# WISCONSIN LYME DISEASE CASE REPORT FORM

## PATIENT/PHYSICIAN INFORMATION

| Patient’s Name: | Date reported to HD: __/__/____ (mm/dd/yyyy) |
| Street Address: | Provider Name: |
| City: | Provider Phone: |
| Patient Phone: | Provider Address: |

## DEMOGRAPHICS

<table>
<thead>
<tr>
<th>State of residence:</th>
<th>County of residence:</th>
<th>Zip code:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex:</td>
<td>Date of birth: <strong>/</strong>/____</td>
<td>Hispanic Ethnicity: yes no unknown</td>
</tr>
<tr>
<td>male</td>
<td>female</td>
<td>unknown</td>
</tr>
</tbody>
</table>

## LABORATORY FINDINGS

### EIA/IFA
- (Please circle: IgM, IgG, total)
- Specimen collection date: __/__/____
- IgM: positive equivocal negative not done

### Western Blot (WB)-
- (If not serum, specify): __/__/____
- IgM: positive negative not done
  - 41kDa (FlaB) 39 kDa (BmpA) 21-25 kDa (OspC)
- IgG: positive negative not done
  - 93 kDa 66 kDa 58 kDa 45 kDa 41 kDa
  - 39 kDa 30 kDa 28 kDa 21 kDa 18 kDa

## CLINICAL SIGNS AND SYMPTOMS AND EXPOSURE

<table>
<thead>
<tr>
<th>Did a physician diagnose this patient with Lyme disease? yes no</th>
<th>Date of Lyme disease diagnosis: <strong>/</strong>/____</th>
</tr>
</thead>
<tbody>
<tr>
<td>Confirmatory signs and symptoms yes no unknown</td>
<td>Non-confirmatory signs and symptoms (check all that apply):</td>
</tr>
<tr>
<td>EM rash (&gt; 5 cm in diameter)</td>
<td>Arthralgias</td>
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<tr>
<td>Arthritis (objective episodes of joint swelling)</td>
<td>Myocarditis</td>
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<tr>
<td>Bells palsy or other cranial neuritis</td>
<td>Bundle branch block</td>
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<tr>
<td>Encephalomyelitis*</td>
<td>Neck pain</td>
</tr>
<tr>
<td>Lymphocytic meningitis</td>
<td>Cognitive impairment</td>
</tr>
<tr>
<td>Radiculoneuropathy</td>
<td>Other rash</td>
</tr>
<tr>
<td>2nd or 3rd degree atrioventricular block</td>
<td>Encephalopathy</td>
</tr>
<tr>
<td></td>
<td>Palpitations</td>
</tr>
<tr>
<td>*If encephalomyelitis is checked, CSF titer must be higher than serum titer</td>
<td></td>
</tr>
</tbody>
</table>

### Exposure
- If EM is present, was the patient exposed to wooded, brushy or grassy areas in a Lyme disease endemic county ≤ 30 days before onset? yes no unknown
- If yes, where: County(s) __________ State(s) __________
- If the patient had EM, was there: A single EM or multiple EM rashes

## SUPPLEMENTAL INFORMATION

- Was the patient pregnant at the time of illness? yes no unknown
- Was the patient hospitalized for this illness? yes no unknown
- Antibiotics used for this illness (check all that apply): doxycycline azithromycin
- Combined duration of antibiotics for this illness: <1 month 1 - 3 months >3 months

### FOR HEALTH DEPARTMENT USE ONLY

#### Confirmed Case
- EM rash in a Wisconsin resident or
- At least one late manifestation that has laboratory evidence of infection that meets criteria (see next page)

#### Probable Case
- Physician-diagnosed Lyme disease with non-confirmatory signs and symptoms and laboratory evidence of infection that meets criteria (see next page)

#### Suspect Case
- At least one late manifestation but only has a positive IgG EIA/IFA result or only has a total antibody result (i.e. IgM or IgG was not specified) or
- Any positive laboratory test with no clinical information available (e.g. a laboratory report without a case report form)
Clinical description: a systemic, tickborne disease with protean manifestations, including dermatologic, rheumatologic, neurologic, and cardiac abnormalities. The best clinical marker for the disease is the initial skin lesion (i.e., erythema migrans (EM)) that occurs in 60%-80% of patients.

Surveillance case definition: this surveillance case definition was based on the revised national case definition effective January 1, 2008. It is developed for national reporting of Lyme disease and not intended to be used in clinical diagnosis.

Case classifications:

Confirmed case:
- EM with a potential exposure in a Lyme disease endemic county <30 days before illness (as defined below), or
- At least one late manifestation that has laboratory evidence of infection that meets criteria.

Probable:
- Physician-diagnosed Lyme disease that has laboratory evidence of infection with non-confirmatory signs and symptoms.

Suspect:
- At least one late manifestation but only has a positive IgG EIA/IFA result or only has a total antibody result (i.e. IgM or IgG was not specified), or
- Any positive laboratory test with no clinical information available (e.g. a laboratory report).

Not a Case:
- Any case report that does not meet the confirmed, probable, or suspect category.

Definitions and Clarifications:

Erythema migrans (EM). For purposes of surveillance, EM is defined as a skin lesion that typically begins as a red macule or papule and expands over a period of days to weeks to form a large round lesion, often with partial central clearing. A single primary lesion must reach greater than or equal to 5 cm in size. Secondary lesions also may occur. Annular erythematous lesions occurring within several hours of a tick bite represent hypersensitivity reactions and do not qualify as EM. For most patients, the expanding EM lesion is accompanied by other acute symptoms, particularly fatigue, fever, headache, mildly stiff neck, arthralgia, or myalgia. These symptoms are typically intermittent. The diagnosis of EM must be made by a physician. Laboratory confirmation is recommended for persons with no known exposure.

Confirmatory late manifestations. Late signs and symptoms include any of the following when an alternate explanation is not found:

1. Musculoskeletal system. Recurrent, brief attacks (weeks or months) of objective joint swelling in one or a few joints, sometimes followed by chronic arthritis in one or a few joints. Manifestations not considered as criteria for diagnosis include chronic progressive arthritis not preceded by brief attacks and chronic symmetrical polyarthritis. Additionally, arthralgia, myalgia, or fibromyalgia syndromes alone are not criteria for musculoskeletal involvement.

2. Nervous system. Any of the following, alone or in combination: lymphocytic meningitis; cranial neuritis, particularly facial palsy (may be bilateral); radiculoneuropathy; or, rarely, encephalomyelitis. Encephalomyelitis must be confirmed by demonstration of antibody production against B. burgdorferi in the CSF, evidenced by a higher titer of antibody in CSF than in serum. Headache, fatigue, paresthesia, or mildly stiff neck alone are not criteria for neurologic involvement.

3. Cardiovascular system. Acute onset of high-grade (2nd-degree or 3rd-degree) atrioventricular conduction defects that resolve in days to weeks and are sometimes associated with myocarditis. Palpitations, bradycardia, bundle branch block, or myocarditis alone are not criteria for cardiovascular involvement.

Non-confirmatory. Other non-confirmatory signs and symptoms include:
- Fever, sweats, chills, fatigue, neck pain, arthralgias, myalgias, fibromyalgia syndromes, cognitive impairment, headache, paresthesias, visual/auditory impairment, peripheral neuropathy, encephalopathy, palpitations, bradycardia, bundle branch block, myocarditis, or other rash.

Disease endemic to county. A county in which Lyme disease is endemic in which at least two confirmed cases have been acquired in the county or in which established populations of a known tick vector are infected with B. burgdorferi. For the purposes of surveillance, all Wisconsin counties are considered as endemic.

Exposure. Exposure is defined as having been (less than or equal to 30 days before onset of EM) in wooded, brushy, or grassy areas (i.e., potential tick habitats) in a county in which Lyme disease is endemic. A history of tick bite is not required.

Laboratory evidence. For the purpose of surveillance, the definition of a qualified laboratory assay is (1) a positive culture for B. burgdorferi, (2) two-tier testing* with IgM immunoblot seropositive result for specimens collected within 30 days of onset date, or (3) single-tier IgG immunoblot seropositive interpreted using established criteria. Additional assays may be added based on periodic review of the scientific literature and strong evidence of comparable or better performance than qualifying assays.

* Two-tier testing includes an initial screen by enzyme immunoassay (EIA) or indirect immunofluorescence assay (IFA), followed by a Western immunoblot on any equivocal or positive EIA or IFA results.