Lyme Disease Multiplex Testing for Dogs

Background on Lyme disease and Lyme diagnostics in dogs

Lyme disease is induced by the spirochete *B. burgdorferi*. Spirochetes are transmitted to dogs by infected ticks. Similar to humans, dogs are incidental, dead-end hosts for *B. burgdorferi*. Typical clinical signs in dogs are sporadic fever, acute arthritis, arthralgia, lameness, and glomerulopathy. Clinical signs of lameness often develop 2-5 months after infection. *B. burgdorferi* can persist for at least 1 year in clinically recovered, untreated dogs. In a report from 1992, clinical signs of Lyme disease were estimated to occur in 5-10% of seropositive animals. This percentage likely underestimates the current Lyme disease prevalence for dogs in endemic areas due to a higher awareness of the disease in owners and veterinarians and improved diagnostic methods.

Serum antibodies to different antigens of *B. burgdorferi* are commonly used to identify dogs that were exposed to the pathogen and to diagnose Lyme disease in dogs with compatible clinical signs. Former ELISA based diagnostic tests identified antibodies as early as 4-6 weeks after infection. The new Lyme Multiplex assay can identify antibodies to *B. burgdorferi* by 3 weeks after infection. High antibody levels were found in serum of experimentally infected dogs for at least 17 months and are likely maintained for much longer times, i.e. as long as *B. burgdorferi* persists in the dog.

How does the Canine Lyme Multiplex Assay work?

The Canine Lyme Multiplex Assay was developed at the Animal Health Diagnostic Center at Cornell University. It detects antibodies to three *B. burgdorferi* antigens in canine serum (Fig. 1). The test is based on fluorescent beads and allows the simultaneous measurement of antibodies to all three *B. burgdorferi* antigens in a single sample.

**Figure 1:** The Canine Lyme Multiplex Assay detects of antibodies to *B. burgdorferi* in canine serum. Outer surface protein (Osp) specific antibodies in the dog’s serum bind to OspA, OspC or OspF multiplex beads, are detected by a fluorescent conjugate and are evaluated in a multiplex reader. The assay values are expressed as median fluorescent intensities (MFI).

Which *B. burgdorferi* antigens are used and how is the test interpreted?

The Canine Lyme Multiplex Assay is based on three antigens, called outer surface proteins (Osp), of *B. burgdorferi*. Various research studies have shown that Osp antigen expression changes on the bacterial surface in response to tick feeding and again after infection of a warm-blooded host, such as dogs, horses, or humans (Fig. 2). In response to infection,
dogs develop antibodies to these Osp proteins and testing for antibodies to specific Osp antigens can assist in the diagnosis of infection and Lyme disease.

**Figure 2**: *B. burgdorferi* regulates its outer surface protein expression depending on its environment. In the tick gut, OspA is expressed. During tick feeding, the bacteria leave the tick’s mid gut and express OspC on their surface. OspC expression is maintained early during infection. In response to the environment in the dog’s body, the bacteria again change their surface expression – OspC disappears and OspF is expressed in the chronic infection stage.

**Interpretation of Lyme multiplex results**: The Lyme Multiplex Assay is a fully quantitative test. It results in a numeric antibody value for each of the three *B. burgdorferi* antigens tested. An interpretation of each value is submitted with the test report. In addition, the antibody profile gives an advanced interpretation on the infection and vaccination status of the dog. Antibodies to OspA serve as markers for vaccination and those to OspC and OspF as markers for infection (Fig. 3). In infected dogs, quantitative antibody values can then be used to follow-up on treatment success.

1. **OspA** – positive values for antibodies to OspA are typically observed in vaccinated dogs. OspA is expressed while *B. burgdorferi* persists in the tick mid-gut and also while the bacteria are cultured *in-vitro*. During infection of mammalian hosts, the bacteria down-regulate OspA. Therefore, antibodies to OspA are almost undetectable after natural infection in non-vaccinated dogs. Very low positive, transient antibody values to OspA can sometimes be detected 2-3 weeks after infection.

2. **OspC** – is a valuable indicator for early infection with *B. burgdorferi*. Antibodies to OspC are detected as early as 2-3 weeks after infection. Antibodies to OspC decline after 7-11 weeks and become undetectable by 4-5 months after infection.

3. **OspF** – is an indicator of chronic infection. Antibodies to OspF are detectable by 5-8 weeks after infection and are maintained at high levels afterwards. Researchers at Cornell observed a very high agreement between antibodies to OspF and C6 as robust markers for infection in dogs in a blinded study. Dogs with positive antibody values to OspF and negative antibody values to OspC have been infected with *B. burgdorferi* for at least 5 months (Fig. 3).

**Figure 3**: Antibody responses to OspA, OspC and OspF in infected dogs. Dogs were not treated and were unvaccinated. Data obtained from Wagner et al. 2012.

**Advantages of the Canine Lyme Multiplex Assay (AHDC, Cornell University)**

The Canine Lyme Multiplex Assay is available only at the Animal Health Diagnostic Center at Cornell University. It combines the results obtained by previous ELISA and Western blot testing and exceeds the information obtained by tests that are based on a single antigen of *B. burgdorferi* such as C6. The Lyme Multiplex Assay provides information whether the dog got infected with *B. burgdorferi* and when the infection occurred. The
test results are fully quantitative and appropriate to follow-up on treatment success or response to vaccination. The advantages of the Lyme Multiplex Assay compared to other Lyme tests are:

- **assay results distinguishing between:**
  1. **early and chronic stages of infection:** Antibodies to OspC and OspF serve as markers for infection. Combined antibody values for OspC and OspF can also identify the stage of infection with *B. burgdorferi* and can determine when the dog was infected. In humans, early infection stages have a better prognosis and increased success rates after antibiotic treatment than chronic infection stages. The same trends have been observed in dogs. A more rapid decline of antibodies, indicating successful treatment, was observed in many dogs that were treated in the early infection stage.
  2. **vaccination and infection:** The assay provides a separate quantitative value for antibodies to OspA, as a marker for vaccination, and to determine the dog’s vaccination status.

- **increased specificity and sensitivity:** Improved test performance results in earlier detection of antibodies to *B. burgdorferi* (as early as 2-3 weeks after infection) and a higher accuracy of the Lyme multiplex assay compared to conventional ELISA assays.

- **quantitative measurement of individual antigens:** The Lyme Multiplex Assay has a much broader linear quantification range than ELISAs. Antibody values are precisely measured in a broad concentration range. Quantification of antibodies and evaluation of treatment success can be performed based on declining antibody values. Successful antibiotic treatment decreases the bacterial load. The reduced or missing antigenic stimulus causes the immune system to produce fewer antibodies which leads to a decrease in serum antibody values. The decline of antibodies is slower than the bacterial clearance caused by the antibiotic treatment. After successful treatment, IgG antibodies to *B. burgdorferi* decline according to their half-life of around 21 days. A decrease of a positive antibody value by at least 50% of the original value in the time frames mentioned below is considered an indicator of treatment success. After successful treatment, antibodies will continue to drop without further treatment and will become negative over time if no re-infection occurs. The best timing for follow-up testing depends on the stage of infection. For example:

  - **For chronic infection stages** (OspC-/OspF+): Follow-up testing with the Lyme Multiplex Assays can be performed as early as 3 months after starting treatment. It should not be performed earlier to give the antibodies time to decline. A successful treatment is indicated by a clear drop in OspF antibody values (≥ 50% of the pretreatment value obtained 3 months earlier). A treatment failure
infected or non-infected dogs. Antibodies to OspF and C6 provide robust markers of infection in dogs. However, the multiplex assay provides additional information on the dog’s infection stage (early or chronic) and vaccination status.

Sample submission

For detection of antibodies to *B. burgdorferi*, 2ml of dog serum needs to be submitted. Serum should be collected in a red top blood tube. The entire red blood tube or isolated serum should be shipped by overnight shipment on an ice pack to the Animal Health Diagnostic Center at Cornell University. For submission forms and shipping address go to the Animal Health Diagnostic Center website (http://ahdc.vet.cornell.edu).

Samples are tested every day (Mon-Fri) and results are available 1-2 days after the sample arrives at the laboratory. Consultation on Canine Lyme Multiplex testing and on result interpretation is available by calling the Serology/Immunology laboratory at the Animal Health Diagnostic Center at Cornell University 607.253.3900.

Special considerations for vaccinated dogs

The Canine Lyme Multiplex Assay can distinguish between vaccinated and infected dogs. All currently available vaccines for dogs induce antibodies to OspA. Therefore, the results on antibodies to OspA should be used to determine the dog’s vaccination status. However, some of the available vaccines also induce antibodies to other Osp antigens. To provide our clients with the best interpretation for each animal, we need information on the vaccine used. Please include the name of the vaccine and the date when the dog was last vaccinated on the submission form if a sample of a vaccinated dog is submitted for testing.

How can the Lyme Multiplex Assay be compared to other serological Lyme assays?

Researchers at the Animal Health Diagnostic Center at Cornell University have compared the former ELISA/Western blot procedure and the commonly used C6-based assays with the Canine Lyme Multiplex Assay. Lyme Multiplex Assay OspF and C6 results highly correlate for the identification of (persistent infection) or reinfection after treatment is indicated by OspF antibody increases, no or minor decreases or alternating antibody values (minor drop followed by increase after re-testing). In case of a re-infection, OspC antibodies will also become positive.

For *early infection stages* (OspC+/OspF+ or OspC+/OspF−): Treatment follow-up can be performed as early as 6 weeks after beginning of the treatment. Early in infection, antibodies have been observed to decline more rapidly. This is likely due to a higher amount of IgM antibodies during early infection and a more rapid decline of IgM compared to IgG. A successful treatment is indicated by a clear drop in OspC (and if positive also OspF) antibody values to ≥50% of the pretreatment value obtained 6 weeks earlier. A treatment failure (persistent infection) or reinfection after treatment is indicated by OspC and/or OspF antibody increases, no or minor decreases or alternating antibody values (minor drop followed by increase after re-testing).

References


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