Overview

Useful For
Aids in the diagnosis of leptospirosis

Highlights
This is test is intended to be used as an aid for the diagnosis of acute or recent leptospirosis due to infection with *Leptospira* species.

This is a qualitative immunodot test for detection of IgM-class antibodies to *Leptospira* species.

A negative result by this assay does not exclude the possibility of leptospirosis and all results must be correlated with clinical presentation and exposure history.

Method Name
Enzyme-Linked Immunoassay Dot (Immunodot)

NY State Available
Yes

Specimen

Specimen Type
Serum

Specimen Required
Container/Tube:

Preferred: Serum gel

Acceptable: Red top

Specimen Volume: 0.3 mL

Collection Instructions: Serum should be collected according to standard practices. Acute and convalescent specimens obtained to determine seroconversion should be collected 2 or more weeks apart.

Forms
If not ordering electronically, complete, print, and send a Microbiology Test Request (T244) with the specimen.

Specimen Minimum Volume
0.1 mL

Reject Due To

<table>
<thead>
<tr>
<th>Gross hemolysis</th>
<th>Reject</th>
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</thead>
<tbody>
<tr>
<td>Gross lipemia</td>
<td>Reject</td>
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Specimen Stability Information
Clinical and Interpretive

Clinical Information

Leptospirosis is a zoonotic disease of worldwide prevalence, though the majority of infections occur in warm, tropical climates. Wild mammals, typically rodents, are the primary, natural reservoir for pathogenic strains of *Leptospira*, however, domestic animals (eg, dogs) also represent a major source of human infection. *Leptospira* are Gram-negative spirochetes with at least 20 different species in the genus. Of these, at least 9 species are considered pathogenic, including the most common agent of leptospirosis, *Leptospira interrogans*.

Transmission occurs through indirect human contact (eg, via mucous membranes or abraded skin) with water, food, or soil contaminated with animal urine containing the *Leptospira* spirochetes. Following infection, the incubation period can range from 3 to 30 days depending on the inoculum dose and immune status of the individual.

The clinical manifestations of leptospirosis can vary, ranging from a mild, flu-like illness (eg, headache, malaise, fever, arthralgia, fatigue) to fulminant disease, with severe liver and kidney involvement. The latter manifestation was previously referred to as Weil disease. *Leptospira* organisms may be found in the blood at the onset of disease and can persist for approximately 1 week. Subsequently, spirochetes may be found in the urine and can persist for 2 to 3 months; however, shedding may be intermittent and the numbers of organisms present may be low.

While *Leptospira* can be grown in culture, this is a fastidious organism and requires immediate transport to the laboratory. Additionally, detectable growth requires prolonged incubation (1-6 weeks), limiting the utility of culture for acute diagnosis. For this reason, serologic detection for antibodies to *Leptospira* remains the method of choice for rapid diagnosis. IgM-class antibodies to this spirochete are detectable by day 6 of illness and remain detectable for 2 to 3 months following symptom onset.

Reference Values

Negative

Interpretation

Positive: IgM antibodies to *Leptospira* species detected suggesting recent infection. Antibody presence alone cannot be used to definitively diagnose acute infection, as antibodies from a prior exposure or infection may remain detectable for a prolonged period of time.

Borderline: Result should be interpreted with caution. Additional testing of a second, convalescent specimen is recommended. If the specimen remains borderline reactive, a second serological method should be considered if leptospirosis infection is still suspected.

Negative: No IgM antibodies to *Leptospira* detected. Since antibodies may not be present or may be present at undetectable levels during early disease, repeat testing of a convalescent sample collected in 2 to 3 weeks is recommended.

Cautions

The temporal IgM immune response can vary among patients. Therefore, a single negative result by this assay
should not be used to exclude diagnosis, especially in patients with symptoms suggestive of leptospirosis who have an appropriate exposure history.

This test does not distinguish between acute or past infection. Clinical correlation is required. Patients may remain seropositive for months to possibly years following resolution of disease; therefore, this test cannot be used to establish cure or response to therapy.

**Supportive Data**

**Accuracy:**

A total of 40 previously characterized serum samples tested by the Focus Diagnostics Inc. Leptospirosis Indirect Hemagglutination Assay (IHA) (30 were positive and 10 were negative) were evaluated by the GenBio *Leptospira* IgM ImmunDOT assay. A summary of the results is provided below:

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<tr>
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<th>Focus IHA Neg</th>
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<tbody>
<tr>
<td>ImmunoDOT Pos</td>
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<td>ImmunoDOT Neg</td>
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Positive Agreement: 100% (30/30); 95% Confidence Interval (CI) 86.5%-100%

Negative Agreement: 100% (10/10); 95% CI 67.9%-100%

Overall Agreement: 100% (40/40); 95% CI 89.5%-100%

**Clinical Reference**


3. Package Insert: ImmunoDot *Leptospira* IgM, GenBio, San Diego, CA; Version 4.0

**Performance**

**Method Description**

The GenBio IgM Immunodot Leptospirosis test utilizes an enzyme-linked immunoassay (EIA) dot technique for the detection of IgM antibodies. *Leptospira biflexa*, serovar Patoc 1 strain antigens are dispensed as discrete dots onto a solid membrane. After adding the test specimen to a reaction cuvette, an assay strip is inserted, allowing patient antibodies reactive with the test antigens to bind to the strip's solid support member. Alkaline phosphatase conjugated goat antihuman IgM antibodies are allowed to react with bound patient antibodies. Finally, the strip is transferred to an enzyme substrate reagent, which reacts with bound alkaline phosphatase to produce an easily seen, distinct spot. (Package insert: ImmunoDOT Leptospora IgM, San Diego, CA)
Test Definition: LEPDT
Leptospira, IgM, S

No

Day(s) and Time(s) Test Performed
Monday, Wednesday, Friday; 9 a.m.

Analytic Time
Same day/1 day

Maximum Laboratory Time
5 days

Specimen Retention Time
14 days

Performing Laboratory Location
Rochester

Fees and Codes

Fees
- Authorized users can sign in to Test Prices for detailed fee information.
- Clients without access to Test Prices can contact Customer Service 24 hours a day, seven days a week.
- Prospective clients should contact their Regional Manager. For assistance, contact Customer Service.

Test Classification
This test has been cleared, approved or is exempt by the U.S. Food and Drug Administration and is used per manufacturer's instructions. Performance characteristics were verified by Mayo Clinic in a manner consistent with CLIA requirements.

CPT Code Information
86720

LOINC® Information

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