National Institute for Health and Care Excellence

Draft for Consultation

Lyme disease: diagnosis and management

[J] Evidence review for the management of lymphocytoma

NICE guideline

Evidence review

September 2017

Draft for Consultation

This evidence review was developed by the National Guideline Centre



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Management (lymphocytoma)

1.1 Review question: What is the most clinically and cost-2 effective treatment for people with lymphocytoma related 3 to Lyme disease? 4

1.2 Introduction 5

6 Borrelial lymphocytoma is an early but rare skin manifestation of Lyme disease. It most commonly occurs on the earlobe, breast or scrotum as a bluish, erythematous well-defined 7 patch of skin. It may or may not be present with an erythema migrans (EM) rash at another 8 9 site.

1.3 PICO table 10

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11 For full details, see the review protocol in appendix A.

Table 1: PICO o	characteristics of review question
Population	Adults (18 years and over), young people (12 to 17 years) and children (under 12 years) with symptoms consistent with lymphocytoma related to Lyme disease
Interventions	Antimicrobials, including but not limited to: Penicillins Amoxicillin (oral, IV) Ampicillin (oral, IV) Benzylpenicillin sodium / Penicillin G (IV) Including Augmentin (Amoxicillin and clavulanic acid; oral, IV) Phenoxymethylpenicillin / Penicillin V (oral) Tetracyclines Doxycycline (oral) Minocycline (oral) Cephalosporins Cefotaxime (IV) Ceftriaxone (IV) Ceftroxime axetil (oral) Macrolides Azithromycin (oral, IV) Fluoroquinolones Ciprofloxacin (oral, IV) Levofloxacin (oral, IV) Nalidixic acid (oral) Norfloxacin (oral, IV) Rifampicin (oral, IV) Rifampicin (oral, IV)
	Steroids (corticosteroids)
Comparisons	Any type of intervention compared to each other
	∘ If data are available consider:
	- Type of agent (within class or between class)

	- Route of administration		
	- Duration of treatment: 1 month versus longer		
	Monotherapy versus polytherapy (any combination)		
	Antimicrobial treatment or steroids compared to no treatment / placebo		
Outcomes	Critical:		
	Quality of life (any validated measure)		
	Cure (resolution of lymphocytoma symptoms)		
	Reduction of lymphocytoma symptoms		
	Relapse of lymphocytoma symptoms		
	Important:		
	5. Adverse events		
Study design	• RCTs		
	Cohort studies (if no RCT evidence is found)		

1 1.4 Clinical evidence

2 1.4.1 Included studies

- 3 No relevant RCTs and cohort studies comparing the effectiveness of antibiotics versus each
- 4 other or placebo as treatment for people with lymphocytoma related to Lyme disease were
- 5 identified.
- 6 See also the study selection flow chart in appendix C.

7 1.4.2 Excluded studies

8 See the excluded studies list in appendix I.

9 1.4.3 Summary of clinical studies included in the evidence review

10 No relevant clinical studies were identified.

11 1.4.4 Quality assessment of clinical studies included in the evidence review

12 No relevant clinical studies were identified.

1.5 Economic evidence

2 1.5.1 Included studies

- 3 No relevant health economic studies were identified.
- 4 See also the health economic study selection flow chart in appendix G.

5 1.5.2 Excluded studies

No relevant health economic studies were identified and excluded.

≥1.5.3 Unit costs

The following unit costs were presented to the committee to aid consideration of cost-effectiveness.

Table 2: UK costs of antimicrobials

Class	Drug	Age	Preparation	Mg/unit	Cost/unit (£)	Units/day	Course duration (days)	Cost per course (£)
Penicillins	Penicillins Amoxicillin	7 days-11 months	125 mg/1.25 ml oral suspension paediatric	125	0.20	3	14–28	8.35–16.70
		1-4 years	250 mg/5 ml oral suspension	250	0.06	3	14–28	2.37–4.75
		>5 years	capsules	500	0.06	3	14–28 (g)	2.54-5.08
Penicillins	Phenoxymethy lpenicillin	Adults (a)	tablets	250	0.04	4	10	1.49
Tetracyclines	Doxycycline	>12 years	capsules	100	0.11	2	10-28 (h)	2.18-6.09
Cephalosporins	Cefuroxime axetil	>3 months	tablets	250	1.27	4	14–28 (g)	70.88–141.76
Macrolide	Clarithromycin	>1 month	tablets	500	0.16	2	14–21	4.42-6.63
Macrolide	Azithromycin	<12 years	40 mg/1 ml oral suspension	40	0.27	10 mg/kg	9 (i)	Weight dependent
		Adults	tablets	500	0.42	1	9 (i)	3.75
Cephalosporins	Cefotaxime	Adults (b)	2 g powder for solution for injection vials (IV)	2,000	3.75	3	10	112.50
Cephalosporins	Ceftriaxone	>9 years (c)(d)	2 g powder for solution for injection vials (IV) (e)	2,000	1.03	1	14–21	14.42–21.63
Penicillins	Benzylpenicilli n sodium	Adults (f)	600 mg powder for solution for injection vials (IM)	600	2.73	2	3	16.38

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Sources: Unit costs from NHS Electronic Drug Tariff January 2017,¹¹⁷ except cefotaxime from BNF, January 2017²⁰ and ceftriaxone from EMIT March 2017;³⁷ dosage from BNF and BNF for Children January 2017,^{20,21} exceptions below:

- (a) Source of dosage from RCT in adults with ECM: Steere 1983, 164 dosage for Lyme disease not available from BNF or BNF for children.
- (b) Source of dosage from RCT in adults with neuroborreliosis: Pfister 1989¹²⁹ and Pfister 1991, 130 dosage for Lyme disease not available from BNF or BNF for children. 20,21
- (c) For disseminated Lyme borreliosis.
- (d) Dose for neonate and child up to 11 years (body weight <50 kg) 50-80mg/kg once daily for 14-21 days. BNF for children January 2017.²¹
- (e) Administration can vary in adults and children >1 month: IV infusion over 30 mins or IV injection over 5 mins or deep muscular injection (doses over 1 g divided between more than 1 site): 2 g per day for 14-21 days BNF January 2017.²⁰
- (f) Source of dosage from RCT in adults with Lyme arthritis: Steere 1985:¹⁶³ 1.2 million U injected in each buttock weekly intramuscularly. Duration 3 weeks. Dosage for Lyme disease not available from BNF or BNF for children. ^{20,21}
- (g) Course duration for early Lyme 14-21 days; 28 days for Lyme arthritis. BNF January 2017.²⁰
- (h) Course duration for early Lyme 10-14 days; 28 days for Lyme arthritis. BNF January 2017.²⁰
- (i) Course dose and duration for adults: 500 mg once daily for 3 days, for 3 weeks. For children under 12 years: 10 mg/kg once daily for 3 days for 3 weeks. Committee expert opinion.

The cost of intravenous antibiotics will vary depending on where these are administered and by whom. These costs will include some of the following cost components:

- antibiotic
- nursing time (for example, Band 6 nurse, £44 per hour, PSSRU 2016⁴⁰)
- clinic space and clerical time (for outpatient administration)
- travel time (for home administration)
- hospital bed (for inpatient administration)
- consumables (for example, cannula, needles, syringes, dressing, IV giving set and glucose or sodium chloride solution).

A large proportion of the total cost of intravenous antibiotics is likely to be the cost of administration rather than the drug itself. As a result, intravenous drugs that have multiple doses administered per day will be more costly than those administered once daily. This was explored in a detailed costing analysis conducted for the NICE CG102 (Meningitis [bacterial] and meningococcal septicaemia in under 16s). 114 In this analysis, they found that ceftriaxone was the cheapest antibiotic when compared to cefotaxime and benzylpenicillin. This was due to savings in staff time associated with once daily dosing, which offset the higher cost of the drug itself.

Inpatient administration

Intravenous antibiotics administered in an inpatient setting will incur the cost of an inpatient stay, which is assumed to include intravenous antibiotics treatment as part of the unit cost. The weighted average unit cost of non-elective inpatient stays and day cases for infectious disease in adults and children are summarised estimated in the table below using the NHS reference costs 2015/2016.⁴⁵

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Table 3: Unit costs of inpatient administration

Schedule	Currency description	Currency codes	Weighted average unit costs (per day)
Day-case adults	Standard/major/complex infectious diseases with/without single/multiple interventions, with/without CC	WJ01B, WJ01D, WJ01E, WJ02B, WJ02C,WJ02D, WJ02E, WJ03A, WJ03B, WJ03C, WJ03D, WJ03E, WJ03F, WJ03G	£352
Day-case paediatrics	Paediatric minor/major/intermediate infections with/without CC	PW01A, PW01B, PW01C, PW16A, PW16B, PW16C, PW16D, PW16E, PW17D, PW17E, PW17F, PW17G	£448
Non-elective inpatient short-stay adults	Standard/major/complex infectious diseases with/without single/multiple interventions, with/without CC	WJ01A, WJ01B, WJ01C, WJ01D, WJ01E, WJ02A, WJ02B, WJ02C,WJ02D, WJ02E, WJ03A, WJ03B, WJ03C, WJ03D, WJ03E, WJ03F, WJ03G	£432
Non-elective inpatient short-stay paediatrics	Paediatric minor/major/intermediate infections with/without CC	PW01A, PW01B, PW01C, PW16A, PW16B, PW16C, PW16D, PW16E, PW17D, PW17E, PW17F, PW17G	£521
Non-elective inpatient long-stay adults	Standard/major/complex infectious diseases with/without single/multiple interventions, with/without CC	WJ01A, WJ01B, WJ01C, WJ01D, WJ01E, WJ02A, WJ02B, WJ02C,WJ02D, WJ02E, WJ03A, WJ03B, WJ03C, WJ03D, WJ03E, WJ03F, WJ03G	£473
Non-elective inpatient long-stay paediatrics	Paediatric minor/major/intermediate infections with/without CC	PW01A, PW01B, PW01C, PW16A, PW16B, PW16C, PW16D, PW16E, PW17D, PW17E, PW17F, PW17G	£699

Source: NHS reference costs 2015/2016⁴⁵

Outpatient administration

Intravenous antibiotics may also be administered as part of an outpatient parenteral antibiotic therapy (OPAT) service, which is available in some hospitals. This allows for administration in an outpatient clinic or in a home setting by a district nurse and is for people who require parenteral treatment but are otherwise stable and well enough not to be in hospital. There is currently no NHS reference cost for this service.

A UK study by Chapman 2009²⁹ reports that this type of service costs between 41% and 61% of the equivalent inpatient costs. Based on these estimates from Chapman 2009 and the unit cost for an adult day case in Table 3, the cost of OPAT would be approximately £144 to £215 per day. These costs would include the cost of the drug as well as the administration.

1.6 **Resource impact** 1

2 We do not expect recommendations resulting from this review area to have a significant 3

impact on resources.

1.7 **Evidence statements**

1.7.1 Clinical evidence statements 5

No relevant clinical evidence was identified. 6

Health economic evidence statements 7 1.7.2

No relevant economic evaluations were identified. 8

1.8 Recommendations 9

10 No recommendations were made for this review question.

1.9 Rationale and impact 11

12 1.9.1 Why the committee did not make any recommendations

Lymphocytoma is a rare early presentation of Lyme disease. The guideline committee 13 agreed not to make a recommendation for the antibiotic management of lymphocytoma 14 because no evidence was identified, and the committee agreed that a person presenting with 15 lymphocytoma only was likely to require specialist investigation of lesions to establish the 16 17 diagnosis in most cases. For people with a clear supportive history and other symptoms suggesting Lyme disease as the likely diagnosis, the committee agreed that they would 18

receive treatment appropriate for their other symptoms.

1.9.2 20 Impact of the recommendations on practice

21 No recommendations were made for this review question.

1.10 The committee's discussion of the evidence 22

1.10.1 Interpreting the evidence 23

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24 1.10.1.1 The outcomes that matter most

25 The guideline committee considered quality of life, cure or the resolution of lymphocytoma 26 symptoms, the reduction of lymphocytoma symptoms, and the relapse of lymphocytoma symptoms to be critical outcomes. Adverse events as a result of treatment were considered 27

28 to be an important outcome.

29 No evidence for any of the outcomes was identified for this review.

30 1.10.1.2 The quality of the evidence

No evidence was identified for this review. 31

1 1.10.1.3 Benefits and harms

2 No evidence was identified for this review.

3 1.10.2 Cost effectiveness and resource use

No health economic evidence was identified. Unit costs of antimicrobials were presented to the committee.

6 1.10.3 Other factors the committee took into account

Lymphocytoma is a harmless pseudolymphoma that can be mistaken for a cutaneous lymphoma. Although most cases cannot be attributed to a specific cause, lymphocytoma has also been associated with Lyme disease.

The committee agreed that a lymphocytoma related to Lyme disease is a very rare condition. People with Lyme disease would not present with a lymphocytoma alone, but also with other signs and symptoms suggestive of Lyme disease. The committee therefore agreed that these people should receive treatment appropriate for their symptoms constellation. The committee considered investigation and management of people with lymphocytoma only was a matter for specialist practice and that including a recommendation in this guideline would be misleading. The committee made a general research recommendation for antibiotic management of Lyme disease but did not prioritise this area for a separate research recommendation.

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 - 185. Vazquez M, Sparrow SS, Shapiro ED. Long-term neuropsychologic and health outcomes of children with facial nerve palsy attributable to Lyme disease. Pediatrics. 2003; 112(2):e93-97
 - 186. Wahlberg P, Granlund H, Nyman D, Panelius J, Seppala I. Treatment of late Lyme borreliosis. Journal of Infection. 1994; 29(3):255-261
 - 187. Weber K, Neubert U, Thurmayr R. Antibiotic therapy in early erythema migrans disease and related disorders. Zentralblatt fur Bakteriologie, Mikrobiologie, und Hygiene Series A, Medical Microbiology, Infectious Diseases, Virology, Parasitology. 1987; 263(3):377-388
- 43 188. Weber K, Preac-Mursic V, Neubert U, Thurmayr R, Herzer P, Wilske B et al.
 44 Antibiotic therapy of early European Lyme borreliosis and acrodermatitis chronica
 45 atrophicans. Annals of the New York Academy of Sciences. 1988; 539:324-345

1 2 3	189.	Weissenbacher S, Ring J, Hofmann H. Gabapentin for the symptomatic treatment of chronic neuropathic pain in patients with late-stage lyme borreliosis: a pilot study. Dermatology. 2005; 211(2):123-127
4 5 6	190.	White B, Seaton RA, Evans TJ. Management of suspected lyme borreliosis: experience from an outpatient parenteral antibiotic therapy service. QJM. 2013; 106(2):133-138
7 8 9	191.	Zochling N, Mullegger RR, Schluepen EM, Soyer HP, Hodl S, Wienecke R et al. Minocycline in early Lyme Borreliosis. Acta Dermatovenerologica Alpina, Panonica et Adriatica. 1996; 5(3-4):163-168
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Appendices

Appendix A: Review protocols

Table 4: Review protocol for the management of lymphocytoma related to Lyme disease

Question number: 4.6

Relevant section of Scope: management

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Field	Content
Review question	What is the most clinically and cost-effective treatment for people with lymphocytoma related to Lyme disease?
Type of review question	A review of health economic evidence related to the same review question was conducted in parallel with this review. For details, see the health economic review protocol for this NICE guideline.
Objective of the review	The review questions on the condition-specific management of Lyme disease aim to identify the most effective treatment in different clinical scenarios. The questions have been developed in a way to identify the evidence for all potential populations and scenarios, even if clinical presentations are more diverse. The population for this review consists of people with lymphocytoma related to Lyme disease.
Eligibility criteria – population / disease / condition	Adults (18 years and over), young people (12 to 17 years) and children (under 12 years) with symptoms consistent with lymphocytoma related to Lyme disease
Eligibility criteria – intervention(s)	Antimicrobials, including but not limited to: Penicillins Amoxicillin (oral, IV) Ampicillin (oral, IV) Benzylpenicillin sodium / Penicillin G (IV) Including Augmentin (Amoxicillin and clavulanic acid; oral, IV) Phenoxymethylpenicillin / Penicillin V (oral) Tetracyclines Doxycycline (oral) Minocycline (oral) Cephalosporins Cefotaxime (IV) Ceftriaxone (IV) Ceftriaxone (IV) Cefuroxime axetil (oral) Macrolides Azithromycin (oral, IV) Fluoroquinolones Ciprofloxacin (oral, IV) Nalidixic acid (oral, IV) Nalidixic acid (oral) Norfloxacin (oral) Norfloxacin (oral)

Field	Content
	Ofloxacin (oral, IV)Rifampicin (oral, IV)
	Steroids (corticosteroids)
Eligibility criteria – comparator(s)	 Any type of intervention compared to each other If data are available, consider: Type of agent (within class or between class) Route of administration Duration of treatment: 1 month versus longer Monotherapy versus polytherapy (any combination) Antimicrobial treatment or steroids compared to no treatment / placebo
Outcomes and prioritisation	Critical: 1. Quality of life (any validated measure) 2. Cure (resolution of lymphocytoma symptoms) 3. Reduction of lymphocytoma symptoms 4. Relapse of lymphocytoma symptoms Important: 5. Adverse events
Eligibility criteria – study design	RCTsCohort studies (if no RCT evidence is found)
Other inclusion exclusion criteria	Date limits for search: none Language: English only Setting: all settings in which NHS care is provided or commissioned The following interventions will not be considered for inclusion: • Metronidazole • Trimethoprim
Proposed sensitivity / subgroup analysis, or meta-regression	 The following groups will be considered separately if data are available (strata): Children (under 12 years); young people and adults (12 years and over) Onset of specific symptoms less than 6 weeks; 6 weeks to 6 months; over 6 months Subgroups (to be investigated if heterogeneity is identified): Pregnant women People who are immunocompromised People in whom a previous course of antimicrobial treatment or steroid treatment has failed
Selection process – duplicate screening / selection / analysis	Studies will be sifted by title and abstract. Potentially significant publications obtained in full text will then be assessed against the inclusion criteria specified in this protocol.
Data management (software)	Pairwise meta-analyses will be performed using Cochrane Review Manager (RevMan5). GRADEpro will be used to assess the quality of evidence for each outcome Bibliographies, citations, study sifting and reference management will be managed using EndNote. Data extractions will be performed using EviBase, a platform designed and maintained by the National Guideline Centre (NGC)
Information sources –	Clinical searches

Field	Content
databases and dates	Medline, Embase, The Cochrane Library all years
	Health economic searches
	Medline, Embase, NHS Economic Evaluation Database (NHS EED), Health Technology Assessment (HTA) all years
Identify if an update	Not applicable
Author contacts	https://www.nice.org.uk/guidance/indevelopment/gid-ng10007
Highlight if amendment to previous protocol	For details, please see section 4.5 of Developing NICE guidelines: the manual.
Search strategy – for one database	For details, please see appendix B
Data collection process – forms / duplicate	A standardised evidence table format will be used, and published as appendix D of the evidence report.
Data items – define all variables to be collected	For details, please see evidence tables in Appendix D (clinical evidence tables) or H (health economic evidence tables).
Methods for assessing bias at outcome / study level	Standard study checklists were used to appraise individual studies critically. For details, please see section 6.2 of Developing NICE guidelines: the manual The risk of bias across all available evidence will be evaluated for each
	outcome using an adaptation of the 'Grading of Recommendations Assessment, Development and Evaluation (GRADE) toolbox' developed by the international GRADE working group http://www.gradeworkinggroup.org/
Criteria for quantitative synthesis	For details, please see section 6.4 of Developing NICE guidelines: the manual.
	Meta-analysis will be conducted wherever possible (that is, where similar studies can be combined)
	In the absence of clinically established MIDs, standard MIDs for dichotomous (25% risk reduction or risk increase) and continuous outcomes (+/-0.5 standard deviation) will be used
	If heterogeneity is found, the influence of subgroups will be examined
Methods for quantitative analysis – combining studies and exploring (in)consistency	For details, please see the separate Methods report for this guideline.
Meta-bias assessment – publication bias, selective reporting bias	For details, please see section 6.2 of Developing NICE guidelines: the manual.
Confidence in cumulative evidence	For details, please see sections 6.4 and 9.1 of Developing NICE guidelines: the manual.
Rationale / context – what is known	For details, please see the introduction to the evidence review.
Describe contributions of authors and guarantor	A multidisciplinary committee developed the evidence review. The committee was convened by the NGC and chaired by Saul Faust in line with section 3 of Developing NICE guidelines: the manual.
	Staff from the NGC undertook systematic literature searches, appraised the evidence, conducted meta-analysis and cost-effectiveness analysis where appropriate, and drafted the evidence review in collaboration with the committee. For details, please see Developing NICE guidelines: the manual.
Sources of funding / support	The NGC is funded by NICE and hosted by the Royal College of Physicians.
Name of sponsor	The NGC is funded by NICE and hosted by the Royal College of

Field	Content
	Physicians.
Roles of sponsor	NICE funds the NGC to develop guidelines for those working in the NHS, public health and social care in England.
PROSPERO registration number	Not registered

Table 5: Health economic review protocol

alth economic review protocol
All questions – health economic evidence
To identify health economic studies relevant to any of the review questions.
 Populations, interventions and comparators must be as specified in the clinical review protocol above.
 Studies must be of a relevant health economic study design (cost-utility analysis, cost-effectiveness analysis, cost-benefit analysis, cost-consequences analysis, comparative cost analysis).
 Studies must not be a letter, editorial or commentary, or a review of health economic evaluations. (Recent reviews will be ordered although not reviewed. The bibliographies will be checked for relevant studies, which will then be ordered.)
 Unpublished reports will not be considered unless submitted as part of a call for evidence. Studies must be in English.
A health economic study search will be undertaken using population-specific terms and a health economic study filter – see appendix B below.
Studies not meeting any of the search criteria above will be excluded. Studies published before 2001, abstract-only studies and studies from non-OECD countries or the US will also be excluded.
Each remaining study will be assessed for applicability and methodological limitations using the NICE economic evaluation checklist which can be found in appendix H of Developing NICE guidelines: the manual (2014). 115
Inclusion and exclusion criteria
• If a study is rated as both 'Directly applicable' and with 'Minor limitations', then it will be included in the guideline. A health economic evidence table will be completed and it will be included in the health economic evidence profile.
• If a study is rated as either 'Not applicable' or with 'Very serious limitations', then it will usually be excluded from the guideline. If it is excluded, then a health economic evidence table will not be completed and it will not be included in the health economic evidence profile.
• If a study is rated as 'Partially applicable', with 'Potentially serious limitations' or both, then there is discretion over whether it should be included.
Where there is discretion
The health economist will make a decision based on the relative applicability and quality of the available evidence for that question, in discussion with the guideline committee if required. The ultimate aim is to include health economic studies that are helpful for decision-making in the context of the guideline and the current NHS setting. If several studies are considered of sufficiently high applicability and methodological quality that they could all be included, then the health economist, in discussion with the committee if required, may decide to include only the most applicable studies and to exclude the remaining studies selectively. All studies excluded based on applicability or methodological limitations will be listed with explanation in the excluded health economic studies appendix below.

The health economist will be guided by the following hierarchies. *Setting:*

- UK NHS (most applicable).
- OECD countries with predominantly public health insurance systems (for example, France, Germany, Sweden).
- OECD countries with predominantly private health insurance systems (for example, Switzerland).
- Studies set in non-OECD countries or in the US will be excluded before being assessed for applicability and methodological limitations.

Health economic study type:

- Cost–utility analysis (most applicable).
- Other type of full economic evaluation (cost–benefit analysis, cost-effectiveness analysis, cost–consequences analysis).
- Comparative cost analysis.
- Non-comparative cost analyses including cost-of-illness studies will be excluded before being assessed for applicability and methodological limitations.

Year of analysis:

- The more recent the study, the more applicable it will be.
- Studies published in 2001 or later but that depend on unit costs and resource data entirely or predominantly before 2001 will be rated as 'Not applicable'.
- Studies published before 2001 will be excluded before being assessed for applicability and methodological limitations.

Quality and relevance of effectiveness data used in the health economic analysis:

 The more closely the clinical effectiveness data used in the health economic analysis match with the outcomes of the studies included in the clinical review the more useful the analysis will be for decision-making in the guideline.

Appendix B: Literature search strategies

The literature searches for this review are detailed below and complied with the methodology outlined in Developing NICE guidelines: the manual 2014, updated 2017

https://www.nice.org.uk/guidance/pmg20/resources/developing-nice-guidelines-the-manual-pdf-72286708700869

For more detailed information, please see the Methodology Review.

B.1 Clinical search literature search strategy

The search for this review was constructed using population terms. An excluded studies filter was applied where appropriate.

Table 6: Database date parameters and filters used

Database	Dates searched	Search filter used
Medline (OVID)	1946 – 03 July 2017	Exclusions
Embase (OVID)	1974 – 03 July 2017	Exclusions
The Cochrane Library (Wiley)	Cochrane Reviews to 2017 Issue 7 of 12 CENTRAL to 2017 Issue 6 of 12 DARE, and NHSEED to 2015 Issue 2 of 4 HTA to 2016 Issue 4 of 4	None

11 Medline (Ovid) search terms

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1.	exp Borrelia Infections/
2.	exp Lyme disease/
3.	Erythema Chronicum Migrans/
4.	(erythema adj3 migrans).ti,ab.
5.	lyme*.ti,ab.
6.	(tick* adj2 (bite* or bitten or biting or borne)).ti,ab.
7.	acrodermatitis chronica atrophicans.ti,ab.
8.	exp lxodidae/
9.	(borreliosis or borrelia* or neuroborreliosis or ixodid or ixodidae or ixodes or b burgdorferi or b afzelii or b garinii or b bissettii or b valaisiana or b microti).ti,ab.
10.	(granulocyctic anaplasmosis or babesia or babesiosis).ti,ab.
11.	or/1-10
12.	letter/
13.	editorial/
14.	news/
15.	exp historical article/
16.	Anecdotes as Topic/
17.	comment/
18.	(letter or comment*).ti.
19.	or/12-18
20.	randomized controlled trial/ or random*.ti,ab.
21.	19 not 20

22.	animals/ not humans/
23.	exp Animals, Laboratory/
24.	exp Animal Experimentation/
25.	exp Models, Animal/
26.	exp Rodentia/
27.	(rat or rats or mouse or mice).ti.
28.	or/21-27
29.	11 not 28
30.	limit 29 to English language

1 Embase (Ovid) search terms

1.	exp Borrelia Infection/
2.	exp Lyme disease/
3.	Erythema Chronicum Migrans/
4.	(erythema adj3 migrans).ti,ab.
5.	lyme*.ti,ab.
6.	(tick* adj2 (bite* or bitten or biting or borne)).ti,ab.
7.	acrodermatitis chronica atrophicans.ti,ab.
8.	exp lxodidae/
9.	(borreliosis or borrelia* or neuroborreliosis or ixodidae or ixodes or b burgdorferi or b afzelii or b garinii or b bissettii or b valaisiana or b microti).ti,ab.
10.	(granulocyctic anaplasmosis or babesia or babesiosis).ti,ab.
11.	or/1-10
12.	letter.pt. or letter/
13.	note.pt.
14.	editorial.pt.
15.	(letter or comment*).ti.
16.	or/12-15
17.	randomized controlled trial/ or random*.ti,ab.
18.	16 not 17
19.	animal/ not human/
20.	Nonhuman/
21.	exp Animal Experiment/
22.	exp Experimental animal/
23.	Animal model/
24.	exp Rodent/
25.	(rat or rats or mouse or mice).ti.
26.	or/18-25
27.	11 not 26
28.	limit 27 to English language

2 Cochrane Library (Wiley) search terms

#1.	MeSH descriptor: [Borrelia Infections] explode all trees	
#2.	MeSH descriptor: [Lyme Disease] explode all trees	
#3.	MeSH descriptor: [Erythema Chronicum Migrans] explode all trees	
#4.	(erythema near/3 migrans):ti,ab	

#5.	lyme*:ti,ab
#6.	(tick* near/2 (bite* or bitten or biting or borne)):ti,ab
#7.	acrodermatitis chronica atrophicans:ti,ab
#8.	MeSH descriptor: [Ixodidae] explode all trees
#9.	(borreliosis or borrelia* or neuroborreliosis or ixodidae or ixodes or ixodid or b burgdorferi or b afzelii or b garinii or b bissettii or b valaisiana or b microti):ti,ab
#10.	(granulocyctic anaplasmosis or babesia or babesiosis):ti,ab
#11.	#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10

B.2 Health Economics literature search strategy

Health economic evidence was identified by conducting a broad search relating to Lyme disease population in NHS Economic Evaluation Database (NHS EED – this ceased to be updated after March 2015) and the Health Technology Assessment database (HTA) with no date restrictions. NHS EED and HTA databases are hosted by the Centre for Research and Dissemination (CRD). Additional searches were run on Medline and Embase for health economics, economic modelling and quality of life studies.

Table 7: Database date parameters and filters used

Database	Dates searched	Search filter used
Medline	1946 – 03 July 2017	Exclusions Health economics studies Health economics modelling studies Quality of life studies
Embase	1974 – 03 July 2017	Exclusions Health economics studies Health economics modelling studies Quality of life studies
Centre for Research and Dissemination (CRD)	HTA - Inception – 03 July 2017 NHSEED - Inception to March 2015	None

Medline (Ovid) search terms

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1.	exp Borrelia Infections/
2.	exp Lyme disease/
3.	Erythema Chronicum Migrans/
4.	(erythema adj3 migrans).ti,ab.
5.	lyme*.ti,ab.
6.	(tick* adj2 (bite* or bitten or biting or borne)).ti,ab.
7.	acrodermatitis chronica atrophicans.ti,ab.
8.	exp Ixodidae/
9.	(borreliosis or borrelia* or neuroborreliosis or ixodid or ixodidae or ixodes or b burgdorferi or b afzelii or b garinii or b bissettii or b valaisiana or b microti).ti,ab.
10.	(granulocyctic anaplasmosis or babesia or babesiosis).ti,ab.
11.	or/1-10
12.	letter/

13.	editorial/
14.	news/
15.	exp historical article/
16.	Anecdotes as Topic/
17.	comment/
18.	(letter or comment*).ti.
19.	or/12-18
20.	randomized controlled trial/ or random*.ti,ab.
21.	19 not 20
22.	animals/ not humans/
23.	exp Animals, Laboratory/
24.	exp Animal Experimentation/
25.	exp Models, Animal/
26.	exp Rodentia/
27.	(rat or rats or mouse or mice).ti.
28.	or/21-27
29.	11 not 28
30.	limit 29 to English language
31.	Economics/
32.	Value of life/
33.	exp "Costs and Cost Analysis"/
34.	exp Economics, Hospital/
35.	exp Economics, Medical/
36.	Economics, Nursing/
37.	Economics, Pharmaceutical/
38.	exp "Fees and Charges"/
39.	exp Budgets/
40.	budget*.ti,ab.
41.	cost*.ti.
42.	(economic* or pharmaco?economic*).ti.
43.	(price* or pricing*).ti,ab.
44.	(cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab.
45.	(financ* or fee or fees).ti,ab.
46.	(value adj2 (money or monetary)).ti,ab.
47.	or/31-46
48.	exp models, economic/
49.	*Models, Theoretical/
50.	*Models, Organizational/
51.	markov chains/
52.	monte carlo method/

53.	exp Decision Theory/
54.	(markov* or monte carlo).ti,ab.
55.	econom* model*.ti,ab.
56.	(decision* adj2 (tree* or analy* or model*)).ti,ab.
57.	or/48-56
58.	quality-adjusted life years/
59.	sickness impact profile/
60.	(quality adj2 (wellbeing or well being)).ti,ab.
61.	sickness impact profile.ti,ab.
62.	disability adjusted life.ti,ab.
63.	(qal* or qtime* or qwb* or daly*).ti,ab.
64.	(euroqol* or eq5d* or eq 5*).ti,ab.
65.	(qol* or hql* or hqol* or h qol* or hrqol* or hr qol*).ti,ab.
66.	(health utility* or utility score* or disutilit* or utility value*).ti,ab.
67.	(hui or hui1 or hui2 or hui3).ti,ab.
68.	(health* year* equivalent* or hye or hyes).ti,ab.
69.	discrete choice*.ti,ab.
70.	rosser.ti,ab.
71.	(willingness to pay or time tradeoff or time trade off or tto or standard gamble*).ti,ab.
72.	(sf36* or sf 36* or short form 36* or shortform 36* or shortform36*).ti,ab.
73.	(sf20 or sf 20 or short form 20 or shortform 20 or shortform20).ti,ab.
74.	(sf12* or sf 12* or short form 12* or shortform 12* or shortform12*).ti,ab.
75.	(sf8* or sf 8* or short form 8* or shortform 8* or shortform8*).ti,ab.
76.	(sf6* or sf 6* or short form 6* or shortform 6* or shortform6*).ti,ab.
77.	or/58-76
78.	30 and 47
79.	30 and 57
80.	30 and 77

Embase (Ovid) search terms

1.	exp Borrelia Infection/
2.	exp Lyme disease/
3.	Erythema Chronicum Migrans/
4.	(erythema adj3 migrans).ti,ab.
5.	lyme*.ti,ab.
6.	(tick* adj2 (bite* or bitten or biting or borne)).ti,ab.
7.	acrodermatitis chronica atrophicans.ti,ab.
8.	exp lxodidae/
9.	(borreliosis or borrelia* or neuroborreliosis or ixodidae or ixodes or b burgdorferi or b afzelii or b garinii or b bissettii or b valaisiana or b microti).ti,ab.
10.	(granulocyctic anaplasmosis or babesia or babesiosis).ti,ab.

11.	or/1-10
12.	letter.pt. or letter/
13.	note.pt.
14.	editorial.pt.
15.	Case report/ or Case study/
16.	(letter or comment*).ti.
17.	or/12-16
18.	randomized controlled trial/ or random*.ti,ab.
19.	17 not 18
20.	animal/ not human/
21.	Nonhuman/
22.	exp Animal Experiment/
23.	exp Experimental animal/
24.	Animal model/
25.	exp Rodent/
26.	(rat or rats or mouse or mice).ti.
27.	or/19-26
28.	11 not 27
29.	limit 28 to English language
30.	health economics/
31.	exp economic evaluation/
32.	exp health care cost/
33.	exp fee/
34.	budget/
35.	funding/
36.	budget*.ti,ab.
37.	cost*.ti.
38.	(economic* or pharmaco?economic*).ti.
39.	(price* or pricing*).ti,ab.
40.	(cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab.
41.	(financ* or fee or fees).ti,ab.
42.	(value adj2 (money or monetary)).ti,ab.
43.	or/30-42
44.	statistical model/
45.	exp economic aspect/
46.	44 and 45
47.	*theoretical model/
48.	*nonbiological model/
49.	stochastic model/
50.	decision theory/

51.	decision tree/
52.	monte carlo method/
53.	(markov* or monte carlo).ti,ab.
54.	econom* model*.ti,ab.
55.	(decision* adj2 (tree* or analy* or model*)).ti,ab.
56.	or/46-55
57.	quality adjusted life year/
58.	"quality of life index"/
59.	short form 12/ or short form 20/ or short form 36/ or short form 8/
60.	sickness impact profile/
61.	(quality adj2 (wellbeing or well being)).ti,ab.
62.	sickness impact profile.ti,ab.
63.	disability adjusted life.ti,ab.
64.	(qal* or qtime* or qwb* or daly*).ti,ab.
65.	(euroqol* or eq5d* or eq 5*).ti,ab.
66.	(qol* or hql* or hqol* or h qol* or hrqol* or hr qol*).ti,ab.
67.	(health utility* or utility score* or disutilit* or utility value*).ti,ab.
68.	(hui or hui1 or hui2 or hui3).ti,ab.
69.	(health* year* equivalent* or hye or hyes).ti,ab.
70.	discrete choice*.ti,ab.
71.	rosser.ti,ab.
72.	(willingness to pay or time tradeoff or time trade off or tto or standard gamble*).ti,ab.
73.	(sf36* or sf 36* or short form 36* or shortform 36* or shortform36*).ti,ab.
74.	(sf20 or sf 20 or short form 20 or shortform 20 or shortform20).ti,ab.
75.	(sf12* or sf 12* or short form 12* or shortform 12* or shortform12*).ti,ab.
76.	(sf8* or sf 8* or short form 8* or shortform 8* or shortform8*).ti,ab.
77.	(sf6* or sf 6* or short form 6* or shortform 6* or shortform6*).ti,ab.
78.	or/57-77
79.	29 and 43
80.	29 and 56
81.	29 and 78

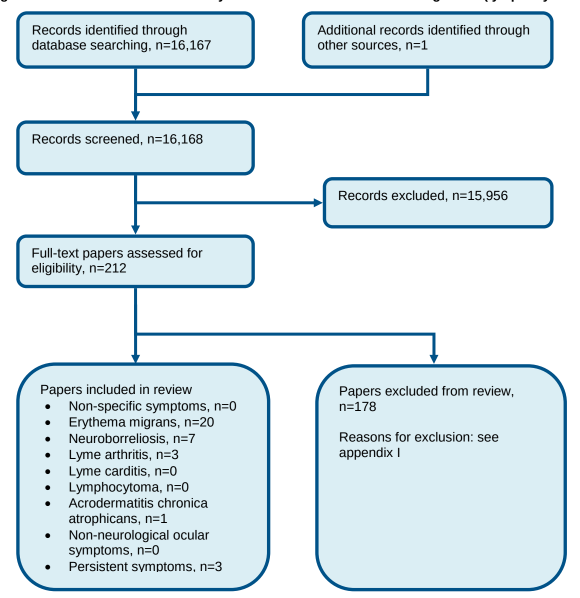
NHS EED and HTA (CRD) search terms

	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
#1.	MeSH DESCRIPTOR Borrelia Infections EXPLODE ALL TREES IN NHSEED,HTA
#2.	MeSH DESCRIPTOR Erythema Chronicum Migrans EXPLODE ALL TREES IN NHSEED,HTA
#3.	((erythema adj3 migrans)) IN NHSEED, HTA
#4.	(lyme*) IN NHSEED, HTA
#5.	((tick* adj2 (bite* or bitten or biting or borne))) IN NHSEED, HTA
#6.	(acrodermatitis chronica atrophicans) IN NHSEED, HTA
#7.	MeSH DESCRIPTOR Ixodidae EXPLODE ALL TREES IN NHSEED,HTA
#8.	((borreliosis or borrelia* or neuroborreliosis or ixodidae or ixodes or b burgdorferi or b afzelii or b garinii or b bissettii or b valaisiana or b microti)) IN NHSEED, HTA
#9.	((granulocyctic anaplasmosis or babesia or babesiosis)) IN NHSEED, HTA
#10.	MeSH DESCRIPTOR Lyme Disease EXPLODE ALL TREES IN NHSEED,HTA

#11. #1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10

2 Appendix C: Clinical evidence selection

Figure 1: Flow chart of clinical study selection for the review of management (lymphocytoma)

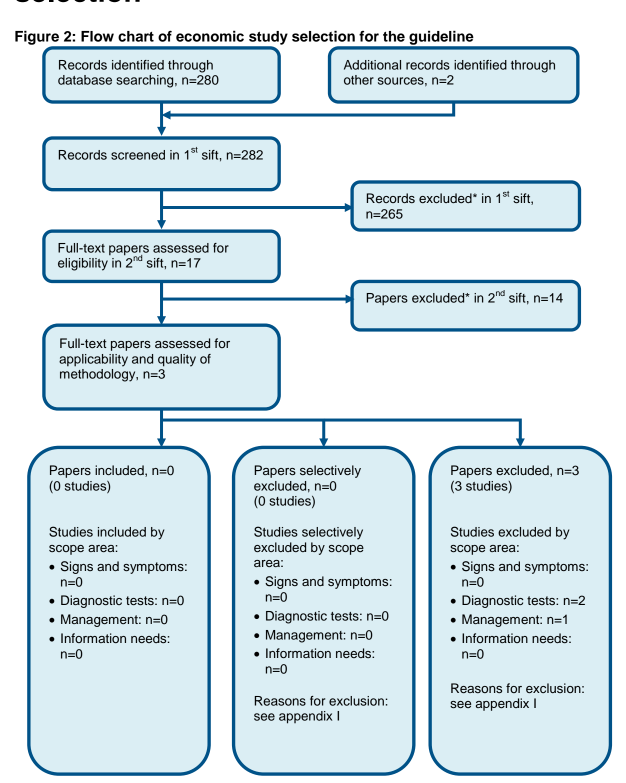


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Appendix E: Forest plots

Appendix F:GRADE tables

Appendix G: Health economic evidence selection



^{*} Non-relevant population, intervention, comparison, design or setting; non-English language

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Appendix H: Health economic evidence tables

Appendix I: Excluded studies

I.1 Excluded clinical studies

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Table 8: Studies excluded from the clinical management reviews

lable 8: Studies excluded from the clinica	
Reference	Reason for exclusion
Aberer 2006 ¹	Excluded due to an incorrect intervention
Abrutyn 1989 ²	Excluded due to an incorrect study design
Agger 1992 ³	Excluded due to an incorrect study design
Agus 1995 ⁴	Excluded due to an incorrect study design
Agwuh 2006 ⁵	Excluded due to an incorrect study design
Ahmed 2005 ⁶	Excluded due to an incorrect study design
Ahmed 2013 ⁷	Excluded due to an incorrect study design
Alarcon 1994 ⁸	Excluded due to an incorrect study design
Andiman 1986 ⁹	Excluded due to an incorrect study design
Anonymous 1991 ¹⁰	Excluded due to an incorrect study design
Arvikar 2015 ¹¹	Excluded due to an incorrect study design
Auwaerter 2004 ¹²	Excluded due to an incorrect study design
Bennet 2003 ¹³	Excluded due to an incorrect study design
Berende 2014 ¹⁴	Excluded due to an incorrect study design
Berger 1988 ¹⁶	Excluded due to an incorrect study design
Berger 1986 ¹⁵	Excluded due to an incorrect study design
Bernardino 2009 ¹⁷	Excluded due to an incorrect study design
Bhate 2011 ¹⁸	Excluded due to an incorrect study design
Bjark 2016 ¹⁹	Not available
Borg 2005 ²²	Excluded due to an incorrect study design
Bratton 2008 ²³	Excluded due to an incorrect study design
Bremell 2014 ²⁴	Excluded due to an incorrect study design
British Infection Association 2011 ²⁵	Excluded due to an incorrect study design
Butler 1978 ²⁶	Excluded due to an incorrect population
Cadavid 2016 ²⁷	Excluded due to an incorrect study design
Canadian Paediatric Society 1992 ²⁸	Excluded due to an incorrect study design
Chen 1999 ³⁰	Excluded due to an incorrect outcome
Choo-Kang 2010 ³¹	Excluded due to an incorrect study design
Christian 1992 ³²	Excluded due to an incorrect study design
Cimmino 1992 ³⁴	Excluded due to an incorrect study design
Cimmino 1997 ³³	Excluded due to an incorrect study design
Cimperman 1999 ³⁵	Excluded due to an incorrect study design
Coblyn 1981 ³⁶	Excluded due to an incorrect study design
Committee on Infectious Diseases 1991 ³⁸	Excluded due to an incorrect study design
Cuisset 2008 ³⁹	Excluded due to an incorrect study design
Dattwyler 1996 ⁴¹	Excluded due to an incorrect comparison
Dattwyler 1987 ⁴²	Excluded due to an incorrect study design
Dattwyler 1988 ⁴³	Excluded due to an incorrect population
Dattwyler 2005 ⁴⁴	Excluded due to an incorrect population

Reference	Reason for exclusion
Dersch 2015 ⁴⁶	Excluded due to an incorrect study design
Dersch 2016 ⁴⁹	Excluded due to an incorrect study design
Dersch 2014 ⁴⁷	Excluded due to an incorrect study design
Dersch 2017 ⁴⁸	Not available
Dhoot 2011 ⁵⁰	Excluded due to an incorrect study design
Dinser 2005 ⁵¹	Excluded due to an incorrect study design
Dotevall 1988 ⁵²	Excluded due to an incorrect study design
Eliassen 2017 ⁵³	Excluded due to an incorrect study design
Eliassen 2017 ⁵⁴	Excluded due to an incorrect intervention
Eppes 2003 ⁵⁵	Excluded due to an incorrect study design
Esposito 2013 ⁵⁶	Excluded due to an incorrect study design
Fallon 1999 ⁵⁸	Excluded due to an incorrect intervention
Fallon 2008 ⁵⁷	Excluded due to an incorrect outcome
Galev 2005 ⁵⁹	Excluded due to an incorrect study design
Garkowski 2017 ⁶⁰	Systematic review
Gasser 1996 ⁶²	Excluded due to an incorrect not available
Gasser 1995 ⁶³	Excluded due to an incorrect study design
Gasser 1995 ⁶¹	Excluded due to an incorrect study design
Gerber 1996 ⁶⁴	Excluded due to an incorrect intervention
Gillies 2015 ⁶⁵	Excluded due to an incorrect study design
Goodwin 1990 ⁶⁶	Excluded due to an incorrect study design
Hansen 1992 ⁶⁷	Excluded due to an incorrect intervention
Hassler 1990 ⁶⁸	Excluded due to an incorrect population
Horton 2017 ⁶⁹	Conference abstract
Hu 2001 ⁷⁰	Excluded due to an incorrect study design
Inboriboon 2010 ⁷¹	Excluded due to an incorrect study design
Kaplan 2003 ⁷²	Excluded due to an incorrect population
Karkkonen 2001 ⁷³	Excluded due to an incorrect study design
Karlsson 1996 ⁷⁴	Excluded due to an incorrect outcome
Kersten 1995 ⁷⁵	Excluded due to an incorrect study design
Kilic Muftuoglu 2016 ⁷⁶	Excluded due to an incorrect study design
Klempner 2013 ⁷⁸	Excluded due to an incorrect study design
Korenberg 1996 ⁷⁹	Excluded due to an incorrect intervention
Kowalski 2010 ⁸¹	Excluded due to an incorrect outcome
Kowalski 2011 ⁸⁰	Excluded due to an incorrect study design
Krbkova 1996 ⁸²	Excluded due to an incorrect comparison
Kuhn 2012 ⁸³	Excluded due to an incorrect study design
Laasila 2003 ⁸⁴	Excluded due to an incorrect population
Lantos 2013 ⁸⁵	Excluded due to an incorrect study design
Lauhio 1994 ⁸⁶	Excluded due to an incorrect population
Lauhio 1991 ⁸⁷	Excluded due to an incorrect population
Lempner 2002 ⁷⁷	Excluded due to an incorrect study design
Liegner 1992 ⁸⁸	Excluded due to an incorrect study design
Lipsker 2002 ⁸⁹	Excluded due to an incorrect study design
Ljostad 2008 ⁹⁰	Study abstract

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Peltomaa 1998 ¹²⁵ Excluded due to an incorrect comparison
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Pena 1999 ¹²⁶ Excluded due to an incorrect study design
Perronne 2015 ¹²⁷ Not available
Pfister 1988 ¹²⁸ Excluded due to an incorrect outcome
Pirila 1951 ¹³¹ Excluded due to an incorrect study design
Plorer 1993 ¹³² Excluded due to an incorrect study design
Plotkin 1991 ¹³³ Excluded due to an incorrect study design
Puchalska 1996 ¹³⁴ Excluded due to an incorrect study design
Puri 2015 ¹³⁵ Excluded due to an incorrect comparison
Puri 2015 ¹³⁶ Excluded due to an incorrect study design
Rebman 2015 ¹³⁷ Excluded due to an incorrect study design
Renaud 2004 ¹³⁸ Excluded due to an incorrect study design
Rohacova 1996 ¹³⁹ Excluded due to an incorrect comparison
Rose 1994 ¹⁴⁰ Excluded due to an incorrect study design

Reference	Reason for exclusion
Rose 1996 ¹⁴¹	Excluded due to an incorrect intervention
Rubin 1992 ¹⁴²	Excluded due to an incorrect study design
Salazar 2005 ¹⁴³	Excluded due to an incorrect intervention
Salazar 1993 ¹⁴⁴	Excluded due to an incorrect study design
Sanchez 2016 ¹⁴⁵	Excluded due to an incorrect study design
Sandstrom 1989 ¹⁴⁶	Excluded due to an incorrect study design
Schmidt 1995 ¹⁴⁷	Excluded due to an incorrect study design
Selby 2008 ¹⁴⁸	Excluded due to an incorrect study design
Shadick 1994 ¹⁴⁹	Excluded due to an incorrect study design
Shadick 1999 ¹⁵⁰	Excluded due to an incorrect study design
Shemenski 2016 ¹⁵¹	Excluded due to an incorrect study design
Shoemaker 2006 ¹⁵²	Excluded due to an incorrect intervention
Sjowall 2012 ¹⁵⁴	Excluded due to an incorrect intervention
Sjowall 2011 ¹⁵³	Excluded due to an incorrect study design
Skogman 2003 ¹⁵⁶	Excluded due to an incorrect intervention
Skogman 2008 ¹⁵⁵	Excluded due to an incorrect study design
Skoldenberg 1988 ¹⁵⁷	Excluded due to an incorrect study design
Smith 2002 ¹⁵⁸	Excluded due to an incorrect study design
Solomon 1998 ¹⁵⁹	Excluded due to an incorrect intervention
Spathling 1992 ¹⁶⁰	Article not in English
Stanek 1999 ¹⁶¹	Excluded due to an incorrect study design
Steere 1980 ¹⁶⁵	Excluded due to an incorrect study design
Steere 1983 ¹⁶⁶	Excluded due to an incorrect study design
Steere 1987 ¹⁶²	Excluded due to an incorrect study design
Steurer 2016 ¹⁶⁷	Article not in English
Stricker 2011 ¹⁶⁸	Excluded due to an incorrect study design
Stricker 2010 ¹⁶⁹	Excluded due to an incorrect study design
Strle 1996 ¹⁷⁰	Excluded due to an incorrect outcome
Strle 1996 ¹⁷¹	Excluded due to an incorrect outcome
Strle 1992 ¹⁷²	Excluded due to an incorrect study design
Strle 1993 ¹⁷³	Excluded due to an incorrect outcome
Stupica 2015 ¹⁷⁵	Excluded due to an incorrect comparison
Stupica 2011 ¹⁷⁴	Excluded due to an incorrect comparison
Suarez-Magdalena 2017 ¹⁷⁶	Not available
Thompson 2012 ¹⁷⁷	Excluded due to an incorrect study design
Thorstrand 2002 ¹⁷⁸	Excluded due to an incorrect study design
Thyresson 1949 ¹⁷⁹	Excluded due to an incorrect study design
Torbahn 2016 ¹⁸⁰	Excluded due to an incorrect study design
Tory 2010 ¹⁸¹	Excluded due to an incorrect comparison
Tseng 2017 ¹⁸²	Excluded due to an incorrect outcome
Valesova 1996 ¹⁸³	Excluded due to an incorrect comparison
Vazquez 2003 ¹⁸⁵	Excluded due to an incorrect study design
Vazquez-Lopez 2016 ¹⁸⁴	Excluded due to an incorrect study design
Wahlberg 1994 ¹⁸⁶	Excluded due to an incorrect intervention
Weber 1988 ¹⁸⁸	Excluded due to an incorrect study design

Reference	Reason for exclusion
Weber 1987 ¹⁸⁷	Excluded due to an incorrect population
Weissenbacher 2005 ¹⁸⁹	Excluded due to an incorrect intervention
White 2013 ¹⁹⁰	Excluded due to an incorrect study design
Zochling 1996 ¹⁹¹	Excluded due to an incorrect study design

I.2 Excluded health economic studies

2 Table 9: Studies excluded from the health economic review

Reference	Reason for exclusion
None	None