An international seroprevalence survey of the IgE sensitisation to the *Dermatophagoides farinae* house dust mite and two of its major allergens (Der f 2, Zen 1) in atopic dogs

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**Background** – Dogs with atopic dermatitis are often immunoglobulin (Ig)E-sensitised to *Dermatophagoides farinae* (Df) house dust mites, yet limited data exist on the sensitisation rates to the individual Df allergens, Der f 2 and Zen 1.

**Objectives** – To determine the IgE sensitisation rates to Df, Der f 2 and Zen 1 in atopic dogs from geographically diverse countries.

**Animals** – Serum was collected from 32 laboratory dogs in Japan, and 837 atopic dogs from 11 countries from five continents: Asia (Japan, Thailand, Taiwan), Europe (Italy, Latvia, the Netherlands, UK), North America (USA), South America (Argentina, Brazil) and Africa (South Africa).

**Methods and materials** – We determined Df-, Der f 2- and Zen 1-specific IgE levels by ELISA. Correlations between the IgE values for these three allergens were calculated.

**Results** – The IgE seropositivity rates for Df varied between 74% (Argentina) and 100% (the Netherlands, Thailand, South Africa), those for Der f 2 between 12% (Argentina) and 88% (South Africa), and for Zen 1 between 70% (Argentina) and 100% (the Netherlands). Apart from the especially low seropositivity rate for Der f 2-specific IgE in Argentina, the percentage of IgE sensitisation varied little between countries. There was significant correlation between the IgE levels to these three allergens which was highest between Df and Zen 1, and lowest between Zen 1 and Der f 2.

**Conclusions and clinical relevance** – The IgE sensitisation to Df is geographically widespread. Der f 2 and Zen 1 are major allergens for dogs in almost all countries where this was evaluated.

**Introduction**

Atopic dermatitis (AD) is a genetically predisposed, pruritic and inflammatory skin disease often associated with immunoglobulin (Ig)E directed against environmental allergens. As IgE antibodies are thought to play a major role in the pathogenesis of AD, it is important to precisely identify disease-exacerbating allergens targeted by these reaginic antibodies. In dogs, house dust mites (HDM) allergens, especially those from *Dermatophagoides farinae* (Df) are considered to be relevant to AD because of the high percentage of positive reactions to intradermal tests with Df, as well as the high level of serum Df-specific IgE in atopic dogs. Additional arguments in support of the relevance of these HDM allergens for canine AD is that they are present in most households with atopic dogs, and treatment of the dog’s environment with acaricides improves the clinical signs in some dogs sensitised to Df. Furthermore, the isolation of atopic dogs in cleaned cages in a veterinary hospital for two weeks results in a marked clinical improvement in dogs with detectable IgE against allergens (mainly Df) from the environment. Importantly, the long-documented skin reactivity and highly specific IgE to Df in healthy dogs, confirms that the mere sensitisation to Df does not always correlate with the presence of atopic signs.
As of early 2021, there were 36 DF proteins considered as allergens for humans (www.allergen.org; page last accessed April 7, 2021). In dogs, the first characterised DF major allergens – that is those recognised by >50% of sera from patients sensitised to the parent extract – were the high-molecular weight (HMWW) Der f 15 and Der f 18 chitinases.9,10 The low-molecular weight (LMWW) Der f 2 was initially shown to represent a major DF allergen for dogs in Japan,11,12 and a similar status was later confirmed in Spain, the UK and Malaysia.13,16 More recently, Zen 1, another HMW allergen of yet-unknown function, was also determined to be a major DF allergen in atopic dogs from multiple countries.13,15,16 Among these four allergens, only Der f 2 has been confirmed to be of clinical importance, as two studies established the high treatment success rate of a commercially available Der f 2 mono-allergen immunotherapy (Allermune HDM, Zenoaq; Koriyama City, Fukushima prefecture, Japan) in dogs with AD.17,18

The goal of this study was to expand the knowledge on the seroprevalence of IgE specific to DF and two of its allergens (Der f 2 and Zen 1) in countries spanning multiple continents. Such a survey is important in preparation for the development of immunotherapy with specific molecular allergens. The data presented herein will clearly establish Der f 2 and Zen 1 as major DF allergens for dogs, worldwide.

Methods and materials

Study population

We assembled sera from 837 dogs in 11 countries: Argentina (50 dogs), Brazil (181), Italy (50), Japan (102), Latvia (30), the Netherlands (52), South Africa (42), Taiwan (100), Thailand (56), the UK (59) and the USA (115). Dogs were selected irrespective of their breed, sex and age, by veterinarians specialised in dermatology. The diagnosis of AD was made after exclusion of ectoparasitic and skin infections, following widely accepted guidelines.18 Care was taken not to enrol atopic dogs whose signs fully responded to an elimination diet. To harmonise selection, dogs should have fulfilled at least five of eight criteria for the diagnosis of AD.20

For the establishment of enzyme-linked immunosorbent assay (ELISA) positive thresholds, we selected 32 healthy, adult laboratory beagles (19 females, 13 males; mean age 24 months), living in a climate-controlled, mite-poor, helminth-free facility in Japan. This cohort was chosen as a way to obtain a “DF-non-sensitised” nonatopic control group owing to its limited exposure to HDMs. The rearing environment was tested beforehand to be negative for DF allergens (Mitye Checker, Sumika Environmental Science; Osaka, Japan).

For all dogs, blood was collected with the owners’ consent and after approbation of the local animal care and use committee, where applicable and necessary.

Sample collection and handling

After collection, sera were separated by centrifugation and kept frozen in standard freezers immediately thereafter. Allergen-specific IgE levels were measured by ELISA as described below. To reduce inter-operator fluctuations between measurements of samples from different geographical regions and different collection times, all samples were sent to a central research laboratory in Japan (Zenoaq) by courier at −20°C for IgE levels to be determined using the same method and operator.

ELISA for allergen-specific IgE levels

ELISA for the determination of serum levels of DF, Der f 2- and Zen 1-specific IgE antibodies were carried out as described previously with minor modifications.12,13 Briefly, lyophilised recombinant Der f 2, native Zen 1 and a whole (crude) extract of DF (Greer Laboratories; Lenoir, NC, USA) were respectively reconstituted to concentrations of 50 μg/mL, 0.1 μg/mL and 1 μg/mL with sodium carbonate buffer (pH 9.6) to prepare the solid-phase antigen, and 100 μL of the resulting solution was added to each well of a 96 well flat-bottomed microtitre plate (Thermo Fisher Scientific; Tokyo, Japan). Following incubation for 1 h at 37°C, the solid-phase antigen was removed by a wash buffer and 100 μL ELISA buffer (25% Block Ace, cat. no. UK-B80, Megmilk Snow Brand; Hokkaido, Japan) was added into each well for another 1 h of incubation as a blocking agent. A negative control and standard sample defined as a positive control (Zenoaq), and test serum samples at 1:10 or 1:50 dilution and ELISA buffer alone (as a blank) were added to the appropriate wells in duplicate, and the plates were incubated for 1 h at 37°C. After washing, a peroxidase-conjugated goat anti-dog IgE antibody (A40-125P, Bethyl Laboratories; Montgomery, TX, USA) diluted up to 10,000-fold with ELISA buffer was added as the secondary antibody and incubated for 1 h at 37°C. Of importance is that, by immunoelectrophoresis and ELISA, this antisera is known to specifically recognise canine IgE with >0.01% reactivity with purified dog IgG1, IgG2, IgA and IgM. The colour development was initiated by adding 3,3′,5,5′-tetramethylbenzidine (TMB) substrate solution (Kirkegaard and Perry Laboratories; Gaithersburg, MD, USA), as instructed by the manufacturer, with the plate protected from light for 30 min. The optical density (OD) was measured at a 450 nm wavelength using a microtitre plate reader (Infinite M200 Pro, Tecan; Mannedorf, Switzerland) after the enzyme reaction had been stopped by the addition of 0.5 M sulfuric acid.

We set the threshold for the seropositivity for each of the three tested allergens as the mean optical density (OD) plus two standard deviations (SD) of the allergen-specific IgE levels in the sera from the 32 healthy laboratory dogs.

Statistics

In the absence of a predefined hypothesis, and owing to the descriptive nature of our objectives, statistical comparisons of the allergen-specific IgE seroprevalence between countries and regions were not performed. However, we calculated the correlation between the OD values for the three allergens, two at a time, using the Spearman rank correlation test (P < 0.01; Prism 9, Graphpad Software; La Jolla, CA, USA).

Results

Study population

Among the 837 dogs enrolled, 51% were female and 49% male; 7% were <1-year-old, 27% 1–3-year-old, 43% 3–8-year-old and 22% >8-year-old. The most frequently represented breeds were shi tzu (48), French bulldog (44), shiba inu (41), Labrador retriever (42), poodle (33), golden retriever (32), Lhasa apso (28), German shepherd dog (22), Maltese (21), boxer (20), dachshund (19), pug (19), beagle (17), toy poodle (17), American Staffordshire terrier (16), Yorkshire terrier (16) and West Highland white terrier (15).

Determination of positivity thresholds

In the 32 healthy laboratory dogs, the mean (SD) OD values for serum IgE specific for Der f 2, Zen 1 and the DF whole extract were 0.243 (0.028), 0.117 (0.039) and 0.129 (0.029), respectively. Setting the seropositivity levels as the means ±2 SD, the calculated ELISA positivity thresholds were 0.299 for Der f 2- and 0.187 for Zen 1-specific IgE.

Seroprevalence of IgE sensitisation to Dermatophagoides farinae, Der f 2 and Zen
In all 837 dogs taken together, the seropositivity rates for Df, Der f 2 and Zen 1 were 93%, 68% and 89%, respectively.

The individual IgE serum levels for all dogs are represented on Figure 1a, b and c. Using the selected thresholds, only one, two and one laboratory dogs had marginally positive IgE levels against Df, Der f 2 and Zen 1, respectively.

The seroprevalence of IgE against the three allergens is summarised in Table 1. With the selected thresholds, the overall IgE seropositivity rates for Df in these atopic dogs varied between 74% (Argentina) and 100% (the Netherlands, Thailand, South Africa), that for Der f 2 between 12% (Argentina) and 88% (South Africa), and that for Zen 1 between 70% (Argentina) and 100% (the Netherlands). Outside of the especially low seropositivity rate for Der f 2-specific IgE in Argentina, the percentages appeared to vary little between countries.

In order to determine if there would be a variation in allergen-specific IgE seroprevalence between different climates, we selected the USA in which veterinarians from nine clinics spanning different regions with various climates had collected samples. Individual OD values from all dogs are shown on Figure 2a, b and c, while the simplified map of the main continental US climates and clinic locations is depicted on Figure 2d. Overall, we did not find any apparent variations in the OD values and positivity rates between US regions, indicating that the sensitisations to Df and its allergens were fairly uniform and geographically widespread.

In an attempt to determine the relationship between the IgE sensitisation to these three allergens, we assessed if they were correlated. The serum IgE values between the allergens, calculated two at a time, were significantly correlated (Figure 3a, b and c). The highest correlation coefficient was between Df and Zen 1 (Spearman $r = 0.88$), and the lowest was between Der f 2 and Zen 1 ($r = 0.34$).

Finally, we applied to this study the currently accepted benchmark for the definition of major allergens (i.e. ≥50% of dogs sensitised to the parent allergen extract (Df) having detectable IgE against the single allergen). As shown in Table 1, both Der f 2 and Zen 1 reached a major allergen status in dogs from all countries, except for Der f 2 in Argentina (16%).

### Discussion

This study’s results strongly suggest that a majority of atopic dogs from multiple countries and continents are similarly sensitised, not only to a whole-body Df extract, but also to two of its allergens, Der f 2 and Zen 1.

The results of this geographically widespread survey expand those of local seroprevalence studies in France, Japan, Malaysia, Spain, Switzerland, UK and USA for Der f 2, and France, Malaysia, Switzerland, UK and USA for Zen 1.11–16

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**Table 1.** Immunoglobulin (IgE) seroprevalence to Dermatophagoides farinae (Df), Der f 2 and Zen 1 allergens.

|                | As a % of all collected sera | As a % of Df-positive sera |
|----------------|-----------------------------|----------------------------|
|                | Whole Df | Der f 2 | Zen 1 | Der f 2 | Zen 1 |
| Americas       |          |         |       |         |       |
| Argentina      | 74%      | 12%     | 70%   | 16%     | 95%   |
| Brazil         | 91%      | 77%     | 90%   | 86%     | 95%   |
| USA            | 89%      | 81%     | 77%   | 86%     | 93%   |
| Europe         |          |         |       |         |       |
| Italy          | 94%      | 48%     | 94%   | 51%     | 100%  |
| Latvia         | 93%      | 63%     | 97%   | 64%     | 100%  |
| Netherlands    | 100%     | 85%     | 100%  | 88%     | 100%  |
| UK             | 98%      | 86%     | 97%   | 89%     | 98%   |
| Asia           |          |         |       |         |       |
| Japan          | 88%      | 56%     | 83%   | 63%     | 91%   |
| Taiwan         | 94%      | 67%     | 83%   | 71%     | 88%   |
| Thailand       | 100%     | 75%     | 86%   | 75%     | 86%   |
| Africa         |          |         |       |         |       |
| South Africa   | 100%     | 88%     | 98%   | 88%     | 98%   |
| Average        | 93%      | 67%     | 89%   | 71%     | 94%   |
| Median         | 94%      | 75%     | 90%   | 75%     | 95%   |
| Minimum        | 74%      | 12%     | 70%   | 16%     | 83%   |
| Maximum        | 100%     | 88%     | 100%  | 89%     | 100%  |

Bold percentages indicate a major allergen status (i.e. ≥50% of sera with IgE to the whole extract of *D. farinae* also had IgE against this single allergen).
Our results confirm that most atopic dogs worldwide are IgE-sensitised to the Df HDM, a finding long ago established (and reviewed4), and which was confirmed more recently in the USA16 and UK;15 it is surprisingly lower in Malaysia.13 That an IgE sensitisation to this particular mite species is widespread is remarkable, as it occurs even when another mite species, Dermatophagoides pteronyssinus (Dp), predominates in the environment of the same geographical areas, such as the São Paolo Brazilian metropolis, Malaysia and the UK.21 In such situations, the detected IgE sensitisation to Df probably represents, a cross-reactivity to another house dust (Dp or other) or storage mite species, or to the Toxocara canis (Tc) nematode (see below for further discussion).22,23,24 In Malaysia, the low rate of IgE seropositivity to Df likely reflects its scarcity in the local environment.22,25

Der f 2, an MD2-like lipid (ML)-binding protein is a major allergen in humans sensitised to Df.26 Although at first it was believed to be of lesser importance in dogs with AD,27,28 a high IgE seroprevalence to this allergen was later confirmed in atopic dogs in Japan,11,12 Spain,14

Figure 2. Levels of Dermatophagoides farinae, Der f 2 and Zen 1-specific immunoglobulin (Ig)E in atopic dogs from the USA. (a) Optical densities for D. farinae-specific IgE; (b) optical densities for Der f 2-specific IgE; (c) optical densities for Zen 1-specific IgE; (d) simplified map of the climates of the continental USA with the location of clinics that provided atopic dog sera. Colours: orange, California (clinic 1); olive, the Southwest (clinics 2 and 3); light blue, central region (clinics 4 and 9); light green, the Southeast (clinics 5, 6 and 7); and dark blue, northern region (clinic 8). Image courtesy of Wikimedia Commons. Each dot represents a single dog, and the red bars highlight median optical densities.

Figure 3. Correlations between Dermatophagoides farinae, Der f 2 and Zen 1. (a) The correlation between Df and Zen 1 is substantial, and is higher than that observed between Df and Der f 2 (b) or between Der f 2 and Zen 1 (c). Spearman’s rank (t) correlation coefficients are indicated. All correlations were significant at a P < 0.0001.
IgE sensitisation rates to *D. farinae*, Der f 2 and Zen 1

Malaysia\textsuperscript{13} and the UK.\textsuperscript{15} In one of our previous papers,\textsuperscript{16} we reported that the rate of positive serum IgE detection to this allergen was low in atopic dogs from the USA, France and Switzerland. In retrospect, we believe that this low seropositivity rate likely resulted from an artificially high positive threshold established for our ELISA to this allergen. Indeed, we had used a mean +3 SD of the Der f 2-IgE levels in 10 laboratory dogs. As only one of these 10 dogs had any IgE above our assay’s detection limit to this allergen, the established threshold was probably too high to reflect biological reality. Looking back at the original dataset, nine of 17 (53%), eight of 12 (67%) and four of 33 (12%) atopic dogs did, in fact, have detectable Der f 2-specific IgE in France, Switzerland and USA (North Carolina), respectively; this would make Der f 2 also a major allergen in Europe and not in North Carolina.\textsuperscript{16} Altogether, the results of the present study add to the existing body of evidence that Der f 2 now can be considered a major allergen for dogs sensitised to Df in almost every country where such investigations have been undertaken recently. A notable exception to such status is the very low IgE seroprevalence to Der f 2 in Argentina (12%, this study). Homes in Argentina are known to harbour mixed populations of Df and Dp.\textsuperscript{29} so dogs should have been exposed to the Df HDM in their local environment. The presence of detectable IgE to Df and Zen 1 in the same atopic dogs from different Argentinian cities suggests that a degradation of serum IgE had not occurred during shipment of the sera to the laboratory. The reason(s) behind the uniquely low IgE seroprevalence to Der f 2 in Argentine atopic dogs for now will remain an unanswered question.

Our study establishes the unequalled IgE seropositivity rate to the novel Df allergen, Zen 1, in all countries where this had been investigated. This observation is similar to that previously reported for France, Switzerland, the USA\textsuperscript{16} and UK.\textsuperscript{15} In Malaysia, the positivity rate is substantially lower (30%),\textsuperscript{13} possibly reflecting the scarcity of Df in the local environment.\textsuperscript{25} Of interest is that Zen 1, a still uncharacterised protein, is not yet a recognised Df allergen for atopic humans. By contrast, in our previous study reporting an identical advance of these HMW allergens could be questioned, as a simple bioengineering systems. Nevertheless, the relevance of IgE against Df and any of its major allergens remains to be established unequivocally. Notwithstanding the latter statement, Der f 2 is likely to represent an important allergen, as a Der f 2 mono-immunotherapy has been shown to lead to a rapid and substantial improvement in clinical signs in dogs sensitised to Df.\textsuperscript{17,18}

A limitation of our work is that we also did not study the IgE seropositivity rate for the two other HMW Df proteins considered to be major allergens for dogs in the USA, Der f 15 and Der f 18.\textsuperscript{9,10} The main reason for this lack of testing was the absence of commercially available recombinant proteins and their notoriously unusual O-glycosylation pattern that prevents their production in simple bioengineering systems. Nevertheless, the relevance of these HMW allergens could be questioned, as a previous study reported an identical — if not higher — prevalence of IgE directed against HMW Df allergens in healthy and atopic dogs.\textsuperscript{34}
This study’s results highlight the need for further research studies. As Der f 15 and Der f 18 only have been shown to be major Df allergens in atopic dogs in the USA and Spain,14 expanding the seroprevalence data beyond these two countries is needed. Furthermore, owing to the high IgE seropositivity to Df observed in normal dogs,7 determining the IgE seropositivity rates to the four major Df allergens in healthy dogs is indispensable. Research on the IgE cross-reactivity between Df O-glycosylated HMW allergens among themselves and those of Tc also is critical. Finally, clinical trials on Zen 1 immunotherapy would help confirm the clinical relevance and effectiveness of this unique allergen.

In conclusion, this study unequivocally establishes the geographically widespread IgE-sensitisation to Df, Der f 2 and Zen 1 in atopic dogs from countries on five continents.

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Author contributions

Claude Favrot: Conceptualization, Data curation, Formal analysis, Visualization, Writing-original draft. Thierry Olivry: Data curation, Validation, Visualization, Writing-original draft. Toshiroh Iwasaki: Data curation, Writing-original draft.

References

1. Halliwell R. Revised nomenclature for veterinary allergy. Vet Immunol Immunopathol 2006; 114: 207–208.
2. Halliwell RE, Deboer DJ. The ACVD task force on canine atopic dermatitis (III): the role of antibodies in canine atopic dermatitis. Vet Immunol Immunopathol 2001; 81: 159–168.
3. Pucheau-Haston CM, Bzikova P, Eisenschink MNC, et al. Review: The role of antibodies, autoantigens and food allergens in canine atopic dermatitis. Vet Dermatol 2015; 26: 115–e130.
4. Hill PB, Deboer DJ. The ACVD task force on canine atopic dermatitis (IV): environmental allergens. Vet Immunol Immunopathol 2001; 81: 169–186.
5. Swinnen C, Vroom M. The clinical effect of environmental control of house dust mites in 60 house dust mite-sensitive dogs. Vet Dermatol 2004; 15: 31–36.
6. Fujimura M. The study of canine atopic dermatitis involving the isolation of dogs. PoI J Vet Sci 2011; 14: 273–277.
7. Layne EA, DeBoer DJ. Allergen-specific IgE in nonatopic dogs. Vet Dermatol 2019; 30: 79–79.
8. Lian TM, Halliwell RE. Allergen-specific IgE and IgGd antibodies in atopic and normal dogs. Vet Immunol Immunopathol 1998; 66: 203–223.
9. McCall H, Hunter S, Stedman K, et al. Characterization and cloning of a major high molecular weight house dust mite allergen (Der f 15) for dogs. Vet Immunol Immunopathol 2001; 78: 231–247.
10. Weber E, Hunter S, Stedman K, et al. Identification, characterization, and cloning of a complementary DNA encoding a 60-kd house dust mite allergen (Der f 118) for human beings and dogs. J Allerg Clin Immunol 2003; 112: 79–86.
11. Masuda K, Tsujimoto H, Fujisawa S, et al. IgE sensitivity and cross-reactivity to crude and purified mite allergens (Der f 1, Der f 2, Der p 1, Der p 2) in atopic dogs sensitive to Dermatophagoides mite allergens. Vet Immunol Immunopathol 1999; 72: 303–313.
12. Yasahista K, Fujisawa C, Azuma R, et al. Determination of antigenic proteins of housedust mites in 90 dogs suffering from atopic dermatitis. J Vet Med Sci 2002; 64: 673–676.
13. Chan WY, Selvarajah GT, Ajat M, et al. The detection of house dust mite Dermatophagoides farinae, Der f 2 and Zen-1 allergen-specific immunoglobulin E antibodies in dogs with atopic Dermatophagoides farinae. Vet Immunol Immunopathol 2019; 212: 43–49.
14. Moya R, Carnes J, Sinovas N, et al. Immunoproteomic characterization of a Dermatophagoides farinae extract used in the treatment of canine atopic dermatitis. Vet Immunol Immunopathol 2016; 180: 1–8.
15. Patel A, Curtis CF, Cerundolo R. Incidence of anti-Der f 2 and anti-Zen-1 specific immunoglobulin E antibodies in atopic dogs from South-East England. Vet Rec 2019; 184: 317–322.
16. Olivry T, Dunston SM, Favrot C, et al. The novel high molecular weight Dermatophagoides farinae protein Zen-1 is a major allergen in North American and European mite allergic dogs with atopic dermatitis. Vet Dermatol 2017; 28: 177-e38.
17. Fischer N, Tarapata N, Leidi F, et al. An open study on the efficacy of a recombinant Der f 2 (Dermatophagoides farinae) immunootherapy in atopic dogs in Hungary and Switzerland. Vet Dermatol 2018; 29: 337-e118.
18. Kawano K, Mizuno T. A pilot study of the effect of pullulan-conjugated Der f 2 allergen-specific immunotherapy on canine atopic dermatitis. Vet Dermatol 2017; 28: 583-e141.
19. Hensel P, Santoro D, Favrot C, et al. Canine atopic dermatitis: detailed guidelines for diagnosis and allergen identification. BMC Vet Res 2015; 11: 196.
20. Favrot C, Steffan J, Seewald W, et al. A prospective study on the clinical features of chronic canine atopic dermatitis and its diagnosis. Vet Dermatol 2010; 21: 23–31.
21. Thomas WR. Geography of house dust mite allergens. Asian Pac J Allergy Immunol 2010; 28: 211–224.
22. Saridomichelakis MN, Marsella R, Lee KW, et al. Assessment of cross-reactivity among five species of house dust and storage mites. Vet Dermatol 2008; 19: 67–76.
23. Fischer N, Rostaher A, Zwicki L, et al. A Toxocara canis infection influences the immune response to house dust mite allergens in dogs. Vet Immunol Immunopathol 2018; 202: 11–17.
24. Zwicki LLMN, Joekel DE, Fischer NM, et al. Total and Toxocara canis larval excretory/secretory antigen- and allergen-specific IgE in atopic and non-atopic dogs. Vet Dermatol 2018; 29: 222-e80.

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25. Mariana A, Ho TM, Gendez BS, et al. First report on sensitization to allergens of a house dust mite, *Suidasia pontifica* (Acari: Saprolegophilidae). *Southeast Asian J Trop Med Public Health* 2000; 31: 722–723.

26. Batard T, Baron-Bodo V, Martelet A, et al. Patterns of IgE sensitization in house dust mite-allergic patients: implications for allergen immunotherapy. *Allergy* 2016; 71: 220–229.

27. Noli C, Bernadina WE, Willems T. The significance of reactions to purified fractions of *Dermatophagoides pteronyssinus* and *Dermatophagoides farinae* in canine atopic dermatitis. *Vet Immunol Immunopathol* 1996; 52: 147–157.

28. Nuttall TJ, Hill PB, Bansignor E, et al. House dust and forage mite allergens and their role in human and canine atopic dermatitis. *Vet Dermatol* 2006; 17: 223–235.

29. Neffen HE, Fernández-Caldas E, Predolini N, et al. Mite sensitivitiy and exposure in the city of Santa Fe, Argentina. *J Investig Allergol Clin Immunol* 1996; 6: 278–282.

30. Thomas WR. House dust mite allergens: new discoveries and relevance to the allergic patient. *Curr Allergy Asthma Rep* 2016; 16: 69.

31. Thomas WR, Smith W-A, Hales BJ, et al. Characterization and immunobiology of house dust mite allergens. *Int Arch Allergy Immunol* 2002; 129: 1–18.

32. Maizels RM. *Toxocara canis*: molecular basis of immune recognition and evasion. *Vet Parasitol* 2013; 193: 365–374.

33. Haim SA, Carlsson MC, Madsen CB, et al. Glycoproteomic analysis of seven major allergenic proteins reveals novel post-translational modifications. *Mol Cell Proteomics* 2015; 14: 191–204.

34. Nuttall TJ, Lamb JR, Hill PB. Characterisation of major and minor *Dermatophagoides* allergens in canine atopic dermatitis. *Res Vet Sci* 2001; 71: 51–57.

Résumé

**Contexte** – Les chiens atopiques ont souvent des immunoglobulines (IgE) dirigées contre les acariens de poussière de maison *Dermatophagoides farinae* (Df) alors que des données limitées existent sur les taux de sensibilisation aux allergènes Df individuels, Der f 2 et Zen 1.

**Objectifs** – Déterminer les taux d’IgE spécifiques de Df, Der f 2 et Zen 1 chez les chiens atopiques issus de pays de différentes régions.

**Sujets** – Le sérum a été collecté chez 32 chiens de laboratoire au Japon et 837 chiens atopiques de 11 pays des cinq continents: Asie (Japon, Thaïlande et Taiwan), Europe (Italie, Lettonie, Pays Bas, Royaume Uni), Amérique du Nord (USA), Amérique du Sud (Argentine, Brésil) et Afrique (Afrique du Sud).

**Matériaux et méthodes** – Nous déterminons les taux d'IgE spécifiques de Df, Der f 2 et Zen 1 par ELISA. Les corrélations entre les valeurs d’IgE pour ces trois allergènes ont été calculées.

**Résultats** – Les taux de séroconversion d’IgE pour Df variaient entre 74% (Argentine) et 100% (Pays Bas, Thaïlande, Afrique du Sud), ceux pour Der f 2 entre 12% (Argentine) et 88% (Afrique du Sud) et ceux pour Zen 1 entre 70% (Argentine) et 100% (Pays Bas). A part pour le taux de séroconversion particulièrement faible pour les IgE spécifiques de Der f 2 en Argentine, le pourcentage de sensibilisation des IgE variait peu entre les pays si l’on avait une corrélation significative entre les taux d’IgE à ces trois allergènes qui était plus élevée entre Df et zen 1, et plus faible entre Zen 1 et Der f 2.

**Conclusions et importance clinique** – La sensibilisation des IgE à Df est géographiquement étendue. Der f 2 et zen 1 sont des allergènes majeurs pour le chien dans presque tous les pays évalués.

Resumen

**Introducción** – Los perros con dermatitis atópica suelen estar sensibilizados con inmunoglobulina (Ig) E a los ácaros del polvo doméstico *Dermatophagoides farinae* (Df), aunque existen datos limitados sobre las tasas de sensibilización a los alérgenos Df individuales, Der f 2 y Zen 1.

**Objetivos** – determinar las tasas de sensibilización de IgE a Df, Der f 2 y Zen 1 en perros atópicos de países geográficamente diversos.

**Animales** – se recogió suero de 32 perros de laboratorio en Japón y 837 perros atópicos de 11 países de los cinco continentes: Asia (Japón, Tailandia, Taiwán), Europa (Italia, Letonia, Países Bajos, Reino Unido), América del Norte (USA.), América del Sur (Argentina, Brasil) y África (Sudáfrica).

**Métodos y materiales** – determinamos los niveles de IgE específicos de Df, Der f 2- y Zen 1 mediante ELISA. Se calcularon las correlaciones entre los valores de IgE para estos tres alérgenos.

**Resultados** – las tasas de seroposividad de IgE para Df varían entre 74% (Argentina) y 100% (Países Bajos, Tailandia, Sudáfrica), las de Der f 2 entre 12% (Argentina) y 88% (Sudáfrica), y para Zen 1 entre el 70% (Argentina) y el 100% (Holanda). Aparte de la tasa de seroposividad especialmente baja para la IgE específica de Der f 2 en Argentina, el porcentaje de sensibilización a la IgE varió poco entre los países. Hubo una correlación significativa entre los niveles de IgE con estos tres alérgenos, que fue más alta entre Df y Zen 1, y más baja entre Zen 1 y Der f 2.

**Conclusiones y relevancia clínica** – la sensibilización por IgE a Df está muy extendida geográficamente. Der f 2 y Zen 1 son alérgenos importantes para los perros en casi todos los países en los que se evaluó.

Zusammenfassung

**Hintergrund** – Hunde mit atopischer Dermatitis haben oft sensibilisierte Immunglobulin (Ig) E auf die Hausstaubmilbe *Dermatophagoides farinae* (Df); allerdings gibt es nur limitierte Daten über die Sensibilisierungsverläufe auf die individuellen Df Allergene, Der f 2 und Zen 1.

**Ziele** – Eine Feststellung der Sensibilisierungsverläufe auf Df, Der f 2 und Zen 1 bei atopischen Hunden aus geographisch unterschiedlichen Ländern.
Tiere – Es wurde Serum von 32 Laborhunden aus Japan sowie von 837 atopischen Hunden aus 11 Ländern und fünf Kontinenten genommen: Asien (Japan, Thailand, Taiwan), Europa (Italien, Lettland, Niederlande, UK), Nordamerika (USA), Südamerika (Argentinien, Brasilien) und Afrika (Südafrika).

Methoden und Materialien – Wir bestimmten Df-, Der f 2- und Zen 1- spezifische IgE Werte mittels ELISA. Es wurden die Korrelationen zwischen den IgE Werten für diese drei Allergene kalkuliert.

Ergebnisse – Die IgE Seropositivitätsraten für Df variierten zwischen 74% (Argentinien) und 100% (Niederlande), jene für Der f 2 zwischen 12% (Argentinien) und 88% (Südafrika), und für Zen 1 zwischen 70% (Argentinien) und 100% (Niederlande). Außer der sehr niedrigen Seropositivität für Der f 2- spezifisches IgE in Argentinien, varierte der Prozentsatz der IgE Sensibilisierung zwischen den Ländern wenig. Es bestanden signifikante Korrelationen zwischen den IgE Werten gegenüber diesen drei Allergenen, wobei die Korrelation zwischen Df und Zen 1 am höchsten und zwischen Zen 1 und Der f 2 am niedrigsten war.

Schlussfolgerungen und klinische Bedeutung – Die IgE Sensibilisierung auf Df ist geografisch weit verbreitet. Der f 2 und Zen 1 stellen Hauptallergene für Hunde in fast allen Ländern dar, wo diese evaluier wurden.

要旨
背景 – アトピー性皮膚炎の犬は、Dermatophagoides farinae (Df) イエダニに対して免疫グロブリン (Ig) E感作していることが多い、個々のDFアレルゲンであるDer f 2およびZen 1に対する感作率のデータは限られている。
目的 – 本研究の目的は、地理的に多様な国のアトピー性犬におけるDf、Der f 2およびZen 1に対するIgE感作率を測定することであった。
供試動物 – 日本の32頭の実験犬および56大陸11カ国837頭のアドピー犬から血清を採取した。アジア（日本、タイ、台湾）、ヨーロッパ（イタリア、ラトビア、オランダ、イギリス）、北米（アメリカ）、南米（アルゼンチン、ブラジル）、アフリカ（南アフリカ）の5大陸11カ国のアトピー犬837頭から血清を採取した。
材料と方法 – ELISA法により、Df、Der f 2、Zen 1-特異的IgE値を測定した。これら3つのアレルゲンに対するIgE値の相関関係を算出した。
結果 – DfのIgE陽性率は74%（アシヘン）から100%（オランダ、タイ、南アフリカ）、Der f 2のIgE陽性率は12%（アシヘン）から88%（南アフリカ）、Zen 1のIgE陽性率は70%（アシヘン）から100%（オランダ）とばらつきがあった。アシヘンのDer f 2特異的IgEの陽性率が特に低かったことを除けば、IgE感作の割合は国による違いはほとんどなかった。これ3つアレルゲンに対するIgEレベルには有意な相関関係があり、DfおよびZen 1間で最も高く、Zen 1およびDer f 2間で最も低かった。
結論と臨床的妥当性 – Dfに対するIgE感作は地理的に広範囲に及んでいる。Der f 2およびZen 1は、評価対象となったほとんどすべての国の犬にとって主要なアレルゲンである。

要約
背景 – 特性地皮炎犬通常にと呼ばれるアガネのアラルゲン（Df）過敏性、但对单一Df过敏原、Der f 2和Zen 1的致敏率数数据有限。
目的 – 确定无分地区国家的特性原对Df、Der f 2和Zen 1的IgE致敏率。
动物 – 从日本的32只实验室犬和来自5大洲11个国家的837只特性病犬中采集血清：亚洲（日本、泰国、台湾）
、欧洲（意大利、拉脱维亚、荷兰、英国）、北美（美国）、南美（阿根廷、巴西）和非洲（南非）。
方法和材料 – 我们通过ELISA法测定了Df、Der f 2和Zen 1-特异的IgE水平，计算这两种过激原的IgE值之间
的相关性。
结果 – DfのIgE血清陽性率を74%（阿根廷）と100%（荷兰、泰国、南非）之间变化、Der f 2在12%（阿根廷）
と88%（南非）之间变化，Zen 1在70%（阿根廷）と100%（荷兰）之间变化。除阿根廷Der f 2特異的IgEの血清
陽性率特別低下、各国之间IgE致敏百分比差異很小，这两种过敏原的IgE水平之间存在显著相关性，Df和Zen
1之间最高、Zen 1とDer f 2之间最低。
结论和临床相关性 – IgE对Df的致敏作用在地理上广泛存在。在几乎所有进行评价的国家中，Der f 2和Zen 1
是犬的主要过敏原。

Resumo
Contexto – Cães com dermatite atópica apresentam comumente sensibilização mediada por IgE para os ácaros da poeira doméstica Dermatophagoides farinae (Df). Entretanto, poucos dados estão disponíveis a respeito da frequência de sensibilização aos alérgenos individuais de Df, Der f 2 e Zen 1.
Objetivos – Determinar a frequência de sensibilização de IgE para Df, Der f 2 e Zen 1 em cães atópicos de países geograficamente diversos.
Animais – O soro de 32 cães de laboratório no Japão, e 837 cães atópicos de 11 países de cinco continentes: Ásia (Japão, Tailândia, Tailândia), Europa (Itália, Lettonia, Holanda, Reino Unido), América do Norte (USA), América do Sul (Argentina, Brasil) e África (África do Sul).
Métodos e materiais – Determinamos os níveis de IgE específicos para Df, Der f 2 e Zen 1 por ELISA. Foram calculadas as correlações entre os valores de IgE para esses três alérgenos.
Resultados – As taxas de soropositividade de IgE para Df variaram entre 74% (Argentina) e 100% (Holanda, Tailândia, África do Sul). As taxas para Der f 2 variaram entre 12% (Argentina) e 88% (África do Sul), e para Zen 1 entre 70% (Argentina) e 100% (Holanda). Além da taxa de soropositividade especialmente baixa de IgE específica para Der f 2 na Argentina, houve pouca variação na porcentagem de sensibilização por IgE entre os países. Houve correlação significativa entre os níveis de IgE para esses três alérgenos, que foi maior entre Df e Zen 1 e menor entre Zen 1 e Der f 2.

Conclusões e relevância clínica – A sensibilização por IgE ao Df é geograficamente difundida. Der f 2 e Zen 1 são os principais alérgenos para cães em quase todos os países onde isso foi avaliado.