Utility of the combined use of 3 serologic markers in the
diagnosis and monitoring of chronic enteropathies in dogs

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Abstract
Background: Dogs with chronic enteropathies (CE) displayed elevated IgA seropositivity
against specific markers that can be used to develop a novel test.
Objective: To assess a multivariate test to aid diagnosis of CE in dogs and to monitor
treatment-related responses.
Animals: One hundred fifty-seven dogs with CE/inflammatory bowel disease (IBD),
24 dogs non-IBD gastrointestinal disorders, and 33 normal dogs.
Methods: Prospective, multicenter, clinical study that enrolled dogs with gastrointesti-
nal disorders. Serum sample collected at enrollment and up to 3 months follow-up
measuring OmpC (ACA), canine calprotectin (ACNA), and gliadin-derived peptides
(AGA) by ELISA.
Results: Seropositivity was higher in CE/IBD than normal dogs (66% vs 9% for ACA;
55% vs 15% for ACNA; and 75% vs 6% for AGA; P < .001). When comparing CE/IBD
with non-IBD disease, ACA and ACNA displayed discriminating properties (66%, 55%
vs 12.5%, 29% respectively) while AGA separated CE from normal cohorts (54% vs
6%). A 3-marker algorithm at cutoff of ACA > 15, ACNA > 6, AGA > 60 differentiates
CE/IBD and normal dogs with 90% sensitivity and 96% specificity; and CE/IBD and
non-IBD dogs with 80% sensitivity and 86% specificity. Titers decreased after treat-
ment (47%-99% in ACA, 13%-88% in ACNA, and 30%-85% in AGA), changes that
were concurrent with clinical improvements.
Conclusion and Clinical Importance: An assay based on combined measurements of
ACA, ACNA, and AGA is useful as a noninvasive diagnostic test to distinguish dogs
with CE/IBD. The test also has the potential to monitor response to treatment.

KEYWORDS
calprotectin, CE/IBD, GI monitoring, gliadins, OmpC

1 | INTRODUCTION

Gastrointestinal conditions characterized by abdominal pain, vomiting,
and diarrhea are common signs in dogs. Chronic enteropathies (CE) in
dogs are a hetrogenous group of gastrointestinal (GI) disorders
characterized by clinical signs that persist for at least 3 weeks or
longer when other intestinal conditions such as parasitism or neopla-
sia have been ruled out.¹² CE in dogs is perceived to be a common
presentation in all types of veterinary clinics with a prevalence reaching 18% based on databases obtained from general veterinary practices and insurance.3,4 The most common clinical signs for CE in dogs are vomiting and diarrhea, although additional clinical signs including dysrexia and weight loss can manifest. All these signs are nonspecific and overlap with other GI conditions, resulting in a significant challenge with a definitive diagnosis.5

Current methodologies to diagnose CE in dogs require relatively costly, labor intensive and invasive clinical, radiographic, endoscopic, and histological techniques.6 Although these more specialized diagnostic techniques remain the gold standard for diagnosis of CE/inflammatory bowel disease (IBD), the high prevalence of CE in dogs means that the development of novel, noninvasive, easily repeatable approaches to aid in diagnosis and monitoring treatment response would be beneficial.

Serological markers are used as clinical tools for diagnosis of gastrointestinal conditions in humans for decades,7,8 and there are multiple iterations of diagnostic panels incorporating an increasing number of both serologic and genetic markers.9 Seropositivity to markers such as perinuclear antineutrophilic cytoplasmic (pANCA10) and OmpC11 have been linked to certain manifestations of ulcerative colitis (UC) or Crohn’s disease (CD) respectively, both different subtypes of IBD in humans. While anti-pANCA as a stand-alone marker lacks sensitivity in CD and UC, its addition to a panel that includes ANCA, ASCA-IgG, and ASCA-IgA (anti-Saccharomyces cerevisiae antibodies) improves sensitivity of identifying CD and UC in 65% and 74% of children, respectively.12 The search for serological markers in dogs to diagnose CE and differentiate it from other GI conditions with overlapping signs has become an important focus in veterinary medicine. Dog serum with antibody responses to pANCA were characterized in dogs with GI disease populations13 and were associated with CE/IBD dogs with specificities ranging between 76% and 94% when compared to dogs with other chronic GI disorders.14 There are specific serological markers that have differential titers in CE/IBD cohorts when compared to cohorts with predominantly acute enteropathies or cohorts with no discernable GI disease.15 Seropositivity to OmpC is highly prevalent in IBD dog cohorts.

The primary aim of this study was to evaluate the use a multivariate 3-assay biomarker test consisting a combination of 2 serologic markers (ie, anti-OmpC IgAs, ACA; antigladin derived peptide IgAs, AGA) and autoantibodies (ie, anticanine calprotectin IgAs, ACNA) that were previously selected in a development study with highly characterized dog cohorts accrued from a limited number of veterinary referral centers. Based on the results of the development study, we set out to conduct a prospective field study where an expanded dog cohort would be enrolled from a broader set of veterinary centers. We assessed the performance of the multivariate test as a noninvasive, convenient adjunctive tool to aid in the diagnosis of CE/IBD under conditions resembling regular practices and procedures when confronted with potential GI cases. Furthermore, based on short half-life of IgA-based serology, a secondary aim of the study was to assess the potential to use these markers to monitor disease response to commonly utilized therapeutics.

2 MATERIALS AND METHODS

2.1 Field multicenter study and protocol

Eleven veterinary centers (4 referral and 7 general practitioners) participated in the study. The following centers were part of the VCA animal hospital network: West Los Angeles, Lakewood, Rossmoor-El Dorado, Emergency Animal Hospital and Referral Ctr., West Bernardo, Animal Specialty Group and Animal Med Ctr El Cajon. In addition, the following 4 independent centers were involved: Aloha AH, Bressi Ranch Pet Hospital, Midland Animal Clinic, and Palomar AH. The field study design was reviewed and approved for VCA sites by the Clinical Review Committee. The approved protocol was provided to all centers and monitored on a regular basis for compliance. Dogs (n = 214) > 6 months age were prospectively enrolled in the field study. The owners were required to sign written consents and were also informed about the option to conduct endoscopy/biopsy as a follow-up diagnostic procedure.

All dogs underwent routine diagnostic evaluations including several (or all) of CBC, chemistry profile, urinalysis, and additional diagnostics to rule out other causes of CE such as endoparasites, exocrine pancreatic deficiencies, and hypoadrenocorticism, or both, utilizing appropriate commercially available tests. Dogs were also given a Canine Inflammatory Bowel Disease Activity Index score.16 A blood (serum) sample was obtained at initial study enrollment to determine baseline titers of markers before any treatment. A study form evaluating the dog’s overall health, diet, and medication history was completed upon enrollment by the attending clinician.17 After enrollment, management of each dog, including diet, antibiotic, anti-inflammatory therapies or combinations thereof, as further described below and in the longitudinal studies, was at the discretion of the attending veterinarian. Diet modification was comprised of elimination, and hydrolyzed diets such as hydrolyzed soy protein or antigen-restricted diet of salmon and rice or sweet potato. Some dogs were administered antibiotics, primarily metronidazole and tylosin, and a smaller number were treated with enrofloxacin, amoxicillin, and clarithromycin. About 20% of the enrolled dogs were treated with immunosuppressive agents, mainly prednisone and budesonide, which were supplemented with other immunosuppressants as determined by the attending veterinarian. Owners were instructed to come back for follow-up visit on average 21 days after initiating treatment. A repeat blood (serum) sample was collected, and all updates on the clinical condition and treatments reported in the follow-up forms.

Inclusion criteria for eligible dogs for CE/IBD (n = 157) cohort were vomiting, diarrhea, excessive flatulence, anorexia, weight loss, or some combination of these signs, either continuous or intermittent for a minimum of 1 month. Dogs with a history of recurrence of clinical signs after treatment or chronic antibiotic responsive diarrhea were also included as long as they presented with no other diagnosed enteropathic disease (eg, parasites, giardiasis, microbial infection, or others) that had been untreated. For a subset of dogs in the CE/IBD cohort, serum samples were collected at multiple visits and were tested for the markers.
Dogs designated as non-IBD cohort (n = 24) included dogs with chronic GI disease (3 weeks to months) comprising 2 dogs with exocrine pancreatic insufficiency (confirmed by serum trypsin-like immunoreactivity and pancreatic lipase immunoreactivity testing), 9 with hypoadrenocorticism (confirmed by ACTH stimulation test result), and 10 with pancreatitis confirmed by the canine pancreatic lipase test (SNAP cPL) further supported by amylase and lipase activities in serum. This cohort also included 2 of the dogs with pancreatitis that had a high suspicion of lymphoma or other GI neoplasia based on ultrasound (eg, discernable mass, disruption of normal architecture like loss of layering and likely enlargement of the mesenteric lymph nodes) or 1 of them with lymphoma by confirmed histopathology.

Exclusion criteria for participation was gastroenteritis associated with the swallowing of foreign bodies or with a known dietary indiscretion, GI-related conditions treated for more than 2 weeks, endoparasites, any uncontrolled medical problem which could impair the study safety, and neoplasia unrelated to lymphoma or GI lymphoma or dogs that were currently receiving active chemotherapy or any other immunosuppressive therapy such as prednisone or budesonide.

The normal cohort included asymptomatic dogs (n = 33) with no prior history of chronic GI disease and no active acute GI disease presenting for a regular checkup, dental prophylaxis, or other routine procedures.

Gastroduodenoscopy was performed on a subset of dogs whose owners consented to biopsy with tests results suggestive of CE, primarily suspected CE/IBD. Biopsy samples from the stomach, duodenum, and colon were collected with flexible endoscopy biopsy forceps between 2 and 10 days of the initial serum sample collection and processed for histopathology by Colorado State University. Full-thickness biopsies or endoscopy biopsies were immediately placed in ice-cold phosphate-buffered saline and 4% buffered paraformaldehyde solution until processed. All tissue samples were processed and graded by a board-certified anatomic pathologist (Dr. Barbara E. Powers, Colorado State University) in accordance with the World Small Animal Veterinary Association (WSAVA) guidelines. Multiple morphological variables (ie, epithelial injury, crypt distension, luteal dilatation, mucosal fibrosis) and inflammatory histological variables (such as plasma cells, lamina propria lymphocytes, eosinophils, and neutrophils) were scored, and the resulting final scores were subdivided into histological severity groups: WSAVA score of 0 = normal, 1 to 6 = mild, 7 to 12 = moderate, >13 = severe.

2.2 Sample collection and analysis

Serum samples were collected from all enrolled dogs, stored on site at −20°C and shipped frozen to the laboratory within 1 month. Samples were analyzed within 72 hours upon arrival. And the remaining sample stored at −80°C. Whole blood was used for routine hematology, serum was used to determine IgA antibody concentrations that bind OmpC (ACA), calprotectin (ACNA), and small gliadin-derived peptides Glu-Gli (AGA) using direct ELISA assays. Both antigen targets and reagents were selected, developed and optimized specifically for dog IgA measurements.

Ninety-six well ELISA plates were coated with selected antigens for at least 16 hours and then washed. Plates were then blocked with blocking solution for at least 1 hour. Standards and samples were prepared in a 1:100 or 1:50 dilution in blocking solution. Standards and samples were added, in duplicate, to the blocked plate for 1 hour, and unbound material was washed away. Horseradish peroxidase-conjugated antibodies (HRP)-conjugated-goat-anti-dog secondary antibody was added to each well and the plate was incubated for 1 hour. Unbound antibody was then washed away, 3,3',5,5'-tetramethylbenzidine (TMB) HRP substrate was added, and the colorimetric reaction was stopped with an acid. The absorbance (A450) or optical density (OD450) was determined by reading the plate at 450 nm on a plate reader spectrophotometer. The standard curve was fitted using a 4-parameter equation and used to estimate the antibody titers in the samples.

2.3 Data analyses

Antibody concentrations were determined relative to a standard/calibration reference obtained from a dog with a positive signal. Analysis of the data was performed using the SoftMax Pro Enterprise (Molecular Devices). The standard curve was plotted as absorbance at 450 nm on the y-axis versus the log ELISA Units of standard on the x-axis. The samples concentrations expressed in ELISA Units (EU/mL) were calculated from the absorbance values at 450 nm. The mean, the SD, and the coefficient of variation (%CV) were obtained for all standards and samples from the raw absorbance values at 450 values and from the calculated values.

Statistical analysis was conducted using R (2016, R Foundation for Statistical Computing, Vienna, Austria) or Microsoft Office Excel (2013, Microsoft, Redmond, Washington) and GraphPad Prism (GraphPad software, San Diego, California). Data were analyzed by using Kruskal-Wallis or Fisher's exact test depending on the comparison data cohorts. A P value <.05 was considered significant. In addition, receiver operating characteristics (ROC) curve analysis was performed for each of the markers (univariate analysis) by ROC curves with the true positive rate (sensitivity) against the false-positive rate (1-specificity) in comparing the multiple cohorts. AUC > 0.5 represented a discriminatory performance of the plotted marker against the selected cohorts.

Field study populations

A total of 214 dogs were enrolled in the study. The characteristics of the cohorts are provided in the tables below. Dogs of varying ages,
sex, and breeds that presented to multiple veterinary centers with chronic gastrointestinal conditions underwent a diagnosis work-up as described above. Dogs \( n = 214 \) enrolled in the study were from more than 30 breeds, with the majority represented by the following breeds: Labrador retriever (10), Golden retriever (9), Chihuahua (12), Yorkshire terrier (10), Border terrier mix (8), and German shepherds (11). A total of 157 dogs that presented with chronic, recurrent GI symptoms (primarily diarrhea, vomiting, or both), or both, and they were enrolled and grouped under the CE/IBD cohort. All dogs enrolled in this cohort were well known to the attending veterinarians with lengthy and comprehensive clinical profiles. All dogs were offered to perform endoscopy/biopsy as part of the work up, and 38 of the owners consented to the procedure. Thirty-six out of 38 dogs that underwent endoscopy were confirmed IBD by histopathology, with a further breakdown in mild (16; 44%), moderate (16; 44%), and severe manifestations (4; 12%). Predominant histopathological diagnosis was lymphoplasmacytic enteritis (10; 28%), and enteritis with eosinophils (21; 58%); followed by neutrophils (3; 8%) and suppurative enteritis (2; 6%).

Thirty-six (36%) of the dogs also presented with gastritis/gastroenteritis as a co-morbidity. The histopathology for the other 2 dogs exhibited normal stomachs with slightly swollen villous structures and occasional dilated lacteals, and also showed some mucous hyperplasia in the colon. The confirmed IBD cohort was comparable with their signalment and clinical history (ie, age, weight, clinical signs, disease chronicity/duration, and marker titers) to the rest of the “suspected”

| TABLE 1 | Characteristics of the 3 cohorts at baseline and treatments |
|---------|---------------------------------------------------------------|
| **Variables** | **CE/IBD (n = 157)** | **non-IBD (n = 24)** | **Normal (n = 33)** |
| **Sex** | | | |
| % Female | 49.0 (77) | 54.2 (13) | 54.5 (18) |
| % Neutered | 43.9 (69) | 41.7 (10) | 39.4 (13) |
| % Spayed | 46.5 (73) | 50.0 (12) | 51.5 (17) |
| **Age (years): median, range** | 7, 0.7–15 | 8, 0.9-14.83 | 4.5, 0.8–10 |
| **Treatment (%)** | | | |
| Diet Modification | 44.6 (70) | 20.8 (5) | NA |
| Antibiotics | 36.9 (58) | 29.2 (7) | NA |
| Metronidazole | 33.1 (52) | 25.0 (6) | NA |
| Tylosin | 5.7 (9) | 0.0 | NA |
| Others | 5.7 (9) | 8.3 (2) | NA |
| Antiparasitics\(^a\) | 8.3 (13) | 4.2 (1) | NA |
| Antinflammatory\(^b\) | 15.9 (25) | 20.8 (5) | NA |
| Antiemetics | 31.2 (49) | 33.3 (8) | NA |
| Probiotics | 24.8 (39) | 8.3 (2) | NA |
| Supplement | 5.7 (9) | 0.0 | NA |
| **Signs (%)** | | | |
| General condition (decreased or lethargy) | 36.3 (57) | 50.0 (12) | NA |
| Vomiting | 61.8 (97) | 45.8 (11) | NA |
| Frequency of defecation (%) | | | |
| Normal or only slightly increased | 70.1 (110) | 58.3 (14) | NA |
| Moderately to severely increased | 13.4 (21) | 20.8 (5) | NA |
| Fecal volume (percent) | | | |
| Normal to Increased | 76.4 (120) | 62.5 (15) | NA |
| Often decreased | 8.9 (14) | 8.3 (2) | NA |
| Appetite disorder (inappetence or anorexia) | 45.2 (71) | 87.5 (21) | NA |
| Abdominal discomfort | 31.8 (50) | 45.8 (11) | NA |
| Regurgitation | 4.5 (7) | 8.3 (2) | NA |
| Presence of mucus | 26.8 (42) | 8.3 (2) | NA |
| Blood in stool | 34.4 (54) | 16.7 (4) | NA |
| Tenesmus (abdominal pain) | 17.8 (28) | 0.0 | NA |

\(^a\)Refer to treatment of ectoparasites such as flea, ticks, and ear mites.
\(^b\)Principally steroids and immunosuppressants as described in the text.
CE/IBD cohort. We combined these groups and referred to as CE/IBD cohort. The non-IBD cohort was composed of 24 dogs that presented with GI signs confirmed by the attending clinicians to have a wide array of underlying causes (see above). The normal cohort consisted of 33 apparently healthy dogs presenting with no relevant signs of GI disease at the time of visit, no known history of gastroenteritis recurrences, and were admitted for regular visits or surgical procedures.

The CE/IBD cohort included 77/157 (49.0%) females and 80/157 (51.0%) males. Sex compositions in the non-IBD cohort were represented by 11/24 (46%) males and 13/24 (54%) females, while the normal cohort included 15/33 (46%) males and 18/33 (54%) females. The mean age (±SD) was 6.96 ± 3.89 years (0.7-15 years); 7.46 ± 3.49 years (range, 0.29-14.83 years); and 5.03 ± 2.99 years (range, 0.8-10.0 years) for the CE/IBD, non-IBD, and normal cohorts, respectively. There were no statistically significant differences between ages (by Kruskal-Wallis test) and sex (by Fisher’s exact test) distribution among the cohorts.

A comprehensive list of characteristics of the 3 cohorts at admission and follow-ups is reported in Table 1. The most frequent clinical findings observed in CE included frequency of defecation +/- increase in volume of feces (110/157, 70% for CE/IBD vs 14/24, 58% non-IBD) and vomiting (97/157, 62% for CE/IBD vs 11/24, 46% non-IBD). While presence of mucus (42/157, 27%), hematochezia (54/157, 34.4%), and tenesmus (28/157, 17.8%) were predominantly associated with the CE/IBD cohort, anorexia in 21/24 (88%) was a predominant clinical sign associated with the non-IBD cohort (Table 1). Long-term antibiotics use and corticosteroid use was recorded in 58/157 (36.9%) and 4/24 (16%) of CE/IBD and non-IBD cases respectively. As part of the field study, the attending veterinarians reported on the treatments their practices considered appropriate for the CE/IBD cohorts based on clinical presentation. While dietary treatments and antibiotics were administered more often in the CE/IBD cohort than the non-IBD cohort (70/157, 44.6% vs 5/24, 20.8%, and 58/157, 36.9% vs 7/24, 29% respectively), treatment with anti-inflammatories was used less frequently in the CE/IBD cohort (25/157, 15.9%) than the non-IBD (5/24, 20.8%) cohort. Probiotics were part of the treatment protocol in 39/157 (24.8%) of the CE/IBD cohort.

### 3.2 Seropositivity of selected markers in the groups

Serum samples were collected from all dogs enrolled during their first visit, and their titers against selected markers determined by ELISA. Single-marker based results are illustrated in Figure 1. The overall anti-IgA titers against ACA, ACNA, and AGA were higher in the CE/IBD cohort when compared to normal and non-IBD (Figure 1). The overall mean titers (mean ± SD, expressed in EU/mL) of these selected markers in the CE/IBD cohort (ACA, 86.68 ± 22.73; ACNA, 33.17 ± 10.97; AGA, 303.23 ± 82.12) were higher than the non-IBD cohort mean titers (ACA, 10.78 ± 2.59; ACNA, 7.14 ± 1.19; AGA, 176.64 ± 80.84) as well as higher than the normal cohort mean titers (ACA, 9.30 ± 5.18; ACNA, 6.04 ± 1.35; AGA, 32.23 ± 9.96) as graphically displayed in Figure 1, with these differences being statistically significant (P values results for the 3 markers less than 0.0001 by the Kruskal-Wallis test).

In this study, seropositivity was defined as the percentage of dogs of a given cohort exhibiting a serum titer of IgA antibodies against the specific marker above predetermined cut-off values as described by Estruch et al. In this respect, serum from dogs with CE/IBD exhibited the highest seropositivity titer for ACA (104/157, 66.2%), ACNA (86/157, 55%), and AGA (118/157, 75.2%). For the normal cohort, seropositivity rates of 9% (3/33), 15% (5/33), and 6% (2/33) for ACA, ACNA, and AGA, respectively, were the lowest. The non-IBD cohort presented intermediate titers, with 3/24 (13%) seropositive for ACA, 7/24 (29%) for ACNA, and 13/24 (54%) for AGA.

### 3.3 ROC and 3D scatter plots to define discriminating markers among cohorts

All 3 selected markers were discriminating when comparing CE/IBD with normal cohorts, with seropositivity’s to ACA and AGA being the most statistically significant with AUCs of 0.89 and 0.91, respectively (Figure 2A). Both ACA and AGA also showed discriminating properties between non-IBD and normal cohorts (AUCs of 0.73 and 0.82, respectively, Figure 2B). By contrast, the ACNA marker offers no discriminatory property between non-IBD and normal. When comparing CE/IBD with non-IBD cohort seropositivity to ACA (larger AUC, 0.79) and ACNA were key distinguishing markers between CE/IBD and non-IBD cohorts (Figure 2C).

Serum titers of the 3 markers can be combined to create a multivariate profile for the dogs of different cohorts and are represented graphically in a 3D plot (Figure 3). Serum titers for ACA, ACNA, and AGA were low in normal cohorts clustering close to the intersection of the 3 axes. A comparable 3D plot for non-IBD displays significant AGA titers while maintaining low values for the other markers. In the CE/IBD cohort, the data set is more spread out along the 3 axes. All serum marker data in the 3 cohorts of the study are represented in Figure 3 demonstrating graphically how multivariate (assay) tests can offer a robust approach to distinguish between normal, CE/IBD and non-IBD cohorts. Algorithms work as a decision tree using as inputs the 3 marker titers for a given sample, and cut-offs values calculated from the ROC curves shown in Figure 2. Specifically, 1 configuration is to set the cutoff for ACA, ACNA, and AGA at 15, 6, and 60 EU/mL. Under this configuration, samples with higher titers would be classified as CE/IBD; with lower as normal; and non-IBD would display combinations in-between. At these cutoff values, the algorithm can discriminate CE/IBD from normal cohorts with 90% sensitivities and 96% specificity; and CE/IBD from non-IBD with 80% sensitivities and 86% specificity.

Given the experimental design discussed above (ie, 1 gate vs 2 gates), while relevant to compare diseased cohorts, the specificity and sensitivity might be overestimated when comparing diseased with normal cohorts. Our experimental design attempts to mitigate such effect by including disease cohorts that are mostly represented by...
samples from dogs with mild (44.4%) or moderate manifestation of the gastrointestinal disease (44.4%), but the potential effect of using different protocols to sample dogs from disease cohorts vs normal cohorts should be taken into consideration. In terms of potential bias due to correlation of the markers, we would note that the overall

FIGURE 1 Scatter graph representations of the titer of seropositivity to OmpC (ACA, A), calprotectin (ACNA, B), and gliadin-derived peptides (AGA, C) measured in serum samples from multiple cohorts of dogs. Dog sera from normal, non-IBD, and CE/IBD were tested by ELISA as described in Section 2. Mean of ELISA values (EU/mL) are indicated by thick horizontal bars. Asterisks denote: *P = .05; **P = .01; ***P = .001. Differential values between the cohorts were considered statistically significant at P < .05 using Kruskal-Wallis test. CE, chronic enteropathies; IBD, inflammatory bowel disease

FIGURE 2 Receiver operator characteristics curves for discriminating the normal vs CE/IBD cohorts, A, normal vs non-IBD, B, and non-IBD vs CE/IBD, C, for continuous serological markers and autoantibodies. Area under the curve (AUC) represents the discriminating performance of each marker. All AUC values for each of the markers when tested and their 95% confidence intervals as determined with 2000 stratified bootstrap replicates. CE, chronic enteropathies; IBD, inflammatory bowel disease

samples from dogs with mild (44.4%) or moderate manifestation of the gastrointestinal disease (44.4%), but the potential effect of using different protocols to sample dogs from disease cohorts vs normal cohorts should be taken into consideration. In terms of potential bias due to correlation of the markers, we would note that the overall
estimates for sensitivity and specificity are based on the evaluation of a single prediction derived from the 3-marker cutoffs, not from the sensitivities and specificities of the component markers. Therefore, their potential correlation should not (or minimally) introduce any bias.

3.4 | Serum markers as monitoring tools

As part of the field study, owners were instructed to return for follow-up visits to facilitate longitudinal monitoring of selected markers. Two centers implemented monitoring 17 enrolled CE/IBD dogs for a period of up to 3 months after diagnosis, and in 14 of them the completeness of the records enabled the data presented in this section. Ten dogs were tracked in multiple follow-ups, serum was collected and marker titers for ACA, ACNA, and AGA were determined (Figure 4). All of them underwent antibiotic treatment (except dog 6 marked by asterisk symbol in Figure 4A) and diet modification after initial diagnosis. The serial titers for ACA and AGA markers are shown in Figure 4A,C, respectively. Furthermore, all dogs except dogs 1 and 6 (also marked by asterisk symbol in Figure 4B) received anti-inflammatory treatment either after initial diagnosis (dogs 2, 3, 4, and 5) or after the 1st follow-up session (dogs 2, 3, 4, and 5). The serial titers for ACNA are shown in Figure 4B. The 1st follow-up took place on average 31 days (range 19-58 days) after initial visit. The 2nd follow-up was scheduled on average 34 days after the 1st follow-up (range 19-55 days). In 7 dogs that received treatments, all 3 markers decreased between 47% to 99% for ACA, 13% to 88% for ACNA, and 30% to 85% for AGA. These reductions occurred concurrently with improvements in their clinical presentations leading to resolution of their signs. Three dogs (dogs 4, 5, 6), also with treatments, displayed limited to no clinical improvements at the time of
data cutoff, and the corresponding markers remained high throughout the time-course of the study.

A more detailed longitudinal profile of 4 additional CE/IBD dogs is shown in Figure 5. The marker titers were measured at enrollment and monitored for up to 3 months. Dog A was fed hydrolyzed protein-based diets, treated with enrofloxacin and metronidazole after diagnosis and received no anti-inflammatory medication. Dog B received gluten-free salmon diet at diagnosis and given budesonide treatment after 1st follow-up (54 days after diagnosis). While dogs A, B, and D noticeably improved their clinical presentations, dog C remained diseased. Results reflect the direct read-outs in EU/mL, and for ACA and AGA are printed in the left axis scale, and for ACNA in the right axis scale.

**FIGURE 5** Longitudinal data profiles for 4 dogs enrolled in the field study. Dogs were monitored for 87, 92, 95, and 97 days for dogs A, B, C, and D respectively. Main treatments included diet modification, antibiotics and anti-inflammatories. All 4 dogs received most treatments after diagnosis, and in the case of dog B, anti-inflammatory were administered after 1st visit (Figure 5B). While dogs A, B, and D noticeably improved their clinical presentations, dog C remained diseased. Results reflect the direct read-outs in EU/mL, and for ACA and AGA are printed in the left axis scale, and for ACNA in the right axis scale.

The principal aim of this study was to assess the performance of combination of serum markers as a noninvasive, convenient test to diagnose cohorts of CE/IBD dogs enrolled under field conditions that resemble how primary care veterinary clinics approach the diagnostic work-up of GI cases. The markers included in the test displayed discriminatory properties as single markers in well-defined dog cohorts affected with GI disorders and have relevance to various physiological aspects known to be impaired in CE/IBD, thus leading to disease susceptibility. We documented the presence of antibodies against immunoreactive targets and the differential prevalence of ACA, ANCA, and AGA in multiple cohorts of dogs defined as CE/IBD and non-IBD cohorts afflicted with CE. Furthermore, we operationalized the multivariate assay in the form of an ELISA test and corresponding algorithms yielded higher discriminating values between the cohorts.

The use of serum markers and autoantibody combinations is routinely implemented in human medicine as part of a comprehensive testing tool to diagnose and monitor GI disorders. Testing co-exists with the practice of endoscopic examinations that are the main way to diagnose complex digestive diseases, but, by comparison, are costly and have increased risk associated with general anesthesia. Multimarker serology-based tests are reportedly used to aid in the diagnosis of IBD, differentiate UC from CD, and to reliably predict endoscopic disease activity in IBD. In veterinary medicine, there has been increased focus on the potential application of serological and fecal markers to assist in the diagnosis and management of gastroenteropathies. Dogs with severe signs of GI disease more frequently displayed abnormal titers of markers than dogs with milder forms or no disease. In particular, antibodies against perinuclear antineutrophilic cytoplasmic proteins (pANCA) are mostly associated with dogs suffering IBD types of CE than other types of enteropathies or normal dogs. More recently, the importance of serological markers in dogs with CE/IBD is further documented where seroconversion to antigens such as OmpC, flagellins, polymorphonuclear leukocytes, calprotectin, and gliadin-derived peptides were significantly associated with CE cohorts compared to normal.

The knowledge of whether there is mucosal inflammation associated with digestive disorders in dogs represents a key finding of the diagnostic process, since it guides appropriate therapeutic interventions. Intestinal inflammation is primarily determined with endoscopically obtained biopsies, followed by histopathology. In dogs, there has been an increased focus to explore the use of noninvasive markers, and in this regard, fecal calprotectin in dogs was recently reported that it behaves in ways comparable to humans and can be useful biomarker for mucosal inflammation. This assay incorporates

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**DISCUSSION**

The use of serum markers and autoantibody combinations is routinely implemented in human medicine as part of a comprehensive testing tool to diagnose and monitor GI disorders. Testing co-exists with the practice of endoscopic examinations that are the main way to diagnose complex digestive diseases, but, by comparison, are costly and have increased risk associated with general anesthesia. Multimarker serology-based tests are reportedly used to aid in the diagnosis of IBD, differentiate UC from CD, and to reliably predict endoscopic disease activity in IBD. In veterinary medicine, there has been increased focus on the potential application of serological and fecal markers to assist in the diagnosis and management of gastroenteropathies. Dogs with severe signs of GI disease more frequently displayed abnormal titers of markers than dogs with milder forms or no disease. In particular, antibodies against perinuclear antineutrophilic cytoplasmic proteins (pANCA) are mostly associated with dogs suffering IBD types of CE than other types of enteropathies or normal dogs. More recently, the importance of serological markers in dogs with CE/IBD is further documented where seroconversion to antigens such as OmpC, flagellins, polymorphonuclear leukocytes, calprotectin, and gliadin-derived peptides were significantly associated with CE cohorts compared to normal.

The knowledge of whether there is mucosal inflammation associated with digestive disorders in dogs represents a key finding of the diagnostic process, since it guides appropriate therapeutic interventions. Intestinal inflammation is primarily determined with endoscopically obtained biopsies, followed by histopathology. In dogs, there has been an increased focus to explore the use of noninvasive markers, and in this regard, fecal calprotectin in dogs was recently reported that it behaves in ways comparable to humans and can be useful biomarker for mucosal inflammation.
immunoreactivity against canine calprotectin (ACNA) as one of its read-out targets. Increased titers of ACNA are associated with dog cohorts exhibiting CE/IBD, with high titers and 55% are seropositive. Moreover, we report that ACNA titers decreased in dogs that underwent interventions with anti-inflammatory agents concomitantly with clinical improvements.

Microbial-based serological markers are important in CE because their expression represents the host immune response to exposure of intestinal bacteria as a result of the breakdown of the gut mucosal barrier. This study has shown that ACA response was detected in 66.2% of dogs enrolled in the CE/IBD cohort, while it was present in less than 12.5% and 9.1% of dogs enrolled in the non-IBD and normal cohorts, respectively. Furthermore, ACA seropositivity was also discriminating between cohorts with CE and other cohorts with dogs predominantly displaying acute enteropathies. This study has further confirmed that OmpC seropositivity is a key discriminating serological marker in dogs with CE/IBD, and therefore could a key component of a potential CE-related serological panel for dogs.

Another assay evaluates the presence of antigliadin IgAs (AGA) produced in response to gliadins. Unlike humans, the clinical significance of AGA in companion animals is unclear. Initial studies in dogs failed to demonstrate the presence of antigliadin antibodies. But, this understanding is changing. Epileptoid cramping syndrome in dogs could be part of a syndrome of gluten intolerance consisting of episodes of transient dyskinesia, signs of gastrointestinal disease and dermatological hypersensitivity. IgA antibodies against gliadins are reported in dogs with CE and intestinal T-cell lymphoma, suggesting an association between glutin-induced repetitive inflammation and subsequent intestinal lymphoma in dogs. Prior observations in dogs with GI disease (including CE) displayed increased AGA seropositivity (between 54% to 75% depending on the cohort) compared to just 6% in normal dogs, together with this study further confirms the potential relevance of serological responses to gliadins in the context of CE dogs. An increase in gut permeability in CE dogs would be consistent with greater antigen movement across the intestinal epithelial barrier and thus an increased immune exposure. However, results in this area are variable with some studies done using dogs affected with severe CE showing an increased in gut permeability while others with milder presentations of the disease displayed no significant difference in GI absorptive capacity when compared to asymptomatic dogs.

Using serological markers to monitor disease progression and treatment responses in CE represents a significant unmet need. In humans, use of non-invasive tests enables monitoring disease relapse. The selected serological markers are IgA based because they have a short half-life (3-6 days). When conducting serial measurements in enrolled dogs undergoing treatments, our study showed encouraging results, albeit in a limited subcohort, supporting the use of single-assay read outs as a noninvasive tool to monitor disease progression and response to therapy.

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CONFLICT OF INTEREST DECLARATION
Authors declare no conflict of interest.

OFF-LABEL ANTIMICROBIAL DECLARATION
Authors declare no off-label use of antimicrobials.

INSTITUTIONAL ANIMAL CARE AND USE COMMITTEE (IACUC) OR OTHER APPROVAL DECLARATION
Authors declare no IACUC or other approval was needed.

HUMAN ETHICS APPROVAL DECLARATION
Authors declare human ethics approval was not needed for this study.

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