## Scientific Report

# Diagnosis of Canine Food Sensitivity and Intolerance Using Saliva: Report of Outcomes 

W. Jean Dodds, DVM<br>Hemopet, 11561 Salinaz Avenue, Garden Grove, CA 92843;<br>Phone: 714-891-2022, ext. 115; Fax: 714-891-2123; e-mail: info@hemopet.org

ABBREVIATIONS<br>Inflammatory bowel disease - IBD<br>Gastrointestinal - GI<br>Immunoglobulin A - IgA<br>Immunoglobulin E - IgE<br>Immunoglobulin G - IgG<br>Immunoglobulin M - IgM


#### Abstract

Objective - To assess the efficacy of a novel saliva-based immunoassay of IgA-and IgM-antibodies in predicting canine food sensitivity and intolerance.


Design - Prospective controlled and clinical trial cohort populations.
Animals - Greyhounds from a closed colony, both healthy ( $\mathrm{n}=29$ ) and with inflammatory bowel disease (IBD) ( $\mathrm{n}=10$ ); clinical samples from dogs of various breeds and mixed breeds, classified as healthy without evidence of IBD ( $\mathrm{n}=208$ ); clinically suspected IBD ( $\mathrm{n}=289$ ); and proven IBD ( $\mathrm{n}=98$ ) cases. Second cohort of clinical samples from dogs suspected to have IBD ( $\mathrm{n}=1008$ ).

Procedures - Saliva was collected with a dental cotton rope from dogs that had not eaten for at least 8 hrs , placed in a double-sleeved saliva collection tube, and transported to the laboratory. Salivary antibodies elicited by 24 foods were measured with goat anti-canine IgA and IgM.

Results - Data distinguished healthy, suspect, and proven IBD cases among Greyhounds and 2 large canine clinical case cohorts. Results were stratified as negative, and as intermediate, medium, and strong reactors against 1 or more of the food antigens tested. The 1-4-year and over 10-year age groups had the highest number of positive food reactors, and the German Shepherd Dog was most represented. Clinical
outcome comparisons after eliminating reactive foods ( $\mathrm{n}=50$ ) and follow up saliva re-testing ( $\mathrm{n}=15$ ) demonstrated the clinical accuracy and predictive outcome of this test.

Conclusions and Clinical Relevance - The novel salivarybased food sensitivity and intolerance test described here for canines offers a reliable and clinically predictive alternative to food elimination trials, serum-based food allergy testing, and skin patch testing.

## Introduction

The background and rationale for a novel approach to diagnosis of canine food sensitivity and intolerance using saliva was recently published (1) (a). Basically, delayed, latent, or pre-clinical elaboration of IgA and/or IgM antibodies to specific food antigens can be detected in mucosal fluids such as saliva, feces, sweat, and tears (2-6). These antibodies to foods appear in the mucosal biofluids before the clinical or gastrointestinal (GI) tract biopsy diagnosis is made of intestinal biopsy-confirmed inflammatory bowel disease (IBD) and/or or "leaky gut syndrome." A major cause of the leaky gut is known to stem from release of zonulin which physiologically modulates intestinal barrier function and serves as a biomarker of impaired gut function (2-6). Zonulin release is triggered primarily by the gliadin protein of dietary glutens and gut bacteria in the small intestine, thereby creating gradients for the optimal transport of nutrients and balancing the body's tolerance or immunity to
external antigens, including foods (4). Frequently, IgA or IgM antibodies to food ingredients appear in saliva but are not detectable in serum (7). Salivary antibodies thus serve as an indication of a mucosal immune response and can be induced in people and animals without parallel antibodies being detected in serum (7-9). The same test is also available for cats and horses (1).

## Materials and Methods

## Study Populations

## Saliva Diagnostic Testing Clinical Validation Protocol

Healthy adult Greyhounds ( $\mathrm{n}=29$ ) were adopted after retirement from the racing industry in Arizona, Oklahoma, and Texas. They were either neutered males or spayed females of similar adult age (2.5-5 years) and weight (25-35 kg [55-75 lbs] for females; 30-40 kg [65-90 lbs] for males). All had periodic general health examinations and laboratory screening profiles performed quarterly. Laboratory profiles checked CBC, chemistry profile, thyroid profile, vaccine titers, von Willebrand factor antigen, infectious disease screening, urinalysis, and fecal ova and parasites.

The Greyhounds live in the company's licensed, closed colony facility (Biologics License \# 84), which is inspected annually by the California Department of Food \& Agriculture; this inspection includes a review of the animal blood bank procedures and production program, animal welfare, and inspection of the animal care and laboratory facilities.

## Regular Diet

All dogs at Hemopet are fed the same control diet, cereal kibble with some canned food (b). The food is given twice daily in pre-determined amounts to maintain ideal body weight.

## Diet for Cohort Group with Food <br> Sensitivity/Intolerance ( $\mathrm{n}=10$ )

Any resident Greyhound exhibiting 1 or more classical symptoms or signs of food sensitivity/intolerance (inflammatory bowel disease, diarrhea, constipation, flatulence, abdominal cramping, gastritis, anorexia or poor appetite, and/or low-grade chronic skin disease [folliculitis, pyoderma]) is fed novel protein source foods (c).

## Initial Clinical Case Cohorts Tested

## Against 6 Purified Food Extracts

The initial clinical trials involved veterinary clinics throughout the USA and Hemopet's resident rescued

Greyhounds. There were 29 healthy control dogs and 81 dogs affected with chronic IBD and/or "leaky gut" syndrome; some cases also had pruritus. Subsequent expansion of these trials included a total of 595 cases ( 566 new cases plus the 29 healthy greyhounds): Healthy dogs without evidence of IBD ( $\mathrm{n}=208$, which included $\mathrm{n}=122$ completely healthy and $\mathrm{n}=86$ healthy with minor non-GI or non-pruritus issues), Suspected cases of IBD based upon the submitting veterinarian's clinical diagnosis ( $\mathrm{n}=289$ ), and Proven cases by intestinal biopsy and/ or food elimination trials to have IBD ( $\mathrm{n}=98$ ).

Sex: There were 455 dogs described by their sex: 244 males and 211 females. Of the males, 195 were intact and 49 neutered; and of the females, 74 were intact and 137 were spayed.

Diet: Fifty dogs of the 566 total cases studied ate raw diets exclusively (Healthy=10; Suspect=34; Proven=6). The majority of dogs studied ate commercial kibbled cereal either dry or with some canned foods and treats. Five of the 6 proven IBD cases ate specialized prescription or homemade elimination diets containing novel proteins and treats.

## Larger Clinical Case Cohort Tested Against 24 Purified Food Extracts

Saliva samples submitted by veterinary clinics throughout North America plus some from Australia, Austria, Brazil, France, Germany, Italy, Hong Kong, Japan, Poland, Portugal, Switzerland, and the United Kingdom ( $\mathrm{n}=1008$ ) were tested against 24 affinity-purified (>98\% pure by molecular analysis) lyophilized food extracts (d) from the foods listed below.

The raw ELISA-based absorbance data measured to 4 decimal places and run in duplicate were averaged from each of the canine clinical trials and were then transformed from the ELISA O.D. readings into a readily understandable data set (units/mL). Known values of standards for each of the initial 6 and the eventual 24 purified food extracts were used to create a baseline standard curve for each of the canine IgA and IgM antibodies, and for each food allergen (i.e. a total of 48 standard curves).

Saliva was collected from either or both sides of the mouth onto the same simple dental cotton rope, 5 inches long by $3 / 8$ inch diameter (e). The saliva-soaked cotton rope was placed in the inner plastic tube of a special double-sleeved collection tube (f), and the tube was capped and taped for additional security while in transit. The samples were
then shipped by regular or Air Mail post to the Hemopet laboratory for testing. Saliva samples were stable for at least 30 days in this sealed tube system, as established by mailing samples nationally and internationally from and to Hemopet. After centrifugation, the saliva samples could be tested immediately, refrigerated for up to 30 days, or frozen at $-20^{\circ} \mathrm{C}$ for later assay; in-house quality control testing of these 3 storage temperatures gave comparable results. For the initial parallel studies comparing results for saliva and serum from Hemopet's resident Greyhounds ( $\mathrm{n}=39$; 29 healthy, 10 with IBD), both saliva and blood samples ( 6 mL whole blood) were collected. After clotting, the serum ( $\sim 2.5$ mL ) was harvested and used for parallel food antigen testing using serum anti-IgG.

## Saliva and Blood Collection

Each of the clinical validation and clinical study cohort dogs had saliva collected with the dental cotton rope. This rope allowed for collection of up to 2 mL of saliva.

## Test Methodology

Assays for Salivary anti-IgA and anti-IgM, and Serum anti-IgG were performed using the specific ELISA Food AntigenCoated Plates containing the 24 affinity-purified food antigens manufactured for this purpose (g). Standard ELISA methodology using a robotic immunoassay autoanalyzer (Tecan [h]) was applied to each of the custom-made food antigen-coated plates. The food antigens were: barley, beef, chicken, corn, duck, egg (hen), lamb, lentil, millet, milk (cow), oatmeal, peanut, pork, potato, quinoa, rabbit, rice, salmon, soy, sweet potato, turkey, venison, wheat, and white-colored fish.


Figure 1a: Nutriscan Standard Dilution Curve for Chicken with $\lg A$

Each of the sealed, refrigerated 96 well ELISA food antigencoated plates was tested in turn, in duplicate, along with the diluent buffer blanks.

Three antibody conjugates were used purified goat antidog IgA, goat anti-dog IgG1, and goat anti-dog IgM, all conjugated to alkaline phosphatase (i).

Quality control was performed by the addition of serum or saliva with low, medium, and high titers of antibodies. In addition, plates were studied for the detection of previously established non-specific reactions to the microwell plates. Without the addition of serum or saliva, the plates underwent the complete ELISA procedure to verify that there was no evidence of nonspecific binding. The plates were stored refrigerated.

## Analysis of Results

Results were analyzed initially as a Panel of 6 antigens, and then subsequently as 2 Panels of 12 antigens each. Calibration graphs for anti-canine IgG in serum and anti-canine IgA +IgM in saliva were obtained from the O.D. values resulting from the blank and negative reactor control samples. The O.D. readings were then converted to units $/ \mathrm{ml}$. The concentration values of anti-canine $\operatorname{IgG}$, $\operatorname{IgA}$, or $\operatorname{IgM}$ were compiled for the initial and subsequent sets of dietary antigens for each healthy control dog and canine patient. The degrees of food reactivity were determined from the calibration slopes measured for each of the 24 foods tested and were then converted to units/ mL for ease of reporting and comprehension (see examples of standard curves in Figure 1). The O.D. values of the duplicate assays for both anti-IgA and anti-IgM were considered acceptable if the coefficient of variation (CV) was not more


Figure 1b: Nutriscan Standard Dilution Curve for Potato with IgM
than $15 \%$ between duplicates (most duplicates had CVs below $5 \%$; any samples with values above $15 \%$ were repeated). The sensitivity and specificity of the assay were $95.5 \%$ (range 9399) and $70.7 \%$ (range 69-72\%), respectively. The likelihood ratios ranged from $3.08-5.30 \%$ for positive ratios and $0.63-$ $0.65 \%$ for the negative ratios (10).

Samples that tested below the $0.63 \%$ negative likelihood ratio cut-off level were clearly negative. Values at or above the $5.30 \%$ positive likelihood ratio cut-off level showed varying degrees of reactivity to the foods tested. Because low level antibody concentrations at or just above the cut-off amount of 10 units $/ \mathrm{mL}$ included some mild or equivocal reactor cases, the lower limit was set at 10 units/mL to avoid the potential misclassification of weak (equivocal) samples as being truly positive. In such cases, the recommendation was made to retest the dog's saliva in 4-6 months. Thus, a $10-11.4$ units/mL amount was set as the range for a weak degree of food sensitivity (clinical significance unclear, if any); and any level at or above 11.5 units/mL indicated a
positive reaction. The positive reaction was then further classified by degree as being borderline, intermediate, medium, or strong food sensitivity. This classification paralleled what is typically used for food sensitivity testing of humans in Europe (d).

Statistical analyses of results were determined using the standard statistical paired t test formulas on Microsoft Excel.

## Results

In the initial clinical validation trials involving 29 healthy Greyhounds and 10 with IBD, anti-IgA and anti-IgM food reactivities were recorded to a varying degree in saliva for the 6 foods tested (beef, corn, cow milk, hen egg, soy and wheat; data not shown). By contrast, none of these dogs had detectable anti-IgG levels with any of the 6 foods tested.

Table 1 shows a typical patient report and illustrates the varying levels of food reactivities for the IgA and IgM antibodies of 24 different foods.

Table 1. Sample Patient Report for Saliva-Based Food Sensitivity Test with 24 Food Antigens

| Accession No. | Doctor | Owner |  | Pet Name | Received |
| :---: | :---: | :---: | :---: | :---: | :---: |
| Test 00116 | Sample Report | Sample Report |  | Sample Report | 10/21/13 |
| Species | Breed | Sex | Weight | Pet Age | Reported |
| Canine | Golden Retriever | FS | 45 Lbs | 5 Yrs | 10/31/13 |
| Diet | Medication | Thyroid Medication | How much medication? | How Often? | Post Pill Timing |
| Raw diet | None | No |  |  |  |
| Test Requested |  | Result | Remark | General Range | Units |
| Beef Salivary IgA |  | 9.500 | Negative Reaction | $<10$ | $\mathrm{U} / \mathrm{mL}$ |
| Beef Salivary IgM |  | 8.125 | Negative Reaction | $<10$ | $\mathrm{U} / \mathrm{mL}$ |
| Chicken Salivary IgA |  | 15.698 | Strong reaction; Avoid | $<10$ | $\mathrm{U} / \mathrm{mL}$ |
| Chicken Salivary IgM |  | 15.524 | Strong reaction; Avoid | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Corn Salivary IgA |  | 8.256 | Negative Reaction | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Corn Salivary IgM |  | 9.635 | Negative Reaction | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Duck Salivary IgA |  | 7.456 | Negative Reaction | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Duck Salivary IgM |  | 6.963 | Negative Reaction | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Lamb Salivary IgA |  | 6.235 | Negative Reaction | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Lamb Salivary lgM |  | 4.653 | Negative Reaction | $<10$ | $\mathrm{U} / \mathrm{mL}$ |
| Milk Salivary IgA |  | 8.563 | Negative Reaction | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Milk Salivary IgM |  | 8.523 | Negative Reaction | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Pork IgA |  | 9.636 | Negative Reaction | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Pork IgM |  | 9.356 | Negative Reaction | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Soy Salivary IgA |  | 7.562 | Negative Reaction | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Soy Salivary lgM |  | 6.235 | Negative Reaction | $<10$ | $\mathrm{U} / \mathrm{mL}$ |
| Turkey Salivary lgA |  | 11.569 | Borderline Reaction; Avoid | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Turkey Salivary lgM |  | 12.375 | Intermediate reaction, Avoid | < 10 | $\mathrm{U} / \mathrm{mL}$ |
| Venison Salivary IgA |  | 8.522 | Negative Reaction | $<10$ | $\mathrm{U} / \mathrm{mL}$ |
| Venison Salivary lgM |  | 7.563 | Negative Reaction | < 10 | $\mathrm{U} / \mathrm{mL}$ |

Table 1. Sample Patient Report for Saliva-Based Food Sensitivity Test with 24 Food Antigens - CONTINUED

| Accession No. | Doctor | Owner | Pet Name | Received |  |
| :--- | :--- | :--- | :--- | :--- | :--- |
| Test 00116 | Sample Report | Sample Report |  | Sample Report | 10/21/13 |
| Species | Breed | Sex | Weight | Pet Age | Reported |
| Canine | Golden Retriever | FS | 45 Lbs | 5 Yrs | 10/31/13 |
| Diet | Medication | Thyroid Medication | How much medication? | How Often? | Post Pill Timing |
| Raw diet | None | No |  |  |  |

Reason for testing: food intolerance, scratching, soft stool

| Test Requested | Result | Remark | General Range | Units |
| :---: | :---: | :---: | :---: | :---: |
| Wheat Salivary IgA | 7.652 | Negative Reaction | <10 | U/mL |
| Wheat Salivary IgM | 9.500 | Negative Reaction | $<10$ | U/mL |
| White Fish Salivary IgA | 8.248 | Negative Reaction | $<10$ | U/mL |
| White Fish Salivary IgM | 7.256 | Negative Reaction | $<10$ | U/mL |
| Barley Salivary IgA | 7.125 | Negative Reaction | $<10$ | U/mL |
| Barley Salivary IgM | 6.359 | Negative Reaction | $<10$ | U/mL |
| Egg Salivary IgA | 7.974 | Negative Reaction | $<10$ | U/mL |
| Egg Salivary IgM | 8.252 | Negative Reaction | $<10$ | U/mL |
| Lentil Salivary IgA | 7.154 | Negative Reaction | $<10$ | U/mL |
| Lentil Salivary IgM | 5.235 | Negative Reaction | <10 | U/mL |
| Millet Salivary IgA | 9.256 | Negative Reaction | $<10$ | U/mL |
| Millet Salivary IgM | 10.254 | Weak Reaction | $<10$ | U/mL |
| Oatmeal Salivary IgA | 12.356 | Intermediate reaction, Avoid | $<10$ | U/mL |
| Oatmeal Salivary IgM | 12.457 | Intermediate reaction, Avoid | $<10$ | U/mL |
| Peanut Salivary IgA | 7.281 | Negative Reaction | $<10$ | U/mL |
| Peanut Salivary IgM | 8.643 | Negative Reaction | $<10$ | U/mL |
| Potato Salivary IgA | 10.120 | Weak Reaction | $<10$ | $\mathrm{U} / \mathrm{mL}$ |
| Potato Salivary IgM | 9.625 | Negative Reaction | $<10$ | U/mL |
| Quinoa Salivary IgA | 9.365 | Negative Reaction | <10 | U/mL |
| Quinoa Salivary IgM | 8.453 | Negative Reaction | $<10$ | U/mL |
| Rabbit Salivary IgA | 5.423 | Negative Reaction | $<10$ | U/mL |
| Rabbit Salivary IgM | 4.536 | Negative Reaction | $<10$ | $\mathrm{U} / \mathrm{mL}$ |
| Rice Salivary IgA | 8.451 | Negative Reaction | $<10$ | U/mL |
| Rice Salivary IgM | 8.263 | Negative Reaction | $<10$ | $\mathrm{U} / \mathrm{mL}$ |
| Salmon Salivary IgA | 11.258 | Weak Reaction | $<10$ | U/mL |
| Salmon Salivary lgM | 14.653 | Medium Reaction; Avoid | $<10$ | U/mL |
| Sweet Potato IgA | 7.124 | Negative Reaction | $<10$ | U/mL |
| Sweet Potato IgM | 8.364 | Negative Reaction | $<10$ | U/mL |

## RECOMMENDATIONS

Food reactions were seen to: Chicken, Turkey, Oatmeal and Salmon. A strong reaction was present for Chicken.
Please avoid feeding these foods.
Interpretation: Pet should avoid food or treats containing ingredient(s) showing results of 11.5 or greater.
Recommend rechecking salivary food sensitivity or intolerance levels every 6-12 months.

## Degree of reactivity:

$<10 \mathrm{U} / \mathrm{mL}$ indicates a normal food antigen tolerance level = negative result.
$10-11.4 \mathrm{U} / \mathrm{mL}$ indicates a weak reaction; clinical significance unclear
11.5-11.9 $\mathrm{U} / \mathrm{mL}$ indicates an borderline reaction

12-12.9 U/mL indicates an intermediate reaction
13-14.9 U/mL indicates a medium reaction
$>/=15 \mathrm{U} / \mathrm{mL}$ indicates a strong reaction
Differences between antibodies to $\lg A$ and $\lg M$ : Antibodies to $\lg A$ measure the secretory immunity from body secretions (tears, saliva, feces, urogenital tract). They act as a mechanical barrier or the "first line of defense" to help protect the bowel from invasion by foreign substances, infectious agents, chemicals, and certain foods that it cannot or poorly tolerate. Antibodies to lgM measure the body's primary immune response to a recent exposure within the last 6 months or so (e.g. to a certain food ingredient).

Table 2 summarizes the age and health status for food sensitivity of 345 dogs from the initial 566 clinical case cohort (ages for the remaining cases were either not stated or unknown). The data show that most of the healthy dogs were between $1-2$ yrs of age with $2-3$ yrs being the next highest age group. For the suspect IBD cases, most of the dogs were between $1-4 \mathrm{yrs}$ of age or over 10 yrs old, whereas in the proven IBD cases, most dogs were over 10 yrs of age.

Table 2. Summary of 345 Cases by Age and Health Status for Food Sensitivity

| Age (yrs) | Heath Status for Food Sensitivity |  | Totals |  |
| :---: | :---: | :---: | :---: | :---: |
|  | Healthy | Suspect |  |  |
|  | $\mathbf{n = 1 0 3}$ | $\mathbf{n}=\mathbf{1 6 5}$ | $\mathbf{n}=\mathbf{7 7}$ |  |
| $<1$ | 3 | 9 | 2 | 14 |
| $1-2$ | 28 | 30 | 8 | 66 |
| $2-3$ | 16 | 27 | 6 | 49 |
| $3-4$ | 10 | 20 | 7 | 37 |
| $4-5$ | 8 | 11 | 7 | 26 |
| $5-6$ | 9 | 12 | 4 | 25 |
| $6-7$ | 3 | 12 | 6 | 21 |
| $7-8$ | 3 | 11 | 7 | 21 |
| $8-9$ | 5 | 5 | 8 | 18 |
| $9-10$ | 8 | 4 | 5 | 17 |
| $>10$ | 10 | 24 | 17 | 51 |

Table 3 summarizes the breed type of 420 dogs from the initial 566 clinical case cohort (breeds for the remaining cases were unknown). The highest number of cases were in breeds stated on the submission form (111 cases), or in miscellaneous breeds where there were fewer than 10 cases each (99). The German Shepherd Dog had more cases (48) than any other affected breed, and 13 of them were of the white German Shepherd variety. Golden Retrievers (32) ranked second, followed by Labrador Retrievers (22) and mixed breeds (20). There were also 39 Hemopet Greyhounds included in the study. The data for the larger second clinical case cohort ( $\mathrm{n}=1008$ ) showed no differences between the intermediate, medium, and strong anti-IgA or anti-IgM antibody reactivity levels for the following food antigens: lamb, oatmeal, potato, quinoa, rabbit, turkey, wheat, or white-colored fish, so these data were combined for further analyses. For other food antigens, namely barley, beef, hen egg, and venison, anti-IgA reactivity levels were observed higher than those of anti-IgM. Similarly, for the food antigens chicken, corn, cow milk, millet, peanut, rice, and soy, anti-IgM reactivity levels were observed higher than those of anti-IgA.

## Veterinary Botanical Medicine Association

the owly protessional cssocidion for velerinary botan
medicine - our members are the best informed in the field!
Members enjoy the following:

- Interactive and informative listserv
- Professional journal
- Herbal wiki (a Wikipedia for herbs)
- Professional listing in our online referrals list
-Helpful and hard-to-find client handouts
- Discounts on specialized CE
- Ecotours to exotic locales

Integrating science and tradition * Expanding treatment options - Providing professional herbal resources

## VBMA.org I Office@VBMA.org



Table 3. Summary of 420 Cases Tested by Breed

| Breed | Number of Cases |
| :--- | :---: |
| Breed Not Stated | 111 |
| German Shepherd Dog [13 = White GSD] | 48 |
| Golden Retrievers | 32 |
| Labrador Retrievers | 22 |
| Mixed Breed | 20 |
| Bernese Mountain Dog | 14 |
| Standard Poodle | 14 |
| Doberman Pinscher | 11 |
| German Shorthaired Pointer | 10 |
| Greyhounds (pre-selected, Hemopet) | 39 |
| Miscellaneous Breeds (less than 10 cases) | 99 |

The intermediate, medium, and strong anti-IgA or anti-IgM antibody reactivity levels of the 1008 cases, when analyzed per 100 cases to permit more direct relative comparisons of the number of positive reactions, showed the following: highest number of reactions $=$ white-colored fish (18); turkey (15); venison (13); corn (12); and hen's egg (11).

The lowest reacting foods for these positive reacting cases were: wheat (5); peanut (4); rice and lamb (each 3); and beef (2).

In comparison to the intermediate, medium, and strong anti-IgA or anti-IgM antibody positive reactivity levels, negative or weak antibody reactivity levels when analyzed per 100 cases showed the following: highest number = wheat (50); lamb, peanut, and rice (each 48); potato (47); beef, chicken, oatmeal, quinoa, and salmon (each 46). The lowest reacting foods for the non-reactive cases were: turkey and venison (each 43), and white-colored fish (41).

Table 4 summarizes the clinical outcome comparisons of 50 cases selected in sequence from the reported data set both before and after eliminating the reactive foods. This group consisted of a wide spectrum of breed types and sizes, and ages varied from 5 months to 14 years of age. The clinical outcomes after removing the reactive foods, based upon follow up interviews with the client and/or submitting veterinarian, varied from good (2 cases), very good (14 cases) to excellent ( 33 cases), with one dog showing no improvement (Table 4).

Table 4. Clinical Outcomes Before \& After Eliminating Reactive Foods (50 Cases)

| Case Breed | Age (yrs) | Sex | Clinical <br> History |  | Initial Results* (Reactive Foods) | Follow Up Results $\dagger$ Clinical Outcome $\ddagger$ |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Lhasa Apso | 2 | FS |  | X | Q, SP | E |  |
| Great Dane | 1 | F | X |  | BA, P, Q, RI, V, WF | E |  |
| Newfoundland | 1.5 | M | X | $X$ | BA, MI, O, P, PO, Q, RA, RI, SA | E |  |
| Wire Fox Terr. | 8 | MN |  | X | BA, MI, O, P, Q, RA, S | G |  |
| Terrier Mix | 3 | FS |  | X | $B A, E, M I, O, P, P E, Q, R A, R 1, S$ | VG |  |
| Min. Aussie. | 2.5 | MN | X |  | BA, C, CO, M, O, P, RA, SA, T, V, W, WF | E |  |
| Rottweiler | 2.5 | FS | X |  | $\mathrm{BA}, \mathrm{CH}, \mathrm{CO}, \mathrm{M}, \mathrm{MI}, \mathrm{O}, \mathrm{PE}, \mathrm{Q}, \mathrm{RI}, \mathrm{SO}, \mathrm{T}, \mathrm{V}, \mathrm{W}$ | E |  |
| Toy Fox Terr. | 14 | MN | X |  | BA, BE, CH, CO, L, M, MI, O, P, PE, Q, SO, T, V, W, WF | E |  |
| Havanese | 3 | MN | X | X | CH, M, V | E |  |
| York. Terrier | 8 | MN | X |  | CO, M, T, V, WF | E |  |
| Std. Poodle | 0.75 | MN | X | X | BE, CH, W | VG |  |
| Tibetan Terr. | 2.5 | FS | X |  | CO, V, Q | VG |  |
| Tibetan Terr. | 1.5 | FS | X |  | CH, CO, M, T, V, W, WF | VG |  |
| Shih Tzu | 5 | MN | X |  | CO, P, Q, RI |  | X |
| Shih Tzu | 4 | FS | X |  | CO, Q, RI, SA, V | VG |  |
| Basenji | 8 | FS |  | X | CH, CO, M, T, V, W, WF | E |  |
| Greyhound | 11 | MN |  | X | BA, CH, CO, M, MI, O, P, Q, RA, RI, SA, T, V, WF | E |  |
| Cairn Terrier | 5 | M |  | X | CH, O, P, SA | E |  |

Table 4. Clinical Outcomes Before \& After Eliminating Reactive Foods (50 Cases) - CONTINUED

| Case Breed | Age (yrs) | Sex | Clinical History GI Skin |  | Initial Results * (Reactive Foods) | Follow Up Results $\dagger$ Clinical Outcome $\ddagger$ Improved No $\Delta$ |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Greyhound | 4 | FS | X |  | BA, CH, CO, M, P, RA, T, V, W, WF | E |  |
| Glen of Imal Terr. | 3.5 | FS | X |  | BE, M, SO, W | E |  |
| Labradoodle | 4 | M |  | X | BA, CH, MI, O, T, V, WF | E |  |
| PONS | 1.5 | F | X |  | V | E |  |
| Greyhound | 4 | FS |  | X | BE, $\mathrm{CO}, \mathrm{E}, \mathrm{M}, \mathrm{SO}, \mathrm{W}$ | E |  |
| White Boxer | 2.5 | MN |  | X | BE, CH, CO, SO, W, WF | G |  |
| York. Terrier | 7.5 | MN | X |  | CH, CO, M, T, V, W, WF | E |  |
| Lab. Mix | 10.5 | FS | X |  | CH, CO, E, M, MI, P, RI, T, V, W, WF | E |  |
| Gr. Swiss Mtn. Dog | 7.3 | FS | X |  | BA, BE, CO, E, LE, MI, O, P, Q, R, RI, SA, SP, V, WF | E |  |
| Belg. Tervuren | 3.25 | M | X |  | BA, BE, CH, M, O, SA, SO, T, V, W, WF | E |  |
| Siberian Husky | 8.9 | M |  | X | BE, CH, M, P, T, V | E |  |
| Boxer/Catahoula Leopard Dog | 6 | FS | X |  | O, Q, RI, V, W, WF | E |  |
| NSDTR | 1 | F |  | X | BE, CO, M, SO, W | E |  |
| Lab/Pointer | 12.9 | M |  |  | MI, O, P, Q, RA, SA, V, WF | VG |  |
| Goldendoodle | 2 | MN | X |  | BA, CO, LE, MI, O, P, SA, V, W, WF | E |  |
| Beagle Mix | 6 | MN | X |  | V, WF | VG |  |
| Belg. Malinois | 1.5 | MN | X |  | PO, V, WF | E |  |
| Wire Fox Terr. | 4.5 | MN |  | X | M, PO, T, V, WF | VG |  |
| Brittany Sp. | 6.5 | MN | X |  | CH, CO, M, T, V, WF | E |  |
| Flat Coat Retr. | 7.7 | MN | X |  | BA, D, E, LE, M, MI, O, PE, PO, Q, RI, SA, SO, SP, T, V, W, WF | E |  |
| York. Terrier | 4.9 | MN | X |  | CH, T, V, WF | E |  |
| French Bulldog | 0.45 | M | X | $X$ | BA, D, LE, M, MI, O, P, PE, PO, RA, RI, SA, T, V, W, WF | E |  |
| German Shepherd | 3.75 | MN |  | X | CH, CO, E, M, MI, O, PE, PO, RA, SA, T, V, W, WF | VG |  |
| Golden Retr. | 7 | FS | X | X | CH, CO, M, SO, T, V, W, WF | VG |  |
| Irish Setter | 11 | MN |  | X | CH, CO, MI, O, P, PO, Q, RA, SA, V, W, WF | E |  |
| Irish Setter | 9 | M | X | X | BA, CH, CO, LE, T, V | VG |  |
| German Shepherd | 2.6 | FS | X |  | WF | VG |  |
| Scottish Terr. | 1.5 | FS |  | X | CO, E, LE, O, Q, RA, T, WF | VG |  |
| Eng. Bulldog | 2 | M | X | X | BE, CH, CO, PO, RA, SA, T, V, W | E |  |
| Border Collie | 9.5 | MN | X |  | E, SP, T, V | VG |  |
| Bernese Mtn. Dog | 3.25 | F | X | X | BE, CH, CO, MI, P, RI, SO, T, V, W, WF | E |  |
| Golden Retr. | 5.75 | M | X |  | BA, CH, CO, M, SO, T, V, W, WF | E |  |

* $\mathrm{BA}=$ barley, $\mathrm{BE}=$ beef, $\mathrm{CH}=$ chicken, $\mathrm{CO}=$ corn, $\mathrm{D}=$ duck, $\mathrm{E}=$ egg, $\mathrm{L}=$ lamb, $\mathrm{LE}=$ lentil, $\mathrm{M}=$ milk, $\mathrm{MI}=$ millet, $\mathrm{O}=$ oatmeal, $\mathrm{PE}=$ peanut, $\mathrm{PO}=$ pork, $\mathrm{P}=$ potato, $\mathrm{Q}=$ quinoa, $\mathrm{R}=$ rabbit, $\mathrm{RI}=$ rice, $\mathrm{SA}=$ salmon, $\mathrm{SO}=$ soy, $\mathrm{SP}=$ sweet potato, $\mathrm{T}=$ turkey, $\mathrm{V}=$ venison, and $\mathrm{W}=$ wheat, $\mathrm{WF}=$ white-colored fish.
* $\dagger$ Reactive Foods Removed. $\ddagger \mathrm{E}=$ excellent, $\mathrm{VG}=$ very good, $\mathrm{G}=$ good, $\Delta=$ change

Table 5 describes results for the 15 cases for which salivary food diagnostic testing was repeated by the owners several months later. Owners of the other cases from Table 4 elected not to retest their dogs because they stated them to be clinically improved upon removing the prior reactive foods from the diet. A variety of breeds and dog sizes were represented here with ages varying from 10 months to 8.75
years. The follow up saliva-based food test results and clinical outcomes, based again upon follow up interviews with the client and/or submitting veterinarian, varied from very good ( 5 cases) to excellent ( 10 cases). The initially reactive foods were mostly non-reactive on retesting, although some newly reactive foods also were identified upon retesting.

Table 5. Initial and Follow Up Test Results After Eliminating Reactive Foods (15 Cases)

| Case Breed | $\begin{aligned} & \text { Age } \\ & \text { (yrs) } \end{aligned}$ | Sex | Clinical <br> History <br> GI Skin |  | Initial Results * <br> (Reactive Foods) | Follow Up Results $\dagger$ (Reactive Foods) | Clinical <br> Outcome $\ddagger$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Boston Terrier | 8.7 | M |  | X | BE, CH, CO, M, SO, T, W | CH, T | VG |
| Boston Terrier | 6 | M |  | X | BA, CH, CO, M, O, P, T, V, W, WF | BE | E |
| German Shepherd | 3 | M | X | X | BE, CH, E, Q, RA, SA, T, V | BE, CH, CO, M, PO, T, V, WF | E |
| German Shepherd | 5 | FS | X | $X$ | BE, CO, M, SO, W | CO, E, M, MI, SA, WF | E |
| Irish Setter | 5 | M | X |  | BA, CH, CO, E, M, MI, O, PE, PO, P, Q, RA, RI, SA, SO, T, V, W, WF | $B A, B E, C O, D, L E, M, ~ O, ~ P E, ~ P O, ~ P, ~ Q, ~ R I, ~ S O, ~ T, ~ V, ~ W, ~ W F ~$ | E |
| Border Collie X | 0.8 | FS | X |  | $B A, B E . C H, C O, D, E, L E, M, M I, O, P E, P O, P, Q, R A, R I, S A, S O, S P$, T, V, W, WF | CH, CO, T, WF | E |
| Gr. Dane/Dogo | 3.75 | M | X |  | $B A, B E, C H, C O, D, E, L A, L E, M, M I, O, P E, P O, P, Q, R A, R I, S A, S O$, $S P, T, V, W, W F$ | BA, E, LE, MI, 0, PE, PO, Q, RA, RI, SA, SP | E |
| Labrador Retr. | 4.25 | MN |  | X | CO, T, V | T, WF | VG |
| Doberman Pin. | 5.5 | FS | $X$ |  | CO, M, SO, T, V, WF | V, WF | E |
| Min. Poodle | 6.5 | M | X |  | BE, M, W | V, WF | VG |
| Glen of Imal Terr. | 3.5 | FS | X |  | BE, M, SO, W | None | E |
| Basset Hound | 7.75 | F | X | X | CO, E, LE, M, MI, P, RI, SP, T, V, W | BE, M, W | E |
| English Setter | 7 | M | X |  | BE, M, W § | CH, CO, M, MI, O, SO, T, V, W, WF | VG |
| English Setter | 7 | FS | X |  | BE, M, W § | CH, CO, M, O, SO, T, V, W, WF | VG |
| NSDTR | 2 | FS |  | X | CH, MI, SO, V, WF | CO, WF | E |

${ }^{*} \mathrm{BA}=$ barley, $\mathrm{BE}=$ beef, $\mathrm{CH}=$ chicken, $\mathrm{C} O=$ corn, $\mathrm{D}=$ duck, $\mathrm{E}=$ egg, $\mathrm{L}=\operatorname{lamb}, \mathrm{LE}=$ lentil, $\mathrm{M}=$ milk, $\mathrm{MI}=$ millet, $\mathrm{O}=$ oatmeal, $\mathrm{PE}=$ peanut, $\mathrm{PO}=$ pork, $\mathrm{P}=$ potato,
$\mathrm{Q}=$ quinoa, $\mathrm{R}=$ rabbit, $\mathrm{RI}=$ rice, $\mathrm{SA}=$ salmon, $\mathrm{SO}=$ soy, $\mathrm{SP}=$ sweet potato, $\mathrm{T}=$ turkey, $\mathrm{V}=$ venison, and $\mathrm{W}=$ wheat, $\mathrm{WF}=$ white-colored fish.
$\dagger$ Reactive Foods Removed; then 2-6 months retesting, reactions were lower or negative.
$\ddagger \mathrm{E}=$ excellent, $\mathrm{VG}=$ very good $\S$ Only 6 foods tested $(\mathrm{BE}, \mathrm{CO}, \mathrm{E}, \mathrm{M}, \mathrm{SO}, \mathrm{W})$

Figures 2 and 3 illustrate the physical differences in 2 dogs shown before and after offending foods identified by the saliva testing had been removed from their diets. These
remarkable beneficial effects were clearly seen within 2 weeks of the diet changes, and the original issues were completely resolved within a month.



Figure 2a: Cattle Dog Mix One Week After Nutriscan \& Reactive Food Removal


Figure 2a: Cattle Dog Mix One Month After Nutriscan \& Reactive Food REmoval RevalPicture2


Figure 2b: Cattle Dog Mix Before Nutriscan


Figure 2b: Cattle Dog Mix One Month After Nutriscan \& Reactive Food Removal


Figure 3a: Aussie Before Nutriscan


Figure 3b Aussie Skin \& Muscle Before Nutriscan

## Discussion

Saliva is a source of body fluid for detection of an immune response to bacterial, food, and other antigens present in the oral cavity and GI tract (1, 7-9, 11, 12). Indeed, salivary antibody induction has been widely used as a model system to study secretory responses to ingested material, primarily because saliva secretion is simple and easy to collect and analyze (1-3, 7, 12-14).

The results presented here using a novel saliva-based test which quantified the IgA and IgM antibody responses to 24 affinity-purified food antigens convincingly demonstrated the clinical predictability, utility, and efficacy of the assay (Tables 4 and 5; Figures 2 and 3). As shown in Table 5, the assay was repeated once more in 15 cases, 2-6 months after the identified offending foods had been completely removed from the diet. In each case, the clinical outcome was stated to be very good ( 5 cases) and excellent ( 10 cases). Even though some of the initially reactive foods were still quantified as reactive on retesting (at or above 11.5 units $/ \mathrm{mL}$ ), the degree of reactivity was lower. It is interesting that on repeat testing, some other food antigens were quantified as reactive, suggesting that these dogs could be especially prone to developing food intolerances. Several possible explanations for the ongoing but lowered reactivity of initially reactive


Figure 3a Aussie Before Nutriscan


Figure 3b Aussie After Nutrscan \& Reactive Food Removal
food antigens and the appearance of additional reactive foods include: initial reactions were actually to residues in the flesh from what the meat or fish ate before becoming a food source; and reactive foods were still present albeit it in presumed smaller amounts in supplements the dog still ate, such as chicken fat, cornstarch, and fish oils (1, 2, 15-18).

Food intolerance is stated to be the third most commonly recognized syndrome in dogs after flea bite sensitivity and atopy (inhalant allergy), and food intolerance makes up an estimated $10-15 \%$ of all allergic skin disease (3). It mimics other skin syndromes. Food intolerance is stated to have no age, sex, or breed predilection, although clinical experience indicates that it can be familial $(2,3)$. In the author's experience, most affected animals had been eating the offending foods for more than 2 years; the major complaint of their owners was bilateral pruritus, and there was often otitis externa. Secondary skin disease such as seborrhea (both dry or oily) and pyoderma was also common (2, 3).

Delayed food sensitivities in people are extremely common and can be manifested by GI, neurological, pulmonary, dermatologic, ear, nose, throat, musculoskeletal, genitourinary, cardiovascular, and endocrine problems (7, 8). For dogs, in addition to the commonly observed GI tract
signs of food sensitivity, the skin is frequently a concurrent or alternate tissue target (1-3).

Creating a healthy acid-base balance within tissues through optimal nutrition should be the goal of case management and therapy (19). When eaten, different foods produce varying metabolic waste products and by-products. Most foods are acid-forming, with the exception of nuts and seeds, so that when people and animals eat more nuts and seeds, the dietary by-products are alkaline and promote health and prevent disease (19-21). Changing the proportions of macro-nutrients and micronutrients in different nutrient and food products is important in obtaining the right tissue and gut balance $(4,19)$. To be effective, diets ideally need to be individualized using nutrigenomic principles (2, 4, 20-23). Studies have indicated that specialized nutrient intake extends and improves life, delays onset and slows progression of disease, and enhances the quality of life of animals (2, 4, 20, 24).

Avoiding additives and supplements, as well as avoiding frequent switching from diet to diet, is important too, as up to $20 \%$ of cases have concurrent other GI tract issues (2-4). Some canine cases have swollen peripheral lymph nodes, although this sign is more common in affected cats. Affected pets may exhibit tension-fatigue, malaise, and dullness. Effects are usually non-seasonal, and the primary disorder is poorly responsive to steroids (3).

The so-called "gold standard" for food sensitivity or intolerance until now has been either diet elimination trials for 3-12 weeks, micronized or hydrolysed prescription diets, skin patch testing considered by clients to be expensive and unsightly, and allergen provocation (20-34). But, even these specialized, limited ingredient diets have been found to contain ingredients not listed on the label, and there is often poor compliance with the diet elimination trial approach (15-17, 25). The alternative diagnostic approach of performing serum allergy tests for food sensitivity is typically based on measuring IgE, IgG, and immune complexes bound to complement; these tests have high sensitivity but lower individual specificity, and measure only more immediate-type reactions (25-34). Dogs with atopic and GI tract disease have higher levels of serum $\operatorname{IgE}$ and IgG antibodies than normal dogs, and the antigen(s) causing the reaction is often contained in the diet (2527). However, there is generally poor correlation between
serum IgE and IgG antibody testing and clinical experience in resolving disease in both humans and dogs (25-34).

Immune complexes containing large food antigens enter the blood from the GI tract and then travel through the liver where most immune complexes are removed. However, if circulating immune complexes pass the liver filtering system, they may cause injury to many body tissues $(7,8$, 12). Malabsorption of food particles from the GI tract can also travel by lymphatic drainage to the body $(4,12)$. The lymph channels in the gut wall converge at the thoracic duct, which drains its contents into the large thoracic veins. This combination of antibody with complement in the blood stream becomes a circulating immune complex. Immune complexes subsequently attach to receptors on red and white blood cells. These altered cells are cleared by the body's liver or spleen (reticuloendothelial system) $(5,7,8,12)$.

Any circulating immune complexes that are not removed by the reticuloendothelial system of the liver (or spleen) can activate the complement cascade. Individuals with more immune complexes on their red blood cells are the ones that can experience chronic food sensitivities or intolerances (5-7). Circulating immune complexes also can damage the integrity of blood vessel capillaries which in turn can trigger inflammatory events $(7,8)$.

Newer testing for food sensitivity has used serum, saliva or feces [j-for people only] in a simple ELISA format or other immunoassay platform (1-3, 31-35). These methods identify IgG, IgA, or immune complexes to foods in serum, and IgA or IgM antibodies to foods in saliva. As antibodies to foods usually appear in saliva several months before the GI tract diagnosis of IBD or the "leaky gut syndrome" (intestinal dysbiosis), saliva testing can thus reveal the latent or pre-clinical form of food sensitivity ( $1-3,5,7,13$, $33,35)$. IgA, especially, but also IgM, are the important antibodies generated by immunological reactions and are expressed as secretory immunity in saliva, as well as other body fluids like tears, sweat, and breast milk $(8,13,14)$. IgE serology has been found to offer no advantage for diagnosis when performing dietary trials because it had a sensitivity of $14 \%$, specificity of $87 \%$, positive predictive value of $40 \%$, and negative predictive value of $61 \%$ (20-22). Thus, this form of serum food allergy testing is clearly inadequate for clinical diagnostic purposes.

## Conclusion

By looking at secretory immune responses to specific food antigens, detected as salivary antibodies to IgA and IgM in

## Endnotes

a. NutriScan ${ }^{\circledR}$, Division of Hemolife Diagnostics, Garden Grove, CA 92843; www.nutriscan.org
b. Foundation Formula, Precise Pet Products, Nacogdoches, TX
c. Wellness Pet Food, WellPet, Tewksbury, MA
d. DST, Diagnostic Systems \& Technologies GmbH, Schwerin, Germany

## References

1. Dodds WJ. Food Intolerance: Diagnostic testing \& dietary management. J Am Hol Vet Med Assoc. 2014;36:36-42.
2. Dodds, WJ, Laverdure, DR. Canine Nutrigenomics: The New Science of Feeding Your Dog for Optimum Health. 2015. DogWise Publishing, Wenatchee, WA.
3. Dodds WJ. Functional foods: The new paradigm based on nutrigenomics. J Am Hol Vet Med Assoc. 2014;36:26-35.
4. Fasano A. Zonulin, regulation of tight junctions, and autoimmune diseases. Ann N Y Acad Sci. 2012;1258(1):25-33.
5. Kiyono H, Kweon MN, Hiroi T, et al. The mucosal immune system: from specialized immune defense to inflammation and allergy. Acta Odntol Scand. 2001;59 (3):145-153.
6. Walker, WA, Isselbacher, KJ. Intestinal antibodies. New Engl J Med. 1977; 297:767-773.
7. Robinson, LE, Reeves, S. EpiCor; Review of sIgA's major role as a first line of immune defense and new indications regarding inflammation and gut health. 2013. Retrieved from http://www.embriahealth.com (last accessed 03/27/14). See pdf of article at: https://www.dropbox.com/ s/3iisl0h54lwdeiq/Science_Report__sIga_E6319CD472E10.pdf?dl=0
8. Mesenteric, J, McGhee, JR, Arnold, RR. Selective induction of an immune response in external secretions by ingestion of bacterial antigen. J Clin Invest. 1987;61:731-737.
9. Challacombe SJ. The induction of secretory IgA responses In: Brostoff J, Challacombe S J, eds. Food allergy and intolerance. Eastborne, England: W. B. Sanders Co, 1987.
10. McGee S. Simplifying likelihood ratios. J Gen Intern Med. 2002;17(8):646-649.
11. Kanda M, Inove H, Fukuizumi T, et al. Detection and rapid increase of salivary antibodies to Staphylococcus lentus and indigenous bacterium in rabbit saliva, through a single tonsillar, application of bacterial cells. Oral Microbiol Immunol. 2001;16:257.
12. Husband AM, Gowens JL. The origin and antigen-dependent distribution of IgA containing cells in the intestine. J Exp Med. 1978;148:1146-1160.
13. Lee, YH, Wong, DT. Saliva: An emerging biofluid for early detection of disease. Am J Dent. 2009;22: 421-428.
14. Miller, CS, Foley, JD, Bailey, AL, et al. Current developments in salivary diagnostics. Biomarker Med. 2010;4:171-189.
15. Ricci R, Granato A, Vascellari M, et al. Identification of undeclared sources of animal origin in canine dry foods used in dietary elimination trials. J Anim Physiol Anim Nutr (Berl). 2013;97(Suppl 1):32-38.
16. Raditic DM. Remillard RL, Tater KC. ELISA testing for common food antigens in four dry dog foods used in dietary elimination trials. J Anim Physiol Anim Nutr. 2011;95: 90-97.
17. Parr JM, Remillard RL. Common cofounders of dietary elimination trials contain the antigens soy, pork, and beef. J Am An Hosp Assoc.

## Patents: Issued US patents:

7,867,720; 7,892,763; 8,450,072; 8,450,074; Canadian patents 2,743,714; 2,771,948; and European patent 2382469.
humans and with the current saliva-based testing in dogs, a direct correlation between results and clinical allergenic reactivity to foods can be demonstrated $(1-3,14,35)$. 委
e. Patterson Dental Supply Inc, St. Paul, MN
f. Starstedt Inc., Newton, NC
g. Oxford Biomedical Corporation, Rochester Hills, MI
h. Tecan Group Ltd, Männedorf, Switzerland
i. Bethyl Laboratories, Montgomery, TX
j. EnteroLab, Dallas, TX

2014;50(5):298-304.
18. Dodds WJ. Adjuvants and additives in human and animal vaccines. Med Res Archives. 2016;2(5):1-8.
19. Remer T. Influence of diet on acid-base balance. Semin Dial. 2000;13(4):221-226.
20. Dodds WJ. Epigenetics: programming for health and longevity. J Am Hol Vet Med Assoc. 2014;37:16-22.
21. German, JB, Roberts, MA, Fay, L, et al. Metabolomics and individual metabolic assessment: the next great challenge for nutrition. J Nutr. 2002;132:2486-2487.
22. Daniel, H. Genomics and proteomics: importance for the future of nutrition research. Brit J Nutr. 2002;87:S305-S311.
23. Fekete, SG, Brown, DL. Veterinary aspects and perspectives of nutrigenomics: a critical review. Acta Vet Hungarica. 2007;55(2):229-239.
24. Swanson, K S, Schook, L B. Canine nutritional model: influence of age, diet, and genetics on health and well-being. Current Nutr Food Sci. 2006;2(2):115-126.
25. Zimmer A, Bexley J, Halliwell RE, et al. Food allergen-specific serum IgG and IgE before and after elimination diets in allergic dogs. Vet Immunol Immunopathol 2011; 144:442-447.
26. Bethlehem S, Bexley J, Mueller RS. Patch testing and allergen-specific serum IgE and IgG antibodies in the diagnosis of canine adverse food reactions. Vet Immunol Immunopathol. 2012;145:582-589.
27. Foster AP, Knowles TG, Hotson MA, et al. Serum IgE and IgG responses to food antigens in normal and atopic dogs, and dogs with gastrointestinal disease. Vet Immunol Immunopathol. 2003;92:113-124.
28. Bahna SL, Furukawa CT. Food allergy: diagnosis and treatment. Ann Allergy. 1983;51:574-580.
29. Kagnoff, MF. Effects of antigen feeding on intestinal and systemic immune responses. I. Priming of precursor cytotoxic T-cells by antigen feeding. J Immunol. 1978;120:395-399.
30. Jeffers, JG, Shanley, KJ, Meyer, EK. Diagnostic testing of dogs for food hypersensitivity. J Am Vet Med Assoc. 1991;198:245-250.
31. Buchanan, BB, Frick, OL. The dog as a model for food allergy. Ann NY Acad Sci. 2002;964:173-183.
32. Day, MJ. The canine model of dietary hypersensitivity. Proc Nutr Soc. 2005;64:458-464.
33. Hall ES, Batt RM. Dietary modulation of gluten sensitivity in a naturally occurring enteropathy of Irish setter dogs. Gut 1992;33(2):198-205.
34. Rinkinen, M, Teppo, AM, Harmoinen, J, et al. Relationship between canine mucosal and serum immunoglobulin A (IgA) concentrations: Serum IgA does not assess duodenal secretory IgA. Microbiol Immunol. 2003;47:155-159.
35. Vojdani A. Detection of IgE, IgG, IgA and IgM antibodies against raw and processed food antigens. Nutr \& Metabol. 2009;6:22-37.

