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Antibiotics for the neurological complications of Lyme disease (Review)

Cadavid D, Auwaerter PG, Rumbaugh J, Gelderblom H

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[Intervention Review]

Antibiotics for the neurological complications of Lyme disease

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ABSTRACT

Background

Various central nervous system-penetrant antibiotics are bactericidal in vitro and in vivo against the causative agent of Lyme neuroborreliosis (LNB), *Borrelia burgdorferi*. These antibiotics are routinely used clinically to treat LNB, but their relative efficacy is not clear.

Objectives

To assess the effects of antibiotics for the treatment of LNB.

Search methods

On 25 October 2016 we searched the Cochrane Neuromuscular Specialised Register, the Cochrane Central Register of Controlled Trials (CENTRAL), MEDLINE, and Embase. We searched clinical trial registers on 26 October 2016. We reviewed the bibliographies of the randomized trials identified and contacted the authors and known experts in the field to identify additional published or unpublished data. There were no language restrictions when searching for studies.

Selection criteria

Randomized clinical trials of antibiotic treatment of LNB in adults and children that compared any antibiotic treatment, including combinations of treatments, versus any other treatment, placebo, or no treatment. We excluded studies of entities considered as post-Lyme syndrome.

Data collection and analysis

We used standard methodological procedures expected by Cochrane.

Main results

We identified seven randomized studies involving 450 European participants with LNB for inclusion in this systematic review. We found no trials conducted in the United States. Marked heterogeneity among these studies prevented meta-analysis. None of the studies included a placebo control on the initial antibiotic treatment, and only one was blinded. None were delayed-start studies. All were active comparator studies, and most were not adequately powered for non-inferiority comparison. The trials investigated four antibiotics: penicillin G and ceftriaxone in four studies, doxycycline in three studies, and cefotaxime in two studies. One study tested a three-month course of oral amoxicillin versus placebo following initial treatment with intravenous ceftriaxone. One study was limited to children. The trials measured efficacy using heterogeneous physician- or patient-reported outcomes, or both. In some cases cerebrospinal fluid analysis was included as an indirect biomarker of disease and outcome. None of the studies reported on our proposed primary outcome, 'Improvement in a measure



of overall disability in the long term (three or more months).' None of the trials revealed any between-group differences in symptom resolution in response to active treatment. In general, treatment was tolerated well. The quality of adverse event reporting, however, was low.

Authors' conclusions

There is mostly low- to very low-quality clinical evidence from a limited number of mostly small, heterogeneous trials with diverse outcome measures, comparing the relative efficacy of central nervous system-penetrant antibiotics for the treatment of LNB. The few existing randomized studies have limited power and lack consistent and well-defined entry criteria and efficacy endpoints. It is not possible to draw firm conclusions on the relative efficacy of accepted antibiotic drug regimens for the treatment of LNB. The majority of people are reported to have good outcomes, and symptoms resolve by 12 months regardless of the antibiotic used. A minority of participants did not improve sufficiently, and some were retreated. These randomized studies provide some evidence that doxycycline, penicillin G, ceftriaxone, and cefotaxime are efficacious in the treatment of European LNB. No evidence of additional efficacy was observed when, in one study, an initial antibiotic treatment with intravenous ceftriaxone was followed by additional longer treatment with oral amoxicillin. There is a lack of evidence identified through our high-quality search strategy on the efficacy of antibiotics for treatment of LNB in the United States.

PLAIN LANGUAGE SUMMARY

Treatment for the neurological complications of Lyme disease

Review question

Are antibiotics effective for the treatment of Lyme disease affecting the nervous system?

Background

In humans, a bacterium called *Borrelia burgdorferi* causes Lyme disease. People become infected when bitten by ticks carrying the bacterium. The person may experience symptoms in the joints, skin, muscles, and nervous system (peripheral nerves (nerves outside the brain and spinal cord), the brain, and the spinal cord). Without antibiotic treatment, neurological Lyme disease either may resolve or cause long-term problems. Neurological Lyme disease differs between Europe and the United States, probably because of differences in *B. burgdorferi*. Limited information exists about which antibiotics are better for the treatment of neurological Lyme disease.

Study characteristics

We found seven trials studying antibiotic treatments for neurological Lyme disease. All but one trial compared different antibiotics. The other trial compared the treatment effects of oral amoxicillin to placebo following initial ceftriaxone treatment. The trials included 450 Europeans. The antibiotics tested were penicillin G, doxycycline, ceftriaxone, and cefotaxime. One of the trials involved children only, while the others included mostly adults. We only selected studies in which treatment allocation was determined by chance (randomly), as such studies provide the best information for comparing the effects of different treatments. Most studies were not blinded (meaning that those taking part and the study staff knew the treatment being given). We could not find any studies of antibiotic treatments for neurological Lyme disease from the United States. No studies assessed the effects of delaying the start of treatment.

Key results and quality of the evidence

The seven studies were too different for their results to be combined, so we analyzed them individually.

None of the studies provided clear evidence that one antibiotic was better than another. One study failed to find evidence that a second and longer treatment with an oral antibiotic (amoxicillin) offered any extra benefit following initial intravenous treatment with ceftriaxone. As none of the other studies used a dummy treatment (placebo), the extra benefit offered by antibiotic treatment over recovery that occurs naturally is unknown. In general, the treatment was tolerated well, although the quality of adverse event reporting in most studies appeared to be low.

The results indicate that treatment with any of the four antibiotics produced similarly good outcomes for treatment of neurological Lyme disease in Europe. A second treatment with amoxicillin does not appear to provide added benefit to ceftriaxone. We found no trials of antibiotics for treatment of neurological Lyme disease in the United States.

The evidence is current to October 2016.



BACKGROUND

Description of the condition

Lyme neuroborreliosis (LNB) is a group of diseases that can affect the central nervous system (CNS) and the peripheral nervous system (PNS), or both, as a result of infection with or the postinfectious consequences of different species of the spirochete bacterium Borrelia burgdorferi. These organisms are transmitted by ixodid ticks in endemic areas in the United States and Europe. Although a multitude of clinical manifestations of LNB have been reported, the most common are radicular pains, facial paralysis, and meningitis, referred to as Bannwarth's syndrome in Europe (Bannwarth 1941; Bannwarth 1944). It was not until 1981 that entomologist Willy Burgdorfer and colleagues in the United States suspected that the cause of Lyme disease was a tick-borne spirochete (Burgdorfer 1982). In the decades since the identification of *B. burgdorferi*, it has become clear that LNB is one of the most common and important complications of Lyme disease. The diagnosis of LNB requires confirmation of infection with B. burgdorferi plus evidence of involvement of the CNS, the PNS, or both. According to the Centers for Disease Control and Prevention, from the 154,405 cases of Lyme disease reported during 2001 to 2010 in the United States, 14% were identified with facial palsy, radiculoneuropathy, meningitis, or encephalitis (CDC 2011a). Looking at Lyme disease occurring in Europe, others have estimated that up to 12% of cases have neurological manifestations (Koedel 2015), and that approximately 5% of individuals with an untreated erythema migrans will develop LNB (Hansen 2013).

Knowledge of the natural course and prognosis of untreated LNB is limited, and both increases in severity of disease and spontaneous remissions may occur. A random review of medical records from the original Lyme disease investigation among people from Connecticut, United States, who were not treated with antibiotics because they were diagnosed before the infectious cause of the disease was known, revealed that when left untreated, LNB can result in long-term sequelae (Kalish 2001). In this report, 31 people who had presented with facial palsy and meningism frequently went on to develop more disseminated manifestations of LNB, with two-thirds being formally diagnosed with lymphocytic meningitis, radiculoneuritis, or both, and 1 in 5 developing atrioventricular block. In a German retrospective study of 72 people with untreated LNB, only 59 went into full remission, whereas 13 developed mild-to-moderate sequelae during 5 to 27 years of follow-up. Importantly, all those participants were eventually judged as "having been cured without antibiotics" (Kruger 1989).

The incidence of LNB varies widely among European countries, with the highest incidences in central European and Scandinavian countries. Population-based annual incidence rates of LNB in central Europe are 30 to 50 per million for acute LNB and less than 0.4 per million for chronic LNB (Hansen 2013).

Description of the intervention

People with the characteristic skin lesion of Lyme disease, erythema migrans, followed by manifestations of infection of the nervous system, referred to as LNB, were successfully treated with antibiotics (penicillin) as early as 1948 (Hollstrom 1951). Treatment with antibiotics capable of crossing the blood-brain barrier is now the standard of care for people diagnosed with LNB. However, no placebo-controlled trials have ever been performed, and the antibiotic of choice, route of administration, dose, and length of treatment for LNB remain controversial.

Why it is important to do this review

At the time that the protocol for this review was conceived, there had been several attempts at producing treatment guidelines for Lyme disease and no high-quality systematic evidence reviews to synthesize the available evidence to feed those reviews. Guidelines have now been produced by the American Academy of Neurology (AAN) (Practice Parameter) (Halperin 2007), the European Federation of Neurological Societies (EFNS) (Mygland 2010), the German Neurological Society (Rauer 2012), and the International Lyme and Associated Diseases Society (ILADS) (Cameron 2014). American Academy of Neurology/American College of Rheumatology/Infectious Diseases Society of America as well as the German Neurological Society are currently working on updated guidelines. This review will continue to synthesize the evidence for the antibiotic treatment of LNB.

OBJECTIVES

To assess the effects of antibiotics for the treatment of Lyme neuroborreliosis.

METHODS

Criteria for considering studies for this review

Types of studies

The review authors only considered quasi-randomized and randomized, prospective, controlled trials of antibiotic treatment for Lyme neuroborreliosis (LNB) for inclusion in this review. We excluded non-randomized and uncontrolled studies. We also excluded single-case reports and case series.

Types of participants

We considered trials that evaluated individuals with clinically diagnosed LNB. The clinical syndrome of LNB included one or more of the following: meningitis, encephalitis, myelitis, radiculitis, cranial neuropathies (including facial nerve palsy or ocular motor palsy, or both), optic neuritis, peripheral neuropathies, and myopathies. We excluded studies of people with post-Lyme disease syndrome, defined as people with persistent symptoms attributed to Lyme disease in the absence of ongoing infection following prior antibiotic treatment. We also excluded trials of LNB prevention through treatment of erythema migrans.

Diagnoses of LNB are supported by a number of approaches, including positive serologic testing (Stiernstedt 1988; Dressler 1993; Anonymous 1995; Engstrom 1995; Wilske 2000). Cerebrospinal fluid (CSF) culture of *B. burgdorferi* was also acceptable as supportive evidence of LNB, as well as a positive CSF Lyme polymerase chain reaction (PCR), the presence of CSF anti-*B. burgdorferi* antibodies, Steere 1990, or CSF pleocytosis, or both (Halperin 1996). Peripheral neuropathy and myopathy required clinically detectable impairment of motor or sensory function confirmed by abnormal electrophysiological tests.

Types of interventions

We considered any antibiotic treatment, including combinations of treatments, versus any other treatment, placebo, or no treatment.

Types of outcome measures

When designing the review we prespecified several primary and secondary outcome measures to assess the efficacy of antibiotics for treatment of LNB, as follows.

Primary outcomes

- 1. Improvement in a measure of overall disability in the long term (three or more months) following treatment.
- 2. Improvement or resolution of the person's presenting neurological deficits in the long term (three or more months) following treatment. In general, we considered improvement as determined and defined by the original authors, provided that they included objective findings as criteria for the outcome.

Secondary outcomes

- 1. Improvement in a measure of overall disability in the short term (two weeks) following treatment.
- 2. Resolution of CSF pleocytosis following treatment.
- For people with peripheral neuropathy or myopathy, improvement in electrophysiological abnormalities following treatment.
- 4. Occurrence of one or more adverse events. We considered serious adverse events, defined as those which required hospitalization or that were life-threatening or fatal, adverse events requiring discontinuation of treatment or substitution of alternative treatment, and any other adverse events as defined and reported by the original authors.

If cost and cost-effectiveness information was available, we planned to include it as part of the effects of the intervention analysis.

Search methods for identification of studies

Electronic searches

On 25 October 2016, we searched the Cochrane Neuromuscular Specialised Register, the Cochrane Central Register of Controlled Trials (CENTRAL) in the Cochrane Register of Studies Online), MEDLINE (1966 to October 2016), and Embase (1980 to October 2016). On 26 October 2016, we also searched trials registers: US National Institutes of Health Ongoing Trials Register ClinicalTrials.gov, the EU Clinical Trials Register (www.clinicaltrialsregister.eu), the World Health Organization Clinical Trials Registry Platform (www.who.int/ictrp/en/), and ISRCTN Registry (www.isrctn.com/).

The detailed search strategies are in the appendices: MEDLINE (Appendix 1), Embase (Appendix 2), CENTRAL (Appendix 3), Cochrane Neuromuscular Specialised Register (Appendix 4), and trials registers (Appendix 5).

Searching other resources

We reviewed the bibliographies of the randomized trials identified, and contacted authors and known experts in the field to identify additional published or unpublished data. We handsearched conference proceedings for additional trials. We applied no language restriction when searching for studies.

Data collection and analysis

Selection of studies

All review authors checked titles and abstracts identified from the searches to determine which studies met the eligibility criteria. When the review authors could not determine eligibility from the title and abstract, they obtained the full text of all potentially relevant studies for independent assessment. Two review authors independently assessed and decided which of the trials identified from the preliminary searches fitted the inclusion and exclusion criteria and graded the risk of bias of the trials. The review authors resolved disagreements about study inclusion by consensus. Two systematic review specialists conducted a duplicate study selection process. The review authors assessed any discrepancies in comparison with their selection.

Data extraction and management

Two review authors independently extracted data from all studies that met the inclusion and exclusion criteria onto a specially designed data extraction form. One of the review authors entered data into the Cochrane Review Manager 5 software (RevMan 2014), and a second review author checked the data extraction. In the case of missing data, the review authors attempted to contact the trial authors. Review authors were not blinded to trial authors, journal, or institution. To assist the review authors, two systematic review specialists conducted an independent data extraction. The Cochrane Neuromuscular Managing Editor created analysis tables and added numerical data to the Results using this data extraction. A review author checked the outcome data entry.

Assessment of risk of bias in included studies

Two review authors independently assessed all of the included studies for risk of bias. In the event of disagreement, all of the review authors achieved consensus through discussion. We used the Cochrane 'Risk of bias' tool to assess risk of bias of the included studies (Higgins 2011). This tool applies the following criteria: random sequence generation; concealment of allocation; blinding of participants and personnel; blinding of outcome assessors; incomplete outcome data (numbers of participants lost to follow-up and use of intention-to-treat analysis); selective reporting; and other sources of bias, such as baseline differences in study populations (other than imbalances caused by inadequate randomization, lack of allocation concealment, or exclusion of participants, which we considered under other criteria). We assessed all included studies under each criterion as at high, low, or unclear risk of bias (we used 'unclear' when there was insufficient information to permit judgement or when what occurred in the study was known but the implications in terms of bias were unclear). To assist the review authors, two systematic review specialists provided by Cochrane conducted an independent 'Risk of bias' assessment, and the review authors addressed any discrepancies in assessments.

Measures of treatment effect

We used Review Manager 5 to calculate risk ratios with 95% confidence intervals for dichotomous outcomes. For continuous data, we reported the mean difference and corresponding 95% confidence intervals. Missing standard deviations were calculated from confidence interval using the Review Manager 5 calculator tool during data entry.



The review authors originally planned to divide the analysis according to whether the LNB studies were North American or European, since infecting strains of Borrelia species as well as clinical manifestations of LNB may differ between the two regions. If the division between European and North American studies did not reveal important differences, we planned to pool all studies. However, we did not identify any eligible study from North America; the review analyses are therefore based only on available European studies until studies from other world areas become available.

Assessment of heterogeneity

Significant heterogeneity of inclusion and exclusion criteria and primary and secondary outcome measures was evident by simple examination. To illustrate this heterogeneity, we have presented a detailed comparison of the study characteristics in Table 1.

Data synthesis

We performed a narrative review of the evidence. In Appendix 6 we provide additional methods relating to the originally planned metaanalysis described in the protocol (Cadavid 2008).

Sensitivity analysis

We did not perform any formal sensitivity analysis because of the small sample size and heterogeneity among the studies.

'Summary of findings' table

We included a 'Summary of findings' table according to recommendations in Chapter 11 of the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). We assessed the evidence using the five GRADE considerations (study limitations, consistency of effect, imprecision, indirectness, and publication bias) to determine the quality of the body of evidence for each outcome, and to draw conclusions about the quality of evidence within the text of the review. We used footnotes to justify any decision to downgrade the quality of evidence. We reported the following outcomes whether or not they were measured or reported in the included studies.

- 1. Improvement in a measure of overall disability in the long term (three or more months) following treatment.
- 2. Improvement or resolution of the person's presenting neurological deficits in the long term (three or more months) following treatment.
- 3. Improvement in a measure of overall disability in the short term (two weeks) following treatment.
- 4. Resolution of CSF pleocytosis following treatment.
- 5. All adverse events.

We presented outcome number two as one outcome, or two if the included studies reported improvement and resolution separately. We created 'Summary of findings' tables for comparisons unless very little data were available for a comparison. We chose to report adverse events in the 'Summary of findings' table as 'All adverse events' with a comment on serious adverse events, as the data were inconsistently reported in the included studies.

RESULTS

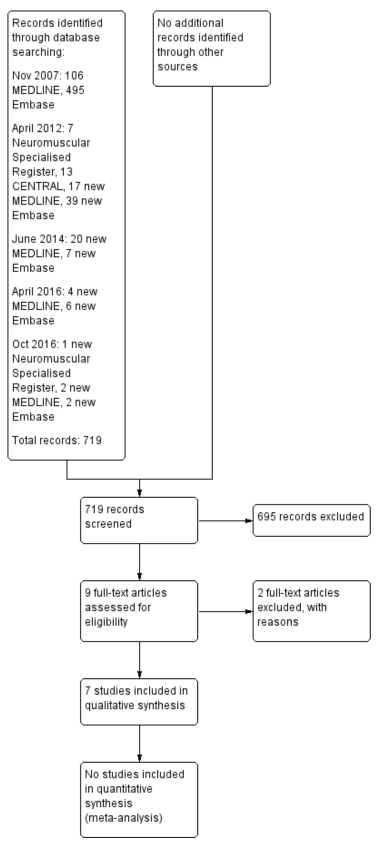
Description of studies

Results of the search

The search strategies listed in the appendices identified 34 studies from the Cochrane Neuromuscular Specialised Register, 35 from CENTRAL, 75 from MEDLINE, and 116 from Embase. The number of references found in each database, the number of studies remaining after removal of duplicates, the number of studies selected for further review, and the number of studies meeting the inclusion criteria are shown in Figure 1. We identified seven trials eligible for inclusion in the review. We identified one ongoing study (NCT02553473).



Figure 1. Study flow diagram.





Included studies

Seven studies fulfilled our predefined inclusion criteria. Table 1 shows an overview of the included studies. All were randomized studies of participants with LNB comparing initial treatment with two different antibiotics, except for one that compared treatment with a second antibiotic to placebo following initial antibiotic treatment. One study only enrolled children. Table 2 shows details of clinical and laboratory eligibility criteria supporting the diagnosis of LNB in the seven studies; all were consistent with current case definitions for LNB (CDC 2011b). Table 3 shows baseline demographics and laboratory findings for enrolled participants by treatment group. The evaluation period in all but one study was a year or less. There was marked heterogeneity in the efficacy assessments used. Baseline characteristics among the seven studies varied widely; disease duration prior to treatment ranged from as short as five days to as long as several months. The number of study participants varied from 22 to 145, with only two studies enrolling over 100 participants. Reporting of concomitant treatment with corticosteroids was incomplete in most studies.

See Characteristics of included studies.

Physician-reported measures of efficacy

In all but one of the included trials, investigators reported neurologic findings on clinical examination, allowing for objective clinical evaluation of efficacy. Physician tools for quantifying efficacy as primary or secondary outcome varied between trials. In the two largest trials, final grading by the treating physician combined objective clinical examination and patient report of subjective symptoms (Oksi 2007; Ljostad 2008). Only the trial by Karlsson 1994 did not report a physician-based judgement.

Patient-reported measures of efficacy

Studies largely used clinical evaluations, which often depended on unspecified symptoms to judge cure, improvement, or failure. These symptoms were typically not systematically tracked. Some studies included people with LNB who were only a subset of participants with disseminated Lyme disease. Subjective patient outcome measures were not usually outlined rigorously but were included within global clinical assessments, or at least inferred. One study, Karlsson 1994, did track specific individual complaints, such as fatigue, during follow-up time points. Ljostad 2008 incorporated subjective symptoms into a clinical composite score, but with a minority weighting compared to objective findings. Oksi 2007 directly incorporated subjective assessments by participants and independent assessments by physicians in a visual analogue scale to assess the success or failure of antibiotic therapy. Pfister 1989 individually tracked radicular pain in all participants.

Table 4 shows an overview of the clinical efficacy assessments used in each study and whether any treatment difference was reported.

CSF parameters

Most trials measured CSF parameters, although there was significant heterogeneity in the percentage of participants whose CSF was examined, the timing of the CSF sampling, and the parameters that were reported, as well as the specific interventions compared (Table 5). The CSF parameter initially prespecified as an outcome measure for this review was resolution of CSF pleocytosis at three or more months following treatment. Of the included trials, Ljostad 2008, Karlsson 1994, Pfister 1989, and Pfister 1991 provided meaningful information on this prespecified parameter. No data were available to determine the effect of any of the antibiotics versus placebo on CSF parameters, except for retreatment with amoxicillin in Oksi 2007. Only two studies looked at the ability to achieve minimum inhibitory concentrations of antibiotics in the CSF.

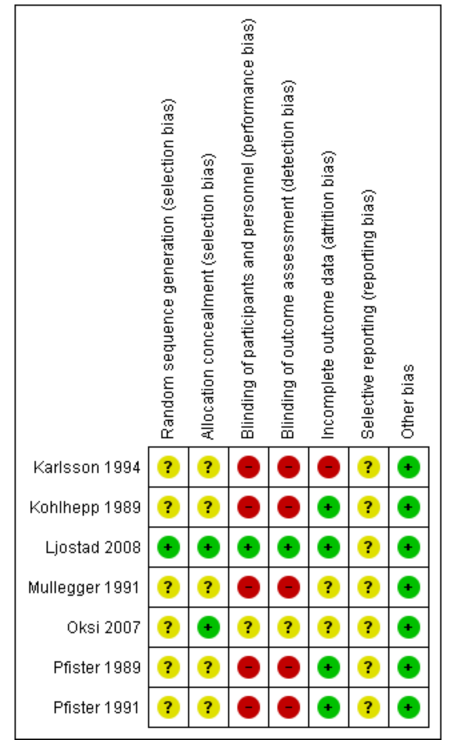
Excluded studies

The Characteristics of excluded studies section refers to relevant randomized studies that were not included in the review and the reasons for their exclusion. We excluded a total of seven studies, one because it tested a non-antibiotic intervention and six that did not specifically address the LNB population.

Risk of bias in included studies

Figure 2 summarizes the review authors' judgements for each 'Risk of bias' domain in the seven included studies.





With the exception of Ljostad 2008 and Oksi 2007 (amoxicillin phase only), all trials were unblinded.

All but one of the studies were at an overall high risk of bias (i.e. at high risk of bias in at least one domain). Only the study by Ljostad 2008 was at low risk of bias, although as no protocol was available, we judged the risk of selective reporting unclear. The risk of bias in Oksi 2007 was mostly unclear, as the information provided on the subgroup of participants with LNB was quite limited. The remaining five included studies were at high risk of bias due to lack of blinding of study participants and outcome assessments. All studies suffered from poor or incomplete reporting in at least one domain, which prevented an adequate assessment of the risk of bias affecting in particular random sequence generation, allocation concealment, and selective reporting in all but Ljostad 2008.



Allocation

Ljostad 2008 was the only study at low risk of selection bias.

Blinding

Ljostad 2008 and Oksi 2007 were the only two studies at low risk of performance and detection bias.

Incomplete outcome data

This domain had the lowest general risk of bias; only Karlsson 1994 was judged to have incomplete outcome data.

Selective reporting

We could not evaluate the risk of bias for this domain for any of the included studies, as we did not have final study protocols or statistical analyses plans, or both for any of them.

Other potential sources of bias

As the reporting of efficacy and safety followed the standard of medical care in the various countries, and none of the studies was identified as having potential financial or other conflicts of interest, we did not consider any of the seven included studies to have other potential sources of bias.

Effects of interventions

Oral antibiotic therapy with amoxicillin versus placebo after previous treatment with 21 days of ceftriaxone for disseminated LNB

Studied in Oksi 2007. See Table 6.

This was the only study to compare amoxicillin treatment with placebo after a three-week course of ceftriaxone. The study report did not distinguish between participants with disseminated Lyme disease and those with LNB. We contacted the trial authors in an effort to obtain separate information on the LNB group and they supplied the supplementary information.

This study provides low- or very low-quality evidence. Despite author transparency, the study remains at high risk of bias overall (unclear risk of bias in most domains), and data for amoxicillin and placebo groups were not reported separately.

Improvement in a measure of overall disability in the long term (three or more months)

Not reported.

Improvement or resolution of the person's presenting neurological deficits in the long term (three or more months)

Oksi 2007 investigated whether 100 days of oral antibiotic therapy with amoxicillin provided any additional benefit after previous treatment with 21 days of ceftriaxone for disseminated Lyme disease. Investigators enrolled 152 participants with definite (74%) or possible (26%) Lyme disease, of whom five were withdrawn due to discontinuation of the study drug and two due to diagnosis other than disseminated Lyme disease. Sixty-two of those enrolled had LNB (Jarmo Oksi, supplementary information). Trialists measured objective efficacy using an investigator-rated visual analogue scale (VAS) that ranged from 0 to 100, administered at baseline (prior to randomization), after completion of ceftriaxone treatment, and at 1, 3, 6, and 12 months thereafter. A value of 50 was attributed to average baseline severity, 0 for full remission of symptoms, and 100 for a certain poor outcome. A similar VAS completed by participants was used for assessment of efficacy. Only in the largest center were participants rated by the same physician throughout the whole study. The degree to which subjective symptoms influenced the investigator VAS judgement was not documented. Also, non-neurologists judged severity of symptoms at some sites. Furthermore, participants with definite and possible LNB were combined. Participants in the definite LNB group had objective signs, making remission of objective neurological signs a more relevant criterion for efficacy assessment in this subgroup. For the disease severity rating, the outcome was categorized as "excellent or good," "controversial," or "poor," using VAS values plus information obtained from participant medical records, but without any standardization.

In participants with definite Lyme disease, the outcome was excellent or good in 49 (92.5%) amoxicillin-treated participants and 47 (87%) placebo-treated participants; the outcome was poor in 3 (5.7%) amoxicillin-treated and 6 (11.1%) placebo-treated participants. In participants with possible Lyme disease treated with amoxicillin, the outcome was excellent or good in 11 (55%) participants and poor in 4 participants (20%); in the placebo group with possible Lyme disease, the outcome was excellent or good in 8 participants (44.4%) and poor in 4 participants (22.2%). In the whole group of 145 participants, the mean differences (MDs) in patient-reported VAS at 3, 6, and 12 months for a moxicillin versus placebo were 4.20 (95% confidence interval (CI) 3.39 to 5.01), -0.50 (95% CI -1.38 to 0.38), and 0.60 (95% CI -0.21 to 1.41). The corresponding MDs for investigator-rated VAS at 3, 6, and 12 months were 0.50 (95% CI -0.28 to 1.28), -2.40 (95% CI -3.18 to -1.62), and -0.40 (95% CI -1.13 to 0.33) (Analysis 1.1). Comparison of all VAS values did not differ significantly between participant and investigator assessments, including the subset of participants with definite LNB (62 of the total 145 participants). The results showed no clinically important differences between amoxicillin and placebo groups on the participant or investigator VAS scores in either the whole group (Analysis 1.2), or the definite Lyme disease subgroup. The report does not provide separate results for the possible LNB subgroup. The trialists reported that there were no statistically significant differences in the definite LNB subgroup between amoxicillin and placebo groups in mean patient or investigator VAS scores at 0, 1, 3, 6, and 12 months. We did not obtain numerical data for analysis.

The risk ratio (RR) for improvement of symptoms (excellent or good according to investigator VAS) at 12 months with amoxicillin versus placebo was 1.06, 95% CI 0.93 to 1.21 in the 107 participants with definite Lyme disease (Analysis 1.3). Investigators concluded that additional amoxicillin therapy was not beneficial, with most participants (59/62) in the LNB subgroup having an excellent or good response regardless of treatment arm. Data from participants with LNB were not reported for amoxicillin and placebo groups separately. Less than 5% of the LNB subgroup ended up with a controversial or poor response on the investigator ratings. The investigators did not find any correlation between the clinical outcomes and persistence without decline of *B. burgdorferi*-specific antibodies in the serum.

A comparison of the overall response to the additional treatment between the definite and possible LNB subgroups showed that the response was better for amoxicillin in the subgroup with definite



LNB than in the subgroup with possible LNB, but this was not the case for placebo.

Improvement in a measure of overall disability in the short term (two weeks)

Not reported.

Resolution of CSF pleocytosis following treatment

Not reported.

Other CSF parameters

Antibody titers

Dr Oksi provided additional information for the number of participants in the LNB subgroup with a decline in *B. burgdorferi*-specific antibodies at 12 months; however, the text is unclear whether serum or CSF levels were reported. Among participants who had LNB, a strong antibody decline (a decrease of over 50%) occurred in 17/30 in the amoxicillin group and 18/32 in the placebo group (RR 1.01, 95% CI 0.65 to 1.56), with a partial decline in 3/30 participants in the amoxicillin group and 2/32 participants in the placebo group (RR 1.60, 95% CI 0.29 to 8.92).

Improvement in electrophysiological abnormalities following treatment

Not reported.

Adverse events

No serious adverse effects of antibiotic treatment occurred in any of the 145 trial participants (Analysis 1.4). Diarrhea was reported in 33 participants (22.8%) during intravenous ceftriaxone treatment and in 19 participants (13.1%) during the second phase of oral treatment, higher with amoxicillin than placebo (15/73 versus 4/72, respectively; RR 3.70, 95% CI 1.29 to 10.61). The diarrhea was usually mild and resolved spontaneously over about two weeks. No participant had to discontinue treatment due to diarrhea. Clostridium difficile colitis was reported in two participants, one in the placebo group and one after "discontinuation of study drug" (amoxicillin or placebo). Cholecystitis or biliary sludging was not observed in any participant. None of the participants developed urticaria or other allergic reactions. Seventeen episodes of fever were reported in 15 participants. Of these, 14 episodes were not attributable to an infection other than *B. burgdorferi* (2 on placebo, 3 on amoxicillin, and 9 on ceftriaxone). One participant developed facial paresis three days after the onset of ceftriaxone treatment, and about half of the participants reported transient intensification of symptoms during ceftriaxone treatment.

Oral doxycycline versus intravenous ceftriaxone for LNB (acute and chronic)

Studied in Ljostad 2008. See Table 7.

The investigators included 118 consecutive participants from nine hospitals in coastal areas of Southern Norway.

Improvement in a measure of overall disability in the long term (three or more months)

Not reported.

Improvement or resolution of the person's presenting neurological deficits in the long term (three or more months)

The primary outcome was a custom-made composite score measured four months after randomization. To calculate this score, the severity of 27 multiple objective neurologic signs (maximum score of 54) and six subjective symptoms (maximum score of 12) were graded from 0 to 2 (total maximum score of 66). Secondary outcomes were full recovery (composite score = 0) at four months and reduction of the composite score two weeks after randomization.

There was no significant difference in reduction in clinical score at 4 months (MD 0.10, 95% CI -1.20 to 1.40; N = 102; Analysis 2.1); the RR for complete resolution of symptoms at 12 months favored doxycyline, but the result was imprecise, with CI including no difference (RR 1.44, 95% CI 0.89 to 2.35; N = 102; moderate-quality evidence; Analysis 2.2).

Improvement in a measure of overall disability in the short term (two weeks)

Not reported.

Resolution of CSF pleocytosis following treatment

In Ljostad 2008, investigators obtained CSF at inclusion and at 13 days and 4 months after the start of antibiotic treatment. No significant difference was found between oral doxycycline and intravenous ceftriaxone for reduction in CSF cell count at 13 days (P = 0.89) or 4 months (P = 0.56) after the start of treatment (data not provided; low-quality evidence).

Improvement in electrophysiological abnormalities following treatment

Not reported.

Adverse events

The safety population included 113 participants with available data. The RR of adverse event between the two groups favored doxycycline, but the data were very imprecise and allowed for the possibility of no difference (RR 0.79, 95% CI 0.51 to 1.23; N = 113; moderate-quality evidence; Analysis 2.3). Three participants discontinued ceftriaxone treatment due to adverse events: one with cholecystitis, one with stomatitis and proctitis, and one with allergy. There were no other serious adverse events. There was one serious adverse event but no withdrawals in the doxycycline group. Results for adverse events leading to discontinuation (RR 0.14, 95% CI 0.01 to 2.71; N = 118; Analysis 2.4) and serious adverse events (RR 0.33, 95% CI 0.04 to 3.05; N = 113; Analysis 2.5) also favored doxycycline but with serious imprecision. Diarrhea, nausea, and urticaria were reported for 19, 15, and 3 participants, respectively; all were generally mild. Emergence of new symptoms compatible with LNB or intensification of symptoms during treatment was not reported.

Ceftriaxone versus penicillin G for acute LNB in children

Studied in Mullegger 1991.

Improvement in a measure of overall disability in the long term (three or more months)

Not measured.



Improvement or resolution of the person's presenting neurological deficits in the long term (three or more months)

No documentation was provided on how clinical status was assessed or whether subjective complaints were incorporated into the assessment in Mullegger 1991. All children (N = 23) initially had objective neurologic findings and recovered completely. Median recovery times differed between interventions: 20 days for penicillin and 33 days for ceftriaxone, with no other statistical information provided. The authors concluded that their study did not show any differences in efficacy between the two treatments.

Improvement in a measure of overall disability in the short term (two weeks)

Not measured.

Resolution of CSF pleocytosis following treatment

This study provided no useful information on CSF parameters with regard to the effect of antibiotic treatment in children, because it did not routinely collect CSF for analysis of parameters after treatment; some were studied but not all reported.

Improvement in electrophysiological abnormalities following treatment

Not reported.

Adverse events

In the ceftriaxone group, transient elevation of serum transaminases (once) and drug-induced toxic skin reaction (twice) were seen. No side effects were noticed in the penicillin G group. Intensification of symptoms or emergence of new symptoms compatible with LNB during treatment was not reported.

Intravenous penicillin G versus oral doxycycline for LNB (acute and chronic)

Studied in Karlsson 1994. See Table 8.

Improvement in a measure of overall disability in the long term (three or more months)

Not reported.

Improvement or resolution of the person's presenting neurological deficits in the long term (three or more months)

Karlsson 1994 compared a 14-day treatment with penicillin G to 14 days of oral doxycycline in 54 participants with LNB. The study included only participants with objective findings and a positive serology or evidence of abnormal CSF. Investigators used a rating of subjective and objective findings on a Likert scale from 0 to 3 (no symptoms, mild symptoms, moderate or severe symptoms) for primary efficacy. The RRs for improvement and resolution with penicillin G versus oral doxycycline at 12 months were 1.0 (95% CI 0.92 to 1.08) and 0.95 (95% CI 0.77 to 1.18), respectively (N = 51; low-quality evidence; Analysis 3.1; Analysis 3.2). Participants were followed for 12 months, with no difference found between the two treatment arms except for the fact that more participants treated with doxycycline reported vertigo at the end of treatment but not at one month. One participant in each treatment group was retreated because of residual symptoms. Subjective symptoms were completely absent at 12 months except for 1 penicillin G participant with neuromuscular pain and hypoesthesia and 1 doxycycline participant with arthralgia. Selection bias is a concern in this study due to a considerable imbalance in the number of participants randomized to each treatment arm. The report provided few statistics and did not allow a distinction to be made between participant- and physician-based judgements. Objective judgement of findings at the end of follow-up was implied but not well documented, thus this study did not use a well-characterized objective measure of efficacy by a physician.

Improvement in a measure of overall disability in the short term (two weeks)

Not reported.

Resolution of CSF pleocytosis following treatment

CSF was obtained at inclusion, at 13 days, and in some participants at one year after the start of treatment. All participants had positive *B. burgdorferi*-specific antibodies in serum, CSF, or both, or had a positive CSF culture (one participant had a positive culture but no specific antibodies) at study entry. The CSF cell count in all 9 participants in the penicillin group and 18 of the 20 participants in the doxycycline group had returned to normal at 1 year (RR 0.93, 95% CI 0.75 to 1.15; N = 29; very low-quality evidence; Analysis 3.3).

Other CSF parameters

All participants in both treatment arms were negative for immunoglobulin M antibodies in the CSF at one year. *B. burgdorferi*-specific CSF immunoglobulin G antibodies were negative in only 5/9 penicillin participants and 9/20 doxycycline participants (56% versus 45%, difference not significant). No significant differences were found between oral doxycycline and intravenous penicillin for any CSF parameter.

Improvement in electrophysiological abnormalities following treatment

Not reported.

Adverse events

Emergence of new symptoms compatible with LNB during treatment was not reported. There was intensification of symptoms during treatment in one participant. One penicillin G-treated participant had a transitory rise of temperature and increased vertigo during treatment. Two participants had thrombophlebitis. Two doxycycline-treated participants had skin rash at the end of treatment; another two reported transient diarrhea (RR for adverse events 1.01, 95% CI 0.25 to 4.08; N = 54; very low-quality evidence; Analysis 3.4).

Intravenous doxycycline versus intravenous penicillin G for LNB (acute and chronic)

Studied in Kohlhepp 1989. See Table 9.

Improvement in a measure of overall disability in the long term (three or more months)

Not reported.

Improvement or resolution of the person's presenting neurological deficits in the long term (three or more months)

Kohlhepp 1989 randomized a clinically well-defined cohort of 75 participants with predominantly acute (n = 67) but also chronic (n = 67)



8) LNB to a 10-day course of intravenous doxycycline or intravenous penicillin G. Follow-up was 12 months, but for cases with residual symptoms the follow-up was three years. The primary outcome was the treating physicians' categorical grading of the clinical status as "no remission," "partial remission," or "full remission," based on objective and subjective signs and symptoms with no specification given. At the end of treatment, over 80% of participants had responded to some degree in both groups. Early responders were usually asymptomatic after six months. Pain, meningitic symptoms, and acute cranial neuritis began to remit within days. A slower improvement was observed in symptoms of radiculitis, myelitis, encephalitis, and peripheral neuropathy. According to data in Figure 2 of the study report, at six months the RR for "partial remission" (improvement) showed no clear difference between interventions (RR 1.10, 95% CI 0.95 to 1.28), whereas the RR for "full remission" (resolution) favored doxycycline, but with the possibility of no effect (RR 1.42, 95% CI 0.83 to 2.42; lowquality evidence; Analysis 4.1 and Analysis 4.2). At 12 months, the RRs for "partial remission" and "full remission" were 0.98, 95% CI 0.80 to 1.21 and 0.96, 95% CI 0.70 to 1.31, respectively (low-quality evidence; Analysis 4.1 and Analysis 4.2). Of the 22 participants with only partial remission after 6 months, 10 chose to receive retreatment with penicillin G, 6 from the penicillin arm and 4 from the doxycycline arm. Three years after randomization, the recovery rate was 94% in the doxycycline and 91% in the penicillin G group if the retreatment group was excluded. In the "partial remission" group, 7/10 participants who chose retreatment recovered completely, compared to 7/12 of those who did not choose retreatment (no significant difference). Participants with partial remission had central nervous system involvement, a disseminated clinical picture, and/or a longer disease duration. The authors concluded that there was no clinically relevant difference between doxycycline and penicillin G. The number of participants with chronic LNB was too low for any subgroup analysis. In addition, the majority of these chronic LNB cases were also treated with immunosuppressants.

Improvement in a measure of overall disability in the short term (two weeks)

Not reported.

Resolution of CSF pleocytosis following treatment

Kohlhepp 1989 did not report resolution of CSF pleocytosis with sufficient detail for reporting in the review.

Improvement in electrophysiological abnormalities following treatment

Not reported.

Adverse events

None of the participants experienced serious side effects such as a Jarisch-Herxheimer reaction or developed an allergic reaction. Intensification of symptoms or emergence of new symptoms compatible with LNB during treatment was not reported.

Intravenous cefotaxime versus intravenous penicillin G for acute LNB

Studied in Pfister 1989. See Table 10.

Improvement in a measure of overall disability in the long term (three or more months)

Not reported.

Improvement or resolution of the person's presenting neurological deficits in the long term (three or more months)

Pfister 1989, in an open-label study, randomized 21 participants with acute LNB (Bannwarth's syndrome, meningitis) to a 10day treatment with either cefotaxime or penicillin G. Neurologic examination was performed daily during treatment or at follow-up (mean 7.7 +/- 2.4 months). In addition, the study authors scored radicular pain daily during therapy. Most participants improved by day 3 or 4 of therapy. Eight of the 10 participants in the penicillin G group and 9/11 in the cefotaxime group had complete remission at follow-up, with slight residual findings like mild radicular hypoesthesia or mild paresis in the rest (RR 1.02, 95% CI 0.67 to 1.55; N = 21; low-quality evidence; Analysis 5.1). Pain subsided in all participants within nine days, with the exception of recurrent radicular pain lasting five weeks in one penicillintreated participant. While on therapy, four participants developed facial palsy, radicular symptoms, or bilateral proximal arm paresis, which subsided by the end of therapy. The authors stated that there was no difference in neurological examinations or pain scorings between groups during treatment and at follow-up, with no statistics given.

Improvement in a measure of overall disability in the short term (two weeks)

Not reported.

Resolution of CSF pleocytosis following treatment

CSF was obtained prior to treatment and 8 to 10 days after the start of treatment (treatment duration was 10 days). There was no significant difference between the two treatment arms for abnormal CSF on repeated lumbar puncture on any of the CSF parameters, including pleocytosis, protein elevation, oligoclonal bands, or positive culture.

CSF cell counts normalized in almost all participants (10/10 in the penicillin G group and 10/11 in the cefotaxime group; RR 0.92, 95% CI 0.71 to 1.18; N = 21; very low-quality evidence; Analysis 5.2), while CSF oligoclonal bands persisted in 6 participants from each treatment group, and intrathecal antibody production was still present in 1 participant from each group.

Other CSF parameters

CSF cefotaxime concentrations reached the minimum inhibitory concentration (MIC) in all participants, while none of the participants treated with penicillin G had CSF concentrations above the MIC.

Improvement in electrophysiological abnormalities following treatment

Not reported.

Adverse events

No side effects of antibiotic treatment were observed (N = 21; lowquality evidence). Two participants per group developed a new symptom compatible with LNB during treatment. Intensification of symptoms during treatment was not reported.

Antibiotics for the neurological complications of Lyme disease (Review)

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Intravenous ceftriaxone versus intravenous cefotaxime for acute LNB

Studied in Pfister 1991. See Table 11.

Improvement in a measure of overall disability in the long term (three or more months)

Not reported.

Improvement or resolution of the person's presenting neurological deficits in the long term (three or more months)

Pfister and colleagues compared a 10-day treatment with ceftriaxone to cefotaxime in acute LNB (Bannwarth's syndrome in 28/33 participants, 30 of whom could be evaluated). They reported no clinical difference between groups as judged by objective and subjective neurologic signs and symptoms during treatment or at follow-up (mean 8.1 +/- 1.9 months). In most participants clinical improvement was observed on days 3 to 5 of therapy, while two participants in each group did not respond or deteriorated during treatment. Sixty-three per cent of participants were asymptomatic at follow-up, while 37% remained symptomatic, although all but 1 participant in each arm had improved to mild symptoms; 1 participant in each treatment arm had recurrent radicular pain with either pleocytosis or isolation of *B. burgdorferi* from the CSF. Symptom resolution at follow-up occurred in 8/12 participants receiving ceftriaxone and 9/15 participants receiving cefotaxime (RR 1.11, 95% CI 0.63 to 1.97; low-quality evidence; Analysis 6.1).

Improvement in a measure of overall disability in the short term (two weeks)

Not reported.

Resolution of CSF pleocytosis following treatment

CSF was obtained prior to treatment and very early at 2 to 4 days after the start of treatment (treatment duration was 10 days). No significant difference was found between the two treatment arms for abnormal CSF parameters on repeated lumbar puncture, including pleocytosis, protein elevation, oligoclonal bands, or positive culture. Cell counts normalized in all 10 participants in the ceftriaxone group and in 11/13 participants in the cefotaxime group with only mild pleocytosis, while oligoclonal bands persisted in 5 participants from the ceftriaxone group and 7 from the cefotaxime group, and culture was positive in 1 participant from the ceftriaxone yers cefotaxime was 1.14 (95% CI 0.90 to 1.44; N = 27; very low-quality evidence; Analysis 6.2).

Other CSF parameters

There was no significant difference in the ability of the two antibiotics to reach MIC in the CSF.

Improvement in electrophysiological abnormalities following treatment

Not reported.

Adverse events

Both drugs were generally well tolerated. Intensification of symptoms during treatment was reported for one participant per group. One cefotaxime recipient developed allergic exanthema on day 9, so therapy was stopped. One ceftriaxone recipient developed fever and diarrhea and had mildly elevated liver enzymes on day 5. All symptoms and laboratory abnormalities resolved after his therapy was terminated on day 9. Two cefotaxime recipients had a worsening soon after antibiotic infusion consistent with the Jarisch-Herxheimer reaction. The RR for adverse events with ceftriaxone versus cefotaxime was 0.38 (95% CI 0.04 to 3.26; N = 30; low-quality evidence; Analysis 6.3).

Cost-effectiveness

We found no cost-effectiveness information in any included study.

DISCUSSION

In clinical practice, almost all people with identified symptomatic bacterial infections are treated with antibiotics. This is also true of people diagnosed with Lyme neuroborreliosis (LNB) (Halperin 2015). However, uncertainty exists about the absolute and relative efficacy of the available antibiotic regimens for LNB. This review did not address the prevention of LNB following initial antibiotic treatment of erythema migrans, the optimal duration of treatment with the various antibiotics, or the efficacy of antibiotics for treatment of late neurological manifestations of Lyme disease, sometimes referred to as late- or third-stage LNB. Also, this review did not address the efficacy of antibiotics for treatment of post-Lyme disease syndrome. We included only randomized comparative trials for quality reasons. The majority of the 450 participants from the 7 randomized treatment trials included in this review had typical manifestations of early disseminated (acute or stage II) LNB. Only a minority had objective, defined central nervous system parenchymal involvement consistent with late disseminated (chronic or stage III) LNB (4/145 participants in Oksi 2007 and 6/75 participants in Kohlhepp 1989). In Oksi 2007, three participants had a disease duration of one year or longer.

Summary of main results

We found no high-quality evidence on the absolute or relative efficacy of antibiotics for the treatment of LNB. Although there was some evidence on the relative efficacy of antibiotic treatment for LNB, it was mostly of low or very low quality. Although we selected only randomized studies for inclusion in this review, most lacked consistent, standardized, and well-defined efficacy outcomes. The heterogeneity of eligibility criteria, interventions, and assessment of outcomes meant that we could not perform a meta-analysis.

None of the studies reported on our proposed primary outcome, 'Improvement in a measure of overall disability in the long term (three or more months).' None of the trials reported a significant difference between antibiotic treatments by physician- or patientreported measures of efficacy. In the studies that reported lack of efficacy or partial response separately, there were no differences between randomized treatment regimens. However, differences in eligibility, the assessments used, and the duration of followup among trials made any assessment of differences in outcomes difficult. In all seven trials, the majority of participants were reported as having recovered completely. The lack of a placebo group for the initial antibiotic treatment prevented assessment of the efficacy of antibiotic treatment versus natural recovery.

The majority of participants enrolled in these seven studies presumably had European LNB, predominantly acute LNB, and appear to have had good clinical outcomes (resolution of presenting signs and symptoms) regardless of whether the initial



antibiotic treatment was ceftriaxone, cefotaxime, penicillin G, or oral or intravenous doxycycline. In a single study (Kohlhepp 1989), the need for retreatment (as determined by the incomplete resolution of symptoms) was 29%, but this was not addressed in most studies. Retreatment with antibiotics usually occurred in people with chronic or stage III LNB, supporting the need for antibiotic treatment as early as possible to prevent disease progression to stage III LNB. A single trial that examined the question of prolonged treatment with a second antibiotic found no evidence of additional efficacy when extending ceftriaxone treatment with amoxicillin, where improvement occurred with antibiotic and placebo at the same rate (small numbers) (Oksi 2007). In Oksi 2007, participants with "probable" Lyme disease did not respond to antibiotic treatment as well, in general, as participants with definite LNB, with the implication that some "probable" participants may have had alternative causation for their symptoms.

In summary:

- We found no clinical trials to evaluate the absolute efficacy of initial treatment with antibiotics for LNB. The lack of placebocontrolled studies prevented us from assessing the extent to which antibiotics improve spontaneous recovery from LNB or prevent further complications.
- 2. Seven randomized studies mostly at high risk of bias and of marked heterogeneity provided some relative efficacy data for antibiotic treatment of LNB, only one of which was blinded (Ljostad 2008). All studies were from Europe.
- 3. Marked heterogeneity among the eligible studies in terms of differences in inclusion and exclusion criteria, supportive laboratory diagnostic criteria, primary and secondary outcome measures, treatment regimens, prior antibiotic treatment, and duration of disease and follow-up prevented incorporation of results into a systematic meta-analysis. The quality of the evidence was mostly low to very low; we have formally presented this in 'Summary of findings' tables.
- 4. All studies reported improvement in the majority of participants following antibiotic treatment, and the majority had complete resolution of their symptoms in long-term followup, irrespective of the antibiotic regimen received and in the case of doxycycline regardless of whether it was given orally or intravenously.
- 5. Only three studies provided information on the need for retreatment (Kohlhepp 1989; Karlsson 1994; Ljostad 2008).
- 6. Where measured, objective biomarkers of response (CSF pleocytosis) recovered in almost all participants examined at follow-up.
- 7. Incomplete or poor treatment responses for the efficacy outcomes used in the trials were reported in only a minority of participants, regardless of the antibiotic used. Where no or partial response was recorded, there were no obvious differences in the rates of partial or lack of response between treatments.
- All of the antibiotics studied appear to have been generally well tolerated as judged by all adverse event reporting. Only four studies provided information on discontinuation due to adverse events (Pfister 1989; Mullegger 1991; Pfister 1991; Ljostad 2008), which can have a major impact on the outcome of the treatment.
- 9. The single study conducted in children treated very early in their disease reported full remission in all children. This study had

a high risk of bias, with qualitative outcomes and incomplete follow-up data.

Overall completeness and applicability of evidence

As placebo-controlled studies have never been performed, the extent to which antibiotic treatment contributes to the natural recovery (absolute efficacy) of LNB is not known. Our review was therefore only able to summarize outcome data for the comparative effectiveness of antibiotics (relative efficacy) when used for the initial treatment of LNB. This review did not address the prevention of LNB following initial antibiotic treatment of erythema migrans, and we identified only one randomized controlled trial (RCT) that investigated the efficacy of retreatment following initial antibiotic treatment with ceftriaxone.

The participants in the seven trials included in the review are likely to all have been representative of LNB from Europe. We did not find any randomized studies that included participants with LNB from the United States. LNB is caused by a number of different *Borrelia burgdorferi sensu lato* species. In Europe it is caused by *B. garinii* and *B. afzelii* and only infrequently by *B. burgdorferi sensu stricto*, while in the United States all cases are caused by *B. burgdorferi sensu stricto*. Possible differences in clinical presentation and disease course between Europe and the United States are grounds for caution in applying the findings of this review to the development of guidance for treatment of LNB in the United States. For this review to people with LNB from the United States has limitations.

All studies used diagnostic criteria for LNB consistent with currently accepted case definitions (CDC 2011b). Owing to the paucity of data from late disseminated LNB, the evidence applies mostly to early disseminated (acute or stage II) LNB rather than to late disseminated (chronic or stage III) LNB.

One challenging aspect of LNB therapy concerns the definition of initial antibiotic treatment failure. Lyme immunoglobulin G antibodies postinfection may persist for many years despite complete clinical response, and on its own is not a diagnostic test of active infection with B. burgdorferi or a reliable criterion for treatment failure. A prospective Danish study of 187 consecutive LNB participants with intrathecal B. burgdorferi-specific antibody synthesis who were treated with penicillin G between 1985 and 1990 could not find a single case of treatment failure based on the clinical outcome and normalization of CSF (Hansen 1992). In contrast, in the studies included in this review, there were reported instances of treatment failure or only partial response, mostly for late disseminated (chronic or stage III) LNB. A second course of treatment with a different antibiotic in a single study was associated with a response in no more participants than placebo (Oksi 2007). Objective quantitative measures of treatment 'failure' are few. The resolution of CSF pleocytosis in 99/106 participants in whom measurements were available provide biomarker evidence of a treatment response in most cases. Qualitative symptom reports and measures of cognitive impairments are non-specific and error prone. Proper assessment of the response to antibiotic treatment in LNB requires a uniform definition of diagnostic criteria, a better understanding of the natural and treated history of the disease, and an understanding of what constitutes a treatment failure from both the patient and physician. Inclusion of subjective symptomatology is patient focused but makes it difficult to distinguish symptoms



from other origins. All but one of the seven trials included in this review included both objective and subjective assessments of efficacy, and these correlated relatively well on the improvement from baseline. Further research is needed to prove the link between infection, ongoing versus pre-existing damage, presenting versus residual symptoms, signs, and other comorbid conditions.

Quality of the evidence

We identified limitations in the design of all the included studies, in particular the lack of blinding in the majority of studies. We found a lack of standardization of entry criteria and of efficacy assessments across trials. All studies had a small sample size, with variable duration of follow-up. The confidence in our estimates on the efficacy of the studied antibiotics for LNB is limited, and the true effects may well be substantially different. We thus graded the quality of the evidence as mostly low to very low, with the GRADE definition of 'low' being "further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate" (GRADE Working Group 2004).

Potential biases in the review process

The review process extended over eight years from initial protocol to completion of the systematic review. The review team included specialists in neurology and infectious diseases and experts in the diagnosis, treatment, and pathogenesis of LNB from both the United States and Europe, who worked as unpaid volunteers. To minimize the risk of publication bias, we performed a comprehensive search for studies and also solicited information from investigators of included trials for unpublished data. We also extended requests for unpublished RCT data to known specialists in the field and to leaders of patient organizations to allow their membership to respond (International Lyme and Associated Diseases Society). Outcome measures differed in the trials, and recording of adverse drug reactions was limited, therefore it was difficult for the review authors to interpret the effect of antibiotic intervention positively or negatively except in general terms. The limited number of participants studied in individual trials also meant that this review could not detect less common or rare adverse events.

We revised the scope of the review to focus on antibiotic treatment for LNB, the usual treatment for Lyme disease, which resulted in the exclusion of one RCT that assessed the effect of steroids (Pfister 1988). We revised our methods section to include current Cochrane methodology, including 'Summary of findings' tables, implemented since publication of our protocol (Cadavid 2008),

We attempted to reduce potential biases in the review process and analytical biases by reselecting studies and extracting data with an independent team.

Agreements and disagreements with other studies or reviews

In their recent systematic review, Dersch and colleagues considered both randomized and non-randomized trials investigating all pharmacological treatments and focused on adults with acute LNB (Dersch 2015). We identified one RCT of non-antibiotic treatment (Pfister 1988), which they listed as a reference but did not include. We agree with their conclusion that the heterogeneity among the selected RCTs and the overall bias precludes conclusion on the preference of any antibiotic regimen investigated. However, unlike Dersch and colleagues, we considered the studies to be heterogeneous to a degree that prevented pooling of data from different studies into a meta-analysis. Differing from the study selection of Dersch and colleagues, we excluded two studies, mainly due to the lack of objective evidence of neurological disease (Hassler 1990; Oksi 1998). We did, however, include two studies not included by Dersch and colleagues, one in adults, Oksi 2007, and one in children (Mullegger 1991). We were able to include Oksi 2007 because the trial authors provided separate information on the LNB participants.

AUTHORS' CONCLUSIONS

Implications for practice

For early disseminated (acute or stage II) Lyme neuroborreliosis (LNB), low- to very low-quality evidence suggests that the majority of people with LNB respond to antibiotics with known brain penetration and appropriate activity spectra, namely penicillin G, ceftriaxone, cefotaxime, and doxycycline, either oral or intravenous. There is no accurate estimate of the absolute efficacy of these antibiotics, as there were no appropriate studies from which to infer this. None of the included trials had a placebo control for initial antibiotic treatment, and most had limited power and lacked consistent and well-defined efficacy endpoints, study duration, and entry criteria. Also, the studies employed a range of treatment duration, from 10 to 21 days. All studies were conducted in Europe and none in the United States, thus no direct conclusion can be drawn on the effect of antibiotic treatment for LNB in the United States. The number of participants with late disseminated (chronic or stage III) LNB in the included trials was very low, and no useful separate conclusions can be drawn about its treatment with antibiotics. We observed no evidence of additional efficacy in a trial extending initial ceftriaxone treatment with amoxicillin. No firm conclusions can be drawn on the relative efficacy of the four antibiotics we reviewed. In the majority of cases reported in the seven included studies, European LNB was treatable with antibiotic regimens recommended in national guidelines, with marked improvement from baseline impairment. A minority of people do not improve after initial treatment.

Implications for research

Clinical research on the treatment of LNB lacks standardization of diagnostic, entry, and efficacy outcome criteria. There is a need for randomized and blinded controlled trials of adequate size and power and with proper study design to compare relative efficacy of antibiotics, in terms of route of administration, treatment duration, safety and tolerability, and the need for retreatment. A multiple-treatments meta-analysis may be possible in the future if adequate data are generated. It is unlikely that a standard placebo arm can be ethically included in any future trial designs, and novel trial designs with delayed start or add-on therapy designs will be required. Future research would benefit greatly from standardization of diagnostic and outcome criteria and treatment failure criteria. Diagnostic research criteria should be sensitive enough to include an acceptable range of patients with different presentations, but also be specific enough to include only patients with Lyme disease-in clinical practice the findings of a trial can be broadened. The assessment of trial outcomes is equally important, as trial-based diagnosis and these outcomes should have both internal and external validity. Assessment of subjective complaints and cognitive impairment is feasible and



should be included. It is important to address not only impairment and derivative biomarkers, but also to measure patient-reported disabilities and activity and participation measures. Rasch-built outcome measures should be preferred over those based on classical test theory.

There is an absence of trials in some important areas. Controlled trials of antibiotic treatment of chronic LNB and also including participants with non-European LNB are required, as current epidemiological and clinical studies suggest different disease courses and etiologic agents of LNB in Europe and the United States. More studies in children are also needed.

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The Information Specialist of Cochrane Neuromuscular, Angela Gunn, assisted with development of search strategies and ran the literature searches.

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CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Karlsson 1994

Methods	Prospective, randomized, non-blinded, parallel trial with an active comparator arm			
Participants	54 participants with clinical signs and symptoms of meningoradiculitis, encephalomyelitis, or chronic meningitis and with elevated <i>Borrelia burgdorferi</i> -specific antibody titers in serum or CSF, or both, or with <i>B. burgdorferi</i> cultured from CSF			
	Doxycycline arm (22 women, 10 men); penicillin arm (13 women, 9 men)			
	Other reported metrics comparable			
	Exclusion criteria included age below 12 years, pregnancy, breast-feeding, allergy to treatment com- pounds, and antibiotic treatment within the previous 4 weeks.			
	See Table 2 and Table 3 for diagnostic criteria and additional baseline characteristics.			
Interventions	14-day course of:			
	 intravenous penicillin G (3 g every 6 hours) (N = 23); or oral doxycycline (200 mg once daily) (N = 31) 			
Outcomes	Participant-scoring daily self report form (0 = no symptoms, 1 = mild, 2 = moderate, 3 = severe).			
	CSF analysis, serologic and clinical follow-up for 1 year. It is unclear how symptoms were scored at lat- er follow-up visits. Symptoms scored included malaise, fatigue, nausea, vomiting, vertigo, headache, neck stiffness, muscular pain, arthralgia, visual and hearing disturbances, hypoesthesia or hyperesthe sia, and paresis. CSF and serum samples were analyzed for antibodies against whole-cell sonicate of <i>B</i> . <i>burgdorferi</i> by ELISA.			
	Assessments were done at 14 days, 3, 6, and 12 months.			
Funding	Not disclosed			
Conflicts of interest	Not disclosed			
Notes	Study years 1987 to 1990			
Risk of bias				
Bias	Authors' judgement Support for judgement			

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Cadavid D, Auwaerter P, Aucott J, Rumbaugh J. Treatment for the neurological complications of Lyme disease. *Cochrane Database of Systematic Reviews* 2008, Issue 1. [DOI: 10.1002/14651858.CD006978]

* Indicates the major publication for the study

Karlsson 1994 (Continued)

Random sequence genera- tion (selection bias)	Unclear risk	Participants were randomly allocated to treatment groups, but the report does not describe the method of randomization. Though a small study, 31 par- ticipants in doxycycline arm and 23 in penicillin G arm would suggest a ran- domization problem. A sex imbalance was present, with many more women in the doxycycline group (N = 22) than in the penicillin G group (N = 13).
Allocation concealment (selection bias)	Unclear risk	Not mentioned
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Non-blinded study
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Non-blinded study
Incomplete outcome data (attrition bias) All outcomes	High risk	Participant dropout was minimal (2 participants in penicillin arm, 1 participant in doxycycline arm). However, participants were randomized and dosed be- fore the serology or culture results were available, and 17 were excluded. The timing of dropout was not specified.
Selective reporting (re- porting bias)	Unclear risk	We found no evidence that all outcomes were prespecified; the pre-study pro- tocol was not available for review, and the study was not registered prior to initiation.
Other bias	Low risk	No other identified

Kohlhepp 1989

Methods	Prospective, randomized, parallel-group, open-label study with active comparator, no placebo
Participants	75 participants with acute and chronic LNB. 12% of participants had a disease duration of ≥ 1 year with no statistically significant difference between treatment groups
	Participants had <i>Borrelia burgdorferi</i> -specific antibodies in serum and at least 3 of the following di- agnostic criteria: radicular pain; meningitic symptoms; cranial neuritis; sensory or motor radiculitis, or both; arthritis or carditis or encephalitis or myelitis or peripheral neuritis; tick bite or erythema mi- grans, or both. All participants were required to also have an elevated <i>B. burgdorferi</i> -specific antibody titer in the serum. The following CSF laboratory parameters were analyzed:
	1. <i>B. burgdorferi-</i> specific antibody titer;
	 lymphocytic pleocytosis (abnormal if > 4 cells/mm³);
	3. elevated CSF protein (> 50 mg/dL);
	4. elevated CSF IgM, IgA, and/or IgG index;
	5. CSF oligoclonal banding.
	See Table 2 and Table 3 for diagnostic criteria and additional baseline characteristics.
Interventions	10-day treatment with either:
	 intravenous doxycycline 200 mg per day for 2 days followed by 100 mg also intravenously per day for another 8 days (N = 39); or intravenous penicillin G 20 mega units per day (N = 36) continuously infused over 16 hours.



Kohlhepp 1989 (Continued)		or all of the following were considered a "treatment failure" and were eligible defined) for a second course of treatment with penicillin G (30 mega units per
	day for 10 days):	defined) for a second course of readment with performance (so mega units per
	relapse or progress	ion of symptoms at the end of therapy;
	no improvement in	
	 >2-fold increase of 	<i>B. burgdorferi</i> -specific antibody concentrations in CSF.
	regimen. Participants	re residual symptoms 3 months after therapy were offered the same therapeutic with chronic encephalomyelitis were also treated with intravenous or intrathecal) and cytarabine (N = 3).
Outcomes		nination (first outcome) was performed prior to the intervention, and 0, 5, 6, and tervention. Investigators graded the clinical status as "no remission," "partial re- ssion."
		ncluded CSF findings (cell count, total protein, IgM index, and intrathecal <i>B.</i> production) and <i>B. burgdorferi</i> -specific IgG concentrations in serum.
Funding	Not disclosed	
Conflicts of interest	Not disclosed	
Notes		
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Participants were randomly allocated to treatment groups, but the report does not describe the method of randomization.
Allocation concealment (selection bias)	Unclear risk	Not mentioned, not clear if it was done
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Not documented. We assume that neither participants nor investigators were blinded to treatment allocation.
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Not documented. We assume that outcome assessors were not blinded to treatment allocation.
Incomplete outcome data (attrition bias) All outcomes	Low risk	All participants returned to follow-up at 1 year or longer.
Selective reporting (re- porting bias)	Unclear risk	No evidence that all outcomes were prespecified; the pre-study protocol was not available for review, and the study was not registered prior to initiation.

Ljostad 2008

Other bias

Methods

Double-blind, double-dummy, randomized clinical non-inferiority trial

No other identified

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Low risk

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jostad 2008 (Continued)				
Participants	118 consecutive adult participants in Norway with prospective newly diagnosed LNB were randomized. 102 were evaluable. Entry criteria were neurological symptoms without alternative explanation and 1 or more of the following criteria: CSF WBC > 5 cells/mL, intrathecal <i>Borrelia burgdorferi</i> -specific anti- body production, and/or verified acrodermatitis chronica atrophicans			
	This study included both acute and chronic LNB, but groups were not analyzed separately, although chronic LNB comprised only 8% and 11% of each arm.			
	See Table 3 for addition	nal baseline characteristics.		
Interventions	14-day course of either	· · · ·		
	 oral doxycycline 200 mg daily (N = 54); or intravenous ceftriaxone 2 g daily (N = 48). 			
Outcomes	The primary endpoint was a composite clinical score administered by experienced clinicians at base- line, 13 days, and 4 months after therapy. Secondary endpoints included the number of participants who had full recovery 4 months after treatment, reduction in CSF cell count at 4 months, and both re- duction in CSF counts and clinical score at day 13.			
Funding	Sørlandet Kompetansefond (100%)			
Conflicts of interest	Authors reported no conflicts of interest.			
Notes	A large number of anticipated participants ended up not qualifying for inclusion and were excluded (N = 18). The study was not powered to investigate true differences in side effects.			
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Low risk	Computerized allocation performed in advance in blocks of 4 participants, stratified according to early disease (duration of symptoms < 6 months) and chronic disease (duration of symptoms > 6 months).		
Allocation concealment (selection bias)	Low risk	Generally comparable baseline characteristics by treatment allocation		
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Investigators who participated in randomization were not further involved in study. A double-blind, double-dummy trial design was used.		
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Physicians were unaware of assignment groups.		
Incomplete outcome data (attrition bias) All outcomes	Low risk	7/55 in ceftriaxone arm and 5/59 in doxycycline arm were excluded due to ei- ther loss to follow-up or new diagnosis. All participants who received doxycy- cline completed a 14-day course; 4 in the ceftriaxone arm did not complete (1 owing to late delivery, 3 owing to adverse reactions).		
Selective reporting (re- porting bias)	Unclear risk	No evidence that all outcomes were prespecified; the pre-study protocol was not available for review, and the study was not registered prior to initiation. Relevant outcomes were reported in the paper, but some with insufficient de- tail for numerical analysis.		

Ljostad 2008 (Continued)

Other bias

Low risk

The study provided separate results for cases of definite and probable LNB both at randomization and by efficacy outcomes.

Methods	Prospective, randomize	ed, parallel-group, open-label study with active comparator, no placebo		
Participants	23 children with acute neurologic symptoms of LNB, i.e. 1 or more of the following symptoms: meningism (n = 17), peripheral facial palsy (n = 14), torticollis (n = 1), VI cranial nerve paresis (n = 1), pseudotumor cerebri (n = 1)			
	Inclusion criteria were:			
	1. presence of neurological signs and symptoms indicative of LNB;			
	2. start of antibiotic treatment within 14 days after onset of symptoms;			
	3. detection of <i>Borrelia burgdorferi</i> -specific antibodies or intrathecal synthesis of <i>B. burgdorferi</i> -specific antibodies, or cultivation of <i>B. burgdorferi</i> from the CSF, or a combination of these.			
	All children had to be antibiotic treatment-naïve.			
	See Table 2 and Table 3 for diagnostic criteria and additional baseline characteristics.			
Interventions	14 days of treatment w	ith either:		
	 intravenous penicillin G (400,000 to 500,000 international units/kg body weight) per day given 3 times daily (N = 11); or 			
	 intravenous ceftriaxone 75 mg/kg to 93 mg/kg body weight per day, given once daily (N = 12). 			
Outcomes	1. Development of specific B. burgdorferi serum ELISA IgG antibody titers			
	2. Clinical outcome (i.e. duration of disease from the beginning of therapy until complete clinical recovery and clinical follow-up for at least 6 months)			
	<i>B. burgdorferi</i> -specific IgG in serum was the primary outcome and was measured by ELISA prior to ran- domization, at the end of treatment, and 3, 6, and 12 months after the end of treatment.			
Funding	Not disclosed	Not disclosed		
Conflicts of interest	Not disclosed			
Notes				
Risk of bias				
Bias	Authors' judgement	Support for judgement		
Random sequence genera- tion (selection bias)	Unclear risk	Children were randomly allocated to treatment groups, but the report does not describe the method of randomization.		
Allocation concealment (selection bias)	Unclear risk	Not mentioned, not clear if it was done		
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Neither participants nor investigators were blinded to treatment allocation.		

Mullegger 1991 (Continued)

Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Not documented. We assume that outcome assessors were not blinded to treatment group.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	All children in both groups recovered completely. Some children did not par- ticipate in the final control visit (12 months), which could have resulted in missed detection of late relapse in these children.
Selective reporting (re- porting bias)	Unclear risk	No evidence that all outcomes were prespecified; the pre-study protocol was not available for review, and the study was not registered prior to initiation
Other bias	Low risk	No other bias identified

Oksi 2007

Methods	Prospective, randomized, double-blind, parallel-group multicenter study with placebo			
Participants	152 consecutive adults (145 evaluable) from 3 tertiary hospitals in Finland who had just completed treatment for Lyme disease with ceftriaxone. 62 participants had definite LNB. Criteria for a definite LNB diagnosis were: a classical manifestation (e.g. facial paresis, meningitis, or meningoradiculitis, along with exclusion of other causes) and inflammatory changes in the CSF or <i>Borrelia burgdorferi</i> -specific intrathecal antibodies, or both. Criteria for possible LNB were less common manifestations of LNB and presence of serum <i>B. burgdorferi</i> -specific antibodies. Diagnosis of definite or probable LNB required exclusion of other causes. Among all Lyme disease participants, 52/73 (71.2%) amoxicillin-treated group and 54/72 (75%) placebo group had a definite diagnosis.			
	See Table 2 and Table 3 for diagnostic criteria and additional baseline characteristics.			
Interventions	All participants received:			
	 3 weeks' treatment with intravenous ceftriaxone 2 g per day, followed by oral amoxicillin 1 g twice per day for 100 days (N = 73); or 			
	 3 weeks' treatment with intravenous ceftriaxone 2 g per day, followed by placebo twice per day for 100 days (N = 72). 			
Outcomes	1-year follow-up, with outcome measurement at the end of ceftriaxone treatment and 1, 3, 6, and 12 months later. Outcome measured by 0 to 100 VAS, where 50 = baseline before intravenous treatment, 0 = symptom free, 100 = "definitely poor outcome".			
Funding	Bristol–Myers Squibb provided amoxicillin tablets, and Roche covered part of the costs of the study; any other funding was not disclosed.			
Conflicts of interest	Not disclosed			
Notes	Study conducted 1998 to 2003.			
	Location: Finland			
	The study may have been underpowered to permit a definite conclusion about the lack of efficacy of the adjunctive treatment, as a total of 200 participants would have been needed to show a 10% difference with an 80% power to detect a significant (P < 0.05, 2-sided) difference. Dr. Oksi provided us with unpublished data separating the LNB from the other Lyme disease participants.			
Risk of bias				
Bias	Authors' judgement Support for judgement			

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Oksi 2007 (Continued)

Random sequence genera- tion (selection bias)	Unclear risk	Participants were randomly allocated to treatment group in a pharmacy, but the report does not describe the exact method. Baseline comparison of treat- ment groups was not adequately described.
Allocation concealment (selection bias)	Low risk	The enrolled participants received labeled containers marked with a code. The investigators had no access to the codes before the end of the study.
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	No blinding for the initial antibiotic treatment, but there was blinding for the second. Although the investigators had no access to the codes until the end of the study, the possibility that side effects unblinded participants was not addressed.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	No blinding for the initial antibiotic treatment, only for the second. Although the investigators had no access to the study codes until the end of the study, the possibility that side effects unblinded investigators was not addressed.
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	7 participants were withdrawn and not included in analysis (5 for discontinu- ing the study drug and 2 who received an alternate diagnosis). No intention-to- treat analysis was performed.
Selective reporting (re- porting bias)	Unclear risk	No evidence that all outcomes were prespecified; the pre-study protocol was not available for review, and the study was not registered prior to initiation. Reporting for the LNB group was complete but never published.
Other bias	Low risk	The study provided separate results for cases of definite and probable LNB both at randomization and by efficacy outcomes.

Pfister 1989

Methods	Prospective, randomized, parallel-group, open-label study with an active comparator, no placebo
Participants	Participants had acute painful LNB radiculitis (n = 18) or LNB meningitis (n = 3). The diagnostic criteria included the following ¹ :
	 clinical signs of acute LNB radiculitis (Bannwarth's syndrome) with severe radicular pain and lympho- cytic pleocytosis in the CSF, elevated <i>Borrelia burgdorferi</i>-specific antibody titers, and/or a history of arthropod bite or erythema migrans (n = 18); and
	 LNB meningitis with a history of a tick bite or erythema migrans and elevated <i>B. burgdorferi</i>-specific antibody titers (n = 3).
	See Table 2 and Table 3 for diagnostic criteria and additional baseline characteristics.
Interventions	10-day treatment with either:
	 intravenous penicillin G 20 million units per day (n = 10); or intravenous cefotaxime, 2 g, 3 times per day (n = 11).
Outcomes	Neurologic examination was performed daily. Improvement or resolution in the neurological history and physical exam (cranial nerve palsies, pareses of extremities and abdominal muscles, headache, and sensory disturbances) was recorded on day 10 (early outcome) and on average 7.7 months later (longer-term outcome).
	The severity of radicular pain was scored daily with a 0-to-10 rating system. For evaluation, the medi- ans of the corresponding maximum daily pain scores in the penicillin group (n = 7) were compared with those in the cefotaxime group (n = 8). Trialists recorded the daily dose of analgesics during the 10-day treatment period and measured the total amount of analgesics taken during the 10-day treatment pe- riod. Investigators performed lumbar puncture prior to (n = 21) and on the 8th to 10th day of treatment



Pfister 1989 (Continued)

(n = 17) and quantified CSF-WBC, CSF protein, and intrathecal IgG synthesis. They visualized oligoclonal bands and cultured CSF in BSK media. In addition, the investigators measured *B. burgdorferi*-specific antibody concentration in serum at randomization and follow-up.

Funding	Not disclosed				
Conflicts of interest	Not disclosed				
Notes	¹ At the time of the onset of therapy, radicular pain and headache had already subsided in 3 partici- pants with radiculitis and in 1 participant with meningitis, respectively.				
	6 of 21 participants were seronegative: 6 participants had normal (n = 5) or marginal (n = 1) <i>B. burgdor-feri</i> -specific antibody titers in the serum and normal <i>B. burgdorferi</i> -specific CSF antibody titers. 4 of these 6 participants had a history of erythema migrans, which was still present in 2 participants at the time of hospital admission. The other 2 seronegative participants (1 participant from each treatment group) had no history of erythema migrans but reported multiple "insect bites" and bites by horseflies within a few weeks prior to the onset of the neurologic disease.				
Risk of bias					

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Participants were randomly allocated to treatment groups, but the report does not describe the method of randomization.
Allocation concealment (selection bias)	Unclear risk	Not stated
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Neither participants nor investigators were blinded to treatment allocation.
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Outcome assessors were not blinded.
Incomplete outcome data (attrition bias) All outcomes	Low risk	The majority of participants returned for longer-term follow-up.
Selective reporting (re- porting bias)	Unclear risk	No evidence that all outcomes were prespecified; the pre-study protocol was not available for review, and the study was not registered prior to initiation.
Other bias	Low risk	No other bias identified

Pfister 1991

Methods	Prospective, randomized, parallel-group, open-label study with an active comparator, no placebo
Participants	33 participants with predominantly acute LNB
	32 of the 33 participants had clinical LNB; 1 was asymptomatic. Trialists excluded 3 of the 33 participants because they were never symptomatic (N = 1) or because the symptoms had resolved prior to randomization (N = 2). 28 participants had typical Bannwarth's syndrome with intense radicular pain and lymphocytic pleocytosis in the CSF. These participants usually also had paresis of the extremities or cranial nerve palsies. 4 had lymphocytic meningitis with detectable <i>Borrelia burgdorferi</i> -specific an-



fister 1991 (Continued)	<i>burgdorferi</i> infection, b <i>feri</i> was isolated from 0	r CSF, or both. The remaining participant had no clinical symptoms due to <i>B.</i> but the <i>B. burgdorferi</i> -specific IgG titer in the serum was elevated and <i>B. burgdor</i> CSF. A history of an arthropod bite or typical erythema migrans within 3 months neurologic disease was found in 18 and 16 participants, respectively.
	See Table 2 and Table 3	³ for diagnostic criteria and additional baseline characteristics.
Interventions	10 days' treatment wit	h either:
		tone 2 g per day (n = 17); or ime, 2 g, 3 times per day (n = 16).
Outcomes	symptoms attributable	was the number of participants whose symptoms improved (improvement of to LNB including radicular pain, headache, cranial nerve palsies, pareses of ex- disturbances). Study authors also reported tolerability.
Funding	Not disclosed	
Conflicts of interest	Not disclosed	
Notes		
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Participants were randomly allocated to treatment groups, but the report does not describe the method of randomization.
Allocation concealment (selection bias)	Unclear risk	Not stated
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	Neither study participants nor study personnel were blinded to treatment group.
Blinding of outcome as- sessment (detection bias) All outcomes	High risk	Outcome assessors were not blinded.
Incomplete outcome data (attrition bias) All outcomes	Low risk	The majority of participants returned for follow-up.
Selective reporting (re- porting bias)	Unclear risk	No evidence that all outcomes were prespecified; the pre-study protocol was not available for review, and the study was not registered prior to initiation.
		No other bias identified

BSK media: Barbour-Stoenner-Kelly media CSF: cerebrospinal fluid ELISA: enzyme-linked immunosorbent assay Ig: immunoglobulin LNB: Lyme neuroborreliosis VAS: visual analogue scale WBC: white blood cells



Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion						
Dattwyler 1988	This study focused only on late LB (all types, not specifically neurological).						
Dattwyler 2005a	This open randomized clinical trial compared 14 and 28 days of ceftriaxone treatment in people with LB including LNB. As LNB participants were not separately documented, we excluded the trial from the review.						
Hassler 1990	Excluded for 2 main reasons: clear objective evidence of neurological disease was lacking, and par- ticipants were a mixed population with manifestations of joint involvement with or without periph- eral nerve involvement, which was poorly documented. We contacted the author, who was unable to retrieve study data from an outdated digital storing system for further analysis.						
Massarotti 1992	Participants had erythema migrans, not untreated LNB. Excluded as the objective of the review was not to determine the efficacy of antibiotic treatment of erythema migrans to prevent the development of LNB.						
Oksi 1998	 The manuscript did not allow for the separation of LNB from other cases of disseminated LB, al- though the trial authors claim that most participants had neurological symptoms. Antibiotic pretreatment in a subgroup (the trial author mentions 6 cases in each arm) prevents 						
	determination of efficacy of the initial intervention.						
	3. The lack of intrathecal production of Ab in the majority of participants is surprising for European LNB (the first author himself found intrathecal antibody production in about 50% of LNB participants in a later study). Pleocytosis of the CSF is not mentioned. Being a mixture of different manifestations of stage II and III Lyme disease, the LNB cases were not sufficiently characterized for judgement of antibiotic efficacy. Most participants had CSF PCR performed, but only 1 or 2 in each group were found to be positive, suggesting that many participants may not have had active LNB, but sequelae from previously treated disease. This is consistent with the observed low response rate to antibiotic therapy in both arms, as the participants were likely selected for failure to respond to earlier treatment and by persisting symptoms (fibromyalgia, "arthritis" not otherwise defined).						
	4. Although symptoms are categorized, it is unclear which symptoms were presenting at time of the study or at onset (and before any antibiotic therapy).						
Pfister 1988	This was a randomized study of treatment of LNB with a non-antibiotic intervention (corticos- teroid).						

Ab: antibody

CSF: cerebrospinal fluid LB: Lyme borreliosis LNB: Lyme neuroborreliosis PCR: polymerase chain reaction

Characteristics of ongoing studies [ordered by study ID]

NCT02553473					
Trial name or title	Six versus two weeks treatment with doxycycline in Lyme neuroborreliosis				
Methods	Multicenter, non-inferiority, randomized, penta-blind (participant, caregiver, investigator, out- comes assessor), placebo-controlled study				
Participants	250 adults (18 years and older) with neuroborreliosis (<i>Borrelia burgdorferi</i>) from Norwegian hospi- tals				



NCT02553473 (Continued)	Inclusion criteria: neurological symptoms suggestive of Lyme neuroborreliosis without other obvi- ous causes, one or both of a) cerebrospinal fluid pleocytosis (> 5 leukocytes/mm ³), b) intrathecal <i>B.</i> <i>burgdorferi</i> antibody production
Interventions	Doxycycline 200 mg once daily for 6 weeks versus doxycycline 200 mg once daily for 2 weeks + placebo for 4 weeks
Outcomes	Primary: composite clinical score at 6 months after the end of treatment Secondary: Fatigue Severity Scale (FSS), Patient Health Questionnaire (PHQ-15), 36-Item Short- Form Health Survey (SF-36), and blood and cerebrospinal fluid findings at inclusion and after 6 and 12 months
Starting date	Safety October 2015
Contact information	Sorlandet Hospital
Notes	Estimated completion 2020

DATA AND ANALYSES

Comparison 1. Oral amoxicillin versus placebo after previous treatment with ceftriaxone for disseminated Lyme disease

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Symptoms (patient-rated VAS, scale 0 to 100, higher worse) in all partici- pants (definite and possible Lyme dis- ease)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
1.1 3 months	1	145	Mean Difference (IV, Fixed, 95% CI)	4.20 [3.39, 5.01]
1.2 6 months	1	145	Mean Difference (IV, Fixed, 95% CI)	-0.5 [-1.38, 0.38]
1.3 12 months	1	145	Mean Difference (IV, Fixed, 95% CI)	0.60 [-0.21, 1.41]
2 Symptoms (investigator-rated VAS, scale 0 to 100 higher worse) in all par- ticipants (definite and possible Lyme disease)	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only
2.1 3 months	1	145	Mean Difference (IV, Fixed, 95% CI)	0.5 [-0.28, 1.28]
2.2 6 months	1	145	Mean Difference (IV, Fixed, 95% CI)	-2.40 [-3.18, -1.62]

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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2.3 12 months	1	145	Mean Difference (IV, Fixed, 95% CI)	-0.40 [-1.13, 0.33]
3 Improvement of symptoms (excel- lent or good on investigator VAS) (12 months) in participants with definite Lyme disease	1	107	Risk Ratio (M-H, Fixed, 95% CI)	1.06 [0.93, 1.21]
4 Adverse events (12 months) in all participants (definite and possible Ly-me disease)	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
4.1 Serious adverse events	1	145	Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]
4.2 Diarrhea	1	145	Risk Ratio (M-H, Fixed, 95% CI)	3.70 [1.29, 10.61]

Analysis 1.1. Comparison 1 Oral amoxicillin versus placebo after previous treatment with ceftriaxone for disseminated Lyme disease, Outcome 1 Symptoms (patient-rated VAS, scale 0 to 100, higher worse) in all participants (definite and possible Lyme disease).

Study or subgroup	Am	oxicillin	P	acebo		Mean Difference	Weight	Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		Fixed, 95% CI		Fixed, 95% CI
1.1.1 3 months								
Oksi 2007	73	26.8 (2.4)	72	22.6 (2.6)				4.2[3.39,5.01]
Subtotal ***	73		72				100%	4.2[3.39,5.01]
Heterogeneity: Not applicable								
Test for overall effect: Z=10.1(P<0.000	01)							
1.1.2 6 months								
Oksi 2007	73	22.3 (2.5)	72	22.8 (2.9)		_ _	100%	-0.5[-1.38,0.38]
Subtotal ***	73		72			-	100%	-0.5[-1.38,0.38]
Heterogeneity: Not applicable								
Test for overall effect: Z=1.11(P=0.27)								
1.1.3 12 months								
Oksi 2007	73	18.8 (2.4)	72	18.2 (2.6)		+	100%	0.6[-0.21,1.41]
Subtotal ***	73		72			-	100%	0.6[-0.21,1.41]
Heterogeneity: Not applicable								
Test for overall effect: Z=1.44(P=0.15)								
			Favou	rs amoxicillin	-4	-2 0 2	4 Favours pla	cebo



Analysis 1.2. Comparison 1 Oral amoxicillin versus placebo after previous treatment with ceftriaxone for disseminated Lyme disease, Outcome 2 Symptoms (investigator-rated VAS, scale 0 to 100 higher worse) in all participants (definite and possible Lyme disease).

Study or subgroup	Am	oxicillin	Р	acebo	Mean Difference	Weight	Mean Difference
	N	Mean(SD)	Ν	Mean(SD)	Fixed, 95% CI		Fixed, 95% Cl
1.2.1 3 months							
Oksi 2007	73	22.3 (2.2)	72	21.8 (2.6)		100%	0.5[-0.28,1.28]
Subtotal ***	73		72		-	100%	0.5[-0.28,1.28]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.25(P=0.21)							
1.2.2 6 months							
Oksi 2007	73	18.1 (2.2)	72	20.5 (2.6)	_ 	100%	-2.4[-3.18,-1.62]
Subtotal ***	73		72		•	100%	-2.4[-3.18,-1.62]
Heterogeneity: Not applicable							
Test for overall effect: Z=6(P<0.0001)							
1.2.3 12 months							
Oksi 2007	73	14.3 (2.2)	72	14.7 (2.3)		100%	-0.4[-1.13,0.33]
Subtotal ***	73		72		-	100%	-0.4[-1.13,0.33]
Heterogeneity: Not applicable							
Test for overall effect: Z=1.07(P=0.28)							
Test for subgroup differences: Chi ² =2	7.63, df=	1 (P<0.0001), I ² =	92.76%				
			Favou	rs amoxicillin	-2 -1 0 1 2	Favours pla	cebo

Analysis 1.3. Comparison 1 Oral amoxicillin versus placebo after previous treatment with ceftriaxone for disseminated Lyme disease, Outcome 3 Improvement of symptoms (excellent or good on investigator VAS) (12 months) in participants with definite Lyme disease.

Study or subgroup	Amoxicillin	Placebo	Risk Ratio	Weight	Risk Ratio
	n/N		M-H, Fixed, 95% Cl		M-H, Fixed, 95% CI
Oksi 2007	49/53	47/54		100%	1.06[0.93,1.21]
Total (95% CI)	53	54	-	100%	1.06[0.93,1.21]
Total events: 49 (Amoxicillin),	47 (Placebo)				
Heterogeneity: Not applicable	2				
Test for overall effect: Z=0.92(P=0.36)				
		Favours placebo	1	Favours amoxicillin	

Analysis 1.4. Comparison 1 Oral amoxicillin versus placebo after previous treatment with ceftriaxone for disseminated Lyme disease, Outcome 4 Adverse events (12 months) in all participants (definite and possible Lyme disease).

Study or subgroup	Amoxicillin	Placebo		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95					M-H, Fixed, 95% Cl
1.4.1 Serious adverse events									
Oksi 2007	0/73	0/72							Not estimable
Subtotal (95% CI)	73	72							Not estimable
	Fa	vours amoxicillin	0.02	0.1	1	10	50	Favours placebo	

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Study or subgroup	Amoxicillin	xicillin Placebo			Risk Ratio			Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI						M-H, Fixed, 95% CI	
Total events: 0 (Amoxicillin), 0 (Placeb	o)									
Heterogeneity: Not applicable										
Test for overall effect: Not applicable										
1.4.2 Diarrhea										
Oksi 2007	15/73	4/72				.		100%	3.7[1.29,10.61]	
Subtotal (95% CI)	73	72						100%	3.7[1.29,10.61]	
Total events: 15 (Amoxicillin), 4 (Place	bo)									
Heterogeneity: Not applicable										
Test for overall effect: Z=2.43(P=0.01)							1			
	Fa	vours amoxicillin	0.02	0.1	1	10	50	Favours placebo		

Comparison 2. Oral doxycycline versus intravenous ceftriaxone for Lyme neuroborreliosis (acute and chronic)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Mean reduction in clinical score (4 months)	1	102	Mean Difference (IV, Fixed, 95% CI)	0.10 [-1.20, 1.40]
2 Resolution of symptoms	1	102	Risk Ratio (M-H, Fixed, 95% CI)	1.44 [0.89, 2.35]
3 All adverse events	1	113	Risk Ratio (M-H, Fixed, 95% CI)	0.79 [0.51, 1.23]
4 Adverse events leading to dis- continuation	1	118	Risk Ratio (M-H, Fixed, 95% CI)	0.14 [0.01, 2.71]
5 Serious adverse events	1	113	Risk Ratio (M-H, Fixed, 95% CI)	0.33 [0.04, 3.05]

Analysis 2.1. Comparison 2 Oral doxycycline versus intravenous ceftriaxone for Lyme neuroborreliosis (acute and chronic), Outcome 1 Mean reduction in clinical score (4 months).

Study or subgroup	Dox	ycycline	Ceftriaxone			Mean Difference				Weight M	lean Difference
	Ν	Mean(SD)	Ν	Mean(SD)		Fix	ed, 95%	CI			Fixed, 95% CI
Ljostad 2008	54	4.5 (3.3)	48	4.4 (3.4)						100%	0.1[-1.2,1.4]
Total ***	54		48							100%	0.1[-1.2,1.4]
Heterogeneity: Not applicable											
Test for overall effect: Z=0.15(P=0.88)											
			Favou	rs ceftriaxone	-2	-1	0	1	2	Favours doxycyc	line

Analysis 2.2. Comparison 2 Oral doxycycline versus intravenous ceftriaxone for Lyme neuroborreliosis (acute and chronic), Outcome 2 Resolution of symptoms.

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Study or subgroup	Doxycycline	Ceftriaxone		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95% CI					M-H, Fixed, 95% CI
Ljostad 2008	26/54	16/48						100%	1.44[0.89,2.35]
Total (95% CI)	54	48						100%	1.44[0.89,2.35]
Total events: 26 (Doxycycline), 16	(Ceftriaxone)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.48(P=0.	.14)								
	Fa	vours ceftriaxone	0.2	0.5	1	2	5	Favours doxycycline	

Analysis 2.3. Comparison 2 Oral doxycycline versus intravenous ceftriaxone for Lyme neuroborreliosis (acute and chronic), Outcome 3 All adverse events.

Study or subgroup	Doxycycline	Ceftriaxone		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95% CI					M-H, Fixed, 95% Cl
Ljostad 2008	21/57	26/56		-				100%	0.79[0.51,1.23]
Total (95% CI)	57	56						100%	0.79[0.51,1.23]
Total events: 21 (Doxycycline), 26 (C	Ceftriaxone)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.03(P=0.3)	1								
	Fa	vours doxycycline	0.2	0.5	1	2	5	Favours ceftriaxone	

Analysis 2.4. Comparison 2 Oral doxycycline versus intravenous ceftriaxone for Lyme neuroborreliosis (acute and chronic), Outcome 4 Adverse events leading to discontinuation.

Study or subgroup	Doxycycline	Ceftriaxone		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		M-H, Fixed, 95% CI					M-H, Fixed, 95% CI
Ljostad 2008	0/59	3/59		+				100%	0.14[0.01,2.71]
Total (95% CI)	59	59						100%	0.14[0.01,2.71]
Total events: 0 (Doxycycline), 3 (Ceft	riaxone)								
Heterogeneity: Not applicable									
Test for overall effect: Z=1.3(P=0.19)				i					
	Fa	vours doxycycline	0.005	0.1	1	10	200	Favours ceftriaxone	

Analysis 2.5. Comparison 2 Oral doxycycline versus intravenous ceftriaxone for Lyme neuroborreliosis (acute and chronic), Outcome 5 Serious adverse events.

Study or subgroup	Doxycycline	Ceftriaxone	xone			0		Weight	Risk Ratio
	n/N	n/N		м-н,	Fixed, 9	5% CI			M-H, Fixed, 95% CI
Ljostad 2008	1/57	3/56						100%	0.33[0.04,3.05]
Total (95% CI)	57	56						100%	0.33[0.04,3.05]
	Fa	vours doxycycline	0.005	0.1	1	10	200	Favours ceftriaxone	

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Study or subgroup	Doxycycline	Ceftriaxone		R	isk Rati	io		Weight	Risk Ratio
	n/N	n/N		м-н,	ixed, 9	5% CI			M-H, Fixed, 95% Cl
Total events: 1 (Doxycycline),	3 (Ceftriaxone)								
Heterogeneity: Not applicable	2								
Test for overall effect: Z=0.98(P=0.33)								
	F	avours doxycycline	0.005	0.1	1	10	200	Favours ceftriaxone	

Comparison 3. Intravenous penicillin G versus oral doxycycline for Lyme neuroborreliosis (acute and chronic)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Improvement of symp- toms	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
1.1 3 months	1	53	Risk Ratio (M-H, Fixed, 95% CI)	1.0 [0.93, 1.08]
1.2 6 months	1	52	Risk Ratio (M-H, Fixed, 95% CI)	1.0 [0.93, 1.08]
1.3 12 months	1	51	Risk Ratio (M-H, Fixed, 95% CI)	1.0 [0.92, 1.08]
2 Resolution of symp- toms	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.1 3 months	1	53	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.64, 1.61]
2.2 6 months	1	52	Risk Ratio (M-H, Fixed, 95% CI)	1.67 [1.10, 2.54]
2.3 12 months	1	51	Risk Ratio (M-H, Fixed, 95% CI)	0.95 [0.77, 1.18]
3 Resolution of CSF pleo- cytosis at 1 year	1	29	Risk Ratio (M-H, Fixed, 95% CI)	0.93 [0.75, 1.15]
4 All adverse events	1	54	Risk Ratio (M-H, Fixed, 95% CI)	1.01 [0.25, 4.08]

Analysis 3.1. Comparison 3 Intravenous penicillin G versus oral doxycycline for Lyme neuroborreliosis (acute and chronic), Outcome 1 Improvement of symptoms.

Study or subgroup	Penicillin	Doxycycline	Risk Ratio	Weight	Risk Ratio M-H, Fixed, 95% Cl	
	n/N	n/N	M-H, Fixed, 95% CI			
3.1.1 3 months						
Karlsson 1994	22/22	31/31		100%	1[0.93,1.08]	
Subtotal (95% CI)	22	31		100%	1[0.93,1.08]	
Total events: 22 (Penicillin), 31 (D	oxycycline)					
Heterogeneity: Not applicable						
Test for overall effect: Not applica	able					
3.1.2 6 months						
Karlsson 1994	21/21	31/31		100%	1[0.93,1.08]	
Subtotal (95% CI)	21	31		100%	1[0.93,1.08]	
	Fa	vours doxycycline	1	Favours penicillin		

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Study or subgroup	Penicillin Doxycycline		Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% Cl	
Total events: 21 (Penicillin), 31 (Doxycyc	line)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
3.1.3 12 months						
Karlsson 1994	21/21	30/30		100%	1[0.92,1.08]	
Subtotal (95% CI)	21	30		100%	1[0.92,1.08]	
Total events: 21 (Penicillin), 30 (Doxycyc	line)					
Heterogeneity: Not applicable						
Test for overall effect: Not applicable						
Test for subgroup differences: Not applie	cable					
	Fa	avours doxycycline	1	Favours penicillin		

Favours doxycycline

Favours penicillin

Analysis 3.2. Comparison 3 Intravenous penicillin G versus oral doxycycline for Lyme neuroborreliosis (acute and chronic), Outcome 2 Resolution of symptoms.

Study or subgroup	Penicillin	Doxycycline	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% Cl
3.2.1 3 months					
Karlsson 1994	13/22	18/31		100%	1.02[0.64,1.61]
Subtotal (95% CI)	22	31		100%	1.02[0.64,1.61]
Total events: 13 (Penicillin), 18 (Dox	(ycycline)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.07(P=0.94	4)				
3.2.2 6 months					
Karlsson 1994	17/21	15/31		100%	1.67[1.1,2.54]
Subtotal (95% CI)	21	31		100%	1.67[1.1,2.54]
Total events: 17 (Penicillin), 15 (Dox	(ycycline)				- / -
Heterogeneity: Not applicable					
Test for overall effect: Z=2.41(P=0.0)	2)				
3.2.3 12 months					
Karlsson 1994	18/21	27/30	- <mark></mark>	100%	0.95[0.77,1.18]
Subtotal (95% CI)	21	30		100%	0.95[0.77,1.18]
Total events: 18 (Penicillin), 27 (Dox	(ycycline)				
Heterogeneity: Tau ² =0; Chi ² =0, df=0	0(P<0.0001); I ² =100%				
Test for overall effect: Z=0.45(P=0.6	5)				
Test for subgroup differences: Chi ² =	=5.58, df=1 (P=0.06), I ²	=64.16%			
	Fa	vours doxycycline	0.5 0.7 1 1.5 2	Favours penicillin	

Analysis 3.3. Comparison 3 Intravenous penicillin G versus oral doxycycline for Lyme neuroborreliosis (acute and chronic), Outcome 3 Resolution of CSF pleocytosis at 1 year.

Study or subgroup	Penicillin	Doxycycline	Risk Ratio	Weight	Risk Ratio
	n/N n/N		M-H, Fixed, 95% Cl		M-H, Fixed, 95% CI
Karlsson 1994	18/20	9/9		100%	0.93[0.75,1.15]
	Fa	vours doxycycline	0.5 0.7 1 1.5 2	Favours penicillin	

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Study or subgroup	Penicillin Doxycycline		Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% Cl		M-H, Fixed, 95% CI
Total (95% CI)	20	9	•	100%	0.93[0.75,1.15]
Total events: 18 (Penicillin), 9 (Doxyo	cycline)				
Heterogeneity: Not applicable					
Test for overall effect: Z=0.7(P=0.49)					
	F	avours doxycycline	0.5 0.7 1 1.5 2	Favours penicillin	

Analysis 3.4. Comparison 3 Intravenous penicillin G versus oral doxycycline for Lyme neuroborreliosis (acute and chronic), Outcome 4 All adverse events.

Study or subgroup	Penicillin Doxycycline		Risk Ratio					Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI						M-H, Fixed, 95% Cl
Karlsson 1994	3/23	4/31						100%	1.01[0.25,4.08]
Total (95% CI)	23	31						100%	1.01[0.25,4.08]
Total events: 3 (Penicillin), 4 (Doxycycl	line)								
Heterogeneity: Not applicable									
Test for overall effect: Z=0.02(P=0.99)			1						
		Favours penicillin	0.2	0.5	1	2	5	Favours doxycycline	

Comparison 4. Intravenous doxycycline versus intravenous penicillin G for Lyme neuroborreliosis (acute and chronic)

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Improvement of symptoms ("partial remission")	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
1.16 months	1	75	Risk Ratio (M-H, Fixed, 95% CI)	1.10 [0.95, 1.28]
1.2 12 months	1	75	Risk Ratio (M-H, Fixed, 95% CI)	0.98 [0.80, 1.21]
2 Resolution of symptoms ("full remission")	1		Risk Ratio (M-H, Fixed, 95% CI)	Subtotals only
2.1 6 months	1	75	Risk Ratio (M-H, Fixed, 95% CI)	1.42 [0.83, 2.42]
2.2 12 months	1	75	Risk Ratio (M-H, Fixed, 95% CI)	0.96 [0.70, 1.31]
3 Serious adverse events	1	75	Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]

Analysis 4.1. Comparison 4 Intravenous doxycycline versus intravenous penicillin G for Lyme neuroborreliosis (acute and chronic), Outcome 1 Improvement of symptoms ("partial remission").

Study or subgroup	Doxycycline	Penicillin	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI	
4.1.1 6 months						
Kohlhepp 1989	37/39	31/36		100%	1.1[0.95,1.28]	
Subtotal (95% CI)	39	36	►	100%	1.1[0.95,1.28]	
Total events: 37 (Doxycycline), 31 (Pe	enicillin)					
Heterogeneity: Not applicable						
Test for overall effect: Z=1.26(P=0.21))					
4.1.2 12 months						
Kohlhepp 1989	32/39	30/36		100%	0.98[0.8,1.21]	
Subtotal (95% CI)	39	36	—	100%	0.98[0.8,1.21]	
Total events: 32 (Doxycycline), 30 (Pe	enicillin)					
Heterogeneity: Not applicable						
Test for overall effect: Z=0.15(P=0.88))					
Test for subgroup differences: Chi ² =0	.74, df=1 (P=0.39), I ² =	=0%				
		Favours penicillin	0.5 0.7 1 1.5 2	Favours doxycycline		

Analysis 4.2. Comparison 4 Intravenous doxycycline versus intravenous penicillin G for Lyme neuroborreliosis (acute and chronic), Outcome 2 Resolution of symptoms ("full remission").

Study or subgroup	Doxycycline	Penicillin	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
4.2.1 6 months					
Kohlhepp 1989	20/39	13/36		100%	1.42[0.83,2.42]
Subtotal (95% CI)	39	36		100%	1.42[0.83,2.42]
Total events: 20 (Doxycycline), 1	3 (Penicillin)				
Heterogeneity: Not applicable					
Test for overall effect: Z=1.29(P=	0.2)				
4.2.2 12 months					
Kohlhepp 1989	26/39	25/36	- <mark></mark> -	100%	0.96[0.7,1.31]
Subtotal (95% CI)	39	36		100%	0.96[0.7,1.31]
Total events: 26 (Doxycycline), 2	5 (Penicillin)				
Heterogeneity: Tau ² =0; Chi ² =0, d	lf=0(P<0.0001); I ² =100%				
Test for overall effect: Z=0.26(P=	0.8)				
Test for subgroup differences: Ch	hi²=1.56, df=1 (P=0.21), I²=	-35.73%			
		Favours penicillin	0.2 0.5 1 2 5	Favours doxycycline	

Analysis 4.3. Comparison 4 Intravenous doxycycline versus intravenous penicillin G for Lyme neuroborreliosis (acute and chronic), Outcome 3 Serious adverse events.

Study or subgroup	Doxycycline	Penicillin			Risk Ratio)	Weight		Risk Ratio
	n/N n/N		M-H, Fixed, 95% CI					M-H, Fixed, 95% Cl	
Kohlhepp 1989	0/39	0/39 0/36				Not estimable			
	Fav	ours doxycycline	0.01	0.1	1	10	100	Favours penicillin	



Study or subgroup	Doxycycline	Penicillin			Risk Ratio)		Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% Cl		% CI			M-H, Fixed, 95% CI	
Total (95% CI)	39	36							Not estimable
Total events: 0 (Doxycycline), 0 (Pe	nicillin)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicab	ole								
	Fa	vours doxycycline	0.01	0.1	1	10	100	Favours penicillin	

Comparison 5. Intravenous cefotaxime versus intravenous penicillin G for acute Lyme neuroborreliosis

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Resolution of symptoms (mean 7.7 months' follow-up)	1	21	Risk Ratio (M-H, Fixed, 95% CI)	1.02 [0.67, 1.55]
2 Resolution of CSF pleocytosis	1	21	Risk Ratio (M-H, Fixed, 95% CI)	0.92 [0.71, 1.18]
3 All adverse events (at 2 weeks)	1	21	Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]

Analysis 5.1. Comparison 5 Intravenous cefotaxime versus intravenous penicillin G for acute Lyme neuroborreliosis, Outcome 1 Resolution of symptoms (mean 7.7 months' follow-up).

Study or subgroup	Cefotaxime	Penicillin G		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		М-Н,	Fixed, 95	5% CI			M-H, Fixed, 95% CI
Pfister 1989	9/11	8/10		-	-	-		100%	1.02[0.67,1.55]
Total (95% CI)	11	10		-	\blacklozenge			100%	1.02[0.67,1.55]
Total events: 9 (Cefotaxime), 8 (Penio	cillin G)								
Heterogeneity: Not applicable									
Test for overall effect: Z=0.11(P=0.92))						1		
	Fa	avours penicillin G	0.2	0.5	1	2	5	Favours cefotaxime	

Analysis 5.2. Comparison 5 Intravenous cefotaxime versus intravenous penicillin G for acute Lyme neuroborreliosis, Outcome 2 Resolution of CSF pleocytosis.

Study or subgroup	Cefotaxime	Penicillin G	Risk Ratio	Weight	Risk Ratio
	n/N	n/N	M-H, Fixed, 95% CI		M-H, Fixed, 95% Cl
Pfister 1989	10/11	10/10		100%	0.92[0.71,1.18]
Total (95% CI)	11	10	-	100%	0.92[0.71,1.18]
Total events: 10 (Cefotaxime)	, 10 (Penicillin G)				
Heterogeneity: Tau ² =0; Chi ² =0	0, df=0(P<0.0001); l²=100%				
Test for overall effect: Z=0.68((P=0.49)				
	Fa	vours penicillin G	0.5 0.7 1 1.5 2	Favours cefotaxime	



Analysis 5.3. Comparison 5 Intravenous cefotaxime versus intravenous penicillin G for acute Lyme neuroborreliosis, Outcome 3 All adverse events (at 2 weeks).

Study or subgroup	Cefotaxime	Penicillin G	Risk Ratio	Weight	Risk Ratio	
	n/N	n/N	M-H, Fixed, 95% Cl		M-H, Fixed, 95% Cl	
Pfister 1989	0/11	0/10			Not estimable	
Total (95% CI)	11	10			Not estimable	
Total events: 0 (Cefotaxime),	0 (Penicillin G)					
Heterogeneity: Not applicable	e					
Test for overall effect: Not app	plicable					
	Fa	vours cefotaxime	1	Favours penicillin G		

Comparison 6. Intravenous ceftriaxone versus intravenous cefotaxime for acute Lyme neuroborreliosis

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Resolution of symptoms (mean 8.1 months' follow-up)	1	27	Risk Ratio (M-H, Fixed, 95% CI)	1.11 [0.63, 1.97]
2 Resolution of CSF pleocytosis	1	27	Risk Ratio (M-H, Fixed, 95% Cl)	1.14 [0.90, 1.44]
3 All adverse events	1	30	Risk Ratio (M-H, Fixed, 95% Cl)	0.38 [0.04, 3.26]

Analysis 6.1. Comparison 6 Intravenous ceftriaxone versus intravenous cefotaxime for acute Lyme neuroborreliosis, Outcome 1 Resolution of symptoms (mean 8.1 months' follow-up).

Study or subgroup	Ceftriaxone	Cefotaxime		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		M-H	H, Fixed, 95%	CI			M-H, Fixed, 95% Cl
Pfister 1991	8/12	9/15			-			100%	1.11[0.63,1.97]
Total (95% CI)	12	15			•			100%	1.11[0.63,1.97]
Total events: 8 (Ceftriaxone), 9 (Cefe	otaxime)								
Heterogeneity: Not applicable									
Test for overall effect: Z=0.36(P=0.72	2)			i		i	i		
	Fa	avours cefotaxime	0.01	0.1	1	10	100	Favours ceftriaxone	

Analysis 6.2. Comparison 6 Intravenous ceftriaxone versus intravenous cefotaxime for acute Lyme neuroborreliosis, Outcome 2 Resolution of CSF pleocytosis.

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Study or subgroup	Ceftriaxone	Cefotaxime		Risk Ratio		Weight	Risk Ratio
	n/N	n/N	M-I	H, Fixed, 95% C	I		M-H, Fixed, 95% Cl
Pfister 1991	12/12	13/15				100%	1.14[0.9,1.44]
Total (95% CI)	12	15		•		100%	1.14[0.9,1.44]
Total events: 12 (Ceftriaxone), 13 (C	Cefotaxime)						
Heterogeneity: Not applicable							
Test for overall effect: Z=1.08(P=0.2	8)						
	Fa	vours cefotaxime	0.1 0.2 0	0.5 1 2	5 10	Favours ceftriaxone	

Analysis 6.3. Comparison 6 Intravenous ceftriaxone versus intravenous cefotaxime for acute Lyme neuroborreliosis, Outcome 3 All adverse events.

Study or subgroup	Ceftriaxone	Cefotaxime		Risk Ratio			Weight	Risk Ratio	
	n/N	n/N		M-H, F	ixed, 9	95% CI			M-H, Fixed, 95% CI
Pfister 1991	1/14	3/16				-		100%	0.38[0.04,3.26]
Total (95% CI)	14	16				-		100%	0.38[0.04,3.26]
Total events: 1 (Ceftriaxone), 3	8 (Cefotaxime)								
Heterogeneity: Tau ² =0; Chi ² =0	, df=0(P<0.0001); l ² =100%								
Test for overall effect: Z=0.88(F	P=0.38)			1					
	Fa	avours ceftriaxone	0.002	0.1	1	10	500	Favours cefotaxime	

Antibiotics for the neurological complications of Lyme disease (Review) Copyright © 2016 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd. ADDITIONAL TABLES

Table 1. Overview of studies included in the review

Study	Popula- tion limited to LNB	Length of follow-up	Interventions	Antibiotic naïve	Clinical remission measure- ment	Time to remission	Pa- tient-re- ported outcomes	CSF remission measured	Serology response mea- sured
Kohlhepp 1989	Yes	Up to 3 years	Penicillin Doxycycline	Unknown	Complete/partial/no	No	No	Yes	No
Pfister 1989	Yes	Average of 7 months	Penicillin Cefotaxime	Unknown	Yes/no	No	Yes: VAS, 0 to 10	Yes	No
Mullegger 1991	Yes (chil- dren only)	>6 months, <12 months	Penicillin Ceftriaxone	Yes	Yes/no	Yes; from treatment onset to complete remission	No	No	Yes
Pfister 1991	Yes	Average of 7.7 months	Ceftriaxone Cefotaxime	Unknown	Yes/no	No	No	Yes	No
Karlsson 1994	Yes	12 months	Penicillin Doxycycline	Past 4 weeks	Yes/no; by specific sign/symp- tom	No	Yes: Lik- ert-like scale, 0 to 3	Yes, at 2 weeks and 12 months	Yes
Oksi 2007	No (but large sub- group of LNB en- rolled)	12 months	Ceftriaxone fol- lowed by amoxicillin or placebo	Past 1 month	Physician VAS 0 to 100; scored as excellent/good, poor/none, controversial	No	Yes: VAS	Lumbar puncture and measurement of CSF antibody levels and PCR for <i>Borrelia burgdorferi</i> was repeated in se- lected cases during or after treatment.	Yes: scored as strong decline, mild, none
Ljostad 2008	Yes	Up to 4 months	Ceftriaxone Doxycycline	Past 14 days	No/mild/more than mild; also change in baseline deficits in past 3 months us-	No	Yes: 6 items, each scored 0 to 2	Yes	No

Table 1. Overview of studies included in the review (Continued)

ing own composite clinical score

CSF: cerebrospinal fluid LNB: Lyme neuroborreliosis PCR: polymerase chain reaction VAS: visual analogue scale

Table 2. Clinical and laboratory criteria for diagnosis of Lyme neuroborreliosis

Criteria	Kohlhepp 1989	Pfister 1989	Mullegger 1991	Pfister 1991	Karlsson 1994	Oksi 2007	Ljostad 2008	
	(N = 75)	(N = 21)	(N = 30) (N = 23)		(N = 54)	(N = 145; 72% to 75% definite, 25% to 27% possible ^a)	(N = 102)	
Clinical	Radicular pain, meningitic symp- toms, cranial neuritis, senso- ry and/or mo- tor radiculitis, arthritis, cardi- tis, myelitis or pe- ripheral neuri- tis, tick bite and/ or erythema mi- grans	Radicular pain (15/21), headache (2/21), facial palsy (8/21), unilateral VI palsy (1/21), lower limb muscle weak- ness (9/21), sensory disturbance (12/21)	Presence of neurologi- cal signs and symptoms indicative of LNB	Radiculopa- thy (motor or sensory, or both), cranial neuropathy (facial palsy, ocular motor)	Headache (71% to 74%), subjective stiff neck (65%), paresis (55% to 57%) including fa- cial palsy in 35% to 43%	Lymphocytic meningitis without radi- culitis in 18 (all definite), meningo- radiculitis (16 definite) or radiculi- tis (11 definite), paresis in 5, en- cephalomyelitis in 4, encephalopathy in 6, facial paresis in 21, sudden deaf- ness in 6, tinnitus in 8, other cranial nerve involvement in 13, peripheral neuritis in 6, and other peripheral ner- vous system manifestations (9 periph- eral mononeuropathy or polyneuropa- thy, 15 paresthesia, 39 with headache without meningitis, 29 with dizziness or vertigo, and 11 with memory im- pairment)	25% to 33% Bannwarth's syndrome, 19% to 22% facial palsy, 24% to 38% radiculopa- thy, various others (other cranial neu- ropathies, ataxia, myelopathy, limb pare- sis, paresthe- sias, cognitive deficits)	
Laboratory	<i>B. burgdorferi</i> - specific antibody titer in serum, <i>B.</i> <i>burgdorferi</i> -spe- cific antibody titer in CSF, lym- phocytic pleo- cytosis, elevat- ed CSF protein (>50 mg/dL), el- evated CSF IgM-,	Elevated <i>B. burgdor- feri</i> -specific IgG and IgM antibody titers in serum (1:64 to 1:512): found in 11, of whom 4 had both elevated IgG and IgM, 6 had only ele- vated IgG, and 1 had only elevated IgM.	1 or more of the following specific CSF laboratory parameters: elevated <i>B.</i> <i>burgdorferi</i> - specific IgG antibody titer, intrathecal- ly produced	Elevated <i>B.</i> <i>burgdorferi</i> - specific IgG and IgM anti- body titers in serum: found in 22 and 8, respectively. 13 had pos- itive <i>B.</i>	Elevated <i>B.</i> burgdorferi- specific IgM or IgG con- centration, or both in 83% to 90%; all had positive serology or <i>B. burgdor-</i> <i>feri</i> -specific	Only 3 of the 145 study participants were seronegative. Presence of inflammatory changes in the CSF or intrathecal antibodies against <i>B. burgdorferi</i> , or both sup- ported a diagnosis of definite LNB; 124/145 participants had lumbar puncture performed at diagnosis.	Intrathecal production of <i>B. burgdor- feri</i> -specific antibodies or <i>B. burgdor- feri</i> -specific antibodies in serum, or both were re-	

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Table 2. Clinical and laboratory criteria for diagnosis of Lyme neuroborreliosis (Continued)

ettineat ana taboratory	cificilia for alagitosis	or Lynne mearo		inucu)	
IgA-, and/or IgG- index, oligoclonal bands in CSF. On- ly group-level in-	4 were seroposi- tive in CSF but not in blood; 6 had neg- ative serology in	<i>B. burgdor- feri</i> -specific antibodies, and/or direct	<i>burgdorferi-</i> specific CSF/ serum anti- body index.	CSF antibod- ies, except for 1 partici- pant who had	quired for en- rollment.
formation is giv- en.	both serum and CSF (seronegative LNB), but 4 had EM and 2 a history of insect bites.	cultivation of <i>B. burgdor-</i> <i>feri</i> from the CSF in a mod- ified Bar- bour-Stoen- ner-Kelly medium		a positive CSF culture.	

^{*a*}LNB considered possible if clinical presentation was an uncommon manifestation, but serum antibodies against *Borrelia burgdorferi* were positive and other causes were excluded.

Abbreviations:

B. burgdorferi: Borrelia burgdorferi

CSF: cerebrospinal fluid

EM: erythema migrans

Ig: immunoglobulin

LNB: Lyme neuroborreliosis

Study	Kohlhep 1989 Peni- cillin	p Kohlhep 1989 Doxy- cy- cline	p Pfister 1989 Peni- cillin G	Pfister 1989 Cefo- taxime	Pfister 1991 Ceftri- axone	Pfister 1991 Cefo- taxime	Mul- legger 1991 Peni- cillin G	Mul- legger 1991 Ceftri- axone	Karls- son 1994 Peni- cillin G	Karls- son 1994 Doxycy- cline	Oksi 2007 Amox- icillin post- ceftriax- one	Oksi 2007 Placebo post- ceftriax- one	Ljostad 2008 Doxy- cy- cline	Ljostad 2008 Ceftri- axone
Number of par- ticipants (evalu- able)	36	39	10	11	14	16	11	12	23	31	73	72	54	48
Age mean (SD) un- less specified	Men 55 (12.6); women 54.1 (16.3)	Men 49.6 (14); women 55.7 (14.3)	56.7 (15)	55.4 (10.8)	58.7 (19.5)	53.7 (16.8)	8.1 (3.1)		Medi- an 55 (range 16 to 88)	Medi- an 49 (range 18 to 74)	Mean 52.3, range 19 to 87	Mean 50.5, range 16 to 80	54 (13)	52 (13)

Table 3. Baseline characteristics of the participants in the seven included studies

Percentage males	44%	51%	50%	64%	64%	44%	36%	42%	44%	29%	48%	50%	52%	65%
History of ery- thema migrans	36%	31%	80%	45%	50%	56%	Not repo	orted	61%	42%	26% (pro	bable)	31%	10%
Mean (SD) time from onset of LNB to treat- ment	5.2 (13.6) months	4.1 (11.1) months	28.7 (33.8) days	23.5 (16.3) days	64.5 (84.7) days	38.6 (23.1) days	All includ dren wer ted to th tal withi days from of sympt	re admit- le hospi- n 5 ± 1.8 m onset	3.5 weeks (1 week to 25 months)	4 weeks (1 week to 18 months)	Unknow	n	10 (19) weeks	8 (13) weeks
Previous treat- ment with an- tibiotics	11%	8%	Not repo	orted	Not re- ported	Not re- ported (men- tioned for 1 partici- pant)	Was an e criteria	exclusion	None for tl prior to en	ne 4 weeks rollment		th EM, 24 ad- and 14 not	Treatme cephalo: penicillin cycline i days exc criterion	sporin, n, tetra- n past 14 lusion
Concomitant treatment with steroids	28%	26%	None		Not repo	rted	Not repo	orted	Not report	ed	Not reported	I	Not reported	1
CSF leukocytes mean (SD) cells/ uL	186 (75)	145 (58)	280.9 (212)	435.7 (528)	86.4 (128.4)	135.3 (299.2)	Not repo	orted	Median 96, range 6 to 1190	Medi- an 117, range 8 to 910	CSF show	n available wed lympho- ocytosis.	194 (237)	178 (187)
CSF total protein mean (SD) in mg/dL	133 (110)	119 (112)	115 (69)	136 (67.4)	72.7 (42)	79.1 (48.4)	Not repo	orted	Medi- an 110, range 40 to 360	Medi- an 120, range 50 to 580	Not reported	I	120 (70)	130 (80)
Presence of CSF oligoclonal pands	78%	62%	70%	64%	64%	Not report- ed	Not reported	i	Not reported		(41%) pa with defi	d in 24/58 articipants inite LNB examined	Not reported	1

CSF: cerebrospinal fluid EM: erythema migrans

4

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LNB: Lyme neuroborreliosis SD: standard deviation



Study	Tool	Signs assessed	Subjective symp- toms elicited
Oksi 2007	Visual analogue scale	Yes	Possibly
Ljostad 2008	Composite clinical score	Yes	Yes
Mullegger 1991	Change of clinical symptoms	Yes	No
	Disease duration	Yes	No
Kohlhepp 1989	Change of clinical symptoms	Yes	Unclear
	(3-level classification)		
Pfister 1989	Change of clinical symptoms	Yes	Yes
Pfister 1991	Change of clinical symptoms	Yes	Unclear
Karlsson 1994	Change of clinical symptoms	Yes	No

Table 4. Measures of efficacy based on the assessment of signs and symptoms*

*The efficacy of interventions was quantified by diverse tools in each study assessing the change in objective findings (signs) or subjective complaints (symptoms), or both, as reported by participants or judged by the study physician.

Study	Parameter
Oksi 2007	Decrease of <i>B. burgdorferi</i> -specific antibody concentrations at 12 months of at least 20% ("moder- ate decline") or 50% ("strong decline")
Ljostad 2008	Resolution of CSF pleocytosis
Kohlhepp 1989	Cell count, protein, antibody index, B. burgdorferi-specific antibody production
Pfister 1989	Abnormal CSF on repeated lumbar puncture ¹
Pfister 1991	Abnormal CSF on repeated lumbar puncture ²
Karlsson 1994	Cell count, B. burgdorferi-specific antibody production
Mullegger 1991	Changes in intrathecally produced specific antibodies against B. burgdorferi

Table 5. Measures of efficacy based on cerebrospinal fluid analyses in the included studies

¹One or more of lymphocytic pleocytosis, protein elevation, oligoclonal bands, *B. burgdorferi*-specific antibody production. ²One or more of lymphocytic pleocytosis, protein elevation, oligoclonal bands, culture positive for *B. burgdorferi*. Abbreviations:

B. burgdorferi: Borrelia burgdorferi

CSF: cerebrospinal fluid

Table 6. Summary of findings table: oral amoxicillin versus placebo for people previously treated with ceftriaxone for Lyme neuroborreliosis (acute and chronic)

Oral amoxicillin versus placebo for people previously treated with ceftriaxone for Lyme neuroborreliosis (acute and late)

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Table 6. Summary of findings table: oral amoxicillin versus placebo for people previously treated with ceftriaxone for Lyme neuroborreliosis (acute and chronic) (Continued)

Patient or population: people previously treated with ceftriaxone for disseminated Lyme neuroborreliosis (acute and chronic)¹ Settings: Finland, hospital-based neurology/internal medicine, outpatient Intervention: oral amoxicillin Comparison: placebo

Outcomes	Illustrative compar	rative risks* (95% CI)	Relative – effect	No of Par- ticipants	Quality of the evi-	Comments
	Assumed risk or score/value	Corresponding risk or score/value	(95% CI)	(studies)	dence (GRADE)	
	Placebo	Oral amoxicillin	-			
Improvement in a measure of over- all disability in the long term (3 or more months) fol- lowing treatment	See comment	See comment	Not es- timable	-	See com- ment	Not report- ed
Improvement or resolution of the person's present- ing neurological deficits in the long term (3 or more months) following treatment ²	62), but not for inter subgroup was provid request. 59/62 partic experiencing improv rological deficits at r sessment: 'excellent	Separate information on the LNB subgroup (N = 62), but not for intervention groups within this subgroup was provided at the review authors' request. 59/62 participants were classified as experiencing improvement of presenting neurological deficits at month 12 (dichotomous assessment: 'excellent or good' based on investigator VAS values and medical record information). ³			Low ⁵	
Improvement in a measure of over- all disability in the short term (2 weeks) following treatment ⁴	See comment	See comment	Not es- timable	-	See com- ment	Not report- ed
Resolution of CSF pleocytosis follow- ing treatment	See comment	See comment	Not es- timable	-	See com- ment	Not mea- sured
All adverse events - 12 months	ticipants, mostly dia need for discontinua events reported. Att to either pretreatme	r all 145 Lyme disease par- rrhea and fever with no ation. No serious adverse ribution of adverse events ent with ceftriaxone, or to po, or both, is unclear.	Not es- timable	145 (1 study)	Very low ⁶	

*The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% CI) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI:** confidence interval; **CSF:** cerebrospinal fluid; **LNB:** Lyme neuroborreliosis; **RR:** risk ratio; **VAS**: visual analogue scale

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.



Table 6. Summary of findings table: oral amoxicillin versus placebo for people previously treated with ceftriaxone for Lyme neuroborreliosis (acute and chronic) (Continued)

Very low quality: We are very uncertain about the estimate.

Evidence based on randomized controlled trials begins as high-quality evidence, but confidence in the evidence was decreased for several reasons, including the following.

- Study limitations
- Inconsistency of results
- Indirectness of evidence
- Imprecision
- Reporting bias

¹Subpopulation with LNB (N = 62) within study of people with definite or possible disseminated Lyme borreliosis (N = 145).

²Month 12: dichotomous outcome: excellent or good, based on investigator VAS values and medical record information.

 3 No transparency on the influence of subjective symptoms on investigator VAS values, no standardization of inclusion of medical record information on LNB subgroup (N = 62). No transparency of date in publication. Only in the larger center did the same physician rate participants during the whole study.

⁴Trialists reported no statistically significant differences in VAS values between amoxicillin and placebo groups at 0, 1, 3, 6, and 12 months without providing numerical data for analysis.

⁵Downgraded twice: for study limitations (unclear risk of bias for all domains) and imprecision (small study size). We did not downgrade the quality of evidence for indirectness as, although flawed, the measure is likely to reflect clinical reality.

⁶Downgraded three times: twice for study limitations (lack of blinding and adverse events not ascribed to interventions) and once for indirectness (participants not limited to those with LNB; separate data not available for the LNB subgroup of 62 participants). In the absence of comprehensive adverse event reporting in the included trials, the table presents 'all adverse events' with a comment on severe adverse events when these data are presented in the trial.

Table 7. Summary of findings table: oral doxycycline compared to intravenous ceftriaxone for Lyme neuroborreliosis (acute and chronic)

Oral doxycycline compared to intravenous ceftriaxone for Lyme neuroborreliosis (LNB) (acute and chronic)

Patient or population: Lyme neuroborreliosis (acute and chronic) Settings: Southern Norway, hospital Intervention: oral doxycycline Comparison: intravenous ceftriaxone

Outcomes	Illustrative co risks* (95% Cl	•	Relative effect - (95% CI)	No of Par- ticipants (studies)	Quality of the evi- dence	Comments	
	Assumed risk or score/value	Correspond- ing risk or score/value	- (557661)	(studies)	(GRADE)		
	Intravenous ceftriaxone	Oral doxycy- cline	-				
Improvement in a mea- sure of overall disabili- ty in the long term (3 or more months) following treatment	See com- ment	See comment	Not es- timable	-	See com- ment	Not reported	
Resolution of the per- son's presenting neu- rological deficits in the long term (3 or more	333 per 1000	480 per 1000 (297 to 783)	RR 1.44 (0.89 to 2.35)	102 (1 study)	Moderate ¹	Symptom resolution; composite clinical score of neurological signs and symptoms at 12 months ²	



Table 7. Summary of findings table: oral doxycycline compared to intravenous ceftriaxone for Lyme

neuroborreliosis (acute and chronic) (Continued) months) following

treatment

Improvement in a mea- sure of overall disabil- ity in the short term (2 weeks) following treat- ment	See com- ment	See comment	Not es- timable	-	See com- ment	Not reported
Resolution of CSF pleo- cytosis following treat- ment	participants no significar difference w days and 4 n	lyzed in 88/102 ; authors state that at between-group as present at 13 nonths, but data able for verifica-	Not es- timable	88 (1 study)	Low ³	Resolution of CSF pleocy- tosis in all participants
All adverse events	464 per 1000	367 per 1000 (237 to 571)	RR 0.79 (0.51 to 1.23)	113 (1 study)	Moderate ⁴	48 adverse events in all participants randomized to study drug. 3 partici- pants on ceftriaxone and 1 on doxycycline expe- rienced serious adverse events (as defined by tri- al authors); RR 0.33 (95% CI 0.04 to 3.05).

*The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% CI) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI:** confidence interval; **CSF**: cerebrospinal fluid; **RR:** risk ratio

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

Evidence based on randomized controlled trials begins as high-quality evidence, but confidence in the evidence was decreased for several reasons, including the following.

- Study limitations
- Inconsistency of results
- Indirectness of evidence
- Imprecision
- Reporting bias

¹Downgraded once for imprecision (small study). We did not downgrade the quality of evidence for indirectness as, although flawed, the measure is likely to reflect clinical reality.

²Participants had predominantly acute LNB, although people with an acute or chronic course of LNB were eligible. Investigators included subjective symptoms in the overall neurologic deficit assessment but with a higher maximum score for objective neurologic findings. No long-term assessment was performed, however worsening after near-resolution in the majority of participants is unlikely.

³Downgraded twice: for indirectness of pleocytosis as an outcome measure and imprecision (small study).

⁴Downgraded once for imprecision (few events, small study). In the absence of comprehensive adverse event reporting in the included trials, the table presents 'all adverse events' with a comment on severe adverse events when these data are presented in the trial.

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Table 8. Summary of findings table: intravenous penicillin G compared to oral doxycyline for Lyme neuroborreliosis (acute and chronic)

Intravenous penicillin G compared to oral doxycyline for Lyme neuroborreliosis (acute and chronic)

Patient or population: Lyme neuroborreliosis (acute and chronic) Settings: Southern Sweden, hospital Intervention: intravenous penicillin G Comparison: oral doxycycline

Outcomes	Illustrative risks* (95%	comparative Cl)	Relative effect - (95% CI)	No of Par- ticipants (studies)	Quality of the evi- dence	Comments
	Assumed risk or score/val- ue	Correspond- ing risk or score/value		(statics)	(GRADE)	
	Oral doxy- cycline	Intravenous penicillin G	-			
Improvement in a measure of overall disability in the long term (3 or more months) following treatment	See com- ment	See comment	Not es- timable	-	See com- ment	Not measured
Improvement of the person's pre- senting neurological deficits in	1000 per 1000	1000 per 1000	RR 1.0 (0.92 to	51 (1 study)	Low ²	Investigators rating symp-
the long term (3 or more months) following treatment 1		(920 to 1000)	1.08)			tom composite on Likert scale from 1 to 3 (no, mild, moderate to severe) ²
Resolution of the person's pre- senting neurological deficits in the long term (3 or more months) following treatment ¹	900 per 1000	855 per 1000 (693 to 1000)	RR 0.95 (0.77 to 1.18)	51 (1 study)	Low ²	Investigators rating symp- tom composite on Likert scale from 1 to 3 (no, mild, moderate to severe) ²
Improvement in a measure of overall disability in the short term (2 weeks) following treatment	See com- ment	See comment	Not es- timable	-	See com- ment	Not reported
Resolution of CSF pleocytosis fol- lowing treatment Follow-up: 1 year	1000 per 1000	930 per 1000 (750 to 1000)	RR 0.93 (0.75 to 1.15)	29 (1 study)	Very low ³	
All adverse events	129 per 1000	130 per 1000 (32 to 526)	RR 1.01 (0.25 to 4.08)	54 (1 study)	Very low ⁴	

*The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% CI) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI:** confidence interval; **CSF**: cerebrospinal fluid; **RR:** risk ratio

GRADE Working Group grades of evidence



Table 8. Summary of findings table: intravenous penicillin G compared to oral doxycyline for Lyme neuroborreliosis

(acute and chronic) (Continued)

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

Evidence based on randomized controlled trials begins as high-quality evidence, but confidence in the evidence was decreased for several reasons, including the following.

- Study limitations
- Inconsistency of results
- Indirectness of evidence
- Imprecision
- Reporting bias

¹Measured at 3, 6, and 12 months; reported here at 12 months.

²Downgraded twice: for study limitations (unclear risk of selection bias and lack of blinding) and imprecision (small number of participants). Although judgement of objective findings is implied, the assessment approach does not allow a distinction between participant- and physician-based judgement on the basis of subjective and objective findings. We did not downgrade the quality of evidence for indirectness as, although flawed, the measure is likely to reflect clinical reality.

³Downgraded three times: for study limitations (unclear risk of selection bias and incomplete outcome data), imprecision (small number of participants), and indirectness of pleocytosis as an outcome measure.

⁴Downgraded three times: twice for study limitations (unclear risk of selection bias and lack of blinding) and once for imprecision (small number of participants, few events, and wide CI). In the absence of comprehensive adverse event reporting in the included trials, the table presents 'all adverse events' with a comment on severe adverse events when these data are presented in the trial.

Table 9. Summary of findings table: intravenous doxycycline compared to intravenous penicillin G for Lyme neuroborreliosis (acute and chronic)

Intravenous doxycycline compared to intravenous penicillin G for Lyme neuroborreliosis (acute and chronic)

Patient or population: Lyme neuroborreliosis (acute and chronic) Settings: Southern Germany, hospital Intervention: intravenous doxycycline Comparison: intravenous penicillin G

Outcomes	Illustrative o risks* (95% (•	Relative effect – (95% CI) –	No of Par- ticipants (studies)	Quality of the evi- dence	Comments	
	Assumed risk or score/val- ue	Correspond- ing risk or score/value		(studies)	(GRADE)		
	Intra- venous penicillin G	Intravenous doxycycline					
Improvement in a measure of overall disability in the long term (3 or more months) fol- lowing treatment	See com- ment	See com- ment	Not es- timable	-	See com- ment	Not reported	

Table 9. Summary of findings table: intravenous doxycycline compared to intravenous penicillin G for Lyme neuroborreliosis (acute and chronic) (Continued)

lieuloboli ellosis (acute allu cili		eu)				
Improvement of the person's presenting neurological deficits in the long term (3 or more months) following treatment ¹	833 per 1000	817 per 1000 (667 to 1000)	RR 0.98 (0.80 to 1.21)	75 (1 study)	Low ²	Clinical findings were classified as no remission, par- tial remission, or full remission.
Resolution of the person's pre- senting neurological deficits in the long term (3 or more months) following treatment ¹	694 per 1000	667 per 1000 (486 to 910)	RR 0.96 (0.70 to 1.31)	75 (1 study)	Low ²	Clinical findings were classified as no remission, par- tial remission, or full remission.
Improvement in a measure of overall disability in the short term (2 weeks) following treat- ment	See com- ment	See com- ment	Not es- timable	-	See com- ment	Not reported
Resolution of CSF pleocytosis following treatment	See com- ment	See com- ment	Not es- timable	-	-	Measured but not reported in detail
All adverse events ³	See com- ment	See com- ment	Not es- timable	75 (1 study)	-	'Adverse events' not reported. No serious adverse events occurred.

*The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% CI) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI:** confidence interval; **CSF**: cerebrospinal fluid; **RR:** risk ratio

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

Evidence based on randomized controlled trials begins as high-quality evidence, but confidence in the evidence was decreased for several reasons, including the following.

- Study limitations
- Inconsistency of results
- Indirectness of evidence
- Imprecision
- Reporting bias

¹Measured at 6 and 12 months; 12-month results reported here.

²Downgraded twice: for study limitations (unclear risk of selection bias, lack of blinding) and imprecision (small sample size). We did not downgrade the quality of evidence for indirectness as, although flawed, the measure is likely to reflect clinical reality.

³In the absence of comprehensive adverse event reporting in the included trials, the table presents 'all adverse events' with a comment on severe adverse events when these data are presented in the trial.

Table 10. Summary of findings table: intravenous cefotaxime compared to intravenous penicillin G for Lyme neuroborreliosis (acute)

Intravenous cefotaxime compared to intravenous penicillin G for Lyme neuroborreliosis (acute)

Patient or population: Lyme neuroborreliosis (acute) Settings: Southern Germany, hospital Intervention: intravenous cefotaxime Comparison: intravenous penicillin G

Outcomes	Illustrative o risks* (95% (Relative effect - (95% CI)	No of Par- ticipants (studies)	Quality of the evi- dence	Comments
	Assumed risk or score/val- ue	Correspond- ing risk or score/value	- (5576 Cl)	(statics)	(GRADE)	
	Intra- venous penicillin G	Intravenous cefotaxime	-			
Improvement in a measure of overall disability in the long term (3 or more months) follow- ing treatment	See com- ment	See comment	Not es- timable	-	See com- ment	Not reported
Resolution of the person's pre- senting neurological deficits in the long term (3 or more months) following treatment ¹	800 per 1000	816 per 1000 (536 to 1000)	RR 1.02 (0.67 to 1.55)	21 (1 study)	Low ²	Investigators' nonstandard- ized judgement of improvement or resolution of symptoms re- ported at 7.7 months.
Improvement in a measure of overall disability in the short term (2 weeks) following treat- ment	See com- ment	See comment	Not es- timable	-	See com- ment	Not reported
Resolution of CSF pleocytosis following treatment	1000 per 1000	920 per 1000 (710 to 1000)	RR 0.92 (0.71 to 1.18)	21 (1 study)	Very low ³	Follow-up: mean 7.7 months
All adverse events	See com- ment	See comment	Not es- timable	21 (1 study)	Low ⁴	No adverse events occurred.

*The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% CI) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI:** confidence interval; **CSF**: cerebrospinal fluid; **RR:** risk ratio

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

Antibiotics for the neurological complications of Lyme disease (Review)

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Table 10. Summary of findings table: intravenous cefotaxime compared to intravenous penicillin G for Lyme

neuroborreliosis (acute) (Continued)

Evidence based on randomized controlled trials begins as high-quality evidence, but confidence in the evidence was decreased for several reasons, including the following.

- Study limitations
- Inconsistency of results
- Indirectness of evidence
- Imprecision
- Reporting bias

¹Improvement in all participants at an average of 7.7 months.

²Downgraded twice: for study limitations (unclear risk of selection bias and lack of blinding) and imprecision (small sample size). We did not downgrade the quality of evidence for indirectness as, although flawed, the measure is likely to reflect clinical reality.

³Downgraded three times: for study limitations (unclear risk of selection bias), imprecision (small sample size), and indirectness (limitation to acute Lyme neuroborreliosis, indirectness of pleocytosis as an outcome measure). One participant in the cefotaxime group had mild residual pleocytosis. Resolution reported in 20/21 participants.

⁴Downgraded twice: for study limitations (unclear risk of selection bias and lack of blinding) and imprecision (small sample size). In the absence of comprehensive adverse event reporting in the included trials, the table presents 'all adverse events' with a comment on severe adverse events when these data are presented in the trial.

Table 11. Summary of findings table: intravenous ceftriaxone compared to intravenous cefotaxime for Lyme neuroborreliosis (acute)

Intravenous ceftriaxone compared to intravenous cefotaxime for Lyme neuroborreliosis (acute)

Patient or population: acute Lyme neuroborreliosis Settings: Southern Germany, hospital Intervention: intravenous ceftriaxone Comparison: intravenous cefotaxime

Outcomes	Illustrative ((95% CI)	comparative risks*	Relative effect - (95% CI)	No of Par- ticipants (studies)	Quality of the evi- dence	Comments	
	Assumed risk or score/val- ue	Corresponding risk or score/ value		(studies)	(GRADE)		
	Intra- venous ce- fotaxime	Intravenous cef- triaxone	-				
Improvement in a measure of over- all disability in the long term (3 or more months) following treatment	See com- ment	See comment	Not es- timable	-	See com- ment	Not mea- sured	
Resolution of the person's present- ing neurological deficits in the long term (3 or more months) following treatment	600 per 1000	666 per 1000 (378 to 1000)	RR 1.11 (0.63 to 1.97)	27 (1 study)	Low ¹	Outcome reported at a mean of 8.1 months' follow-up.	
Improvement in a measure of over- all disability in the short term (2 weeks) following treatment	See com- ment	See comment	Not es- timable	-	See com- ment	Not report- ed	

Antibiotics for the neurological complications of Lyme disease (Review)

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Table 11. Summary of findings table: intravenous ceftriaxone compared to intravenous cefotaxime for Lyme

neuroborreliosis (acute) (Continued)

Resolution of CSF pleocytosis fol- lowing treatment	867 per 1000	988 per 1000 (780 to 1000)	RR 1.14 (0.90 to 1.44)	27 (1 study)	Very low ²
All adverse events	188 per 1000	71 per 1000 (7 to 611)	RR 0.38 (0.04 to 3.26)	30 (1 study)	Low ³

*The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI).

CI: confidence interval; CSF: cerebrospinal fluid; RR: risk ratio

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

Very low quality: We are very uncertain about the estimate.

Evidence based on randomized controlled trials begins as high-quality evidence, but confidence in the evidence was decreased for several reasons, including the following.

- Study limitations
- Inconsistency of results
- Indirectness of evidence
- Imprecision
- Reporting bias

¹Downgraded twice: for study limitations (unclear risk of selection bias and lack of blinding) and imprecision (small sample size). We did not downgrade the quality of evidence for indirectness as, although flawed, the measure is likely to reflect clinical reality.

²Downgraded three times: for study limitations (unclear risk of selection bias), imprecision (small sample size), and indirectness of pleocytosis as an outcome measure.

³Downgraded twice: for study limitations (unclear risk of selection bias and lack of blinding) and imprecision (small sample size, few events). In the absence of comprehensive adverse event reporting in the included trials, the table presents 'all adverse events' with a comment on severe adverse events when these data are presented in the trial.

APPENDICES

Appendix 1. MEDLINE (OvidSP) search strategy

Database: Epub Ahead of Print, In-Process & Other Non-Indexed Citations, Ovid MEDLINE(R) Daily and Ovid MEDLINE(R) <1946 to Present> Ovid MEDLINE(R) 1946 to October Week 2 2016

Search Strategy:

1 randomized controlled trial.pt. (433771) 2 controlled clinical trial.pt. (91846) 3 randomized.ab. (374509) 4 placebo.ab. (180527) 5 clinical trials as topic.sh. (180437) 6 randomly.ab. (265901) 7 trial.ti,ab. (458646) 8 groups.ab. (1652249) 9 or/1-8 (2538867)



10 exp animals/ not humans.sh. (4331351) 11 9 not 10 (2156855) 12 Borrelia/ or borrelia.tw. (9635) 13 lyme.mp. (12202) 14 erythema migrans.mp. (1126) 15 erythema chronicum migrans.mp. (944) 16 or/12-15 (15327) 17 Lyme Neuroborreliosis/ or lyme neuroborreliosis.tw. (820) 18 Brain/ or brain.tw. (977583) 19 Meningitis/ or meningitis.tw. (52969) 20 Radiculopathy/ or radiculitis.tw. (4629) 21 Facial Paralysis/ (11282) 22 (facial palsy or facial paralysis).mp. (14025) 23 Myelitis/ or myelitis.mp. (5359) 24 Encephalitis/ or encephalitis.tw. (42001) 25 encephalopathy.mp. (44315) 26 Peripheral Nervous System Disease/ or neuropathy.mp. (73365) 27 Neuritis/ or Neuritis.tw. or neuro\$.mp. (1820199) 28 nerve damage.mp. (4781) 29 nerve involvement.mp. (2351) 30 Muscle Diseases/ or myopathy.mp. (17574) 31 Uveitis/ or uveitis.mp. (19924) 32 white matter.mp. (45516) 33 cranial nerve diseases/ or cranial nerve pals\$.mp. (5673) 34 Bannwarth\$1.mp. (165) 35 or/17-34 (2497544) 36 Anti-Bacterial Agents/ (279571) 37 antibiotics.mp. (191436) 38 ceftriaxone.mp. or Ceftriaxone/ (10329) 39 cephalosporin.mp. or Cephalosporins/ (22734) 40 doxycycline.mp. or Doxycycline/ (14221) 41 Tetracycline/ or tetracycline.mp. (37783) 42 erythromycin.mp. or Erythromycin/ (24037) 43 azithromycin.mp. or Azithromycin/ (7134) 44 penicillin.mp. or Penicillins/ (74810) 45 Amoxicillin/ or amoxicillin.mp. (16952) 46 or/36-45 (485048) 47 11 and 16 and 35 and 46 (77) 48 remove duplicates from 47 (75)

Appendix 2. Embase (OvidSP) search strategy

Database: Embase <1980 to 2016 Week 43> Search Strategy:

1 crossover-procedure.sh. (53448) 2 double-blind procedure.sh. (135943) 3 single-blind procedure.sh. (26362) 4 randomized controlled trial.sh. (456203) 5 (random\$ or crossover\$ or cross over\$ or placebo\$ or (doubl\$ adj blind\$) or allocat\$).tw,ot. (1332905) 6 trial.ti. (212018) 7 or/1-6 (1488225) 8 (animal/ or nonhuman/ or animal experiment/) and human/ (1699334) 9 animal/ or nonanimal/ or animal experiment/ (3720880) 10 9 not 8 (3019629) 11 7 not 10 (1375833) 12 Borrelia/ or borrelia.mp. (14679) 13 lyme.mp. (15502) 14 erythema migrans.mp. (1364) 15 erythema chronicum migrans.mp. (2210) 16 or/12-15 (21043) 17 Lyme Neuroborreliosis/ or lyme neuroborreliosis.mp. (12645)



18 Brain/ or brain.mp. (1747986) 19 Meningitis/ or meningitis.mp. (78071) 20 Radiculopathy/ or radiculitis.mp. (9952) 21 Facial Paralysis/ (9009) 22 (facial palsy or facial paralysis).mp. (9192) 23 Myelitis/ or myelitis.mp. (8571) 24 Encephalitis/ or encephalitis.mp. (58216) 25 encephalopathy.mp. (66605) 26 Peripheral Nervous System Disease/ or neuropathy.mp. (161107) 27 Neuritis/ or Neuritis.mp. (18566) 28 nerve damage.mp. (6280) 29 nerve involvement.mp. (3163) 30 Muscle Diseases/ or myopathy.mp. (35040) 31 Uveitis/ or uveitis.mp. (26998) 32 white matter.mp. (70624) 33 cranial nerve diseases/ or cranial nerve pals\$.mp. (6212) 34 Bannwarth\$1.mp. (219) 35 or/17-34 (2120234) 36 Anti-Bacterial Agents/ (163846) 37 antibiotics.mp. (202847) 38 ceftriaxone.mp. or Ceftriaxone/ (48176) 39 cephalosporin.mp. or Cephalosporins/ (51075) 40 doxycycline.mp. or Doxycycline/ (44244) 41 Tetracycline/ or tetracycline.mp. (86047) 42 erythromycin.mp. or Erythromycin/ (72161) 43 azithromycin.mp. or Azithromycin/ (27929) 44 penicillin.mp. or Penicillins/ (133479) 45 Amoxicillin/ or amoxicillin.mp. (79682) 46 or/36-45 (607536) 47 11 and 16 and 35 and 46 (119) 48 remove duplicates from 47 (116)

Appendix 3. CENTRAL (CRSO) search strategy

Search run on Tue Oct 25 2016

#1 borrelia:TI,AB,KY 92
#2 (erythema near3 migrans):TI,AB,KY 40
#3 lyme:TI,AB,KY 117
#4 #1 OR #2 OR #3 141
#5 (brain or meningitis or radiculopathy or radiculitis):TI,AB,KY 28989
#6 ("facial paralysis" or "facial palsy" or myelitis or encephalitis or encephalopathy):TI,AB,KY 2487
#7 MESH DESCRIPTOR Peripheral Nervous System Diseases 419
#8 (neuropathy or neuritis or "nerve damage" or "nerve involvement" or neuro*):TI,AB,KY 56849
#9 (myopathy or myositis or uveitis or "white matter"):TI,AB,KY 2057
#10 ("cranial nerve disease" or "cranial nerve diseases" or "cranial nerve palsy" or "Bannwarth NEAR syndrome"):TI,AB,KY 41
#11 #5 or #6 or #7 or #8 or #9 or #10 77315
#12 antibiotic* or ceftriaxone or cephalosporin* or doxycycline or tetracycline or erythromycin or azithromycin or penicillin or amoxicillin
25671
#13 #4 AND #11 AND #12 35
#14 sr-neuromusc:cc 5977
#145 WEAR MARCH AND #11 AND #12 45

#15 #13 not #14 11

Appendix 4. Cochrane Neuromuscular Specialised Register (CRS) search strategy

#1 BORRELIA [REFERENCE] [STANDARD]
#2 erythema near3 migrans [REFERENCE] [STANDARD]
#3 lyme [REFERENCE] [STANDARD]
#4 #1 or #2 or #3 [REFERENCE] [STANDARD]
#5 brain or meningitis or radiculopathy or radiculitis [REFERENCE] [STANDARD]
#6 "facial paralysis" or "facial palsy" or myelitis or encephalitis or encephalopathy [REFERENCE] [STANDARD]
#7 MeSH DESCRIPTOR Peripheral Nervous System Diseases [REFERENCE] [STANDARD]
#8 neuropathy or neuritis or "nerve damage" or "nerve involvement" or neuro* [REFERENCE] [STANDARD]



#9 myopathy or myositis or uveitis or "white matter" [REFERENCE] [STANDARD]

#10 "cranial nerve disease" or "cranial nerve diseases" or "cranial nerve palsy" or "Bannwarth NEAR syndrome" [REFERENCE] [STANDARD] #11 #5 or #6 or #7 or #8 or #9 or #10 [REFERENCE] [STANDARD] #12 #4 and #11 [REFERENCE] [STANDARD]

#13 antibiotic* or ceftriaxone or cephalosporin* or doxycycline or tetracycline or erythromycin or azithromycin or penicillin or amoxicillin [REFERENCE] [STANDARD]

#14 #12 and #13 [REFERENCE] [STANDARD]

#15 #12 AND #13 AND (INREGISTER) [REFERENCE] [STANDARD]

Appendix 5. Search strategy for clinical trial registries

Terms used:

Lyme disease

Lyme Borreliosis

Borrelia

Neuroborreliosis

Nervous system Lyme

Sites searched:

clinicaltrials.gov/

www.clinicaltrialsregister.eu/

www.who.int/ictrp/en/

www.isrctn.com/

Appendix 6. Additional methods

If data are available for meta-analysis in the future, we will follow methods outlined in our protocol (Cadavid 2008), updated to meet current Cochrane guidelines.

Types of outcome measures

If studies use different follow-up periods, we will scale results to a standardized follow-up period before we pool them in a meta-analysis.

If cost and cost-effectiveness information is available, we will include it as part of the effects of the intervention analysis.

Data extraction and management

If published trials exclude any randomized participants from analyses, we will contact trial authors and incorporate any available additional data.

Measures of treatment effect

We will use the Cochrane Review Manager 5 software to calculate risk ratios (RRs) with 95% confidence intervals (CIs) for dichotomous outcomes. We will use odds ratios (OR) for logistic regression analysis. For continuous data, we will examine the mean difference (MD) and report corresponding 95% CI when studies use the same measurement tool. Where studies assess an outcome using different measurement scales, we will calculate the standardized mean difference (SMD) with 95% CI.

If studies with different scales are combined, we will ensure that higher scores for continuous outcomes all have the same meaning for any particular outcome. We will explain the direction of interpretation and report when directions are reversed.

Assessment of heterogeneity

We will investigate statistical heterogeneity among studies using I² methodology (Higgins 2011). If significant heterogeneity exists (I² > 50%) for primary outcome measures, we will scrutinize reasons such as disease manifestation and severity, or treatment selection and duration. However, we will bear in mind uncertainty in interpretation when there are few studies.

Data synthesis

We will perform a fixed-effect analysis and will repeat the calculations using the random-effects model, as this is more conservative than a fixed-effect model, which assumes that studies are measuring the same intervention effect.



We will undertake a meta-analysis only if we consider participants, interventions, comparisons, and outcomes to be sufficiently similar to ensure a result that is clinically meaningful.

If we include multi-arm studies, we will analyze multiple intervention groups to avoid omission of relevant groups or double-counting of participants, following guidance in the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011).

Subgroup analysis and investigation of heterogeneity

We will also consider the results of European studies in combination with and separately from results of studies performed in the United States. In Europe, diagnostic guidelines are less standardized, and the causative organisms of neuroborreliosis are different and more varied. We will want to learn what, if any, effect these geographic differences have on treatment outcomes.

Sensitivity analysis

We will perform sensitivity analyses by repeating the analysis excluding unpublished studies and studies at high risk of bias, and examining the effect of large, dominant studies or particular inclusion or exclusion criteria. Since the number of clinical studies of Lyme neuroborreliosis is small, we will consider the criteria used in each of the studies, assign levels of confidence to each of the definitions used, and interpret our meta-analysis in light of this information.

WHAT'S NEW

Date	Event	Description
9 December 2016	Amended	Addition to Declarations of interest

HISTORY

Protocol first published: Issue 1, 2008 Review first published: Issue 12, 2016

Date	Event	Description
2 September 2008	Amended	Converted to new review format.

CONTRIBUTIONS OF AUTHORS

D Cadavid: wrote most sections of the review with assistance from H Gelderblom and PG Auwaerter. D Cadavid liaised with editorial base and co-ordinated the writing and updates to the actual review. D Cadavid prepared several of the tables and final revisions.

P Auwaerter: work on the review of several included studies and on the preparation of the actual review and obtained supplementary data from J Oksi.

J Rumbaugh: liaised with editorial base for the original protocol and co-ordinated contributions from review authors in the planning phase; participated in the initial phases of the review, but not in the later phases.

H Gelderblom: work on the data analysis and preparation of the actual review and contributed to the preparation of the GRADE tables.

DECLARATIONS OF INTEREST

D Cadavid was a full-time paid employee of Biogen during most of the preparatory time for this review. He is currently a full-time employee of Fulcrum Therapeutics. Neither Biogen nor Fulcrum Therapeutics is involved in research on LNB. D Cadavid's work on this review is not related to his employment with Biogen or Fulcrum Therapeutics.

PG Auwaerter has served as a medical-legal expert witness regarding Lyme disease; has been reimbursed for travel expenses related to an update of the Lyme Disease Guideline by the Infectious Diseases Society of America, the American Academy of Neurology, and the American College of Rheumatology (IDSA/AAN/ACR); and has been given honoraria for CME courses regarding Lyme disease.

J Rumbaugh has been reimbursed for travel expenses related to an update of the Lyme Disease Guideline (IDSA/AAN/ACR).

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H Gelderblom: none known.

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Internal sources

• Paul Auwaerter, USA.

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Diego Cadavid, USA.

Diego Cadavid was a full-time employee of Biogen and consultant at the Center for Immunology and Inflammatory Diseases at Massachusetts General Hospital during much of the preparation of this review. He is currently a full-time employee of Fulcrum Therapeutics. His work on this review is not related to his prior employment at Biogen or his current employment at Fulcrum Therapeutics.

External sources

• No sources of support supplied

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

The review has a published protocol (Cadavid 2008). A change in authors occurred: Dr J Aucott withdrew, and Dr H Gelderblom joined the review team. Subsequent to the publication of the protocol, we updated the methodology for the 'Risk of bias' assessment according to the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). Early in 2014 we decided that a meta-analysis was not feasible and changed the focus of the review to a systematic narrative review.

The original protocol listed any treatment, including antibiotics and steroids. However, we changed the title of the review from 'Treatment for the neurological complications of Lyme disease' to 'Antibiotics for the neurological complications of Lyme disease' to focus the review on the usual treatment for this condition. This resulted in exclusion of a single randomized study of oral corticosteroids (Pfister 1988).

The original plan was to discuss non-randomized, open, and uncontrolled studies and case series with more than 10 participants as appropriate, but not to include them in a meta-analysis. We did not assess non-randomised studies in the full review. We were unable to perform meta-analysis; therefore, our focus changed to a narrative review of randomized studies, which mostly provided low or very-low quality evidence. We considered that non-randomized studies were unlikely to provide additional valuable safety and efficacy data of sufficient quality for conclusions to be drawn.

The original protocol proposed to study neuropathy and myopathy. This was not possible in the current review, as the included studies did not provide distinct information on peripheral LNB.

The original protocol described a minimum set of diagnostic criteria for inclusion. We followed this to a large degree, however due to the heterogeneity of the seven studies included in the review, we opted for listing the detailed inclusion criteria in a separate table (Table 2).

The original protocol proposed as primary endpoints for the meta-analysis 'improvement in a measure of overall disability or resolution of the presenting neurological deficits at three or more months following treatment'. This was not possible due to the heterogeneity in efficacy assessments. The same problem occurred with the secondary efficacy endpoints proposed in the protocol. For the review we listed the efficacy assessments as performed in each of the seven included trials and added Table 1, which summarizes efficacy assessments in each trial.

The original protocol proposed to scale appropriately the study duration to a standardized follow-up period before pooling data in a metaanalysis. This was not done. For the review we listed the duration of follow-up for each study and summarized the results in Table 1.

None of the proposed meta-analyses specified in the original protocol were possible.

We transferred methods reported in the protocol relating to meta-analysis to Appendix 6.

INDEX TERMS

Medical Subject Headings (MeSH)

Amoxicillin [therapeutic use]; Anti-Bacterial Agents [*therapeutic use]; Borrelia burgdorferi; Cefotaxime [therapeutic use]; Ceftriaxone [therapeutic use]; Doxycycline [therapeutic use]; Lyme Disease [complications]; Lyme Neuroborreliosis [*drug therapy]; Penicillin G [therapeutic use]; Randomized Controlled Trials as Topic



MeSH check words

Humans