatments such as antihistamines, suggesting contributing factors other than MC mediators. We hypothesized that the increased nerve density histologically found in AD lesions might be responsible for the persistent itch sensation [30]. Therefore, any potential actions of D.p. extracts on nerve elongation were determined by neurite outgrowth bioassay using PC-12 cells. While NGF in
approximately 40% of cells induced neurite outgrowth (Fig. 4e), neither of the D.p. extracts had any effect on the PC-12 cells (Fig. 4e and Additional file 1: Figure S5).

**Discussion**

The present study demonstrates that D.p. has a potent enzymatic impact on the ECM. This may lead to epithelial barrier defects, thereby explaining the high prevalence of cutaneous hypersensitivity to D.p. in humans and animals.

By zymography casein and gelatin digestion capacity could be demonstrated in extracts obtained from both dust-purified or cultured mites. This method has long been used for studies of ECM-degrading enzymes particularly of MMPs with gelatin, the denatured form of collagen [23]. Further, casein is favorably applied to readout of a wide range of serine and cysteine proteases, such as kallikreins [9], in the absence of metal ions. Accordingly, the in vitro gels using purified proteins (i.e., casein and gelatin) represent an experimental approximation to the in vivo situations. Gelatin digestion at high molecular mass bands in the gels which was diminished by both chelators, might be attributed in part to enzymes classified into the MMP family. In contrast, the chelators had much less effect on digestion of bands at a low molecular mass. These findings indicate that D.p. extracts contain several different kinds of enzymes that can degrade protein components of the cutaneous ECM.

The D.p. proteases were optimally active at neutral to alkaline pH. Therefore, the acidic mantle of healthy skin (i.e., at pH 4.5–5.5 in humans) may not only prevent the overgrowth of indigenous pathobionts [31], but also prevent the activation of D.p. proteases. A rise in skin surface pH has been related to skin defects, such as AD [32]. Further, a more recent study indicates that alkalization of skin surface induces abnormal activation of endogenous serine proteases and thereby disrupts the skin barrier function, even before the onset of AD [10]. The temperature of the skin surface is generally lower than the core body at 37 °C, thereby controlling surface-attached pathogens such as Staphylococcus aureus [33]; the fact that casein and gelatin digestion by the D.p. proteases occurred at temperatures lower than 37 °C suggests that they are active at the skin surface. Collectively, these findings raise the possibility that the enzymatic activity of D.p. proteases could be enhanced when the pH of the host skin surface is elevated leading to barrier damage.

This is the first demonstration, to our knowledge, that D.p. extracts exhibit β-HEX activity. β-HEX acts on glucoside- and galactoside- types of GAGs, and functionally overlaps with hyaluronidases, known as HA catabolic enzymes [34], thereby acting as spreading factor in some venom [35]. Most notably, although almost all D.p. allergens are proteins, D.p. extracts induced degranulation of MCs in an IgE-dependent manner, indicating that they possess allergenic potency in parallel to the enzymatic functions. This seems crucially important, given that defected epithelial barrier facilitates secondary sensitization to allergens [7]. In contrast with our results concerning the innate effects on MCs, it has been reported that papain activates human cord blood-derived MCs IgE-independently via protease-activated receptor-2 and its enzymatic activity [6]. This contradiction may be due to low abundance of Der p 1 in the extracts or decreased sensitivity of the canine cells. Collectively, although further investigations are needed, these data support the hypothesis that D.p. may enzymatically inflict damage on protein and also GAG components of the ECM, and subsequently induce allergic responses through an IgE-mediated mechanism.

Finally, commensal organisms of D.p., being determined by various environmental and climatic conditions of mite habitats, may contaminate dust samples and mite cultures. In particular, Aspergillus species, known as ubiquitous saprophytic fungi, are favorable food to D.p. [36], which often generate enzymes to degrade mammalian tissue for nutrient acquisition and invasion [37]. Therefore, to this end we cannot entirely exclude the possible involvement of such microbial symbionts in the observed enzymatic functions, which may explain the remarkable differences between extract I and II from two different mite breed sources, especially in gelatinolysis under acidic pH, and in the thermal susceptibility of β-HEX.

**Conclusions**

The present study highlights the crucial role of D.p. carrying a variety of enzymes with a potential for degrading ECM and acting as allergens. Given that activation of a large proportion of D.p. proteases is abrogated by acidic pH of skin surface, elevated skin pH can be a critical host factor required for really developing disease following D.p. exposure. This novel finding highlights that skin pH regulation is an important strategy for prevention and treatment in this type of dermatitis. In conclusion, our findings demonstrate that enzymatic potency of D.p. in alkaline conditions damages the cutaneous ECM and therefore weakens the host defense in humans and their best friend, the dog.

**Additional file**

Additional file 1: Appendix. Methods. Figure S1. Distinct band pattern of proteins in D.p. extracts. Figure S2. Proteolytic activity of D.p. extract II. Figure S3. The pH optimum of proteolysis by D.p. extract II. Figure S4. Proteolytic activity of recombinant D.p. allergens. Figure S5. Innate effects of D.p. extracts on neurite outgrowth. Table S1. Specification of
Table S2. Specification of canine subjects with CAD. (PPTX 13.0 mb)

**Abbreviations**
- AD: Atopic dermatitis
- ANOVA: Analysis of variance
- CAD: Canine atopic dermatitis
- CBB: Coomassie Brilliant Blue
- D.F.: Dermatophagoides farinae
- ECM: Extracellular matrix
- FBS: Fetal bovine serum
- Fc\(\varepsilon\):
- M.H.: Matrix metalloproteinase
- MW: Molecular weight
- NGF: Nerve growth factor
- N-P-GlcNAc: N-acetyl-β-D-glucosaminide
- OD: Optical density
- Phen: 1,10-phenanthroline
- r: Recombinant
- SDS: Sodium dodecyl sulfate
- SDS-PAGE: SDS-polyacrylamide gel electrophoresis
- SEM: Standard error of the mean

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**Availability of data and material**
Please contact author for data requests.

**Authors’ contributions**
KO and LE performed experiments, analyzed the data, and wrote the original draft; H and LP provided canine serum samples and were responsible for clinical data; YR and SV provided the D.p. extract and recombinant allergens and helped in manuscript writing; GH contributed to cell-based assays and helped in manuscript writing; HM provided MPT-1 cells and knowhow, and helped in manuscript writing; AT and HM contributed to the data analysis and helped in finalizing the manuscript; EJH and KO designed the study and wrote the final paper. All authors read and approved the final manuscript.

**Competing interests**
The authors state no competing interest.

**Consent for publication**
Not applicable.

**Ethics approval and consent to participate**
This study was approved by the institutional ethics committee of the University of Veterinary Medicine Vienna (Vienna, Austria) in accordance with Good Scientific Practice and EU and Japanese national legislation.

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