IgE reactivity to components of fish allergens in Pacific cod (Gadus macrocephalus) in atopic dogs

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Abstract
Background IgE reactivity to fish allergens in atopic dogs, which are used as models for food allergy, has not been elucidated to date. We investigated IgE reactivity to crude and purified Pacific cod (Gadus macrocephalus) allergens in atopic dogs to identify the allergenic components of cod.

Methods Specific IgE to crude cod allergens in the sera of 179 atopic dogs, including 27 dogs with cod allergy, were measured using enzyme-linked immunosorbent assay (ELISA). The allergenic components of crude cod antigen were analyzed by ELISA, immunoblotting, and liquid chromatography-tandem mass spectrometry (LC-MS/MS). IgE reactivity to parvalbumin, collagen, and tropomyosin was evaluated using the sera of atopic dogs that were positive for specific IgE to crude cod allergens.

Results Specific IgE to crude cod allergens were present in 36 (20%) of the 179 atopic dogs and 12 (44%) of the 27 dogs with cod allergy. In allergen component analysis, IgE reactivity to tropomyosin and enolase was observed in the sera of dogs with cod allergy. Among the 36 dogs with IgE reactivity to crude cod extracts, 9 (25%), 14 (39%), and 18 (50%) dogs had specific IgE to parvalbumin, collagen, and tropomyosin, respectively.

Conclusions The dogs exhibited IgE reactivity to the cod allergen components similar to that observed in humans, providing support for the use of atopic dogs with fish allergy as a model for fish allergy in humans.

Background
The prevalence of fish allergy, which affects approximately 0.2% of the world population [1], is over ten times higher in geographic regions where fish is an essential dietary component such as Japan [2, 3]. Fish allergy is typically known to be a life-long condition in contrast to other food allergies [3]. Since clinical cross-reactivity to different fish species is a widely accepted feature of fish allergy, affected individuals have to avoid all fish species for long periods, and inadvertent exposure to fish allergens, and the consequent severe or fatal reactions remain a grave risk for fish-sensitive individuals [3]. Among the fish allergens, parvalbumin is the best characterized major allergen that is found in many species [4–6]. In a previous study, we identified fish gelatin (type I collagen) as an
allergen [7]. More recently, other proteins such as tropomyosin, enolases, and aldolases were also considered as relevant allergens in fish [8]. To resolve the issue of quality of life in fish-allergic patients, it is necessary not only to investigate the mechanisms underlying the acquisition of allergenicity but also to elucidate the approaches to reduce allergenicity.

Animal models are beneficial to test food allergies as they provide a more rapid and extensive evaluation of allergenicity of certain foods [9]. Dogs are recognized as a useful model to study IgE-mediated hypersensitivity [10, 11] as they are one of the few species other than humans in which allergies develop naturally following environmental exposure to a broad spectrum of allergens, including foods [12, 13]. Disease states in dogs include atopic dermatitis, gastroenteric inflammation, and anaphylaxis [14]. Based on the many similarities between canine atopic dermatitis and humans [13], atopic dogs have been utilized as animal models for food allergies to cow’s milk [15], corn [16], and nuts [11].

The allergenic components of atopic dogs in food are assumed to correspond with those of humans [17]. However, Kubota et al. reported that the allergenic components in atopic dogs were different from those in humans [18]. Previous studies reported that atopic dogs were appropriate models for food allergies by demonstrating the production of specific immunoglobulin (Ig) E to crude allergens and positive oral challenges similar to those observed in human subjects [11, 15]. These dogs might be suitable models to elucidate the mechanisms underlying the allergenicity of the food components if the allergenic components in the dogs are analyzed. Therefore, in the present study, we investigated IgE reactivity to crude and purified Pacific cod (Gadus macrocephalus) allergens in atopic dogs and elucidated similarities in fish allergy between humans and dogs to assess the potential of atopic dogs with fish allergy as a suitable animal model.

Methods

Sera of atopic dogs

To examine IgE reactivity in atopic dogs, we obtained sera from 179 dogs that were diagnosed with atopic dermatitis based on the criteria by Willemse [19] and Prelaud et al [20] among dogs visiting Fujimura Animal Hospital (Osaka, Japan). Twenty samples from laboratory dogs were used as negative
controls. The dogs were housed indoors as experimental laboratory animals and had never been exposed to fish antigens. None of the laboratory dogs exhibited signs of atopic dermatitis. Enzyme-linked immunosorbent assay (ELISA) for parvalbumin or collagen from salmon (Atlantic salmon; *Salmo salar*), sardine (Japanese pilchard; *Sardinops melanostictus*), and mackerel (Chub mackerel; *Scomber japonicus*) were performed using individual sera of nine negative control dogs that provided a large amount of serum. All sera were stored at −80°C before use. Oral informed consent was obtained from the dog owners. All experimental procedures were carried out in accordance with Japanese law and approved by the animal care and user committee of Azabu University.

**Food elimination and oral provocation tests**

Among the 179 dogs with atopic dermatitis, 27 were confirmed to have cod reactivity by the oral provocation test after the elimination diet test; consent for food provocation test was obtained from the dog owners. These 27 dogs were fed commercial hydrolyzed protein diets as elimination diets for 6–8 weeks by the dog owners. When the veterinary physician recognized the complete resolution of the clinical signs during the food elimination test, the dog was admitted to the animal hospital and challenged with various cod ingredients, including grilled cod meat and cod-containing dog foods. The cod provocation test was discontinued immediately upon the relapse of the clinical signs including vomiting, diarrhea, erythema, pruritic urticaria, and conjunctival hyperemia [21].

**Preparation of crude cod allergens**

Cod is not only one of the most commonly consumed fish species in Europe and Japan [22] but also one of the most characterized fish species with allergen components [8]. Thus, Pacific cod was purchased from a fish market in Japan to be used in the study. The fresh, raw meat of four fishes (500 µg) was homogenized in 500 µl phosphate-buffered saline (PBS, 10 mM pH 7.2) and rotated overnight at 4°C. After centrifugation at 21500 g for 5 min at 4°C, the supernatant was collected, and the protein was quantified using the BCA protein assay (Bio-Rad, Hercules, CA, USA).
Purification of parvalbumin, collagen, and tropomyosin

Fish parvalbumin [23] and collagen [4] were purified as described previously. Tropomyosin was purified from the ether powder by Bailey’s method with slight modification [19]. Briefly, fish ether powder was stirred in a beaker with 75 ml extraction buffer containing 15 mM Tris HCl pH 7.6 (Sigma Aldrich, St Louis, MO, USA), 1 M KCl (Kanto Kagaku, Tokyo, Japan), and 2 mM dithiothreitol (Sigma Aldrich) overnight at 4°C. The extract was collected by centrifugation at 5400 g for 10 min at 4°C. The supernatant pH was adjusted to 4.5 with 1 N HCl to precipitate tropomyosin, and the precipitation was collected by centrifugation at 5400 g for 10 min at 4°C. The isoelectric precipitation was repeated once, and the precipitated material was dissolved in the extraction buffer. The supernatant after the extraction was collected by centrifugation and fractionated by ammonium sulfate at a concentration of 50%. The sample precipitated by ammonium sulfate was dissolved and dialyzed against PBS. The obtained protein extracts were confirmed by sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis (PAGE).

SDS-PAGE and immunoblotting

Among the 20 atopic dogs with IgE sensitivity to crude cod allergens that did not react to parvalbumin or collagen, sera were used from two dogs (no. 34 and no. 128) that provided a large amount of sera and reacted to crude cod allergens but not cod parvalbumin or collagen, as confirmed with the provocation test using cod-containing foods (Table 1). SDS-PAGE was performed according to the method of Laemmli [24]. Precision plus protein standards (Bio-Rad) were used as molecular-mass markers. Protein components were electrophoretically separated using 5%-20% gradient polyacrylamide gels, and proteins were visualized either by Coomassie brilliant blue R250 (Bio-Rad) or transferring onto polyvinylidene difluoride membranes (GE Healthcare, Chicago, IL, USA). Immunoblotting was performed as described previously [7]. IgE in patient dog sera were used as primary antibodies, which were diluted 1:10 in tris buffered saline containing 0.1% Tween-20 and 5% nonfat dried milk. Mouse monoclonal anti-dog IgE antibodies (0.5 μg/ml) were used as secondary antibodies [21]. Detection was performed using an enhanced chemiluminescence immunoblotting
detection reagent (GE Healthcare).

**Liquid chromatography (LC)-tandem mass spectrometry (MS/MS)**

For LC-MS/MS, the protein bands detected by immunoblotting were excised, and in-gel digestion was performed with 0.5 mg N-tosyl-L-phenylalanine chloromethyl ketone-treated trypsin (Promega, Madison, Wisconsin) at 37°C for 16 h. Tryptic digests were acidified using formic acid (pH < 2.0) and centrifuged at 21500 g for 15 min. The supernatants were analyzed using high-performance LC (Advance System; AMR, Tokyo, Japan) connected to an electrospray ionization triple quadrupole mass spectrometer (4000 QTRAP; AB Sciex, Framingham, MA, USA). Extracts were injected into a reversed-phase column (electrospray ionization column [octa decyl silyl]; particle inner diameter, 75 mm; length, 100 mm; diameter 3 mm; LC Assist, Tokyo, Japan) that was eluted with a 5%-45% gradient of acetonitrile containing 0.1% formic acid for 60 min at 300 nl/min. Ionization was performed using an ion-spray voltage of 2000 V at a capillary temperature of 200°C. The mass spectrometry instrument was operated in the positive ion mode over the range of 450-1200 m/z. The MS/MS spectra were obtained in the enhanced production scan mode, and two higher-intensity peaks in each mass spectrometry scan were chosen for collision-induced dissociation.

The MS/MS data were used to search in entries under the *Liza aurata* category of the UniProt database using the Mascot peptide search engine. An MS tolerance of 1.0 Da for precursor ion and an MS/MS tolerance of 0.8 Da were set as windows of processing parameters for matching peptide mass values.

**Fluorometric ELISA for allergen-specific serum IgE**

Specific IgE levels to cod crude allergens were measured using a fluorometric ELISA as previously described [18]. A microplate (NUNC Immuno Plate Maxisorp F96; Nalge Nunc International, Roskilde, Denmark) was coated with crude cod extracts (10 µg/ml) or the purified allergens (parvalbumin, collagen, or tropomyosin; 1 µg/ml) at 4°C overnight. After washing, the plate was incubated with diluted sera (1:10) at room temperature for 3 h. The plate was then washed and incubated at 4°C
overnight with a mouse monoclonal anti-dog IgE antibody (0.5 µg/ml) [25]. After washing with PBS containing 0.1% (v/v) Tween 20 (PBS-T), the plate was incubated with a biotinylated rat monoclonal anti-mouse IgG₁ (Zymed Laboratories, San Francisco, CA, USA) at room temperature for 1 h. After washing, the plate was incubated with b-D-galactosidase-conjugated streptavidin (Zymed Laboratories) at room temperature for 1 h. After the final washing, the plate was incubated with 0.1 mM 4-methylumbelliferyl- b-D-galactopyranoside (Sigma Aldrich) at 37°C for 2 h. The enzymatic reaction was stopped with the addition of 0.1 M glycine-NaOH (pH 10.2). The fluorescence intensity was measured as fluorescence units (FU) on a microplate fluorescence reader (Fluoroskan; Flow Laboratories, McLean, VA, USA). The absorbance was measured at 355 nm with a 460 nm reference filter. All the washing steps were performed three times for 5 min in PBS-T. The cutoff value was determined as the average + three standard deviations (SDs) of FU in serum samples of 20 dogs used as negative controls. All tests were performed in triplicate.

Results

**Atopic dogs with cod allergy: clinical characteristics and IgE reactivity to crude cod allergens**

The present study included 179 atopic dogs from 34 breeds, and there were 79 males and 100 females (age, 2 months–11 years; mean age, 3.9 years). We first examined specific IgE reactivity to crude cod allergens in the atopic dogs using ELISA and found that 20% (36/179) of the dogs exhibited increased levels of specific IgE to crude cod allergens (Figure 1). Based on diet history, the clinical reactivity to cod meat was confirmed by oral provocation in 27 atopic dogs (15%). Among the 27 dogs that underwent the food provocation test, 8 dogs that were administered grilled cod meat and 19 dogs that were administered cod-containing dog foods were diagnosed with cod allergy (i.e. dogs with cod allergy). IgE reactivity to crude cod allergens was positive in 44% (12/27) of the dogs with cod allergy (Figure 1).

We next performed a field survey of commercial canine dry food products to estimate the difference in fish allergen exposure within and outside Japan and found that 75% (117/157) of the Japanese canine dry food products contained fish. In contrast, 9% (7/82) of the products produced in Australia
and the USA included fish.

**IgE reactivity to cod parvalbumin and collagen among atopic dogs exhibiting IgE reactivity to crude cod allergens**

IgE reactivity to the purified cod allergens parvalbumin and collagen were tested by ELISA using the sera of the 36 dogs with IgE reactivity to crude cod allergens. IgE reactivity to parvalbumin was observed in 25% (9/36) of the dogs, whereas IgE reactivity to collagen was present in 33% (12/36) of the dogs. However, 56% (20/36) of the dogs showed no IgE reactivity to these purified cod allergens (Figure 2A), indicating that reaction occurred to other fish proteins. We also compared IgE reactivity to parvalbumin and collagen among different fish species by analyzing IgE reactivity to parvalbumin and collagen in four fish species in six dogs that were reactive to cod parvalbumin and eight dogs that were reactive to cod collagen. These dogs reacted strongly to crude cod antigen and had sufficient sera to conduct the tests. IgE reactivity to parvalbumin from all fish species excluding mackerel occurred in 100% (6/6) of the dogs (Figure 2B), whereas IgE reactivity to collagen from all fish species occurred in 100% (8/8) of the dogs (Figure 2C).

**Identification of other cod allergen components in atopic dogs with cod allergy**

Two dogs with a documented clinical history of food allergy to cod meat and IgE reactivity to crude cod allergens but not cod parvalbumin or collagen were recruited (Table 1). After the separation of crude cod meat proteins using SDS-PAGE, IgE immunoblotting experiments using total crude cod allergens was performed to detect IgE-reactive protein bands at around 35 and 55 kDa (Figure 3). LC-MS/MS performed for identification of the proteins corresponding to the bands of crude cod allergens in SDS-PAGE showed that the protein band around 35 kDa in weight matched tropomyosin from golden gray mullet (*Liza aurata*), and the protein band around 55 kDa in weight matched an enolase of python (*Python regius*) (Table 2).

We next purified cod tropomyosin from crude cod allergens and confirmed with SDS-PAGE (Figure 3). The IgE levels for fish tropomyosin in the serum of dog no. 34 was measured using ELISA (Figure S1),
which revealed that the IgE level to cod tropomyosin in the serum of dog no. 34 was significantly higher than those in the sera of the 20 control dogs. The levels of specific IgE to other fish tropomyosins were also higher in dog no. 34 compared to the negative controls.

A dog of no. 34 presented with specific IgE and positive intradermal test to crude mite allergen (House dust mite; *D. farinae*) (data not shown). To compare the relationship of allergenicity of mite tropomyosin and IgE reactivity of cod tropomyosin, we evaluated with IgE level for recombinant mite tropomyosin (*Der f 10*) in the serum of dog no. 34 using ELISA (Figure S1), which revealed that the IgE level to mite tropomyosin in the serum of dog no. 34 was also significantly higher than those in the sera of the 20 control dogs.

**IgE reactivity to cod tropomyosin and crude cod allergens in atopic dogs**

Using the sera of 36 atopic dogs with IgE reactivity to crude cod allergens, we determined IgE reactivity to cod tropomyosin using ELISA (Figure 4), which revealed that the IgE reactivity to tropomyosin was present in 50% (18/36) of the dogs. Therefore, 67% (12/18) of the dogs with atopic dermatitis that had high IgE levels to crude cod antigen and tropomyosin. Conversely, 25% (9/36) of the dogs showed no IgE reactivity for any of these allergens (Figure 5).

**Discussion**

Although one study reported that the rate of food allergy or intolerance due to fish was only 1.3% (4/297) in dogs [23], the present study results imply that the prevalence of fish allergy might be higher (Fig. 1). Our field survey of commercial canine dry food products suggested that dogs in Japan might be exposed to fish more frequently compared with dogs in other countries such as USA. These results suggest that atopic dogs might at a higher risk of developing fish allergy due to an increase in the frequency of daily exposure for fish, implicating that atopic dogs mimic the human condition.

Moreover, we showed that the rate of IgE reactivity to crude cod allergens among dogs with cod allergy was 44% (12/27) (Fig. 1), which was comparable to that reported in humans [26, 27]. Atopic dermatitis with food allergy can be a manifestation of an IgE- or a non-IgE-mediated reaction to food [28, 29]. Non-IgE-mediated reactions are more often delayed, in contrast to IgE-mediated reactions
Delayed symptoms associated with reactions were frequently observed during the provocation test in the present study. For further elucidation of these aspects of atopic dermatitis, atopic dogs that are not sensitive to crude cod allergens might be useful as a spontaneous animal model of non-IgE-mediated allergy.

To our knowledge, this is the first study to describe the allergenic potency of parvalbumin and collagen in dogs. Parvalbumin has a higher allergenic potency than collagen in humans with cod allergy [2]. The present study revealed that the rate of collagen allergy was higher in dogs and that collagen elicited a stronger reactivity based on specific IgE levels compared with parvalbumin in these animals (Fig. 2A). One possible interpretation of this discrepancy might be due to a loss of parvalbumin from dog food via physical and chemical steps in food processing, because parvalbumin is a water-soluble protein, unlike collagen. Additionally, the sera of the dogs that exhibited reactivity to crude cod as strongly as collagen and parvalbumin showed reactivity to collagen and parvalbumin from the other fish species tested in the current study, including salmon, mackerel, and sardine (Fig. 2B and 2C). Humans exhibit broad cross-reactivity to parvalbumin and collagen from distinct fish species [22, 30], which we predict might be occurring in dogs with fish allergy as well.

Of note, the level of specific IgE to tropomyosin was higher than that for other allergenic components of cod, i.e., parvalbumin and collagen, in the dogs with cod allergy (Fig. 5). Tropomyosin was demonstrated to be a fish allergen in a study using the sera of human patients with tilapia allergy [31]. Additionally, our comparison of the protein sequence of cod tropomyosin with those of other fish species revealed that cod tropomyosin exhibited 94–99% sequence similarity with tropomyosin of other fish species that are commercially available on the market (Table S1). Tropomyosin is a major allergen that is the cause of many forms of crustacean allergy [32] as well as mite allergy [33] in humans. Although comparison of the tropomyosin protein sequence revealed a low sequence similarity between cod and shrimp (Table S1), fish-shrimp cross-reactivity was previously reported in humans [34, 35]. Additionally, mite-crustacean cross-reactivity was widely reported in humans, and the tropomyosin sequence similarity between the two species is over 90% [36]. In dogs, mite is one of the most frequent sensitizing allergens, and mite tropomyosin is the allergenic component of mite
allergy in canine atopic dermatitis [37]. Additionally, the serum of dog no. 34 exhibited reactivity to mite tropomyosin as strongly as cod tropomyosin (Figure S1). Taken together, these findings raise the possibility that mite allergy in dogs with IgE reactivity to mite tropomyosin might be associated with increased cod allergy in dogs with IgE reactivity to cod tropomyosin.

The results of the current study also revealed enolase is a potential allergen associated with canine fish allergy (Fig. 3 and Table 1). Enolase was recently defined as a fish allergen that exhibited cross-reactivity to chicken in humans and dogs [35, 38]. Numerous fish proteins, other than those purified proteins that are recognized as critical allergenic components in humans, have been registered in the International Union of Immunological Societies allergen database [39]. In some cases, minor allergens in humans can be dominant allergens in dogs [18, 40]. Moreover, in the present study, 25% (9/36) of the dogs with specific IgE to crude cod antigens did not show IgE reactivity to any of the three major purified allergens by ELISA, implying that other purified cod allergens might underlie fish allergy in dogs.

In conclusion, the current study revealed the presence of specific IgE to crude cod allergens as well as parvalbumin, collagen, and tropomyosin in atopic dogs. The observed IgE reactivity also revealed similarities in fish allergens between dogs and humans. Previous studies utilized experimental models of canine atopic dermatitis created in laboratory facilities [9, 11, 15, 16]. However, globally, atopic dermatitis affects approximately 10-20% of the canine population, which often visit animal hospitals [41]. The utility of atopic dogs with reactivity to allergens that correspond with components of human allergies as spontaneous animal models can facilitate and contribute significantly to efforts to elucidate food allergy, and future studies should focus on identification of allergenic components of foods in atopic dogs.

Declarations

Ethics approval and consent to participate

All experimental procedures were carried out in accordance with Japanese law and approved by the animal care and user committee of Azabu University.
**Consent for publication**

All authors read and approved the final manuscript.

**Availability of data and materials**

All data generated or analysed during this study are included in this published article and its supplementary information files.

**Competing interests**

Masahiro Sakaguchi has received research funding from ITEA Inc., whereas other authors declare no conflicts of interest.

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**Author’s contributions**

I. Imanishi made substantial contributions to the study conception and design as well as data acquisition and analysis; he also drafted the article. K. Kurata, J. Kamiie, and M. Fujimura made substantial contributions to performance of the experiments. J. Uchiyama, K. Mizukami, K. Shimakura, and K. Nishifuji assisted in the study design and manuscript preparation. M. Sakaguchi organized all the experimental settings and manuscript editing. He also revised the article critically for important intellectual content.

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Tables

Table 1. Clinical characteristics of dogs with IgE reactivity to crude cod allergen
Table 2. List of suspected allergens detected in two dogs with IgE reactivity to crude cod meat by LC-MS/MS

| Band (kDa) | Accession number | Protein name/species | Theoretical MW (kDa) | PI | Score | Peptide matches |
|------------|------------------|----------------------|---------------------|----|-------|----------------|
| 35         | P84335           | Tropomyosin/ Liza aurata | 32.710              | 4.69 | 111   | 3              |
| 55         | gi17367183       | Alpha enolase/ Python regius | 47.541              | 6.97 | 64    | 1              |

LC-MS/MS, liquid chromatography-tandem mass spectrometry; MW, molecular weight; PI, isoelectric point.

Figures

○ no. 34 dog
○ no. 128 dog
○ other dogs with cod allergy
Immunoglobulin (Ig) E reactivity to cod crude allergens in atopic dogs based on the levels of specific IgE to crude cod meat in 20 negative controls (N.C.). The mean fluorescence units (FU) ± standard deviation (SD) is 457 ± 61 FU, and the cutoff value (mean + 3SD) is 638 FU, shown as the dotted line. IgE reactivity to crude cod allergens among 179 atopic dogs. Thirty-six atopic dogs exhibited specific IgE reactivity to crude cod meat extracts, with IgE levels ranging from 640 to 3483 FU. Blue circle indicates IgE reactivity in dog no. 34. Red circle indicates IgE reactivity in dog no. 128. Gray circles indicate IgE reactivity in other cod allergic dogs.
Figure 2

IgE reactivity to parvalbumin and collagen among atopic dogs exhibiting significant
elevations in cod-specific IgE levels. (A) IgE reactivity to cod parvalbumin and collagen in 36 dogs with IgE reactivity to crude cod allergens was determined using diluted sera (1:10). The dotted line shows the cutoff value, which was calculated using sera from 20 negative controls (N.C.). Based on the levels of specific IgE to parvalbumin and collagen in the negative controls (mean ± SD, 84 ± 67 and 74 ± 25 FU respectively), the cutoff values (mean + 3SD) for specific IgE against parvalbumin and collagen were 286 and 149 FU, respectively. The specific parvalbumin and collagen IgE levels in atopic dogs were 299–1121 and 151–3774 FU, respectively. Blue circle indicates IgE reactivity in dog no. 34. Red circle indicates IgE reactivity in dog no. 128. Specific IgE reactivity to (B) parvalbumin and (C) collagen was measured in the sera of six dogs with specific IgE to parvalbumin and eight dogs with specific IgE to collagen and those of nine healthy dogs (N.C.). The cutoff values (mean + 3SD) of specific IgE were 379, 374, 351, and 372 FU for parvalbumin from cod, salmon, mackerel, and sardine, respectively, which are shown with the dotted lines. The cutoff values (mean + 3SD) of specific IgE were 393, 310, 231, and 372 for collagen from cod, salmon, mackerel, and sardine, respectively.
Figure 3

Immunoblotting for crude cod meat allergen. In the left column, the molecular standard is shown. Lanes 1 and 5 show immunoblotting for cod meat and tropomyosin, respectively, stained by Coomassie brilliant blue R250. Lanes 2, 3, and 4 show immunoblotting using the sera of two atopic dogs (lane 2, dog no. 34; lane 3, dog no. 128) and the serum of a healthy dog serum (lane 4, negative control). The arrow and arrowhead next to lane 1 indicate the bands analyzed by liquid chromatography-tandem mass spectrometry, which corresponded to the band detected by immunoblotting (lanes 2, 3).
Determination of reactivity to cod tropomyosin among 36 dogs with specific IgE to crude cod meat allergen using enzyme-linked immunosorbent assay. The cutoff value (dotted line) calculated from 20 negative control samples. Based on the levels of specific IgE to tropomyosin in negative controls (mean ± SD, 33 ± 67 FU), the cutoff value (mean + 3SD) was determined as 234 FU.
Figure 5

Venn diagram of the number of dogs harboring fish allergens specific IgE.

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