Comments Quality of the (GRADE) BOOOD WEIZY BOWE 12 145 (2 studies) 145 (2 studies) RR 1.3 (0.65 to 2.62) RR 1.5 (0.43 to 5.26) Corresponding risk Single-dose * (95% CI) 260 per 1000 (130 to 524) 318 per 1900 (159 to 642) Single class werean short-cense (3-7 days) antibiods

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Unary ruce infection (UTI) is one of the moor common bacterial infections in infama and children and the moor common bacterial infections in infama and children and the moor common bacterial infections in infama and children and the moor common bacterial infections in infama and children and infection common bacterial infection of the universal presents in intervery are contributed in the universal properties of the universal infection of the kidneys (replacephoins or upper UTI). However, the commonly presenting and in the majority of case can be easily irrated with a course of analohoric theory of case can be easily irrated with a course of analohoric theory of case can be easily irrated with a course of analohoric theory of case can be easily irrated with a course of analohoric theory of case can be easily irrated with a course of analohoric theory of case can be easily irrated with a course of analohoric theory of case can be easily irrated with a course of analohoric theory of case can be easily irrated with a course of analohoric theory of the complete of the course of the

are not underprined by mong evidence and are largely the results of cidical judgment and the hological plantality of funet side of cidical judgment and the hological plantality of funet side. The difficult we obtain accurate estimates of the number of infinite and children who will percent with a lover UTI during childhood. Contract practice in more contract in best of the contract practice in more contract in best of the contract practice. The first of the proposal contract practice in more contract in best of the contract practice. The first own proposal contract practice in the contract practice. The first own proposal contract practice. The first own proposal contract practices are contracted as a contract practice. The first own proposal contract practices are contracted as a contract practice. The first own proposal contract practices are contracted as a contract practice. The contract practices are contracted as a contract practice and the practices are contracted as a contract practice. The contract practices are contracted as a contract practice and the contract practices are contracted as a contract practice. The contract practices are contracted as a contract practice and the contract practices are contracted as a contract practice. The contract practices are contracted as a contract practice and the contract practices are contracted as a contract practice and the contract practices are contracted as a contract practice and the contract practices are contracted as a contract practice and the contract practices are contracted as a contract practice and an advantage of the contract practices are contracted as a contract practice and an advantage of the contract practices are contracted as a contract practice and an advantage of the contract practices are contracted as a contract practice and an advantage of the contract practices are contracted as a contract practice and an advantage and a contract practices are contracted as a contract practice and an advantage and a contract practices are contracted

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- Esclusion criteria

 Chilidre hospitalised for a condition not related to UTI.

 Chilidres with bacteriologically proven UTI and apmynema
 rei gued organization of the principal control principal

- sypes of unterventions

 Antibiotic therapy (tundad course) versus placebo, no therapy, a different antibiotic or alternative non-antibiotic therapy.

 Single-dose (or single-day therapy) versus standard dose. Mode of administration (oral, intravenous or intramuscular)

Electronic searches
The sards traregy was comprehensive and was designed to correct two review being undertaken by the authors, this review and "Insert two reviews being undertaken by the authors, this review and "Insert two reviews being undertaken by the authors, this review and "Insert two reviews being undertaken by the authors that review and "Insert two reviews being undertaken by the authors that review and "Insert two reviews being under the reviews that the standard was a third authors."

Fall details of the search starting are reported in Approximation, the Real Colonger Beautiful Colo

- 1. Quartely tastehen of the Cochana Central Register of Convoiled Tails CENTRAL
 2. Weely searches of MEDINE OVID SP,
 3. Handasseafing of restar-leads possion at the proceedings of major renal conferences.
 4. Searching of the current year of EMBASE OVID SP.
 5. Weely current awareness alres for selected renal-ploratels.
 Conscience of the Immensional Clinical Trials Register
 Conscience of the Immensional Clinical Trials Register
 Conscience of the Immensional Clinical Trials Register
 Studies considered in the specialised register are identified through
 sevent arranging for ECETRAL AMEDINA, and EMBASE Stude
 on the scope of the Cochana Renal Group. Details of these stratepier as well as a list of hundrearched pounds, conference proceedings and current awareness alors are available in the Specialised Register's seeion of information about the Cechrane
 Renal Group.

- Searching other resources

 Reference lists of includogy testbooks, review articles and relevant studies.

 Letters seeking information about unpublished or incomplete studies to investigates shown to be involved in pre

Types of outcome measures

Proisines approars a completion of resument.

Proisines approars a completion of resument.

Proisines. Spinional societaria (> 10³ cfu/mL) and pre-four studies.

Combinations of persistent bacteriaria (> 10³ cfu/mL) and yampinens at completion of resument.

Recurrent symponatic CTTI following measurem.

Provident approaria CTTI following measurem.

Any rend participant diamage on DMSA, four to set to the completion.

Adverse creats.

Adverse creats.

Any changes to antibiotic regimen.

Data cutteraction and management which studies satisfied the inclusion of the completion.

Any changes to antibiotic regimen.

Data cutteraction and management control of the completion of the The rearch strategy described was used to obtain titles and ab-stracts of studies relevant to the review. The titles and abstracts were acreened independently by two authors assessed to determine which studies satisfied the inclusion criteria.

Data extraction and management.
Data extraction was carried our independently by the same suthern using randed data extraction forms. Studier reported in
non-English language journals were translated before assuments.
Where more than one publication of one much existed, reports
were to be grouped together and the most recent or most complete
dataset used. Any descriptory between published versions was to
be highlighted. Diagreements were resolved in consultration with
a third arther.

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Bata synthesis

Data synthesis

Data with continued aurona

Processing the study (desection bia)?

Practicipans and personnel

Outcome aurona

Eventual bian?

Are reperted of the subje (see of sugaration of selective outcome reporting (reporting bias)?

Was the nody appreciably red of char problems that could put it at a risk of bian?

Measures of treatment effect

For dichosomous outcomes (e.g. proponn resolution, persistent bacterius), securem infection) results were expressed at risk ratio of Buscerius's, securem infection fresults were expressed at risk ratio of the secure of th

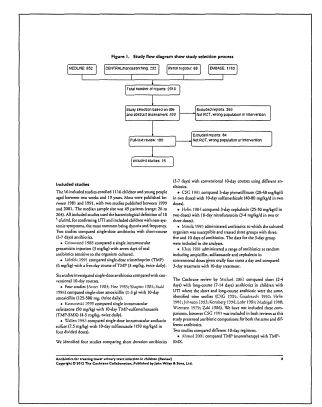


Figure 3. Risk of bias summary: review authors' judgements about each risk of bias item for each included study. Ahmed 2001 7 7 🕏 🚳 🚳 Fine 1985 7 7 7 9 9 9 Grimwood 1988 8 7 7 8 6 Helin 1984 (2) (2) (3) 🛞 🛞 Khan 1981 😗 😗 😮 🛞 Komoroski 1999 🐮 🈗 😝 😗 🔻 Lidefelt 1991 3 2 3 3 6 Kalaka-Zafirul 1984 😗 😗 😵 🍪 Mitnik 1985 (?) (?) (?) 🚱 🚱 Principi 1990 7 2 7 8 8 Sanchez 1990 7 7 7 7 7 Shapiro 1981 7 7 8 7 Stahl 1984 30 30 30 68 3 Wallen 1983 🛞 😗 😗 🚳 🏶 Antibiotics for treating lower urinary tract infection in children (Review)
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Random sequence generation

The quality of reporting of random sequence generation was poor.

The rounders reported using random numbers table to generate random sequence (Coincond) 1998. With 1999 and 1998 and 1998 and 1999 and

- in complete outcome data

 In two studies it appeared that children were randomized to treatment before inclusion and exclusion criteria were applied.

 SCS 1991; 2606 diddient annobiated sevents to follow-up for variety of reasons including nor falfilling inclusion criteria transmet discontinued before streed to the follow-up for a variety of reasons including nor falfilling inclusion criteria transmet discontinued before sheedheld and did not have repear units cultures within the allocated diments and because of the small number and because they were not evenly distributed between 10 to 30 days.

Sanchez 1990 was presented in an abstract and it was not clear whether patients were analysed in groups to which they were randomised.
 In the remaining eight studies, all patients were analysed in groups to which they were randomised.

Alleading of Machine months was very poor. CSG [59]

Effects of interventions.

Examples of interventions.

Examples of interventions.

Effects of interventions.

Examples of interventions.

Examples of interventions.

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Examples of interventions.

Example

small number and Secuse they were not evenly distributed between groups.

• Komosculi 1999: 37% of the chaldren randomised were between groups.

• Komosculi 1999: 37% of the chaldren randomised were between the properties of the properties described by the properties of the propert

Single-dose versus short-course (3-7 days) treatment

Souches 1990 randomied children to one of five antibiotics:
amosicillis: mencellin e seturalusi exide epidaterin. TMPs or
continuated. Beautie of the multi amoster of puricipeus in
the continuated of the multisetural of puricipeus in
the continuated. Beautie of the multisetural of puricipeus in
the continuated of the continuate

TMP (10 days) versus TMP-SMX (10 days)

Single-dose versus short-course. (17-d sps) treatment.

There was no significant difference in person 10-day TMP restriction in the cut on 10-day TMP restriction 10-day TMP restrictio

Short-course (2.7 days) versus long-course (7-10 days)
Treatment

There was no significant difference in persistent because in level several solution course of long-course ankiesis creaments after some significant difference in persistent proposes because and course of long-course ankiesis creaments after some series of long-course ankiesis creaments after some series (1.6 days) versus amplicities (16 days)

There was no significant difference in persistent proposes because and considerable and

Head-to-head studies

There were no significant difference in persistent bacterium because in the control of th

Single-dose versus conventional 16-day treatment
There studies reported outcomes for persistent symptoms at Completion of treatment
There studies reported outcomes for persistent symptoms following resonant and showed no differences between trated and untened groups; all for different ambioric comparisons and durations.

Single-dote versus conventional 10-day treatment
There was no significant difference between single-dote treatment
compared with conventional 10-day treatment (Analysis 2.2 (4 and 6.2), 26 diletters, Ret. 25, 59% C. O.24 o. 21, 18 p. v. 98).

2.8 diletters, Ret. 25, 59% C. O.24 o. 21, 18 p. v. 98).

2.8 diletters Ret. 25, 59% C. O.24 o. 21, 18 p. v. 98).

2.8 diletters Ret. 25, 59% C. O.24 o. 21, 18 p. v. 98).

2.9 diletters Ret. 25, 59% C. O.24 o. 21, 18 p. v. 98).

2.1 four surface compared 3-day to 10-day treatment. Of the series of the single-dote group land pressured symposium compared with 21/10-5 (12.9) of the 10-dot-come group land reconsistent compared with 21/10-5 (12.9) of the long-context compared with 21/10-5 (12.9) of the 10-dot-context compared with 21/10-5 (12.9) of the 10-dot-context context compared with 21/10-5 (12.9) of the 10-dot-context context context compared with 21/10-5 (12.9) of the 10-dot-context context conte

- Mead-to-hand studies

 There were no applicant difference in recurrent symptomatic
 Theorems.

 *TMR 10 days) versum TMP-SMX (10 days) (Analysis 4), 51(1)
 The studies of the

Single-dose versus short-course (1-7 days) treatment
There was no significant difference between single-dose compared
with short-course treatment (Analysis 2.3 (1 study, 45 children);
RR (0.16, 59% Cl.00) to 1.20, shows 1.125 (49%) of the short-course group had a re-infection compared with 5/25 (20%) of the
short-course group.

Adverse conts

Short-course (1-7 days) versus long-course (7-10 days) creatment
There was no significant difference between thort-course models defect (1-8, 12) and the second of the s

and this 1910 CH to the size of the beginning to go and a reference of the property of the beginning and the 1910 CH (1910) one child andomized to single-door. TMP expected varieties of the included unline of the proposed product of the property of the p

Re-infection following treatment

There studies reported outcomes for re-infection (with a different experted outcomes for re-infection (with a different experted) of the wind in the control of t

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Local discomfort from injection sizes was reported in revenuable (Komenetek 1979); Principe (1970).

 State effects were not reprorted in its mades (Grimwood 1988; Klam 1981; Marmil: 1985; Sanche: 1990; Shapiro 1981; Shahim 1981; Marmil: 1985; Sanche: 1990; Shapiro 1981; Sh

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- 66T -

** Very small numbers of patients (50)

**Politics budy reported allocation concealment. One reponds using a random numbers table, the other reported blinding. Neither study used TT analysis.

**Summer of participants is small, <25 in each group across both studies. CI are wide and include 1.

**Summer of participants is small, <25 in each group across both studies. CI are wide and include 1.

**Commerced participants is small, <25 in each group across both studies. CI are wide and include 1.

**Commerced participants is small, <25 in each group across both studies. CI are wide and include 1.

Short (3-7 days) versus long-course (10-14 days) antibiotics for treating lower urinary tract infection in children Patient or population: children with lower urinary tract infection Settings: paediatric department (1); not stated (3) Intervention: short-course (3-7 days)
Comparison: long-course (10-14 days) Illustrative comparative risks* (95% CI) Quality of the evidence Comments (GRADE) Assumed risk Corresponding risk Long-course (10-14 Short-course (3-7 days) RR 1.1 " (0.68 to 1.77) Persistent bacteriuria Study population 265 (3 studies) 186 per 1000 205 per 1000 (126 to 329) 204 per 1000 (126 to 327) 185 per 1000 Study population 363 (4 studies) GOOO very low3.4 (0.7 to 1.86) 127 per 1000 145 per 1000 (89 to 236) Medium risk population 100 per 1000 114 per 1000 (70 to 186)

ADDITIONAL SUMMARY OF FINDINGS (Explanation) Single-dose versus conventional 10-day antibiotic treatment for treating lower urinary tract infection in children Patient or population: children with lower uninary tract infection Settings: outpatient and/or emergency department intervention: single-dose Comparison: conventional 10-day treatment illustrative comparative risks* (95% CI) Assumed risk Corresponding risk Conventional 10-day Single-dose treatment Persistent bacteriuria Study population RR 2.01 (1.06 to 3.8) 228 (6 studies) 104 per 1000 209 per 1000 (110 to 395) Medium risk population 126 per 1000 253 per 1000 (134 to 479) Persistent symptoms Study population RR 0.29 (0.03 to 2.5) 30 (1 study) #200 very low3.4 214 per 1000 62 per 1000 (6 to 535) Medium risk population 214 per 1000 62 per 1000 (6 to 535)

	Study population		RR 1.83	46	⊕○○○ very low ^{5,6}
and symptoms	45 per 1000	82 per 1000 (8 to 848)	(0.18 to 18.84)	(1 study)	very low ^{2,4}
	Medium risk popula	illon			
	46 per 1000	84 per 1000 (8 to 867)			
Recurrence	Study population		RR 1.38	79	⊕000
	158 per 1000	218 per 1000 (87 to 553)	(0.55 to 3.5)	(2 studies)	very low ^{7,8}
	Medium risk population				
	154 per 1000	213 per 1000 (85 to 539)			
Moderate quality: Further	earch is very unlikely to r research is likely to h arch is very likely to h	ave an important impact on	our confidence in the esti	mate of effect and may change t nate of effect and is likely to cha	
reported ITT analyses.	O across all groups	od; no study reported alloca		lies reported blinding; 1/6 studie	s

- 200 **-**

	0 per 1000	0 per 1000 (0 to 0)	
	Medium risk popu	fallon	
	0 per 1000	0 per 1000 (0 to 0)	
*The basis for the a assumed risk in the o GI; Confidence interv	comparison group and the	edian control group risk acro e relative effect of the interve	ss studies) is provided in footnotes. The corresponding risk (and its 95% confidence interval) is based on th ndon (and its 95% CQ.
High quality: Further Moderate quality: Fi Low quality: Further	urther research is likely to research is very likely to 'e are very uncertain abou	have an important impact on I the estimate.	our confidence in the estimate of effect and may change the estimate. our confidence in the estimate of effect and is likely to change the estimate.
10%		alment were not reported. Inv s very wide and crosses 1	estigator blind only. No ITT analysis and loss to follow-up >
10%		•	stigator bind day, to II I analysis and loss to telew-up >
10%		•	sagator blind only. No II I analysis and loss to tolery-up >
10%		•	stigator blind dely, fo II I analysis and loss to telew-up >

10-day celadroxil versu	ıs 10-day amplcillin for I	treating lower urinary tract	infection in children			
Patient or population: c Settings: not stated Intervention: 10-day cel Comparison: 10-day arr	fadroxii	r tract infection				
Outcomes	Illustrative comparati	ive risks* (95% CI)	Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk	Ī			
	10-day ampicillin	10-day cefadroxil				
ersistent bacteriuria Study population		RR 0.33 (0.01 to 7.62)	32 (1 study)	⊕○○○ very low!.2		
	62 per 1000	21 per 1000 (1 to 480)	(0.01 to 7.02)	(i swoy)	very low	
	Medium risk population					
	63 per 1000	21 per 1000 (1 to 450)				
Persistent symptoms	Study population		RR 0.33 (0.01 to 7.62)	32 (1 study)	⊕○○○ very low ^{1,2}	
	62 per 1000	21 per 1000 (1 to 480)	(0.01 to 1.02)	(T Saloy)	very low	
	Medium risk populati	on				
	63 per 1600	21 per 1000 (1 to 480)				

147 per 1000 129 per 1000 (55 to 256) Molicum risk population 154 per 1000 136 per 1000 (68 to 266)	Re-infection	Study population		RR 0.88	211	⊕ 000	
154 per 1000 (55 to 265) 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 assumed risk in the compations group and the relative effect of the intervention (and its 95% CD). Cit. Confidence interval. RR: Risk ratio and the relative effect of the intervention (and its 95% CD). Cit. Confidence interval. RR: Risk ratio and the relative effect of the intervention (and its 95% CD). Cit. Confidence interval. RR: Risk ratio and the relative effect of the intervention (and its 95% CD). Cit. Confidence interval. RR: Risk ratio and the relative effect of the estimate of effect. Risk parallel further research is key not enhanced and the estimate. Low quality. Further research is key view laby by have an important impost on our confidence in the estimate of effect and is likely to change the estimate. Low quality. Further research is key view laby by have an important impost on our confidence in the estimate of effect and is likely to change the estimate. Low quality. Further research is key view laby by have an important impost on our confidence in the estimate of effect and is likely to change the estimate. Very low waster, low was very incertain about the estimate. No study reported blinding. One study was quasi-RCT exing alternation, the other two studies did not report randomisation method. One study reported blinding, One study was quasi-RCT exing alternation, the other two studies did not report randomisation method. One study reported blinding over group and recognition of an wide and cross 1 **Parallel RCT analysis was adequate in one of		147 per 1900		(0.44 to 1.74)	(2 studies)	very low ⁴⁻⁵	
(68 to 268) The basis for the assumed risk (e.g., the median control group risk across studies) is provided in footholes. The corresponding risk (and its 95% confidence interval) is based assumed risk the companions group and the retain's effect of the Intervention (and its 95% CD). Cl. Confidence interval. RR: Risk risk in CDO-CONTROL (and its 95% CD). Cl. Confidence interval. RR: Risk risk risk in the CDO-CONTROL (and its 95% CD). Cl. Confidence interval. RR: Risk risk risk in the CDO-CONTROL (and its 95% CD). Cl. Confidence interval risk rey uniformly to change are confidence in the estimate of effect. Risk grain (in matter cases in its lay to low an important inprove or or confidence in the estimate of effect and is likely to change the estimate. Low quality Firther research is lay by the lay to have an important inprove or our confidence in the estimate of effect and is likely to change the estimate. Low quality Firther research is lay we likely to have an important import on our confidence in the estimate of effect and is likely to change the estimate. Low quality Firther research is lay we should be estimate. No study reported blinding. One study was quasi-RCT esting alternation; the other two studies did not report randomisation method. One study reported blinding. One study was quasi-RCT esting alternation; the other two studies did not report randomisation method. One study reported blinding one study was quasi-RCT esting alternation; the other two studies did not report randomisation method. One study reported blinding one study was quasi-RCT esting alternation; the other two studies did not report randomisation method. One study reported blinding one study was quasi-RCT analyses. Province of the companion of the study o		Medium risk popula	tion				
assumed risk in the comparison group and the relative effect of the Intervention (and its 05% Ci). GRADEC Workship Group grades of ediforce Hold reality facility relative research is labely to change our confidence in the estimate of effect. Moderate quality. Future research is labely to thange our confidence in the estimate of effect and may change the estimate. Moderate quality, refure research is labely to thange our confidence in the estimate of effect and may change the estimate. Very low quality, which we are important import on our confidence in the estimate of effect and sitely to change the estimate. Very low quality, which we very uncertain about the estimate. Very low quality, which we very uncertain about the estimate. Very low quality, which we very uncertain about the estimate. Very low quality, which we very uncertain about the estimate. Very low quality, which we very uncertain about the estimate. Very low quality, which we very uncertain about the estimate. Very low quality, which we very uncertain about the estimate. Very low quality, which we very uncertain about the estimate. Very low quality, which we very uncertain about the estimate. Very low quality, which we very uncertain about the estimate. Very low quality, which we very uncertain about the estimate. Very low quality, which we will be estimated to the estimate and the estimate and the estimate of effect and may change the estimate of effect and estimate the estimate of estimate and may change the estimate the estimate of estimate an		154 per 1000					
GRIJOE Working Group grades of evidence High qualifie, Futther research is very unlikely to change our confidence in the estimate of effect. Moderate qualifie, Futther research is large by to have an important impact on our confidence in the estimate of effect and using change the estimate. Low qualifie, Further research is very lakely in have an important impact on our confidence in the estimate of effect and is likely to change the estimate. Low qualifie, Further research is very lakely in have an important impact on our confidence in the estimate. Very law qualifie in the very investigation for the estimate. In the study reported behalding. One early was quasi-FOT using alternation, the other two studies did not report randomisation method. The study reported behalding, One study was quasi-FOT using alternation. The study reported behalding, One study was quasi-FOT using alternation. The study reported behalding, One study was quasi-FOT using alternation. The study reported behalding, One study was quasi-FOT using alternation. The study reported behalding, One study was quasi-FOT using alternative the other two studies of the other study and the study was resonably until (XES) and Clis are wide and cross 1 **The observation was used to the study of the study	assumed risk in the	comparison group and the	dian control group risk acr relative effect of the interv	oss studies) is provided in foo ention (and its 95% Ci).	notes. The correspon	ding risk (and its 95% confidence interval) i	is based
	Moderate quality: F Low quality: Further Very low quality: W No study reported One study reported	r research is very untikely to urther research is likely to ha research is very likely to ha e are very uncertain about to blinding. One study was qui d allocation concealment ar	nave an important impact of ave an important impact or the estimate. ussi-RCT using alternation; and two studies reported ITT	n our confidence in the estimate n our confidence in the estimate the other two studies did not r analyses.	of effect and is likely	o change the estimate.	
	Moderate quality: Further Very low quality: W 1 No study reported One study reported 2 Number of patients 3 No explanation was 4 Cl crosses 1 5 Randomisation met	r research is very unlikely to unther research is likely to it research is very likely to had e are very uncertain about blinding. One study was qui d allocation concealment ar across groups was reason provided	nave an important impact of ave an important impact of the estimate. casi-RCT using alternation; d two studies reported ITT ably small (265) and Cis as	n our confidence in the estimate our confidence in the estimate the other two studies did not r analyses. re wide and cross 1	of effect and is likely	o change the estimate.	
	Moderate quality: Further Very low quality: W 1 No study reported One study reported 2 Number of patients 3 No explanation was 4 Cl crosses 1 5 Randomisation met	r research is very unlikely to unther research is likely to it research is very likely to had e are very uncertain about blinding. One study was qui d allocation concealment ar across groups was reason provided	nave an important impact of ave an important impact of the estimate. casi-RCT using alternation; d two studies reported ITT ably small (265) and Cis as	n our confidence in the estimate our confidence in the estimate the other two studies did not r analyses. re wide and cross 1	of effect and is likely	o change the estimate.	
	Moderate quality: Further Low quality: Further Very low quality: W 1 No study reported One study reported 2 Number of patients 3 No explanation was 4 Cl crosses 1 5 Randomisalion met	r research is very unlikely to unther research is likely to it research is very likely to had e are very uncertain about blinding. One study was qui d allocation concealment ar across groups was reason provided	nave an important impact of ave an important impact of the estimate. casi-RCT using alternation; d two studies reported ITT ably small (265) and Cis as	n our confidence in the estimate our confidence in the estimate the other two studies did not r analyses. re wide and cross 1	of effect and is likely	o change the estimate.	

10-day trimethoprim ve	rsus 10-day trimethopri	m+sulfamethoxazole for tre	ating lower urinary tra	t infection in children	
Patient or population: c Settings: outpatients de Intervention: 10-day tri Comparison: 10-day tri	partment nethoprim				
Outcomes	Mustrative comparati	ve risks* (95% CI)	Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence Comments (GRADE)
	Assumed risk	Corresponding risk			
	10-day trimethop +suffamethoxazole	rim 10-day trimethoprim			
Persistent bacterioria	Study population		RR 1.93	59	9 000
	69 per 1000	133 per 1000 (26 to 673)	(0.38 to 9.76)	(1 study)	very low ^{1,2}
	Medium risk populati	on.			
	69 per 1000	133 per 1000 (26 to 673)			
Persistent symptoms	Study population		RR 4.84 (0.24 to 96.66)	59 (1 study)	9000 very low ^{1,2}
	0 per 1000	0 per 1000 (0 to 0)	(0.24 to 90.00)	(r suby)	very low
	Medium risk populati	on			
	0 per 1000	0 per 1000 (0 to 0)			naci Tali, in Si Si Ne retive per la saci di Sila. Naci Per
Recurrence	Study population		RR 2.9 (0.12 to 68.5)	59 (1 study)	⊕OOO very low ^{1,2}

where gover peace of externor will be charge our confidence in the estimate of effect.

The first research is even unable to charge our confidence in the estimate of effect and may change the estimate, are qualify. From research is even to large in their a important impact on our confidence in the estimate of effect and is taken to the estimate of effect and is taken to change the estimate. Antibiotics for treating lower urinary tract infection in children (Review)
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The basis for the assumed fit it is a few modes control group (it's arous studies) is provided to honouse. The corresponding trist pool is 55% consumed to the requisition group and the provides of the provi

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Single-dose faslomycin versus single Patient or population: children with k Settings: outpollents department Intervention: single-dose fostomycin Comparison: single-dose nestimicin	Single doze fazionycia versus single-dese netilmiein for teas Patent or population. Children with lower urinary tract infection selestings condemne department or children single dose fazionycia naturendines, single-dose netilmichi.	Single doss followych verna single date entitlede for teating fower unleay test introller in bibliotes the properties all of the properties with over uniny tact intoleo selfuego cuplante opportune opportune or selfuego cuplante opportune or Companies supple do controller.	ry tract infection in ch	ldren		
Outcomes	Mustralive comparative risks* (95%, CI)	ive risks* (95% Ci)	Relative offect (95% CI)	No of Participants (shudies)	Quality of the evidence Comments (GRADE)	Comments
	Assumed risk	Corresponding risk				
	Single-dose netifimicin	n Single-dose fosfomycin				
Persistent bacteriuria	Study population		RR 3.15	135	0000	
	31 per 1088	98 per 1000 (21 to 454)	(0.68 to 14.64)	(1 study)	ve ry low ^{1,2}	
	Medium risk population	10				
	31 per 1000	98 per 1000 (21 to 454)				
Recurrence	Study population		RR 0.63	135	0000	
	156 per 1000	98 per 1000 (41 to 243)	(0.26 to 1.56)	(1 study)	very low ^{1,2}	
	Medium risk population	10				
	156 per 1000	98 per 1000				

Summary of main results

LTIL are relatively common childhood illuses and require in the unitary rans. This review was designed to include all results in the common and the common childhood illuses and require interest ment for hildren with lower UTI. Belaively for malatic investment for hildren with lower UTI. Belaively for malatic investment for hildren with lower UTI. Belaively for malatic investment for hildren with lower UTI. Belaively for malatic investment for hildren with lower UTI. Belaively for malatic investment for hildren with lower UTI. Belaively for malatic investment for hildren with lower UTI. Belaively for malatic investment in common articles of the common artic

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Potential blazes in the review process

The literature search included major international databases and a fairly beautiful search assumed to the search and a fairly beautiful search assumed was utilized to ensure all relevant notice was eliminated. We assume that we identified all RCTs relevant to our review questions, bowever many of the excluded suiside were foreign language anticle and their is the potential other a relevant study was eventobed in the translation process. Consider their search and their search process of the remainded process. On their search and their sear

Contact with any fiveningator.

Agreements and disagreements with other studies or reviews.

A Cochate review comparing alone CA dayly with studied (7.4 day) therapy should be conducted to enablish the studies of the control of

Data on retistant organisma were only reported in one mody. There is incineerin regarding the increasing resistance to anniholotics, specifically to possibility and caphalogoshios. Increasing anniholotics, specifically one possibility and caphalogoshios. Increasing anniholotics related to the control of t

Implications for practice
Implications for practice
In the rise was to the evidence that short-course therapy is an
appropriate therapy for children with lower UTI. While 10 days of
therapy was aignificantly more efficient bean single-done therapy,
there were mo differences in children with releve or Michael 2003
beween about an longer drustion anabilished therapy. Single-done
therapy is not recommended in children with UTI: 10-day rearness was aignificantly more effective in children with UTI: 10-day reartense was aignificantly more effective in children with UTI: 10-day reartense was aignificantly more effective in children with UTI: 10-day reartense was aignificantly more effective in children with UTI: 10-day reartense was aignificant following transment,
however this in due to a paneloy of RCII: rather than demonstratithemselves the control of the children of the

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References to other published versions of this review

Fixegerald 2007
Fixegrald A. Lee CW, Most R. Annibastics for treating uncomplicated unionsy treat infection in children. Cochange Database of Systematic Periors 2007, Issue 4. [DOI: 10.1002/1405188S.CD004857]
*Indicates the major publication for the study

Antibiotics for treating lower urinary tract infection in thildren (Review)
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- 203 -

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Ahmed 2001

TIME EVO		
Methods	Study design: parallel RCT Study period: NS	
Parisipans		nonths and 12 years with signs and symptoms of UTI, 10 ⁵ cfu/mL, and the presence of organisms susceptible collection method not reported.
Interventions	Treatment group • 10-day TMP (monotherap) Control group • 10-day TMP (8 mg/kg/d) •	y; 10 mg/kg/d) in 2 doses (SMX 40 mg/kg/d) in 2 doses
Outcomes	Persistent bacteriuria (16-1) Persistent symptoms (16-15 Recurrence (16-19 days foll	days following treatment)
Notes	Source of funding: NS	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding (performance bias and detection bias) All outcomes	Low risk	Investigators blinded
Incomplete outcome data (attrition bias) All outcomes	High risk	Less than half the randomised patients were analysed, no reason for losses to follow-up given
Selective reporting (reporting bias)	Low risk	Planned outcomes were all analysed

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35

Methods	Study design: parallel RCT Study period: NS	
Participans	Medical centre, Massachuseta G Country, USA Children aged between 2 as inclusion. Children were require frequency, dynamic, or abnormal two midstream clean catch or on Number 19 randomized, 4 Number 19 randomized, 4 O Tieatment group; 24 Control group; 25 Sex (MF): treatment group Eachusion criteria Acutely ill with temperature Acutely ill with temperature	
Interventions	45 kg (3.0 g) Control group	3 kg (1.0 g); 23 to 32 kg (1.5 g); 32 to 45 kg (2.0 g icillin: < 23 kg (125 mg); > 23 kg (250 mg)
Outcomes	Persistent bacteriuria (4 da)	s following treatment)
Notes	Study presents data for chil for children without abnormaliti Source of funding: Hoffma	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	All patients randomised were analysed
Selective reporting (reporting bias)	Low risk	Planned outcomes were all analysed

hagen ountry: Denmark hildren aged 1 to 15 yea	pspiral paediatric departments in and around
arch mid-stream urine si umber: 359 randomised o Treatment group: 90 o Control group: 78 ex (M/F): All female on criteria ntibiotic treatment one onamides; required pates (condition; SCr > 120 µ	re with distalled progresses requiring immediate a defined at 2 10 of Annual, of a single betweening in a mythe. Bug samples of wine-were not succepted 2.64 unalysed; 168 included in this review week prior to inclusion; assiption of allergy to penicilli treatments from severe unitary years malformations; and the severe severe unitary years malformations; or known instrumediated in severe severe unitary years malformations; or known instrumediated in severe severe unitary years malformations; or known instrumediated in severe severe severe severe to known instrumediate or known in the severe or known instrumediated in severe to known instrumediate or known instrumediated to known instrumediate or known instrumediated to known instruments to known in the to known instruments to known in the to known in the to known in the to known in t
ivmecillinam, 20-40 mg Il group	kg/d in 2 doses for 3 days fkg/d in 2 doses for 10 days
essistent bacteriuria (1-1	days following treatment)
nt UTI (52%) nother intervention arm Cochrane review by Mi epeated in this review.	previous UTI (17%), a history of UTI (31%), or was included in this study, 3-day sulfamethizole. that 2003 reports outcomes for this comparison which Medical Research Council (5.52.11.10 and 5.52.14.
rs' judgement	Support for judgement
r risk:	To ensure an equal number of patients in each group, a block randomisation methods was used. Randomisation was in blocks of 6 wishin each of the 10 participating de partment. No detail about the way the block randomisation was performed were reported.
	ex (M/F); All female in oriental unablicite treatment one unablicite treatment one consideration SCV > 120 µm consideration SCV > 120 µm consuperative treatment or tent group treatment or tent group treatment constitution, 20-40 mg/dg proup treatment constitution, 20-40 mg/dg proup treatment bacterium 20-40 mg/dg proup 20-40 m

Allocation concealment (selection bias)	Low risk	Allocation concealed by drawing consect tively numbered sealed envelopes prepare by the manufacturer
Blinding {performance bias and detection bias} All outcomes	Unclear risk	Nor reported
Incomplete outcome data (attrifion bias) All outcomes	Uncterr nisk	Só shaldern dels nos fadili rhedusion cites. Ols baceriotta nos ejunificant. Il provide lorg amplel; treatment vos disconitues in 6 children before schodelded, 32 chi dern did not have unine cultures complete within 10 days from treatment; polite were not evaluated for other exason; il hops were exastlede because of the mu dirribuned benweren group. The fielde ex- fector of the 50 children who were not an yard were included as they received trea- ment.
Selective reporting (reporting bias)	Low risk	Planned outcomes were all analysed
Fine 1985 Methods	Study design: parallel RCT Study period: NS	
Participans	Maryland Hospital Country: USA Female adolescents aged 12 n UTI (frequency, dysuris, heistancy fevor or malaris and ingrificant be mid-stream urine sample Mean age: 16.5 years Number: 34 nandomised, 31 Teatment group: 16 Control group: 15 Sex (M/F): all female Exclusion criteria	nt Adolescent General Medical Clinic, University of 18 years with clinical symptoms of an acute fower lower abdominal pain, urgency, anoreals, flow-goal certaint defined as > 10° cluml. In a clean earth analysed lergy to penicillin or concurrent antibiotic use
	Treatment group	

Outcomes	Persistent bacteriuria (2-5 da Persistent symptoms (2-5 da	
Notes	28/31 participants were sexu Source of funding: Maternal	ally active and Child Health Grant (MCH #000980)
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	Two participants were excluded from analyses because of early pregnancy; one participant did not turn up to the follow-up appointments
Selective reporting (reporting bias)	Low risk	Planned outcomes were all analysed
Grimwood 1988		
Methods	Study design: parallel RCT Study period: NS	
Participants	Country: New Zealand Children aged 2 weeks to 12 cfu/ml. in 2 consecutive clean cat aspiration. Children with cystris tenderness and were without othe Mean age: 4.9 years Number of participants: 45 e Exclusion criteria	children s were also included in this study and wete reported

nterventions	Treatment group • Single intramuscular gentami Control group • 7-day course of appropriate assandard doses (included TMP-SM	ntibiotic depending on culture sensitivity in
Dutcomes	Persistent bacteriuria (1 day f Recurrence (< 1 week followi Re-infection (> 1 week follow	ng treatment)
Rotes	23 children with 3 or more proven UTIs during the preceding 12 months were defined as having a history of recurrent UTIs. Source of funding: National Children's Health Research Foundation	
Risk of bias		
lias	Authors' judgement	Support for judgement
andom sequence generation (selection	Low risk	Random numbers table
allocation concealment (selection bias)	Unclear risk	Not reported
ilinding (performance bias and detection ias) Il outcomes	Unclear risk	Not reported
ncomplere outcome data (attrition bias)	Low risk	All patients randomised were analysed
elective reporting (reporting bias)	Low risk	Planned ourcomes were all analysed
elin 1984		
Sethods	Study design: parallel RCT Study period: NS	
articipans		

Helin 1984 (Continued)	38.5°C, flank pain, elevated ESR a	uggesting upper usinary tract involvement (fev nd leukocytosis); known sensitivity to cephales dder disorder; known structural malformation
Interventions	Treatment group • 3-day cephalexin 25-50 mg/k • Control group • 10-day nitrofurantoin 3-4 mg	
Outcomes	Persistent bacteriuria (4-7 day Recurrence (any time during Re-infection (any time during	follow-up; mean 8 months)
Notes	Source of funding: NS	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	All patients randomised were analyze
Selective reporting (reporting bias)	Low risk	Planned outcomes were all analysed
Khan 1981		
Methods	Study design: Quasi-RCT Study period: NS	
Participans	University of New York, Downstate Country: USA Children aged six months to	15 years with symptoms of cysticis (including rr) and significant bacteriuria defined as > 10 ⁵ c ine samples.

	Sex (M/F): 4/50 Exclusion criteria Younger than 6 months, or abnormal SCr or BUN values.	older than 15 years; urinary trace malformations;
Interventions		es at random for both groups and included ampicillis onventional doses given orally 4 times/day
Outcomes	Persistent bacteriuria (3-7 c Recurrence (> 2 months for	
Notes	Source of funding: NS	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Alternation
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	All patients were analysed in groups to which they were assigned
Selective reporting (reporting bias)	Lowrisk	Planned outcomes were analysed. Data fo re-infection was presented across cystiti pyelonephritis and asymptomatic bacter utia and was not reported for cystitis alon
Komoroski 1999		
Methods	Study design: parallel RCT Study period: NS	
Participants	Inclusion criteria • Steinip/terusiment: outputient and emergency department. Arkansas Children's Hospital • Country: USA • Children aged 1to 19 years with as 1 or more clinical symptoms of cystois	

Komoroski 1999 (Continued)		
	significant bacterium defined > 1 orlo d'unimi defined > 1 orlo d'unimi dato considered positive if 5 10° orlo d'unimi dato considered positive if 5 10° orlo major canada programa, and in vitro sentiririty patrem as the simple clean-catho programa, programa de Numbers 97 annadonised, 55 or Concust organization criteria • Numbers 97 annadonised, 55 or Concust organization criteria • Pregramary antibiotic therap requiring additional antibiote the Preprince to a UTI sign and sp grama programa programa controverbal angle to explain programa controverbal angle to explaination for the programa pr	analysed yin the previous 2 weeks; concomitant infection energy; known resul or sur-logic problems that could improun of probosphatis (ill or unic appearance, one processing = 28 34°C), history or periodilina significant history of gaussinnessing or corteal revorws spreen disease; history of gaussinnessing ouses as a childis a promy or gaussinn who was unable one a family strained in which follow-up our as a family strained in which follow-up our as a child; and the processing of the contraction of th
Interventions	o 27 received ceftriaxone Control group • TMP-SMX 4-5 mg/kg twice	cone 50 mg/kg (to a maximum of 500 mg) (500 mg); 9 received ceftriaxone (250 mg) e daily for 10 days (, 1 patient received amoxicillin because of aulfa
Outcomes	Persistent bacteriuria (10-30 Persistent bacteriuria and sy.	days following treatment) mptoms (10-30 days following treatment)
Note	appears the study was originally of control. The ceftriaxone 500 mg treatment failures were reported. 250 mg group is not reported as groups should be of equal size. It allocated to the ceftriaxone 250 mallocated to the other 2 groups.	
Risk of bias	hadd raddig	
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported

	Unclear risk	Nor reported
Blinding (performance bias and detection bias) All outcomes	High risk	Open label study
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	It appears that children were randomized to treatment before the inclusion/exclusion circins were applied, 25 children had urine cultures that showed no significant growth. 8 children had a laboratory or procedural error occurred (e.g., urinalysis obtained but culture not done, organisms in culture not worked up). 3 children did not return for follow-up assument:
Selective reporting (reporting bias)	Unclear risk	Relapse and recurrence were reported, but not in a format suitable for data extraction for this review
Lidefelt 1991		
Methods	Study design: parallel RCT Study period: 1986-1988	
Puridipus	Stockholm Country: Sweden Claidern aged less than 3 year frequency, dynuria, and painful mic CfulmL in 2 separately voided uri more than 2 perdous UTis, and the the study. Median age: 5 years Number: treatment group (50 Sex (M/F): 13/87 Exclusion criteria	ent (temperature < 38.5°C, absence of loin pain,
Puridipuna	String/secruliment: emerges Stockholm Country Sweden Children aged less than 3 year frequency, dyauria, and painful mie? dufurli. in 2 separately voided uri more than 2 perfosso UTis, and the te study. Median age 5 years Numbers treatment group (50: Sec (MF); 13/87 Eachtsion criteria Signs of upper tract involvem.	so 12 years with symptoms of a UTI (including monitors) and significant bacterium defined as ≥ 10 me tamples. Children were required to have had not emost recent at least 6 months prior to the start of 0; control group (50).

Notes	Source of funding: Swedish Medical Council, grant number 19X765, and th Swedish Society of Medicine.	
Risk of bias		alianosia notal da his
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	All patients were analysed in groups t which they were assigned
Selective reporting (reporting bias)	Low risk	Planned outcomes were analysed
Participants	Inclusion criteria Setting/tercnistment: NS Children aged 8 months to 11.1 year Column. of a single pathogen in 2 consect Number: treatment group (16); cont Exclusion criteria Hypersensitivity to explusiosporins or function, or surceural anomalies	ol group (16)
Interventions	Treatment group • Celeforcial 25 mg/kg once daily for 10 days Control group • Ampicillin 50 mg/kg/d in 4 divided doses for 10 days	
Outcomes	Persistent bacteriuria (10 days following treatment) Persistent symptoms (10 days following treatment)	
Notes	Children with pyelonephricis were also included in this study and were reporte separately. Source of funding: NS	

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	All patients were analysed in group t which they were assigned
Selective reporting (reporting bias)	Low risk	All planned outcomes were analysed
Mitnik 1985		
Methods	Study design: parallel RCT Study period: NS	
Participans	Paediatric Clinic of the Chilean Country. Chile Children aged 2 years to 14 dysuria, urgency, food smelling us bacteriuria defined at ≥ 10° cful supra-public aspiration. Children UTIs, and the most recent at leas Number: treatment group 1 Sex (M/F): 11/87 Exclusion criteria	years with symptoms of a UTI (including frequency rine, enturesis and/or haematurol) and significant mil. in voided urine sample, or \$2 1000 offurth on were required to have had not more than 2 previous of somothe poir or the start of the scut) (1 (27); treatment group 2 (35); control group (36) in; a history of UTI; anatomical abnormalities receive
Interventions	Treatment group 1 • 3-day antibiotics Treatment group 2 • 5-day antibiotics Control group	

Outcomes	Recurrence (at 2-3 months)	
Notes	The 3-day and 5-day intervent to the 10-day control Source of funding: NS	entions were combined into one group and compared
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	All patients analysed in group to which they were assigned
Selective reporting (reporting bias)	Low risk	All planned outcomes were reported
Methods	Study design: parallel RCT Study period: NS	
Participans	Country: Italy Children aged 1 month to defined as > 105 cfu/mL of a single.	ients authors were from various university hospitals 16 years with a lower UTI. Significant bacteriuria $g_{\rm part}$ govern in 2 clean catch or catheterited urine as absence of fever, ESR < 25 mm/L/h and CRP < 20 71); control group (64)
Interventions	Treatment group Single-dose fosfornyein tror Control group Single-dose netilmicin (5 m	nctamol (2 g orally; 1 g in children < 1 year) g/kg intramuscularly)

Outcomes	Persistent bacteriuria (2-4) Recurrence (up to 30 days	
Notes	Source of funding: NS	
Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Low risk	All patients analyzed in group to which the were assigned
Selective reporting (reporting bias)	Low risk	All planned outcomes were reported
Methods	Study design: parallel RCT Study period: NS	
Methods Participants	Study design: parallel RCT Study period: NS Inclusion criteria	
	Serting/recruitment: emerg d'Hebron, Barcelona Country: Spain	ency department, Hospital Materno-Infantil Vall
	 Children aged 8 months to 11.1 years with significant bacteriuria defined as ≥ 1 ⁵ CulmL of a single pathogen in 2 consecutive mid-stream urine samples. Numbers 37 Exclusion criteria Children aged less than 4 months, with a fever of >38.5°C, back pain or mass, 	
	malaise, duration of symptoms I	onger than one week, vorniting, received antibiotics is disease involving immunosuppression, or known
Interventions	Children received amoxicil co-trimoxazole at standard doses	in, amoxicillin + clavulanic acid, cephalexin, TMP or for 7 days.
Outcomes	Persistent bacteriuria (2-3 c	lays following treatment)
Antibiotics for treating lower urinary tract infe		

Notes	This study was published as Source of funding: NS	s an abstract	
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Unclear risk	Not reported	
Allocation concealment (selection bias)	Unclear risk	Not reported	
Blinding (performance bias and desection bias) All outcomes	Unclear risk	Not reported	
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Abstract only, not enough detail provid	
Selective reporting (reporting bias)	Unclear risk	Abstract only, not enough detail provide	
Participants	Inclusion criteria	211 111 115	
Participants		ency department, Children's Hospital of Pittsburgh	
	Country: USA	h symptoms of a UTI (including frequency, dysuria	
	and/or urgency) and significant b	pacteriuria defined as ≥ 105 cfu/mL in 2 clean catch	
	urine samples, or ≥ 1000 cfu/m! • Mean age: 5.6 years	Lon supra-pubic aspiration.	
	Number: 37 randomised, 3	5 analysed	
	o Treatment group: 18 o Control group: 17		
	Exclusion criteria	UT1 (fever > 38°C and/or flank pain); known	
	anatomic or functional urinary to	race abnormality; currently receiving antibiotics;	
	history of penicillin allergy		
	Treatment group		
Interventions		Single-dose amoxicillin 50 mg/kg (to a maximum of 2.5 g) Control group	
Interventions	Control group		
Interventions	Control group	cg/d in 3 divided doses (to a maximum of 500 mg/	

Blas Author' judgment Support for judgment Random sequence generation (telection Unclear risk Nor reported bath) Allaction oncealment (telection bins) Unclear risk Nor reported Blinding (performance bias and detection tow risk bins) All doctormes Uncomplete outcome data (strittion bins) Unclear risk Two children were encluded from analyze the concerned are the exceed utilise callure was neg strice Selective reporting (reporting bins) Low risk All planned outcomes were reported	Blas Author' judgment Support for judgment Blas I Author' judgment Support for judgment Blas I Author' judgment Support for judgment Blas I Author' judgment I Support for judgment Blas I Support for judgment for judgment I Support for judgment I Support for judgment I Support for judgment		Persistent bacteriuria (2 days fi Recurrence (within 3 months	
Blas Authors' judgments Random sequence generation (telection Loclear risk Not reported Minding (performance bias and detection Low risk Discounting (performance bias and detection Low risk Discounting (performance bias and detection Low risk Patient and physician Discounting Di	Blas Author' judgment Support for judgment Random sequence generation (telection Unclear risk Nor reported band) Milection concealment (selection bits) Unclear risk Nor reported Milection concealment (selection bits) Unclear risk Patient and physician bits) All outcomes Incomplete outcome data (stratition bits) Unclear risk Two children were necluded from analyze the concealment (reporting bits) Unclear risk Two children were necluded from analyze the selection of the second uniter culture was neg since Selection exporting (reporting bits) Unow risk All planned outcomes were responsed Selection exporting (reporting bits) Participant **Souly designs parallel BCT** **Souly designs parallel BCT** **Souly proble MS** Participant **Souly designs parallel BCT** **Souly proble MS** **Country USA** **Grid gard 2 to 17 years with program of a UTI (including frequency, dynain, urgency, coparia, superposit, pain, or la heamsturis with prarial and significant between the offices and 2 to 10 years with program of a UTI (including frequency, dynain, urgency, coparia, superposit, pain, or la heamsturis with prarial and significant between the offices and 2 to 10 years with prarial and significant between the offices and 2 to 10 years with prarial and significant between the offices and 2 to 10 years with prarial and significant between the offices and 2 to 10 years with prarial and significant between the offices and 2 to 10 years with prarial and significant between the offices and 2 to 10 years with prarial and significant between the offices and 2 to 10 years and 2 years	Notes	Source of funding: Not reported	
Random sequence generation (telection Unclear risk Not reported bital) Allocazione concealment (telection bital) Unclear risk Patient and physician Low risk Patient and physician Low risk Patient and physician Allocazomes Incomplete outcome data (attrition bital) Unclear risk Two children were excluded from analyze because the second write culture was neg strict Selective reporting (reporting bital) Low risk All planned outcomes were reported **Sordy designs parallel RCT* **Sordy	Random sequence generation (selection Indear risk Nor reported Shal) Allocation concealment (selection bias) Allocation soncealment (selection bias) Allocation soncealment (selection bias) Bridding (selectromance loss and detection Low risk Patient and physician Indoar risk Patient and physician Interpretate Incomplete concerne data (stration bias) All concernes Incomplete concerne data (stration bias) Low risk Two children were enclosed from analyze since Selective reporting (reporting bias) Low risk All planned concounts were reported * Smoly designs parallel RCT * Smoly designs parallel RCT * Smoly designs parallel RCT * Smoly periods NS * Participants Includen criteria * Sensing/retrainment comparients and emergency deparament. Children's Houghtal of Philadelphia and Schrienspher's Houghtal Gord Children, Wilseldsphia and Children, Wilseldsphia and Children, Wilseldsphia and Schrienspher's Houghtal Gord Children, Wilseldsphia and Children, Wilselds	Risk of bias		Additional State Addition
hiad Allocation concealment (orlection bias) Unclear risk Not reported Patient and physician All coatcomes Uncomplex coatcome data (attrition bias) Unclear risk Unclear risk Two children were excluded from analyze because the second write culture was neg street street Selective reporting (reporting bias) Low risk All planned coatcomes were reported Scalal 1984 Methods • Sordy designs parallel RCT • Sordy designs parallel RCT • Sordy retoric NS Inclusion criteria • Senting/retoriuments comparisent and emergency deparament. Children's Hospital of Children, Wilsdelphia. • Country LSS • Country LSS • Country LSS • Median age 4.75 years • Median age 4.75 years • Median age 4.75 years • Number of participants • Teammer group: 10 The Stalant contral • Supro or symptoms of upon time of a single organism in 2 sequential dean cassh with the stalant of this shop of the stalant with proposal or a spital contral or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stalant or a single organism in 2 sequential dean cassh with the stala	hiad Allocation concealment (orlection bias) Unclear risk Not reported Patient and physician All coatcomes Uncomplex coatcome data (attition bias) Unclear risk Unclear risk Two children were enclosed from analyze because the second write culture was neg sive Selective reporting (reporting bias) Low risk All planned coatcomes were reported Sealt 1984 Methods Sondy designs parallel RCT Sondy designs parallel RCT Sondy designs parallel RCT Sondy designs parallel RCT Sondy periods NS Inclusion criteria Senting/recruitments companients and emergency deparament. Children's Houghtal of Philadelphia and SC Inclusiopher's Houghtal for Children, Willadelphia Commy LSS Lower control, Rayman with purposen of a UT1 Goodsing frequency, dynatio, surgery or sentins, lampathelphia of Schildren, or hancomassis deep propial and girlings and screenfast defined at ≥ 10° clulmit. of a single organism in 2 sequencial clean casch universal to the control of the contr	Bias	Authors' judgement	Support for judgement
Blinding (performance bias and desection Low risk Dial) All autocomes Two children were encluded from analyze Incomplete outcome data (stration bias) All autocomes Low risk Two children were encluded from analyze between the second urine culture was neg since Selectrice reporting (reporting bias) Low risk All planned outcomes were reported biash 1984 Methods - Sundy designs parallel BKCT - Sundy periods NS Participants - Sending from immon compatients and emergency department. Children's Houghts of Children's Houghts - Commy 1984 - Gill sign and 2 to 17 years with proposed and 11 for designing from general companies of the proposed of	Blinding (performance bias and desection Low risk Dial) All autocomes Two children were encluded from analyze Incomplete outcome data (stration bias) All autocomes Low risk Two children were encluded from analyze between the second urine culture was neg since Selectrice reporting (reporting bias) Low risk All planned outcomes were reported biash 1984 Methods - Sundy designs parallel BKCT - Sundy periods NS Participants - Sending from immon compatients and emergency department. Children's Houghts of Children's Houghts - Commy 1984 - Gill sign and 2 to 17 years with proposed and 11 for designing from general companies of the proposed of		Unclear tisk	Not reported
blad All occument Incomplete eucome data (amnion bias) Localez rink Versuse the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was not garden experience of the eccord unine culture was not garden experience of the eccord unine culture was not garden experience of the eccord unine culture experience of the eccord experience of the eccord experience of eccord experi	blad All occument Incomplete eucome data (amnion bias) Localez rink Versuse the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was no garden experience of the eccord unine culture was not garden experience of the eccord unine culture was not garden experience of the eccord unine culture was not garden experience of the eccord unine culture experience of the eccord experience of the eccord experience of eccord experi	Allocation concealment (selection bias)	Unclear risk	Not reported
All outcomes Selective reporting (reporting bias) Low risk All planned outcomes were reported Stabl 1984 Methods Study design: parallel RCT Study design: parallel	All outcomes Selective reporting (reporting bias) Low risk All planned outcomes were reported Stabl 1984 Methods Study design: parallel RCT Study design: parallel	bias)	Low risk	Patient and physician
Sachl 1984 ## Sundy designs parallel RCT ## Sundy designs parallel RCT ## Sundy periods NS ## Sundy N	Souls 1984 Methods • Sundy designs parallel RCT • Sundy persion 185 Darkiepanu Annie Control 185 Annie Control 185 Annie Control 185 - Sundy persion to comparison and emergency deparement. Children's Hospital of Whitelephina and Schlichmagher's Hospital for Children's United behavior of Whitelephina and Schlichmagher's Hospital for Children's United behavior of Whitelephina and Schlichmagher's Hospital for Children's United Schlichmagher's Hospital for Children's Indiana and Septiment Interests of Benderal 2010 (Septiment Schlichmagher's Hospital) and septiment Successful of Benderal 2010 (Septiment Schlichmagher's Hospital)		Unclear risk	because the second urine culture was neg
Mediods Sundy design: parallel IKCT Sundy prediot NS Including critical Sundy prediot NS Including critical Sunsing precipitations or comparises and emergency deparament. Children's Hospital of Philadelphia and Sc. Christopher's Hospital for Children, Philadelphia Country, USA Girl aged 2 to 17 years with symptoms of a UTI (including frequency, dysuria, urgency, centrels, suprapole; pain, or the measuria with prediot and applicant uries samples. Median age 4.75 years. Median age 4.75 years. Number of participature 36 anothonicel, 26 analysed Tituraness group; 10 S. Counted group; 10 S. Counted group; 10 S. Spur or symptoms of upper UTI (inexpectator = 3.8.9°C, flink pain, convertebul angles critedness or unite agreements), known read or mulpic, disorder	Methods Sundy designs parallel KCT Sundy periods XS Includence riteria Sundy periods XS Includence riteria Sunsing periods in Continuopher's Hospital for Children, Filluladophia Contempt VSA Gill aged 2a to 17 years wich symptoms of a UTI (including frequency, dysuria, urges; centrels, superposite pain, or Batenamin with printal and significant suries are period of the Contempt VSA Welling aged 2a to 17 years with symptoms of a UTI (including frequency, dysuria, urges; centrels, superposite pain, or Batenamin with printal and significant suries amplets. Welling 2a 473 years Number of participature 36 are domined. 26 analyzed Transmers group: 10 Scienced group 16 Sectioned group 16 Sectioned group 171 (centrelature > 83.9-C. flink pain, conservation and printing conservation and participature of supplementations).	Selective reporting (reporting bias)	Low risk	All planned outcomes were reported
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				er and emergency described. Children's Marsial

Interventions	Control group	mg/kg ozilly (to a maximum of 3 g) sg/d in 3 divided doses (to a maximum of 250 mg/
Outcomes	Persistent bacteriuria (2-4 d Reinfection (> 2 weeks folk	
Notes	both a positive culture more than as recurrence by this review) or a re-infection by this review). The	nor be used from this study as the definition included n 2 weeks following therapy of any organism (defined n infection caused by a different organism (defined as se results were presented together. In Laboratories, Bristol, Tennesse
Risk of blus		
Bizs	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Nor reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	Six girls were lost to follow-up, in 3 girls the 2nd urine culture was negative and 1 girls that received antibiotics within the personal 2 weeks. One girl in the single-dose grounded as amontofills resistant cognism as was prefetched to 10 days TMP-SMX and them followed in the conventional therap group
Selective reporting (reporting bias)	Unclear risk	All planned outcomes were reported
Wallen 1983		
Methods	Study design: parallel RCT Study period: NS	
Participants	Country: USA	tients, The Children's Memorial Hospital, Chicago s with suspected UTI and significant bacteriuria

	samples. Median age: 5.45 years Number: 54 randomised, 49 analys Teatment group: 26 Control group: 25 Exclusion criteria Clinical symptoms of pyelonephrini	us in 2 clean cauch or urine collection bug ed s fincloding fewer > 38,3°C, flank pain, chills cic use during the week prior to the study;	
Interventions	Treatment group • Single-dose intramuscular amikacin Control group • 10-day sulfisoxazole 150 mg/kg/day	sulface 7.5 mg/kg (to a maximum of 240 mg	
Outcomes	Persistent bacteriuria (2-4 days follo Recurrence (30-40 days following t		
Notes :	Re-infection rates were presented, but were only available for the single-dose amiliacin group; these rates have not been reported in this review. Source of funding: NS		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation (selection bias)	Low risk	Random numbers table	
Allocation concealment (selection bias)	Unclear risk	Not reported	
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Not reported	
Incomplete outcome data (attrition bias) All outcomes	High risk	At the 2-4 day follow-up, 6 girls were lor to follow-up. By the 30-40 day follow-up 10 girls were lost to follow-up	
Selective reporting (reporting bias)	Low risk	All planned outcomes were reported	

Study	Reason for exclusion
Adam 1982	Children with pre-existing conditions, and who have symptoms of pyelonephritis are not reported separatel from children with lower UTI
Anttila 1980	Not RCT
Arap 1983	Half of included children had fever and were not reported separately from those without
Arguedas 2009	Children had complicated UTI
Arrieta 2001	Included children had pyelonephritis
Asscher 1973	Not an RCT; screening study only
Bahur 2003	Included children had fever
Bailey 1977	Almost half of the included children had known renal impairment
Baker 2001	Included children were required to be febrile (i.e. systemic illness)
Bukkaloglu 1996	Included children had pyelosephritis
Belet 2004	Prophylaxis for preventing recurrence
Bose 1974	More than half of the included children had pre-existing renal abnormalities
Bourillon 1994	Included children had pyelosiephritis
Caparelli 1983	Some children had pyelonephritis; unclear how many
Carapetis 2001	Most included children had systemic symptoms
Careddu 1987	Study is conducted in symptomatic and asymptomatic children, but proportion of each is unknown. Also, 11 51 children had known renal abnormalities
Carlsen 1985	Prophylactic antibiotics
Chibante 1994	Some children had complicated UTI and were not presented separately from those with lower UTI
Chong 2003	Children had systemic symptoms
Chrapowicki 1975	Included children had pyelonephritis
Clemente 1994	Included children had fever
Dagan 1992	Majority of included children had fever

De Garace 1988	One third of included children had cystopyelitis or pyelonephritis and were not reported separately from those without
Ellerstein 1977	Not enough information reported on symptoms to know whether children had lower UTI; 5/34 had reflux and 3/34 had abdominal pain, but other symptoms were not reported
Elo 1975	Two thirds of included children had renal abnormalities
Feldman 1975	Some children had fever and were not reported separately from those without
Fischbach 1989	Included children had signs of systemic illness (fever)
François 1995	Included children had pyelonephritis
Françoise 1997	Included children had pyelonephritis
Foji 1987	Some children had pyelonephritis and were not reported separately from children without
Gaudreault 1992	Comparison of short versus standard duration antibiotic for lower UTI - included in Michael 2003
Ginsburg 1982	Approximately 1/3 of included children had fever and were not reported separately from those without
Gak 2001	Approximately 1/3 of included children had pyelonephritis and over half had urinary tract abnormalities
Goldberg 1977	Children with fever not reported separately from children without
Gonzalez 1985	35% of included children had fever and were not reported separately from those without
Goos 2006	Not a RCT, or quasi-RCT
Goos 2007	Not RCT
Goszczyk 2000	Children received 3 months antibiotic treatment for preventing recurrence
Gould 1975	Unclear if participants were children. Included participants had prostatitis, acute cystitis, urethritis, and/or trigonitis but results were not reported separately
Granados 1998	Prophylactic antibiotics
Hansen 1981	Approximately half of children presented with fever and were not reported separately from children without fever
Hayashida 1970	Some children had pyelonephritis and were not reported separately from those without
Helin 1981	Comparison of short versus standard duration antibiotic for lower UTI - included in Michael 2003
Hoberman 1999	Included children were required to have a temperature of > 38.3°C

Howard 1978 Just under half of the included children had fever and approximately 65% had symptoms of malaise Johnson 1993 Comparison of short versus standard duration antibiotic for lower UTI - included in Michael 2003 Jojant 1991 Comparison of short versus standard duration antibiotic for lower UTI - included in Michael 2003 Khan 1987 Not RCT Komberg 1994 Comparison of abort versus aundard duration antibiotic for lower UTI - included in Michael 2003 Krepler 1976 Included children had pyelonephritis Comparison of short versus standard duration antibiotic for lower UTI - included in Michael 2003 Lohr 1981 Lubitz 1984 Symptoms not reported. 35% of included children had renal abnormalities Madrigal 1988 Comparison of short versus standard duration antibiotic for lower UT1 - included in Michael 2003 Included children were required to have fever McCracken 1981 > 20% of children had fever, abdominal/flank pain and consoverrebral tenderness indicating pyelonephritis Moe 1977 Not all included children had bacteriologically proven UTI Montini 2003 Children had pyelonephritis Montini 2007 Children had pyelonephritis Half of the included children had fever, loin pain and/or back pain Noorbakhsh 2004 Included children had pyelonephtitis Olbing 1971 Some children had renal abnormalisies; although the results refer to patients with and without abnormalisies, no numbers are included so data cannot be extracted Palcoux 1986 Half of included children had known renal abnormalities More than half of the included children had abdominal pain and/or fever Pylkkunen 1981 Compared 10-day treatment with 42-day treatment in children Children with fever were not analysed separately from children without fever Repetto 1984 Rodriguez 1983 Included children had fever Antiblotics for treating lower utinary trect infection in children (Review) Copyright © 2012 The Cochrane Gollaboration. Published by John Wiley & Sons, Ltd.

52% of children had fever and were not reported separately from those without
Majority of included children had fever
Most children received concomitant surgical therapy
Some patients had fever and/or lumber pain and were not reported separately from patients without
Symptoms not reported. Approximately half of included children had reflux, but grade of reflux was not reported
Most included children had fever
No symptoms of UTI reported. Bacteriological definition of UTI only
As per Michael 2093. Study was excluded because significantly more patients (32/59) with pyelonephitists were Included in the 7-day group compared with 3-day group (11/58) (Chi ² = 15.65, df = 1; P < 0.001), which strongly suggested non-random allocation
Included children were required to have fever
Some children had pyelonephritis but were not reported separately from those without
Included children presented with fever
One third of included children had known renal abnormalities and are not presented separately from those without
Included children had pyelonephritis
Included children had pyelonephritis
More than half of the included children had fever and were not reported separately from those without
Comparison of shore versus standard duration antibiotic for lower UT1 - included in Michael 2003
Comparison of short versus standard duration antibiotic for lower UTI - included in Michael 2003
Comparison of abore versus usundard duration antihionic for lower UTI - included in Mitchael 2003 Hed trial-RCT: UTI - unitarry tract infection

DATA	AND	ANALYSES	

Comparison 1. Single-dose versus conventional 10-day treatment

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
l Persistent bacteriuria	6	228	Risk Ratio (M-H, Random, 95% CI)	2.01 [1.06, 3.80]
1.1 Amoxicilin	4	131	Risk Rario (M-H, Random, 95% CI)	1.97 [0.90, 4.33]
1.2 Other antibiotics	2	97	Risk Rario (M-H, Random, 95% CI)	2.09 [0.71, 6.18]
2 Persistent symptoms	1		Risk Ratio (M-H, Random, 95% Cl)	Totals not selected
5 Recurrence	2	79	Risk Ratio (M-H, Random, 95% CI)	1.38 [0.55, 3.50]
4 Persistent bacteriusia and symptoms	1		Risk Ratio (M-H, Random, 95% CI)	Totals not selected

Comparison 2. Single-dose versus short-course (3-7 days) treatment

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
l Persistent bacteriuria	2	145	Risk Ratio (M-H, Random, 95% CI)	1.30 [0.65, 2.62]
2 Recurrence	2	145	Risk Ratio (M-H, Random, 95% CI)	1.50 [0.43, 5.26]
5 Re-infection	1		Risk Rario (M-H, Random, 95% CI)	Totals not selected

Comparison 3. Short-course (3-7 days) versus long-course (10-14 days) treatment

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Persistent bacteriuria	3	265	Risk Ratio (M-H, Random, 95% CI)	1.09 [0.67, 1.76]
2 Recurrence	4	328	Risk Ratio (M-H, Random, 95% CI)	1.25 [0.74, 2.13]
3 Re-infection	2	211	Risk Ratio (M-H, Random, 95% CI)	0.88 [0.44, 1.74]

Antibiotics for treating lower urinary tract infection in children (Review)

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Comparison 4. Trimethoprim (10 days) versus trimethoprim+sulfamethoxazole (10 days)

Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Persistent bacteriuria	1		Risk Ratio (M-H. Random, 95% CI)	Totals not selected
2 Persistent symptoms	1		Risk Ratio (M-H, Random, 95% CI)	Totals not selected
3 Recurrence	1		Risk Ratio (M-H, Random, 95% CI)	Totals not selected

Comparison 5. Cefadroxil (10 days) versus ampicillin (10 days)

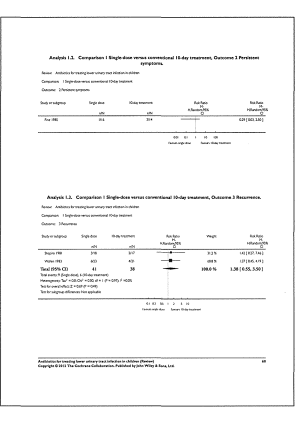
Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
I Persistent bataeriuria	1		Risk Ratio (M-H, Random, 95% CI)	Totals not selected
2 Persistent symptoms	1		Risk Rario (M-H, Random, 95% CI)	Totals not selected

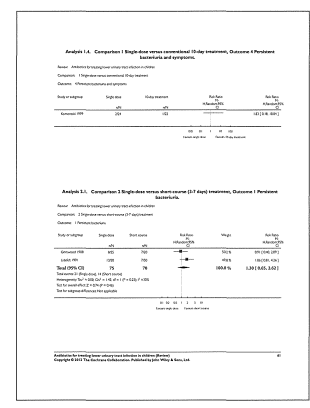
Comparison 6. Single-dose fosfomycin versus single-dose netilmicin

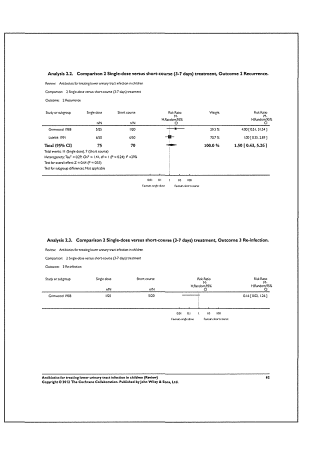
Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Persistent bacteriuria	1		Risk Ratio (M-H, Random, 95% CI)	Totals not selected
2 Recurrence	1		Risk Ratio (M-H, Random, 95% CI)	Totals not selected

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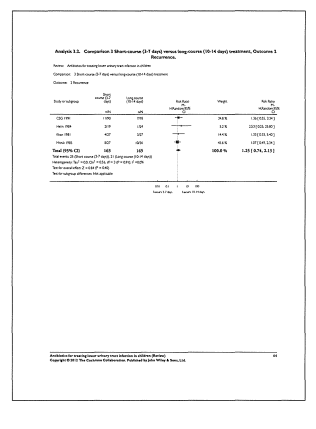
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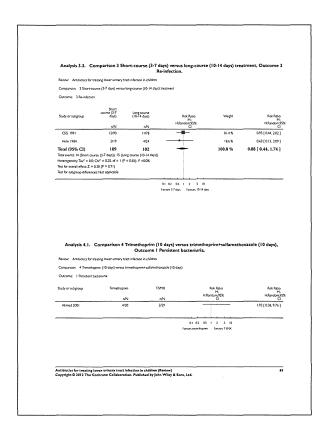


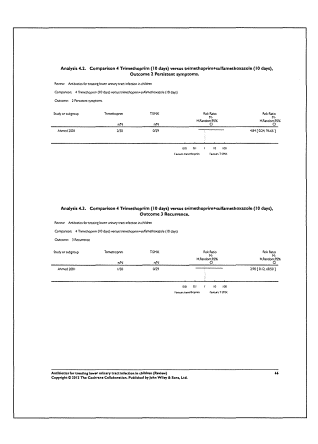


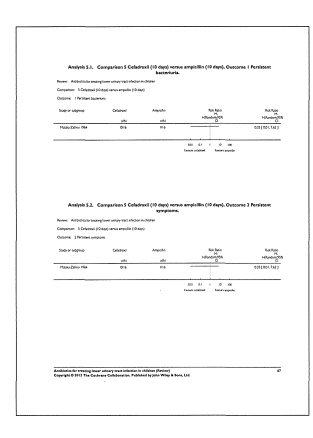


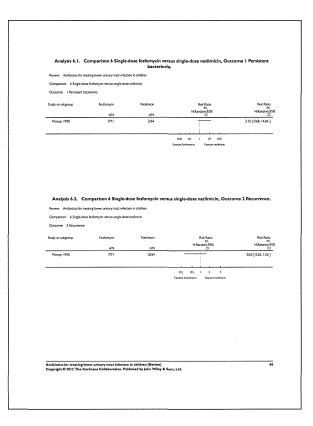
Analysis 3.1. Comparison 3 Short-course (3-7 days) versus long-course (10-14 days) treatment. Outcome 1 Perview Aribidica for tracing linear every trice discions coldate Congruence 1 Discretions (2-7 days) was large course (10-14 days) beamest Outcome 1 Provinces Liberary Sect. S











A P P E N D I C ES Appendix I. Electronic search strategy Database Search terms used CENTRAL 1. child* triabbw 3. (adoloces* or pubmic or necess* or newbom* or toddles*) ni,abbw 4. (peture or periodical-rinabbw 4. (peture or periodical-rinabbw 6. (a) (P and T or peture or pubmic or perpubers* or perpubers* or pivenils* or youth* or teen*) ni, abbw 6. (a) (P and T or peture or pubmic or perpubers* or peture or petu



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Date Event Description 10 July 2008 Amended Converted to new review format.

CONTRIBUTIONS OF AUTHORS

- Writing of protocol and review: AF, RM
 Screening of titles and abstracts: AF, RM
 Astenment for inclusion: AF, RM
 Quality assessment: AF, RM
 Data extraction: AF, RM
 Data entry into RevMan: AF

- Data analysis: AF, RM

DECLARATIONS OF INTEREST

Anita Fingendid Some of this work was undersiden when all authors were employed by, or were advisor's to, the National Collaborating Centre for Women's and Children's Health which received funding from NICE. The views expressed in this publication are token of the authors and one measuraby those of NICE.
 The Anita Lakhangsut I was the Childred Director as the National Collaborating Centre for Women's Health and led the checkponent of the NICE Unlessy Tract Indication Guiddien. I are as longer the Clinical Director but remain on the NICE-WCH beard and I are a NICE Fellow and member of the NISE evidence advisory ratm.

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

- Re-defined the outcome of recurrence to include re-infections we used the definition recurrence (growth of original bacteria) and re-infections (growth or new bacteria).

 Re-defined the protection for new bacteria) in the result of the protection of the p
- We initially defined recurrence as at least three episodes of cystitisflower UTI: however in the included studies any recurrence was reported. We therefore included data on any recurrence.
- Adverse effects were to be tabulated this was not performed.

 Risk of bias assessment tool has replaced the quality assessment checklist.

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INDEX TERMS Medical Subject Headings (MeSH)

Anti-Bacteria Agenta Įdefinisiuration & douge; therapeutic unel; Anti-Infective Agenta, Urinary Įdefinisiuration & douge;
'therapeutic unel; Bucteriur'i Pfung therapyl; Drug Administration Schedule; Infant, Newborn: Randomized Controlled Titals as
Topic Urinary Tates Infection (1"due ferzpy)

MeSH check words

Adolescent; Child; Child, Preschool; Humans; Infant

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