

**SUMMARY OF FINDINGS FOR THE MAIN COMPARISON (Continued)**

Single-dose versus short-course (3-7 days) antibiotics for treating lower urinary tract infection in children	No. of Participants (studies)	Relative effect (RR, CI)	Quality of the evidence (GRADE)	Comment
<b>Outcome:</b> Short-course (3-7 days)				
Illustrative comparative risks <sup>1</sup> (95% CI)				
Assumed risk	Short-course (3-7 days)	Corresponding risk		
Study population	200 per 1000	200 per 1000 (130 to 320)		
Median risk population	245 per 1000 (159 to 447)	318 per 1000 (192 to 497)		
<b>Reference</b>	Study population	RR 1.3 (0.9 to 1.9)	⊕○○○ very low <sup>2,3</sup>	
Study population	100 per 1000	130 per 1000 (82 to 197)		
Median risk population	85 per 1000 (57 to 147)	110 per 1000 (71 to 170)		
<b>Re-infection</b>	Study population	RR 0.16 (0.02 to 1.26)	⊕○○○ very low <sup>4</sup>	
Study population	45 (1 study)			

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200 per 1000	40 per 1000 (5 to 315)
Median risk population	49 per 1000 (5 to 315)
200 per 1000	49 per 1000 (5 to 315)

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

⊕○○○ Confidence Interval, RR, Risk ratio

GRADE Working Group grades of evidence

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

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

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

<sup>1</sup> Either study reported allocation concealment or blinding, one reported adequate randomization method, but the other did not. IT was used in both studies. See notes and dates in following: <sup>2</sup> 100% <sup>3</sup> 100% <sup>4</sup> 100% <sup>5</sup> 100% <sup>6</sup> 100%

<sup>1</sup> Allocation concealment and blinding not reported. Random number table and IT analyses used and no losses to follow-up.

<sup>2</sup> Very small numbers of patients (45)

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**BACKGROUND**

Urinary tract infection (UTI) is one of the most common bacterial infections in infants and children and the most common bacterial infection in infants under three months (Savelyev 2005). Infection of the urinary tract presents in three ways: as covert bacteriuria on screening; non-systemic symptoms and infection limited to the ureters and bladder (epididymitis or lower UTI); or systemic symptoms and infection of the kidneys (pyelonephritis or upper UTI). Lower UTI is the most commonly presenting and in the majority of cases can be easily treated with a course of antibiotic therapy with no further complications. Some children, particularly young infants who are unable to describe how they feel, present with non-specific symptoms making it difficult to diagnose as either lower or upper UTI, whereas older children present with more specific complaints such as dysuria or frequency. The prognosis of lower UTI in childhood is largely unknown. The relationship between UTI, renal scarring and vesicoureteric reflux (VUR) is unclear, as is the progression of lower UTI to pyelonephritis and subsequent damage to the kidneys. Current practice and treatment decisions are not underpinned by strong evidence and are largely the result of clinical judgment and the biological plausibility of future kidney damage.

It is difficult to obtain accurate estimates of the number of infants and children who will present with a lower UTI during childhood. Current practice in most countries is based on historical data from studies conducted in secondary and tertiary referral centres and renal registries. This fails to adequately describe the larger primary care population for whom very limited population-based data of robust quality exist. Of the population-based studies available, variable incidence rates, ranging from approximately 0.1% (Matali 1988) to 3% (Winberg 1974) have been presented. The cumulative incidence of childhood UTI is likely to be somewhere between 5% and 13% (Grosshans 1979; Harrison 1993).

A number of antimicrobials have been used to treat children with lower UTIs; however there is neither agreement on the most effective agent, nor the optimal dosage. This review aims to evaluate the benefits and harms of antibiotics used to treat lower childhood UTI by investigating the alleviation of symptoms and persistence of bacteriuria following treatment, recurrent symptomatic UTI and renal parenchymal damage.

An existing Cochrane review investigates antibiotic therapy for acute pyelonephritis in children (Holton 2007) and another compared short to standard duration oral antibiotic treatment for acute UTI in children (Michael 2003). The latter concluded that a 2-4 day course of oral antibiotics appears to be as effective as 7-14 days in reducing lower tract UTI (epididymitis) in children. Two additional reviews were identified (Feres 2002; Tian 2001) comparing the efficacy of single-dose, short-course and standard course antimicrobial therapy for childhood UTI. This review provides additional data in Cochrane format, particularly on the duration of antibiotic therapy.

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**OBJECTIVES**

This review aimed to summarise the benefits and harms of antibiotics for treating bacteriologically proven, symptomatic, lower UTI in children.

**METHODS**

**Criteria for considering studies for this review**

**Types of studies**

All randomised controlled trials (RCTs) and quasi-RCTs (RCTs in which allocation to treatment was obtained by alternation, use of alternate medical records, date of birth or other predictable methods) in which antibiotic therapy was used to treat lower UTI in children. The first period of randomised cross-over studies was also included.

**Types of participants**

**Inclusion criteria**

- Children aged zero to 18 years with bacteriologically proven UTI (at least one culture of a known urinary pathogen (bacterial culture > 10<sup>5</sup> cfu/ml) in a child with first time or recurrent UTI and who had at least one localised symptom of UTI (such as dysuria or frequency) treated with antibiotics in primary and community health care settings or an outpatient department were included.

- Studies that primarily included children with acute pyelonephritis (at least one culture of a known urinary pathogen (bacterial culture > 10<sup>5</sup> cfu/ml) in a child with at least one symptom or sign of systemic illness such as fever, loin pain or flank pain) or otitis and additional diagnostic criteria as defined by the authors) were excluded. The symptom of fever is a controversial one: for the purposes of this review, we have excluded children who present with fever in an effort to differentiate between children with lower UTI and those with pyelonephritis.

- Children found to have renal abnormalities during the study were included, however if they had known abnormalities prior to the study they were excluded. We included children with low grade (< 2) reflux.
- Studies including patients with lower UTI and either upper UTI or covert bacteriuria were included if the data for the patients with lower UTI could be extracted separately, otherwise these studies were excluded. Any urine collection method was acceptable, including urine collection pad or bag, clean-catch method, catheter, or supra pubic aspiration.

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**Exclusion criteria**

- Children hospitalised for a condition not related to UTI.
- Children with bacteriologically proven UTI and symptoms or signs of systemic illness (including fever, loin pain, toxicity).
- Children with covert bacteriuria (non-symptomatic).
- Children with pre-existing renal abnormalities or known underlying renal conditions (including high grade (3-4) VUR, nephrotic syndrome and reno-genetic bladder).
- Children receiving prophylactic antibiotics for UTI.
- Children receiving antibiotics for any other condition.
- Immunosuppressed children.

**Types of interventions**

- Antibiotic therapy (standard 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).

**Types of outcome measures**

- Persistent symptoms at completion of treatment.
- Pyelonephritis, significant bacteriuria (> 10<sup>5</sup> cfu/ml) at completion of treatment.
- Combinations of persistent bacteriuria (> 10<sup>5</sup> cfu/ml) and symptoms at completion of treatment.
- Recurrent symptomatic UTI following treatment.
- Symptomatic re-infection following treatment.
- Any renal parenchymal damage on DMSA, four to six months following UTI.
- Compliance.
- Adverse events.
- Development of resistant organisms.
- Any changes to antibiotic regimens.

**Search methods for identification of studies**

**Electronic searches**

The search strategy was comprehensive and was designed to cover two reviews being undertaken by the authors, this review and 'Interventions for covert bacteriuria in children' (Fringsdrift 2012). We searched the following databases to identify relevant studies. Full details of the search strategies are reported in Appendix 1. We searched the Cochrane Renal Group Specialised Register (May 2012) through contact with the Trials Search Co-ordinator using search terms relevant to this review. The Cochrane Renal Group Specialised Register contains studies identified from:

- Quarterly searches of the Cochrane Central Register of Controlled Trials CENTRAL.
  - Weekly searches of MEDLINE OVID SP.
  - Handsearching of renal-related journals & the proceedings of major renal conferences.
  - Searching of the current year of EMBASE OVID SP.
  - Weekly current awareness alerts for selected renal-journals.
  - Searches of the International Clinical Trials Register (ICTRP) Search Portal & ClinicalTrials.gov
- Studies contained in the specialised register are identified through search strategies for CENTRAL, MEDLINE and EMBASE based on the scope of the Cochrane Renal Group. Details of these strategies as well as a list of handsearched journals, conference proceedings and current awareness alerts are available in the 'Specialised Register' section of information about the Cochrane Renal Group.

**Searching other resources**

- Reference lists of nephrology textbooks, review articles and reference studies.
- Letters seeking information about unpublished or incomplete studies to investigators known to be involved in previous studies.

**Data collection and analysis**

**Selection of studies**

The search strategy described was used to obtain titles and abstracts of studies relevant to the review. The titles and abstracts were screened independently by two authors assessed to determine which studies satisfied the inclusion criteria.

**Data extraction and management**

Data extraction was carried out independently by the same authors using standard data extraction forms. Studies reported in non-English language journals were translated before assessment. Where more than one publication of one study existed, reports were to be grouped together and the most recent or most complete dataset used. Any discrepancy between published versions was to be highlighted. Disagreements were resolved in consultation with a third author.

**Assessment of risk of bias in included studies**

- The following items were independently assessed by two authors using the risk of bias assessment tool (Higgins 2011) (see Appendix 2).
- Was there adequate sequence generation (selection bias)?
- Was allocation adequately concealed (selection bias)?

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- Was knowledge of the allocated interventions adequately prevented during the study (detection bias)?
  - Outcome assessors
- Were incomplete outcome data adequately addressed (attrition bias)?
  - Are reports of the study free of suggestion of selective outcome reporting (reporting bias)?
  - Was the study apparently free of other problems that could put it at a risk of bias?

**Measures of treatment effect**

For dichotomous outcomes (e.g. symptom resolution, persistent bacteriuria, recurrent infection) results were expressed as risk ratio (RR) with 95% confidence intervals (CI). No continuous scales of measurement were used in the review.

**Dealing with missing data**

Any further information required from the original author was requested by written correspondence and any relevant information obtained in this manner was included in the review.

**Assessment of heterogeneity**

Heterogeneity was analysed using a Chi<sup>2</sup> test on N-1 degrees of freedom (df), with an alpha of 0.05 used for statistical significance and with the I<sup>2</sup> test (Higgins 2003). I<sup>2</sup> values of 25%, 50% and 75% corresponded to low, medium and high levels of heterogeneity.

**Assessment of reporting biases**

If possible, funnel plots were to be used to assess for the potential existence of small study bias (Higgins 2011).

**Data synthesis**

Data was pooled using the random-effects model.

**Subgroup analysis and investigation of heterogeneity**

Subgroup analysis was used to explore possible sources of heterogeneity (e.g. participants, interventions and study quality). Heterogeneity among participants could be related to age and renal pathology. Heterogeneity in treatments could be related to prior agent(s) used and the agent, dose and duration of therapy. Older and newer manuscripts evaluating the same antibiotic was also analysed as a subgroup. Subgroup analyses were also used to explore paediatric sub-populations (infants, toddlers, children and adolescent groups).
 

- Infants: under one year of age
- Toddlers: one to under three years of age
- Children: three to under 12 years of age
- Adolescents: twelve to 18 years of age

**RESULTS**

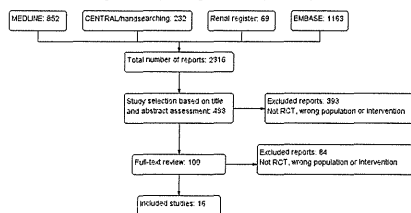
**Description of studies**

See Characteristics of included studies; Characteristics of excluded studies.

**Results of the search**

We initially identified 2316 potentially relevant reports of studies. After evaluating their titles and abstracts we excluded 1823 articles because they were not RCTs, included adult patients, or did not investigate antibiotics for UTI. The full-text articles of the remaining 493 studies were evaluated, with a further 393 excluded, leaving 100 potentially relevant RCTs. After further assessment 16 studies met our inclusion criteria (Figure 1).

Figure 1. Study flow diagram show study selection process



**Included studies**

The 16 included studies enrolled 1116 children and young people aged between two weeks and 19 years. Most were published between 1981 and 1991, with two studies published between 1999 and 2001. The median sample size was 40 patients (range: 26 to 264). All included studies used the bacteriological definition of 10<sup>3</sup> cfu/mL for confirming UTI and included children with no-systemic symptoms, the most common being dysuria and frequency. Two studies compared single-dose antibiotics with short-course (3-7 days) antibiotics.

- Crimwood 1988 compared a single intramuscular gentamicin injection (1 mg/kg) with seven days of oral antibiotics sensitive to the organism cultured.
- Lidfeldt 1991 compared single-dose trimethoprim (TMP) (6 mg/kg) with a five-day course of TMP (3 mg/kg, twice daily).

Six studies investigated single-dose antibiotics compared with conventional 10-day courses.

- Four studies (Aver 1983; Fine 1985; Shapiro 1981; Suli 1984) compared single-dose amoxicillin (1-3 g) with 10-day amoxicillin (125-500 mg, thrice daily).
- Kucurova 1999 compared single intramuscular ceftriaxone (50 mg/kg) with 10-day TMP-sulfamethoxazole (TMP-SMX) (4.5 mg/kg, twice daily).
- Wallen 1985 compared single-dose intramuscular amikacin sulfate (7.5 mg/kg) with 10-day sulfisoxazole (150 mg/kg/d in four divided doses).

We identified four studies comparing short duration antibiotics

(3-7 days) with conventional 10-day courses using different antibiotics.

- Czak 1991 compared 3-day pivmecillinam (20-40 mg/kg/d in two doses) with 10-day sulfamethizole (40-80 mg/kg/d in two doses).
- Jolin 1981 compared 3-day cephalixin (25-50 mg/kg/d in two doses) with 10-day nitrofurantoin (3-4 mg/kg/d in two or three doses).
- Minnik 1985 administered antibiotics to which the cultured organism was susceptible and treated three groups with three, five and 10 days of antibiotics. The data for the 3-day group were included in the analysis.
- Khan 1981 administered a range of antibiotics at random including ampicillin, sulfisoxazole and cephalixin in conventional doses given orally four times a day and compared 3-day treatment with 10-day treatment.

The Cochrane review by Michal 2003 compared short (2-4 days) with long-course (7-14 days) antibiotics in children with UTI where the short and long-course antibiotic were the same. Identified nine studies (CSG 1991; Gaudinaki 1993; Jelin 1981; Johnson 1993; Kember 1994; Lohr 1981; Madagal 1988; Wenzien 1979; Zaki 1988). We have not included these comparisons, however CSG 1991 was included in both reviews as this study presented antibiotic comparisons for both the same and different antibiotics.

Two studies compared different 10-day regimens.
 

- Ahmed 2001 compared TMP (monotherapy) with TMP-SMX.

- Malaka-Zafiri 1984 compared cefadroxil (25 mg/kg, once daily) with ampicillin (50 mg/kg/d in four divided doses).
- One study compared single-dose regimens.
  - Principi 1990 compared oral fosfomicin trometamol (2 g) with intramuscular ceftriaxone (5 mg/kg).
- In Sanchez 1990 (published only as an abstract), children were randomised to one of five antibiotics: amoxicillin, amoxicillin and clavulanic acid, cephalixin, TMP or co-trimoxazole at standard doses for seven days.
  - Included children with pyelonephritis (18)
  - Included children with both pre-existing abnormalities and pyelonephritis (2)
  - Included children with fever or signs of systemic illness (50)
  - Included prophylactic antibiotics (5)
  - Reported in a Cochrane review comparing long and short duration antibiotics (Michal 2003) (9)
  - Included children without bacteriologically proven UTI (1)
  - Significantly more patients were assigned to one group compared with the other suggesting non-random allocation (1)
  - Four articles were excluded for other reasons (4) (see Characteristics of excluded studies)

**Excluded studies**

- We excluded 84 articles.
  - Not RCT (6)
  - Included children with pre-existing renal abnormalities (10)

**Risk of bias in included studies**

See Figure 2 and Figure 3.

Figure 2. Risk of bias graph: review authors' judgements about each risk of bias item presented as percentages across all included studies.

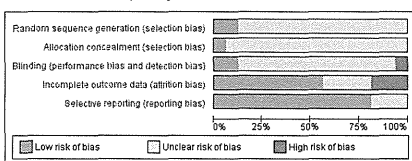


Figure 3. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.

Study	Random sequence generation (selection bias)	Allocation concealment (selection bias)	Blinding (performance bias and detection bias)	Incomplete outcome data (attrition bias)	Selective reporting (reporting bias)
Ahmed 2001	?	?	?	?	?
Aver 1983	?	?	?	?	?
CSO 1991	?	?	?	?	?
Fine 1985	?	?	?	?	?
Grimwood 1988	?	?	?	?	?
Helin 1984	?	?	?	?	?
Khan 1991	?	?	?	?	?
Komoroski 1999	?	?	?	?	?
Lidfeldt 1991	?	?	?	?	?
Malaka-Zafiri 1984	?	?	?	?	?
Minnik 1985	?	?	?	?	?
Principi 1990	?	?	?	?	?
Sanchez 1990	?	?	?	?	?
Shapiro 1981	?	?	?	?	?
Stani 1984	?	?	?	?	?
Wallen 1983	?	?	?	?	?

#### Allocation

#### Random sequence generation

The quality of reporting of random sequence generation was poor. Two studies reported using random numbers tables to generate a random sequence (Greenwood 1988; Wilton 1983) and one study was quasi-randomised using alternation (Khan 1981). CSG 1991 used a block randomisation method to ensure an equal number of patients in each group. Randomisation was in blocks of six within each of the ten participating departments. No details about the way the block randomisation was performed were reported. The remaining 12 studies did not report the randomisation method.

#### Allocation concealment

The quality of allocation concealment was very poor. CSG 1991 reported using sealed envelopes which were prepared by the manufacturer of the study drug. The remaining 15 studies did not report allocation concealment.

#### Blinding

The quality of blinding was also very poor. Shapiro 1991 reported blinding both patients and physicians. Ahmed 2001 reported blinding of the investigators; and Komonoki 1999 was reported to be open label. The remaining 13 studies did not report blinding.

#### Incomplete outcome data

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

- CSG 1991: 26% of children randomised were lost to follow-up for a variety of reasons including not fulfilling inclusion criteria; treatment discontinued before scheduled; and did not have repeat urine cultures within the allocated time. Nineteen boys were excluded from this study because of the small number and because they were not evenly distributed between groups.

- Komonoki 1999: 37% of the children randomised were lost to follow-up. Some urine cultures showed no significant growth; some urine samples were subject to laboratory or procedural errors; and some children did not return for follow-up assessments.

- Ahmed 2001 reported that only 52% of randomised patients were analysed, no reason for attrition were given.
- Four other studies reported losses to follow-up of less than 10% (Pain 1985; Shapiro 1991; Subi 1984; Wilton 1983).

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11

- 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.

#### Selective reporting

Most studies reported all planned outcomes. Komonoki 1999 reported relapse and recurrence, but not in a format suitable for data extraction for this review. Sanchez 1990 was published only as an abstract and it was not clear whether all planned outcomes were reported.

#### Effects of interventions

See Summary of findings for the main comparison. Single-dose versus short-course (3-7 days) antibiotics for treating lower urinary tract infection in children. Summary of findings 2 Single-dose versus conventional 10-day antibiotic treatment for treating lower urinary tract infection in children. Summary of findings 3 Short (3-7 days) long-course (10-14 days) antibiotics for treating lower urinary tract infection in children. Summary of findings 4 10-day trimethoprim versus 10-day trimethoprim-sulfamethoxazole for treating lower urinary tract infection in children. Summary of findings 5 10-day cefaclor versus 10-day ampicillin for treating lower urinary tract infection in children. Summary of findings 6 Single-dose fosfomycin versus single-dose neomycin for treating lower urinary tract infection in children.

#### Persistent bacteriuria at completion of treatment

Fourteen studies reported outcomes for persistent bacteriuria at completion of treatment: five studies completed follow-up urine cultures between two and five days; and one study completed follow-up urine cultures between 10 to 30 days.

#### Single-dose versus conventional 10-day treatment

Conventional 10-day antibiotic treatment significantly increased the number of children free of persistent bacteriuria compared to single-dose treatment (Analysis 1, 1) (6 studies, 228 children; RR 2.01, 95% CI 1.46 to 2.80). No heterogeneity was observed (I<sup>2</sup> = 0%). The test for subgroup differences between the studies using amoxicillin in both arms and studies using other antibiotics did not show any difference (Chi<sup>2</sup> = 0.01, df = 1, P = 0.93, I<sup>2</sup> = 0%).

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

There was no significant difference in persistent bacteriuria between single-dose and short-course antibiotic treatment (Analysis 2, 2) (1 study, 145 children); RR 1.30, 95% CI 0.65 to 2.62; I<sup>2</sup> = 30%.

#### Short-course (3-7 days) versus long-course (7-10 days) treatment

There was no significant difference in persistent bacteriuria between short-course and long-course antibiotic treatment in three studies (Analysis 3, 1) (3 studies, 265 children); RR 1.09, 95% CI 0.67 to 1.76; I<sup>2</sup> = 0%.

#### Head-to-head studies

There were no significant differences in persistent bacteriuria between:

- TMP (10 days) versus TMP-SMX (10 days) (Analysis 4, 1) (1 study, 59 children); RR 1.93, 95% CI 0.38 to 9.70;
- cefaclor (10 days) versus ampicillin (10 days) (Analysis 5, 1) (1 study, 32 children); RR 0.35, 95% CI 0.01 to 7.02; and
- fosfomycin (single-dose) versus neomycin (single-dose) (Analysis 6, 1) (1 study, 135 children); RR 3.15, 95% CI 0.26 to 1.50.

Sanchez 1990 randomised children to one of five antibiotics: amoxicillin, amoxicillin + clavulanic acid, cephalosporin, TMP or co-trimoxazole. Because of the small number of patients in each group (1 to 2) this data was unable to be included in meta-analysis. Following treatment, the number of children free of bacteriuria in each group were 4/5 children who received amoxicillin, 2/7 children who received amoxicillin + clavulanic acid, 2/8 children who received cephalosporin, 2/8 children who received TMP; and 2/9 children who received co-trimoxazole.

#### Persistent symptoms at completion of treatment

Three studies reported outcomes for persistent symptoms following treatment and showed no differences between treated and untreated groups: all for different antibiotic comparisons and durations.

#### Single-dose versus conventional 10-day treatment

There was no significant difference between single-dose treatment compared with conventional 10-day treatment (Analysis 1, 2) (1 study, 30 children); RR 0.29, 95% CI 0.03 to 2.50) where 1/16 of the single-dose group had persistent symptoms compared with 3/14 of the 10-day group (Efte 1995).

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

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12

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

There was no significant difference between 10-day TMP treatment compared with 10-day TMP-SMX treatment (Analysis 4, 1) (1 study, 59 children); RR 4.84, 95% CI 0.24 to 96.60; where 2/30 of the TMP group had persistent symptoms compared with 0/29 of the TMP-SMX group (Ahmed 2001).

#### Cefaclor (10 days) versus ampicillin (10 days)

There was no significant difference in persistent symptoms between 10-day cefaclor treatment compared with 10-day ampicillin treatment (Analysis 5, 2) (1 study, 32 children); RR 0.33, 95% CI 0.01 to 7.62) where 0/16 of the cefaclor group had persistent symptoms compared with 1/16 in the ampicillin group (Makka-Zafrafi 1984).

#### Recurrent symptomatic UTI following treatment

Ten studies reported outcomes for recurrence (with the same organism) following treatment for the initial infection; five studies reported recurrences within one month of antibiotic treatment; four studies reported recurrences between one and three months following antibiotic treatment; and one study reported recurrences at any time with a mean time of eight months.

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

There was no significant difference between single-dose treatment compared with short-course (3-7 days) treatment (Analysis 2, 2) (1 study, 145 children); RR 1.30, 95% CI 0.43 to 3.26; I<sup>2</sup> = 29%, where 11/75 (15%) of the single-dose group had recurrence compared with 7/79 (10%) of the short-course group.

#### Single-dose versus conventional 10-day treatment

There was no significant difference between single-dose treatment compared to conventional 10-day treatment (Analysis 1, 2) (1 study, 79 children); RR 1.38, 95% CI 0.55 to 3.50; I<sup>2</sup> = 0%, where 9/41 (22%) of the single-dose group had recurrent symptomatic UTI compared with 6/38 (16%) in the 10-day group.

#### Short-course (3-7 days) versus long-course (7-10 days) treatment

There was no significant difference between short-course treatment compared with long-course treatment (Analysis 3, 2) (4 studies, 328 children); RR 1.25, 95% CI 0.74 to 2.13; I<sup>2</sup> = 0%. All four studies reported 3-day to 10-day treatment. Of these four studies, 25/163 (15%) of the short-course group had recurrent symptomatic UTI compared with 21/165 (13%) of the long-course group.

#### Head-to-head studies

There were no significant differences in recurrent symptomatic UTI between:

- TMP (10 days) versus TMP-SMX (10 days) (Analysis 4, 1) (1 study, 59 children); RR 2.90, 95% CI 0.12 to 68.50, where 1/30 in the TMP (monotherapy) group had recurrent symptomatic UTI compared with 0/29 in the TMP-SMX groups; and
- fosfomycin (single-dose) versus neomycin (single-dose) (Analysis 6, 2) (1 study, 135 children); RR 0.63, 95% CI 0.26 to 1.56) where 7/71 (10%) in the single-dose fosfomycin group had recurrent symptomatic UTI compared with 10/64 (16%) in the single-dose neomycin group.

#### Re-infection following treatment

Three studies reported outcomes for re-infection (with a different organism) following antibiotic treatment for the initial infection; one study reported re-infection occurring at more than one week, one study reported re-infection at 1-10 days and one study reported re-infection at any time following antibiotic treatment, with a mean time of eight months.

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

There was no significant difference between single-dose compared with short-course treatment (Analysis 2, 1) (1 study, 45 children); RR 0.16, 95% CI 0.02 to 1.20, where 1/25 (4%) of the single-dose group had a re-infection compared with 5/25 (20%) of the short-course group.

#### Short-course (3-7 days) versus long-course (7-10 days) treatment

There was no significant difference between short-course treatment compared with long-course treatment (Analysis 3, 2) (2 studies, 211 children); RR 0.88, 95% CI 0.44 to 1.74; I<sup>2</sup> = 0%, where 14/109 (13%) of the short-course group had a re-infection compared with 15/102 (15%) of the long-course group.

#### Renal parenchymal damage

None of the included studies reported renal parenchymal damage. Two studies (Khan 1983; Subi 1984) reported the use of intravenous osteostethogram or intravenous pyelogram following antibiotic therapy, however the studies were aimed to identify structural abnormalities and VUR and not renal parenchymal damage.

#### Compliance

Five 1985 reported compliance with follow-up assessment. Compliance at the first scheduled follow-up appointment was 100%

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13

in the single-dose treatment group compared to 60% in the 10-day treatment group. Non-returning patients often required several phone calls and letters before returning.

- Three studies reported compliance with antibiotic treatment.
- Fite 1985: 27% of the 10-day treatment group reported perfect compliance with their medication; other patients finished their medication in more than 10 days, less than 10 days, and some had not finished their medication at the time of follow-up.
- Greenwood 1988: side effects prevented two children randomised to amoxicillin from completing their course of antibiotics.
- Heilin 1996: 'optimal patient compliance was achieved in both groups'.

#### Development of resistant organisms

While many included studies reported the antibiotic sensitivity of organisms cultured prior to treatment, most did not report development of resistant organisms during the study period. Subi 1984, comparing single-dose with 10-day amoxicillin, reported that in each of the three single-dose patients who relapsed, the single-dose failed to clear the urinary tract of a sensitive organism, and each of the four conventional therapy patients who relapsed developed organisms resistant to amoxicillin during therapy. All of these patients subsequently responded to additional or different antibiotic therapy.

#### Adverse events

- Patients in five studies did not experience any side effects (Auner 1983; Heilin 1984; Makka-Zafrafi 1984; Wilton 1983).
- CSG 1991: children randomised to 10-day sulfamethoxazole reported no side effects. Two children randomised to 3-day pivmecillinam developed urticarial rash; two children discontinued treatment due to vomiting and abdominal pain; one child had diarrhoea and one child developed irritability and fatigue.
- Lakfelt 1991: one child randomised to single-dose TMP experienced vomiting.
- Prince 1990: two children receiving single-dose fosfomycin experienced mild and transient diarrhoea which disappeared spontaneously; one had nausea; and one had skin rash.
- Data reported in Ahmed 2000) was part of a larger study on children with both UTI and otitis media. Across both TMP and TMP-SMX groups, adverse events included vomiting, abdominal pain, diarrhoea and skin rash, although less than 5% of children experienced these symptoms.
- Fite 1985: *Canalida vaginaria* occurred in three patients receiving 10-day amoxicillin.
- Komonoki 1999: two sexually active females reported vaginal itching after receiving intramuscular ceftriaxone.

- Local discomfort from injection sites was reported in two studies (Komonoki 1999; Prince 1990).
- Side effects were not reported in six studies (Greenwood 1988; Khan 1983; Minni 1985; Sanchez 1990; Shapiro 1991; Subi 1984).

#### Subgroup analyses

We were unable to perform the pre-specified subgroup analyses. The majority of studies did not report results for different practice sub-populations, so this analysis could not be undertaken. Comparing older and newer manuscripts was also not possible because there were not enough 'newer' manuscripts. With the excep-

tion of Ahmed 2001 (which was the only study to compare 10-day TMP with 10-day TMP-SMX) and Komonoki 1999 (single-dose versus 10-day treatment), all studies compared were performed within a 10-year period of each other. Because the method of randomisation was not sufficiently described for most of the studies, we did not perform a sensitivity analysis that excluded quasi-RCTs. Data of randomisation, method of allocation concealment and blinding were not reported by the majority of studies and we were therefore unable to conduct sensitivity analyses comparing higher and lower quality studies.

Formal testing for publication bias using funnel plots was not possible because of the small number of studies identified.

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

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14

\* Very small numbers of patients (59)  
 † Neither study reported allocation concealment. One reported using a random numbers table, the other reported blinding. Neither study used ITT analysis.  
 ‡ Number of participants is small, <25 in each group across both studies. CI are wide and include 1.

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17

**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 urinary tract infection  
 Settings: outpatient and/or emergency department  
 Intervention: single-dose  
 Comparison: conventional 10-day treatment

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk				
	Conventional treatment	10-day Single-dose				
Persistent bacteriuria	Study population		RR 2.01 (1.06 to 3.8)	228 (6 studies)	@○○○ very low <sup>2</sup>	
	104 per 1000	209 per 1000 (110 to 395)				
	Medium risk population	126 per 1000	253 per 1000 (134 to 475)			
Persistent symptoms	Study population		RR 0.29 (0.03 to 2.5)	30 (1 study)	@○○○ very low <sup>4</sup>	
	214 per 1000	62 per 1000 (6 to 535)				
	Medium risk population	214 per 1000	62 per 1000 (6 to 535)			

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15

**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)

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk				
	Long-course (10-14 days)	Short-course (3-7 days)				
Persistent bacteriuria	Study population		RR 1.1 (0.98 to 1.77)	265 (3 studies)	@○○○ very low <sup>2</sup>	
	186 per 1000	205 per 1000 (126 to 329)				
	Medium risk population	165 per 1000	204 per 1000 (126 to 327)			
Recurrence	Study population		RR 1.14 (0.7 to 1.88)	353 (4 studies)	@○○○ very low <sup>4</sup>	
	127 per 1000	145 per 1000 (89 to 236)				
	Medium risk population	100 per 1000	114 per 1000 (70 to 169)			

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18

Persistent bacteriuria and symptoms	Study population		RR 1.83 (0.18 to 18.84)	46 (1 study)	@○○○ very low <sup>4</sup>	
	45 per 1000	82 per 1000 (8 to 848)				
	Medium risk population	48 per 1000	84 per 1000 (8 to 807)			
Recurrence	Study population		RR 1.38 (0.56 to 3.5)	79 (2 studies)	@○○○ very low <sup>4</sup>	
	193 per 1000	218 per 1000 (87 to 553)				
	Medium risk population	154 per 1000	213 per 1000 (85 to 539)			

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

GRADE Working Group grades of evidence  
 High quality: Further research is very unlikely to change our confidence in the estimate of effect.  
 Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.  
 Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.  
 Very low quality: We are very uncertain about the estimate.

- <sup>1</sup> Only 1/6 studies reported randomisation method; no study reported allocation concealment; 1/5 studies reported blinding; 1/5 studies reported ITT analysis.
- <sup>2</sup> Number of patients <250 across all groups
- <sup>3</sup> Randomisation method, allocation concealment, and blinding not reported, ITT analysis not used.
- <sup>4</sup> Very small numbers of patients (51)
- <sup>5</sup> Randomisation method and allocation concealment not reported. Open label study. ITT analysis not used and losses to follow-up > 10%.

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16

0 per 1000	0 per 1000 (0 to 0)
Medium risk population	
0 per 1000	0 per 1000 (0 to 0)

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

**GRADE Working Group grades of evidence**

**High quality:** Further research is very unlikely to change our confidence in the estimate of effect.  
**Moderate quality:** Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.  
**Low quality:** Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.  
**Very low quality:** We are very uncertain about the estimate.

<sup>1</sup> Randomisation method and allocation concealment were not reported. Investigator blind only. No ITT analysis and loss to follow-up > 10%

<sup>2</sup> Very small numbers of patients (59) and CI is very wide and crosses 1

Re-infection	Study population		RR 0.88 (0.44 to 1.74)	211 (2 studies)	⊕○○○ very low <sup>1,5</sup>
	147 per 1000	129 per 1000 (65 to 256)			
	Medium risk population				
	194 per 1000	136 per 1000 (65 to 268)			

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

**GRADE Working Group grades of evidence**

**High quality:** Further research is very unlikely to change our confidence in the estimate of effect.  
**Moderate quality:** Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.  
**Low quality:** Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.  
**Very low quality:** We are very uncertain about the estimate.

<sup>1</sup> No study reported blinding. One study was quasi-RCT using alternation; the other two studies did not report randomisation method.

<sup>2</sup> One study reported allocation concealment and two studies reported ITT analyses.

<sup>3</sup> Number of patients across groups was reasonably small (265) and CIs are wide and cross 1

<sup>4</sup> No explanation was provided

<sup>5</sup> CI crosses 1

<sup>6</sup> Randomisation method and blinding were not reported in either study. Allocation concealment and ITT analysis was adequate in one of the two studies.

**10-day cefadroxil versus 10-day ampicillin for treating lower urinary tract infection in children**

**Patient or population:** children with lower urinary tract infection  
**Settings:** not stated  
**Intervention:** 10-day cefadroxil  
**Comparison:** 10-day ampicillin

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk				
	10-day ampicillin	10-day cefadroxil				
Persistent bacteriuria	Study population		RR 0.33 (0.01 to 7.62)	32 (1 study)	⊕○○○ very low <sup>1,2</sup>	
	62 per 1000	21 per 1000 (1 to 480)				
	Medium risk population					
	63 per 1000	21 per 1000 (1 to 480)				
Persistent symptoms	Study population		RR 0.33 (0.01 to 7.62)	22 (1 study)	⊕○○○ very low <sup>1,2</sup>	
	62 per 1000	21 per 1000 (1 to 480)				
	Medium risk population					
	63 per 1000	21 per 1000 (1 to 480)				

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

**10-day trimethoprim versus 10-day trimethoprim + sulfamethoxazole for treating lower urinary tract infection in children**

**Patient or population:** children with lower urinary tract infection  
**Settings:** outpatients department  
**Intervention:** 10-day trimethoprim  
**Comparison:** 10-day trimethoprim + sulfamethoxazole

Outcomes	Illustrative comparative risks* (95% CI)		Relative effect (95% CI)	No of Participants (studies)	Quality of the evidence (GRADE)	Comments
	Assumed risk	Corresponding risk				
	10-day trimethoprim + sulfamethoxazole	10-day trimethoprim				
Persistent bacteriuria	Study population		RR 1.93 (0.38 to 9.76)	59 (1 study)	⊕○○○ very low <sup>1,2</sup>	
	69 per 1000	135 per 1000 (26 to 673)				
	Medium risk population					
	69 per 1000	133 per 1000 (26 to 673)				
Persistent symptoms	Study population		RR 4.84 (0.24 to 96.66)	59 (1 study)	⊕○○○ very low <sup>1,2</sup>	
	0 per 1000	0 per 1000 (0 to 0)				
	Medium risk population					
	0 per 1000	0 per 1000 (0 to 0)				
Recurrence	Study population		RR 2.8 (0.12 to 68.5)	59 (1 study)	⊕○○○ very low <sup>1,2</sup>	

RR 1.05 (0.99 to 1.11). The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison (and its 95% confidence interval).

GRADE Working Group grades of evidence  
 High quality: further research is very unlikely to change our confidence in the estimate of effect.  
 Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.  
 Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.  
 Very low quality: we are very uncertain about the estimate.

<sup>1</sup> Randomisation method, allocation concealment and blinding not reported.  
<sup>2</sup> Very small number of patients (CI and CrI are very wide and crossed 1).

Single-dose trimethoprim versus single-dose nitrofurantoin for treating lower urinary tract infection in children

Effect of nitrofurantoin: children with lower urinary tract infection  
 Settings: Outpatient department  
 Intervention: single-dose trimethoprim  
 Comparison: single-dose nitrofurantoin

Outcomes	Illustrative comparative risk <sup>a</sup> (95% CI)	Corresponding risk <sup>b</sup>	Relative effect (95% CI)	No. of Participants (studies)	Quality of the evidence (GRADE)	Comments
Persistent bacteriuria	Assumed risk	Single-dose nitrofurantoin				
	Study population	31 per 1000	RR 0.15 (0.08 to 0.24)	115 (1 study)	⊕○○○ very low <sup>c</sup>	
	Median risk population	98 per 1000 (2.7 to 454)				
Recurrence	Assumed risk	Single-dose nitrofurantoin				
	Study population	115 per 1000	RR 0.63 (0.45 to 0.85)	115 (1 study)	⊕○○○ very low <sup>c</sup>	
	Median risk population	185 per 1000 (4.1 to 243)				

RR 1.05 (0.99 to 1.11). The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison (and its 95% confidence interval).

GRADE Working Group grades of evidence  
 High quality: further research is very unlikely to change our confidence in the estimate of effect.  
 Moderate quality: further research is likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.  
 Low quality: further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.  
 Very low quality: we are very uncertain about the estimate.

<sup>1</sup> Randomisation method, allocation concealment and blinding not reported.  
<sup>2</sup> Number of patients is reasonably small (143) and CrI are very wide and crossed 1.

**DISCUSSION**

**Summary of main results**

UTIs are relatively common childhood illness and require antibiotic treatment to alleviate symptoms and clear infection from the urinary tract. This review was designed to include all randomised and quasi-RCTs addressing all aspects of antibiotic treatment for children with lower UTI. Relatively few studies investigated the same comparisons, therefore meta-analyses included only a small number of studies. Nevertheless, we did not identify many instances of heterogeneity. The included studies predominantly examined bacteriological outcomes (persistent bacteriuria, recurrence, re-infection) rather than clinical outcomes (persisting symptoms) as measures of treatment efficacy. Amoxicillin or amoxicillin + clavulanic acid, cephalosporins, nitrofurantoin or TMP-SMX were the most common antibiotics given in the included studies.

Compared to single-dose therapy, 10-day antibiotic treatment was more effective in eliminating bacteriuria (RR 2.01, 95% CI 1.06 to 3.89). When we limited our analysis to only include those studies using the same antibiotic (amoxicillin) in both treatment groups, the differences no longer remained statistically significant (RR 1.97, 95% CI 0.99 to 4.33). It is unclear whether this is attributable to bias considering most of our included studies were of poor quality and sample sizes were too small to detect differences or whether just by chance. Nevertheless, the actual numbers of children with persistent bacteriuria, 18/63 (29%) in the single-dose group compared to 8/68 (12%) of the 10-day group, represents a clinically significant difference. No differences were observed for persistent bacteriuria, recurrence or re-infection following treatment for any other dosing regimen. In this review, there were not enough data to draw conclusions about these results at this stage, the lack of significant differences is more likely due to the inclusion of small, poorer quality studies, rather than demonstrating equivalence between different antibiotic or dosing regimens.

**Overall completeness and applicability of evidence**

While there were 16 RCTs included in this review, the number of children analysed totalled 1116 (1331 randomised), only three studies included more than 100 children. The median sample size was small (49 children), and wide CIs around most of the effect

estimates suggest that studies probably lacked the statistical power to identify such differences. Of the 16 studies included, 13 were conducted between 1981 and 1990. Diagnosis of bacteriuria since this time has not changed significantly, but not all of the antibiotics used in these studies remain available. In most of the meta-analyses carried out, there were few studies included, preventing a thorough investigation of the sources of heterogeneity between study results. In particular, we could not explore the influence of specific sources of bias or methodological quality, and most importantly we could not offer results stratified by age subgroup. Adverse events were reported by some studies, but were not analysed by any included studies; this hindered our efforts to present a full picture of the benefits and harms of antibiotic treatment for lower UTI, which was our intended aim.

**Quality of the evidence**

The quality of the included studies for every comparison was 'very low' according to GRADE criteria (see *Summary of findings table*). The lack of reporting of randomisation method, allocation concealment and blinding in most studies, and the large losses to follow-up in these studies are likely to contribute to issues in the results reported. Despite the inclusion of 16 studies, their methodological weakness and small sample sizes made it difficult to conclude if any of the included antibiotics or regimens were superior to another.

It was unfortunate that no study specifically addressed whether the efficacy of therapies differed according to patients age; we had planned subgroup analyses to investigate this because there are claims that UTIs can lead to long-term damage in younger children (Venoo 1997). In Fine 1985, 90% of the adolescents included were sexually active; this may have caused bias in that sexually active people (particularly women) are known to experience more frequent UTIs than those who abstain (Lehovich 1987). Four studies included young people over the age of 16 years: Shapiro 1981 included children aged 2 to 18 years, but the mean age was 5.6 years; Stahl 1984 included children aged 2 to 17 years, but the mean age was 4.75 years; Kowowski 1999 included children aged 0 to 18 years, but 50% were younger than 16 years. Fine 1985 included female adolescents aged 12 to 18 years with a mean age of 16.3 years. Although a lower tract UTI in an adolescent female is likely a very different condition than lower tract UTI in an infant or young child, post-hoc sensitivity analysis, removing Fine 1985 did not alter the conclusions for persistent bacteriuria.

None of the included studies systematically collected data on the adverse effects of antibiotics; this made our original objective of summarising the benefits and harms of antibiotic treatment difficult. Although it is not possible to compare the benefits and harms of antibiotic therapy from the included studies, from the few adverse antibiotic effects reported it seems unlikely that the expected side-effects course of antibiotics would be significant.



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Georghiou A, Boshniakov V, Jung A. Clinical assessment of Uro-Vaxom in the treatment and prophylaxis of recurrent urinary tract infections in children: preliminary results (Oena klinična preiskava profylakse i terapije Uro-Vaxom u bolesnika s recidivirajućim urinarnim infekcijama djetinjstva). *Medicinski glasnik* 2000;8(4):242-3. [MEDLINE: 1697636]

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**Granados 1998** (published data only)  
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\* Indicates the major publication for the study

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## CHARACTERISTICS OF STUDIES

### Characteristics of included studies (ordered by study ID)

Ahmed 2001

Methods	<ul style="list-style-type: none"> <li>Study design: parallel RCT</li> <li>Study period: NS</li> </ul>	
Participants	<p><b>Inclusion criteria</b></p> <ul style="list-style-type: none"> <li>Setting/recruitment: outpatient department</li> <li>Country: USA</li> <li>Children aged between 6 months and 12 years with signs and symptoms of UTI, significant bacteriuria defined as <math>&gt; 10^5</math> cfu/mL, and the presence of organisms susceptible to TMP and TMP-SMX. Urine collection method not reported.</li> <li>Number: 125 randomised, 59 analysed               <ul style="list-style-type: none"> <li>Treatment group: 30</li> <li>Control group: 29</li> </ul> </li> </ul> <p><b>Exclusion criteria:</b> NS</p>	
Interventions	<p><b>Treatment group</b></p> <ul style="list-style-type: none"> <li>10-day TMP (monotherapy; 10 mg/kg/d) in 2 doses</li> </ul> <p><b>Control group</b></p> <ul style="list-style-type: none"> <li>10-day TMP (8 mg/kg/d) + (SMX 40 mg/kg/d) in 2 doses</li> </ul>	
Outcomes	<ul style="list-style-type: none"> <li>Persistent bacteriuria (16-19 days following treatment)</li> <li>Persistent symptoms (16-19 days following treatment)</li> <li>Recurrence (16-19 days following treatment)</li> </ul>	
Notes	Source of funding: NS	
<b>Risk of bias</b>		
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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Avner 1983

Methods	<ul style="list-style-type: none"> <li>Study design: parallel RCT</li> <li>Study period: NS</li> </ul>	
Participants	<p><b>Inclusion criteria</b></p> <ul style="list-style-type: none"> <li>Setting/recruitment: emergency and outpatient departments, Children's Hospital Medical Centre, Massachusetts General Hospital, and Cambridge Hospital, Boston</li> <li>Country: USA</li> <li>Children aged between 2 and 12 years with acute, lower UTIs were eligible for inclusion. Children were required to have symptoms of abdominal pain, urinary frequency, dysuria, or abnormal urinalyses consisting of hematuria with pyuria, and two midstream clean catch or one suprapubic aspiration urine culture <math>&gt; 10^5</math> cfu/mL.</li> <li>Number: 49 randomised, 49 analysed               <ul style="list-style-type: none"> <li>Treatment group: 24</li> <li>Control group: 25</li> </ul> </li> <li>Sex (M/F): treatment group (2/22); control group (2/23)</li> </ul> <p><b>Exclusion criteria</b></p> <ul style="list-style-type: none"> <li>Acutely ill with temperature <math>&gt; 38^\circ\text{C}</math>, discrete flank pain, rigours, or signs of systemic toxicity; known renal disease or other systemic illness; allergic to penicillin</li> </ul>	
Interventions	<p><b>Treatment group</b></p> <ul style="list-style-type: none"> <li>Single-dose amoxicillin: <math>&lt; 23</math> kg (1.0 g), 23 to 32 kg (1.5 g), 32 to 45 kg (2.0 g); <math>&gt; 45</math> kg (3.0 g)</li> </ul> <p><b>Control group</b></p> <ul style="list-style-type: none"> <li>Conventional 10-day amoxicillin: <math>&lt; 23</math> kg (125 mg); <math>&gt; 23</math> kg (250 mg)</li> <li>Three times daily</li> </ul>	
Outcomes	<ul style="list-style-type: none"> <li>Persistent bacteriuria (4 days following treatment)</li> </ul>	
Notes	<ul style="list-style-type: none"> <li>Study presents data for children with and without known abnormalities. Results for children without abnormalities are presented in this review.</li> <li>Source of funding: Hoffman-La Roche Company</li> </ul>	
<b>Risk of bias</b>		
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 randomized were analysed
Selective reporting (reporting bias)	Low risk	Planned outcomes were all analysed

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CSG 1991

Methods	<ul style="list-style-type: none"> <li>Study design: parallel RCT</li> <li>Study period: NS</li> </ul>	
Participants	<p><b>Inclusion criteria</b></p> <ul style="list-style-type: none"> <li>Setting/recruitment: 10 hospital paediatric departments in and around Copenhagen</li> <li>Country: Denmark</li> <li>Children aged 1 to 15 years with clinical symptoms requiring immediate treatment, significant bacteriuria defined as <math>\geq 10^5</math> cfu/mL of a single bacterium in a clean catch mid-stream urine sample. Bag samples of urine were not accepted</li> <li>Number: 359 randomised, 264 analysed; 168 included in this review               <ul style="list-style-type: none"> <li>Treatment group: 90</li> <li>Control group: 78</li> </ul> </li> <li>Sex (M/F): All female</li> </ul> <p><b>Exclusion criteria</b></p> <ul style="list-style-type: none"> <li>Antibiotic treatment one week prior to inclusion; suspicion of allergy to penicillin or sulfonamides; required parental antibiotic treatment; fever <math>&gt; 39^\circ\text{C}</math> or impaired general condition; <math>\text{SG} &gt; 120</math> <math>\mu\text{mol/L}</math>; known severe urinary tract malformations; immunosuppressive treatment or known immunodeficiency</li> </ul>	
Interventions	<p><b>Treatment group</b></p> <ul style="list-style-type: none"> <li>Pivocillinam, 20-40 mg/kg/d in 2 doses for 3 days</li> </ul> <p><b>Control group</b></p> <ul style="list-style-type: none"> <li>Sulfamethizole, 40-80 mg/kg/d in 2 doses for 10 days</li> </ul>	
Outcomes	<ul style="list-style-type: none"> <li>Persistent bacteriuria (1-10 days following treatment)</li> </ul>	
Notes	<ul style="list-style-type: none"> <li>Children included had no previous UTI (17%), a history of UTI (31%), or recurrent UTI (5%)</li> <li>Another intervention arm was included in this study, 3-day sulfamethizole.</li> <li>A Cochrane review by Michael 2003 reports outcomes for this comparison which is not reported in this review.</li> <li>Source of funding: Danish Medical Research Council (5.52.11.10 and 5.52.14.80)</li> </ul>	
<b>Risk of bias</b>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	To ensure an equal number of patients in each group, a block randomisation method was used. Randomisation was in blocks of 6 within each of the 10 participating departments. No details about the way the block randomisation was performed were reported

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CSG 1991 (Continued)

Allocation concealment (selection bias)	Low risk	Allocation concealed by drawing consecutively numbered sealed envelopes prepared by the manufacturer
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	30 children did not fulfil inclusion criteria (26 bacteriuria not significant, 10 provided bag sample); treatment was discontinued in 6 children before scheduled; 32 children did not have urine cultures completed within 10 days from treatment; 2 children were not evaluated for other reasons; 19 boys were excluded because of the small number and because they were not evenly distributed between groups. The side effects of the 95 children who were not analysed were included as they received treatment
Selective reporting (reporting bias)	Low risk	Planned outcomes were all analysed

Fine 1985

Methods	<ul style="list-style-type: none"> <li>Study design: parallel RCT</li> <li>Study period: NS</li> </ul>	
Participants	<p><b>Inclusion criteria</b></p> <ul style="list-style-type: none"> <li>Setting/recruitment: outpatient: Adolescent General Medical Clinic, University of Maryland Hospital</li> <li>Country: USA</li> <li>Female adolescents aged 12 to 18 years with clinical symptoms of an acute lower UTI (frequency, dysuria, urgency, lower abdominal pain, urgency, anorexia, low-grade fever or malaise) and significant bacteriuria defined as <math>&gt; 10^5</math> cfu/mL in a clean catch mid-stream urine sample</li> <li>Mean age: 16.5 years</li> <li>Number: 34 randomised, 31 analysed               <ul style="list-style-type: none"> <li>Treatment group: 16</li> <li>Control group: 15</li> </ul> </li> <li>Sex (M/F): all female</li> </ul> <p><b>Exclusion criteria</b></p> <ul style="list-style-type: none"> <li>Pregnancy; pyelonephritis; allergy to penicillin or concurrent antibiotic use</li> </ul>	
Interventions	<p><b>Treatment group</b></p> <ul style="list-style-type: none"> <li>Single-dose amoxicillin 3.0 g</li> </ul> <p><b>Control group</b></p>	

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Flue 1985 (Continued)

		<ul style="list-style-type: none"> <li>10-day amoxicillin 250 mg, 3 times/day</li> </ul>
Outcomes		<ul style="list-style-type: none"> <li>Persistent bacteriuria (2-5 days following treatment)</li> <li>Persistent symptoms (2-5 days following treatment)</li> </ul>
Notes		<ul style="list-style-type: none"> <li>28/31 participants were sexually active</li> <li>Source of funding: Maternal and Child Health Grant (MCH #000980)</li> </ul>
<b>Risk of bias</b>		
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	<ul style="list-style-type: none"> <li>Study design: parallel RCT</li> <li>Study period: NS</li> </ul>
Participants	<p><b>Inclusion criteria</b></p> <ul style="list-style-type: none"> <li>Setting/recruitment: outpatients department, Christchurch Hospital</li> <li>Country: New Zealand</li> <li>Children aged 2 weeks to 12 years with significant bacteriuria defined as <math>&gt; 10^5</math> cfu/mL in 2 consecutive clean catch urine samples or any growth on supra-pubic aspiration. Children with cystitis were definite or had fever <math>&gt; 38^\circ\text{C}</math>, no loin pain or tenderness and were without other significant systemic symptoms.</li> <li>Mean age: 4.9 years</li> <li>Number of participants: 45 children</li> </ul> <p><b>Exclusion criteria</b></p> <ul style="list-style-type: none"> <li>Children with pyelonephritis were also included in this study and were reported separately (and excluded from this review).</li> </ul>

Grimwood 1988 (Continued)

Interventions	<p><b>Treatment group</b></p> <ul style="list-style-type: none"> <li>Single intramuscular gentamicin injection 3 mg/kg</li> </ul> <p><b>Control group</b></p> <ul style="list-style-type: none"> <li>7-day course of appropriate antibiotic depending on culture sensitivity in standard doses (included TMP-SMX, amoxicillin, cephalosporins).</li> </ul>	
Outcomes	<ul style="list-style-type: none"> <li>Persistent bacteriuria (1 day following treatment)</li> <li>Recurrence (<math>\leq</math> 1 week following treatment)</li> <li>Re-infection (<math>\leq</math> 1 week following treatment)</li> </ul>	
Notes	<ul style="list-style-type: none"> <li>23 children with 3 or more proven UTIs during the preceding 12 months were defined as having a history of recurrent UTIs.</li> <li>Source of funding: National Children's Health Research Foundation</li> </ul>	
<b>Risk of bias</b>		
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	Low risk	All patients randomized were analysed
Selective reporting (reporting bias)	Low risk	Planned outcomes were all analysed

Helin 1984

Methods	<ul style="list-style-type: none"> <li>Study design: parallel RCT</li> <li>Study period: NS</li> </ul>
Participants	<p><b>Inclusion criteria</b></p> <ul style="list-style-type: none"> <li>Setting/recruitment: NS</li> <li>Country: Sweden</li> <li>Children aged under 15 years with at least 2 clinical symptoms of a UTI (including frequency, dysuria, urgency and enuresis) and significant bacteriuria defined as <math>&gt; 10^5</math> cfu/mL in a clean catch mid-stream urine sample</li> <li>Mean age: 7.2 years</li> <li>Number: treatment group (19); control group (24)</li> <li>Sex (M/F): 1/42</li> </ul> <p><b>Exclusion criteria</b></p>

Helin 1984 (Continued)

		<ul style="list-style-type: none"> <li>Signs or laboratory findings suggesting upper urinary tract involvement (fever <math>&gt; 38.5^\circ\text{C}</math>, flank pain, elevated ESR and leukocytosis); known sensitivity to cephalosporins and nitrofurantoin; neurogenic bladder disorder; known structural malformation of the kidneys</li> </ul>
Interventions	<p><b>Treatment group</b></p> <ul style="list-style-type: none"> <li>3-day cephalosin 25-50 mg/kg/d in 2 doses</li> </ul> <p><b>Control group</b></p> <ul style="list-style-type: none"> <li>10-day nitrofurantoin 3-4 mg/kg/d in 2 or 3 doses</li> </ul>	
Outcomes		<ul style="list-style-type: none"> <li>Persistent bacteriuria (4-7 days following treatment)</li> <li>Recurrence (any time during follow-up; mean 8 months)</li> <li>Re-infection (any time during follow-up; mean 8 months)</li> </ul>
Notes		Source of funding: NS
<b>Risk of bias</b>		
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 randomized were analysed
Selective reporting (reporting bias)	Low risk	Planned outcomes were all analysed

Khan 1981

Methods	<ul style="list-style-type: none"> <li>Study design: Quasi-RCT</li> <li>Study period: NS</li> </ul>
Participants	<p><b>Inclusion criteria</b></p> <ul style="list-style-type: none"> <li>Setting/recruitment: Jewish Hospital and Medical Centre of Brooklyn, State University of New York, Downstate Medical Centre</li> <li>Country: USA</li> <li>Children aged six months to 15 years with symptoms of cystitis (including frequency and dysuria without fever) and significant bacteriuria defined as <math>&gt; 10^5</math> cfu/mL in 2 consecutive clean catch urine samples.</li> <li>Mean age: 5.65 years</li> <li>Number: treatment group (27); control group (27)</li> </ul>

Khan 1981 (Continued)

		<ul style="list-style-type: none"> <li>Sex (M/F): 4/50</li> </ul> <p><b>Exclusion criteria</b></p> <ul style="list-style-type: none"> <li>Younger than 6 months, or older than 15 years; urinary tract malformations; abnormal SCr or BUN values.</li> </ul>
Interventions	<p><b>Treatment group</b></p> <ul style="list-style-type: none"> <li>2-day treatment</li> </ul> <p><b>Control group</b></p> <ul style="list-style-type: none"> <li>10-day treatment</li> </ul> <p>Antimicrobial agents were 'chosen at random' for both groups and included ampicillin, sulfisoxazole and cephalosin in conventional doses given orally 4 times/day</p>	
Outcomes		<ul style="list-style-type: none"> <li>Persistent bacteriuria (3-7 days following treatment)</li> <li>Recurrence (<math>&gt;</math> 2 months following treatment)</li> </ul>
Notes		Source of funding: NS
<b>Risk of bias</b>		
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)	Low risk	Planned outcomes were analysed. Data for re-infection was presented across cystitis, pyelonephritis and asymptomatic bacteriuria and was not reported for cystitis alone

Komarovski 1999

Methods	<ul style="list-style-type: none"> <li>Study design: parallel RCT</li> <li>Study period: NS</li> </ul>
Participants	<p><b>Inclusion criteria</b></p> <ul style="list-style-type: none"> <li>Setting/recruitment: outpatient and emergency department, Arkansas Children's Hospital</li> <li>Country: USA</li> <li>Children aged 1 to 19 years with at 1 or more clinical symptoms of cystitis</li> </ul>

Komoroski 1999 (Continued)

	<p>(including frequency, dysuria, emesis, haematuria, pyuria, suprapubic tenderness) and significant bacteriuria defined as <math>10^5</math> cfu/mL of a single organism from 1 catheterized bladder specimen or <math>&gt;10^4</math> cfu/mL from a nitrite-positive specimen. Urine cultures were also considered positive if <math>&gt; 10^4</math> cfu/mL of a single organism was obtained from a single clean-catch specimen, and the second specimen contained organisms of the same in vitro sensitivity pattern as the first specimen.</p> <ul style="list-style-type: none"> <li>Number: 93 randomized, 59 analysed</li> <li>Treatment group: 36</li> <li>Control group: 23</li> </ul> <p>Exclusion criteria</p> <ul style="list-style-type: none"> <li>Pregnancy antibiotic therapy in the previous 2 weeks; concomitant infection requiring additional antibiotic therapy known renal or urologic problems that could predispose to a UTI; signs and symptoms of pyelonephritis (ill or toxic appearance, flank pain, costovertebral angle tenderness, or temperature <math>\geq 38.3^\circ\text{C}</math>); history of hypersensitivity to cephalosporins or penicillin; a significant history of gastrointestinal, hematologic, hepatic, psychiatric, or central nervous system disease; history of drug or alcohol abuse; history of sexual abuse as a child; a parent or guardian who was unable to understand or follow instructions; a family situation in which follow-up could not be assured; or refusal to obtain a catheterized sample, if necessary.</li> </ul>	
Interventions	<p>Treatment group</p> <ul style="list-style-type: none"> <li>Single intramuscular ceftriaxone 50 mg/kg (to a maximum of 500 mg)</li> <li>27 received ceftriaxone (500 mg); 9 received ceftriaxone (250 mg)</li> </ul> <p>Control group</p> <ul style="list-style-type: none"> <li>TMP-SMX 4.5 mg/kg twice daily for 10 days</li> <li>22 received TMP-SMX; 1 patient received amoxicillin because of sulfa hypersensitivity</li> </ul>	
Outcomes	<ul style="list-style-type: none"> <li>Persistent bacteriuria (10-30 days following treatment)</li> <li>Persistent bacteriuria and symptoms (10-30 days following treatment)</li> </ul>	
Notes	<ul style="list-style-type: none"> <li>Nine patients receiving ceftriaxone 250 mg were excluded from this analysis. It appears the study was originally designed to investigate ceftriaxone 250 mg versus control. The ceftriaxone 500 mg group was added at a later date. When high treatment failures were reported, the 250 mg group was discontinued. The ceftriaxone 250 mg group is not reported as part of this review; using block randomization, the groups should be of equal size. It does not seem logical that only 9 children were allocated to the ceftriaxone 250 mg group when more than double this number were allocated to the other 2 groups.</li> <li>Numbers reported in the text do not match the numbers reported in the tables. Numbers reported in tables have been used for this review.</li> <li>Source of funding: Roche Laboratories Inc.</li> </ul>	
<b>Risk of bias</b>		
Bias	Authors' judgement	Support for judgement
Random sequence generation (selection bias)	Unclear risk	Not reported
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43

Komoroski 1999 (Continued)

Allocation concealment (selection bias)	Unclear risk	Not 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 criteria were applied. 23 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 assessment
Selective reporting (reporting bias)	Unclear risk	Relapse and recurrence were reported, but not in a format suitable for data extraction for this review
<p>Lidefelt 1991</p>		
Methods	<ul style="list-style-type: none"> <li>Study design: parallel RCT</li> <li>Study period: 1986-1988</li> </ul>	
Participants	<p>Inclusion criteria</p> <ul style="list-style-type: none"> <li>Setting/recruitment: emergency department, Sachs' Children's Hospital, Stockholm</li> <li>Country: Sweden</li> <li>Children aged less than 3 years to 12 years with symptoms of a UTI (including frequency, dysuria, and painful micturition) and significant bacteriuria defined as <math>\geq 10^5</math> cfu/mL in 2 separately voided urine samples. Children were required to have had not more than 2 previous UTIs, and the most recent at least 6 months prior to the start of the study.</li> <li>Median age: 5 years</li> <li>Number: treatment group (50); control group (50)</li> <li>Sex (M/F): 1/87</li> </ul> <p>Exclusion criteria</p> <ul style="list-style-type: none"> <li>Signs of upper tract involvement (temperature <math>&lt; 38.5^\circ\text{C}</math>; absence of loin pain, ESR <math>&lt; 20</math> mm/h); more than 2 previous UTIs</li> </ul>	
Interventions	<p>Treatment group</p> <ul style="list-style-type: none"> <li>Single-dose TMP 6 mg/kg</li> </ul> <p>Control group</p> <ul style="list-style-type: none"> <li>5-day TMP 3 mg/kg/12 h</li> </ul>	
Outcomes	<ul style="list-style-type: none"> <li>Persistent bacteriuria (7 days following treatment)</li> <li>Recurrence (&gt; 7 days following treatment)</li> </ul>	
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44

Lidefelt 1991 (Continued)

Notes	<ul style="list-style-type: none"> <li>Source of funding: Swedish Medical Council, grant number 19X765, and the Swedish Society of Medicine.</li> </ul>	
<b>Risk of bias</b>		
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 to which they were assigned
Selective reporting (reporting bias)	Low risk	Planned outcomes were analysed
<p>Malaka-Zafrafi 1984</p>		
Methods	<ul style="list-style-type: none"> <li>Study design: parallel RCT</li> <li>Study period: NS</li> </ul>	
Participants	<p>Inclusion criteria</p> <ul style="list-style-type: none"> <li>Setting/recruitment: NS</li> <li>Children aged 8 months to 11.1 years with significant bacteriuria defined as <math>\geq 10^5</math> cfu/mL of a single pathogen in 2 consecutive mid-stream urine samples</li> <li>Number: treatment group (16); control group (16)</li> </ul> <p>Exclusion criteria</p> <ul style="list-style-type: none"> <li>Hypersensitivity to cephalosporins or penicillins; abnormal hepatic, renal function, or structural anomalies</li> </ul>	
Interventions	<p>Treatment group</p> <ul style="list-style-type: none"> <li>Cefadroxil 25 mg/kg once daily for 10 days</li> </ul> <p>Control group</p> <ul style="list-style-type: none"> <li>Ampicillin 50 mg/kg/d in 4 divided doses for 10 days</li> </ul>	
Outcomes	<ul style="list-style-type: none"> <li>Persistent bacteriuria (10 days following treatment)</li> <li>Persistent symptoms (10 days following treatment)</li> </ul>	
Notes	<ul style="list-style-type: none"> <li>Children with pyelonephritis were also included in this study and were reported separately.</li> <li>Source of funding: NS</li> </ul>	
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45

Malaka-Zafrafi 1984 (Continued)

<b>Risk of bias</b>		
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 to which they were assigned
Selective reporting (reporting bias)	Low risk	All planned outcomes were analysed
<p>Misnik 1985</p>		
Methods	<ul style="list-style-type: none"> <li>Study design: parallel RCT</li> <li>Study period: NS</li> </ul>	
Participants	<p>Inclusion criteria</p> <ul style="list-style-type: none"> <li>Setting/recruitment: Nephrology clinic of the Hospital Roberto del Rio and the Pediatric Clinic of the Chilean Air Force</li> <li>Country: Chile</li> <li>Children aged 2 years to 14 years with symptoms of a UTI (including frequency, dysuria, urgency, foul smelling urine, emesis and/or haematuria) and significant bacteriuria defined as <math>\geq 10^5</math> cfu/mL in voided urine sample, or <math>\geq 1000</math> cfu/mL on supra-pubic aspiration. Children were required to have had not more than 2 previous UTIs, and the most recent at least 6 months prior to the start of the study.</li> <li>Number: treatment group 1 (27); treatment group 2 (53); control group (36)</li> <li>Sex (M/F): 11/87</li> </ul> <p>Exclusion criteria</p> <ul style="list-style-type: none"> <li>Fever <math>&gt; 38^\circ\text{C}</math>; low back pain; history of UTI; anatomical abnormality; received antibiotics in the week prior to the study.</li> </ul>	
Interventions	<p>Treatment group 1</p> <ul style="list-style-type: none"> <li>3-day antibiotics</li> </ul> <p>Treatment group 2</p> <ul style="list-style-type: none"> <li>5-day antibiotics</li> </ul> <p>Control group</p> <ul style="list-style-type: none"> <li>10-day antibiotics</li> </ul> <p>Children were administered a first generation cephalosporin, nitrofurantoin or TMP-SMX depending on the sensitivity of the organism cultured</p>	
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46

Mitsak 1985 (Continued)

Outcomes	• Recurrence (at 2-3 months)	
Notes	• The 3-day and 5-day interventions were combined into one group and compared to the 10-day control • Source of funding: NS	
<b>Risk of bias</b>		
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

Principi 1990

Methods	• Study design: parallel RCT • Study period: NS	
Participants	<b>Inclusion criteria</b> <ul style="list-style-type: none"> <li>• Setting/recruitment: outpatients authors were from various university hospitals</li> <li>• Country: Italy</li> <li>• Children aged 1 month to 16 years with a lower UTI. Significant bacteriuria defined as <math>\geq 10^5</math> cfu/ml of a single pathogen in 2 clean catch or catheterised urine samples. Lower UTI was defined as absence of fever, ESR <math>&lt; 25</math> mm/L/h and CRP <math>&lt; 20</math> pg/ml.</li> <li>• Number: treatment group (71); control group (64)</li> <li>• Sex (M/F): 45/99</li> </ul> <b>Exclusion criteria</b> <ul style="list-style-type: none"> <li>• Renal failure</li> </ul>	
Interventions	<b>Treatment group</b> <ul style="list-style-type: none"> <li>• Single-dose fosfomycin trometamol (2 g orally; 1 g in children <math>&lt; 1</math> year)</li> </ul> <b>Control group</b> <ul style="list-style-type: none"> <li>• Single-dose netilmicin (5 mg/kg intramuscularly)</li> </ul>	

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47

Principi 1990 (Continued)

Outcomes	• Persistent bacteriuria (2-4 days following treatment) • Recurrence (up to 30 days following treatment)	
Notes	• Source of funding: NS	
<b>Risk of bias</b>		
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

Sanchez 1990

Methods	• Study design: parallel RCT • Study period: NS	
Participants	<b>Inclusion criteria</b> <ul style="list-style-type: none"> <li>• Setting/recruitment: emergency department, Hospital Materno-Infantil Vall d'Hebron, Barcelona</li> <li>• Country: Spain</li> <li>• Children aged 8 months to 11.1 years with significant bacteriuria defined as <math>\geq 10^5</math> cfu/ml of a single pathogen in 2 consecutive mid-stream urine samples.</li> <li>• Number: 37</li> </ul> <b>Exclusion criteria</b> <ul style="list-style-type: none"> <li>• Children aged less than 4 months, with a fever of <math>\geq 38.5^\circ\text{C}</math>, back pain or mass, malaise, duration of symptoms longer than one week, vomiting, received antibiotics in the previous 2 weeks, underlying disease involving immunosuppression, or known urinary tract malformation were excluded</li> </ul>	
Interventions	<ul style="list-style-type: none"> <li>• Children received amoxicillin, amoxicillin + clavulanic acid, cephalosin, TMP or co-trimoxazole at standard doses for 7 days.</li> </ul>	
Outcomes	• Persistent bacteriuria (2-3 days following treatment)	

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48

Sanchez 1990 (Continued)

Notes	• This study was published as an abstract • Source of funding: NS	
<b>Risk of bias</b>		
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	Unclear risk	Abstract only, not enough detail provided
Selective reporting (reporting bias)	Unclear risk	Abstract only, not enough detail provided

Shapiro 1981

Methods	• Study design: parallel RCT • Study period: NS	
Participants	<b>Inclusion criteria</b> <ul style="list-style-type: none"> <li>• Setting/recruitment: emergency department, Children's Hospital of Pittsburgh</li> <li>• Country: USA</li> <li>• Girls aged 2 to 18 years with symptoms of a UTI (including frequency, dysuria and/or urgency) and significant bacteriuria defined as <math>\geq 10^5</math> cfu/ml in 2 clean catch urine samples, or <math>\geq 1000</math> cfu/ml on supra-pubic aspiration.</li> <li>• Mean age: 5.6 years</li> <li>• Number: 37 randomised, 35 analysed <ul style="list-style-type: none"> <li>o Treatment group: 18</li> <li>o Control group: 17</li> </ul> </li> </ul> <b>Exclusion criteria</b> <ul style="list-style-type: none"> <li>• Clinical evidence of upper UTI (fever <math>&gt; 38^\circ\text{C}</math> and/or flank pain); known anatomic or functional urinary tract abnormality; currently receiving antibiotics; history of penicillin allergy</li> </ul>	
Interventions	<b>Treatment group</b> <ul style="list-style-type: none"> <li>• Single-dose amoxicillin 50 mg/kg (to a maximum of 2.5 g)</li> </ul> <b>Control group</b> <ul style="list-style-type: none"> <li>• 10-day amoxicillin 40 mg/kg/d in 3 divided doses (to a maximum of 500 mg/dose)</li> </ul>	

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49

Shapiro 1981 (Continued)

Outcomes	• Persistent bacteriuria (2 days following treatment) • Recurrence (within 3 months following treatment)	
Notes	Source of funding: Not reported	
<b>Risk of bias</b>		
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	Patients and physician
Incomplete outcome data (attrition bias) All outcomes	Unclear risk	Two children were excluded from analysis because the second urine culture was negative
Selective reporting (reporting bias)	Low risk	All planned outcomes were reported

Stahl 1984

Methods	• Study design: parallel RCT • Study period: NS	
Participants	<b>Inclusion criteria</b> <ul style="list-style-type: none"> <li>• Setting/recruitment: outpatients and emergency department, Children's Hospital of Philadelphia and St Christopher's Hospital for Children, Philadelphia</li> <li>• Country: USA</li> <li>• Girls aged 2 to 17 years with symptoms of a UTI (including frequency, dysuria, urgency, enuresis, suprapubic pain, or haematuria with pyuria) and significant bacteriuria defined as <math>\geq 10^5</math> cfu/ml of a single organism in 2 sequential clean catch urine samples.</li> <li>• Median age: 4.75 years</li> <li>• Number of participants: 36 randomised, 26 analysed <ul style="list-style-type: none"> <li>o Treatment group: 10</li> <li>o Control group: 16</li> </ul> </li> </ul> <b>Exclusion criteria</b> <ul style="list-style-type: none"> <li>• Signs or symptoms of upper UTI (temperature <math>&gt; 38.9^\circ\text{C}</math>, flank pain, costovertebral angle tenderness or toxic appearance); known renal or urologic disorder; history of penicillin allergy; received antibiotics in the previous 2 weeks</li> </ul>	

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50

Stahl 1984 (Continued)

Interventions	Treatment group • Single-dose amoxicillin 50 mg/kg orally (to a maximum of 3 g) Control group • 10-day amoxicillin 30 mg/kg/d in 3 divided doses (to a maximum of 250 mg/dose)	
Outcomes	• Persistent bacteriuria (2-4 days following treatment) • Re-infection (> 2 weeks following treatment)	
Notes	• Data on re-infection could not be used from this study as the definition included both a positive culture more than 2 weeks following therapy of any organism (defined as recurrence by this review) or an infection caused by a different organism (defined as re-infection by this review). These results were presented together. • Source of funding: Beecham Laboratories, Bristol, Tennessee	
<b>Risk of bias</b>		
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	High risk	Six girls were lost to follow-up, in 3 girls the 2nd urine culture was negative and 1 girl had received antibiotics within the previous 2 weeks. One girl in the single-dose group had an amoxicillin resistance organism and was switched to 10 days TMP-SMX and then followed in the conventional therapy group
Selective reporting (reporting bias)	Unclear risk	All planned outcomes were reported

Wallen 1983

Methods	• Study design: parallel RCT • Study period: NS	
Participants	Inclusion criteria • Setting/recruitment: outpatients, The Children's Memorial Hospital, Chicago • Country: USA • Girls aged 1 year to 12 years with suspected UTI and significant bacteriuria	

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Wallen 1983 (Continued)

	defined as $\geq 10^6$ cfu/ml <i>E. coli</i> organisms in 2 clean catch or urine collection bag samples. • Median age 5.45 years • Number: 54 randomised, 49 analysed ◦ Treatment group: 26 ◦ Control group: 23 Exclusion criteria • Clinical symptoms of pyelonephritis (including fever > 38.3°C, flank pain, chills, ESR $\geq 21$ mm/h) previous UTIs antibiotic use during the week prior to the study, known urinary tract abnormalities	
Interventions	Treatment group • Single-dose intramuscular amikacin sulfate 7.5 mg/kg (to a maximum of 240 mg) Control group • 10-day sulfisoxazole 150 mg/kg/day in 4 divided doses	
Outcomes	• Persistent bacteriuria (2-4 days following treatment) • Recurrence (30-40 days following treatment)	
Notes	• Re-infection rates were presented, but were only available for the single-dose amikacin group; these rates have not been reported in this review. • Source of funding: NS	
<b>Risk of bias</b>		
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 lost 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

BLUN - blood urea nitrogen; CRP - C-reactive protein; ESR - erythrocyte sedimentation rate; NS - not stated; SCR - serum creatinine; SMX - sulfamethoxazole; TMP - trimethoprim; UTI - urinary tract infection

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Characteristics of excluded studies (ordered by study ID)

Study	Reason for exclusion
Adam 1982	Children with pre-existing conditions, and who have symptoms of pyelonephritis are not reported separately from children with lower UTI
Annala 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
Arieta 2001	Included children had pyelonephritis
Aucher 1973	Not an RCT; screening study only
Babar 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)
Bakkaloglu 1996	Included children had pyelonephritis
Belez 2004	Prophylaxis for preventing recurrence
Bose 1974	More than half of the included children had pre-existing renal abnormalities
Bourillon 1994	Included children had pyelonephritis
Caparelli 1983	Some children had pyelonephritis; unclear how many
Carapeta 2001	Most included children had systemic symptoms
Caroddi 1987	Study is conducted in symptomatic and asymptomatic children, but proportion of each is unknown. Also, 11/51 children had known renal abnormalities
Carlen 1985	Prophylactic antibiotics
Chahane 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

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(Continued)

De Gaiser 1988	One third of included children had cystitis 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)
Francois 1995	Included children had pyelonephritis
Francoise 1997	Included children had pyelonephritis
Fuji 1987	Some children had pyelonephritis and were not reported separately from children without
Gandreas 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
Gok 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
Gooss 2006	Not a RCT, or quasi-RCT
Gooss 2007	Not RCT
Goszyk 2000	Children received 3 months antibiotic treatment for preventing recurrence
Gozd 1975	Unclear if participants were children. Included participants had prostatic, acute cystitis, urethritis, and/or vaginitis 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
Hypathidis 1970	Some children had pyelonephritis and were not reported separately from those without
Holin 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

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(Continued)

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 Michalek 2003
Johari 1991	Comparison of short versus standard duration antibiotic for lower UTI - included in Michalek 2003
Kenda 1995	Not RCT
Khan 1987	Not RCT
Kornberg 1994	Comparison of short versus standard duration antibiotic for lower UTI - included in Michalek 2003
Krepler 1976	Included children had pyelonephritis
Lohr 1981	Comparison of short versus standard duration antibiotic for lower UTI - included in Michalek 2003
Lubina 1984	Symptoms not reported. 35% of included children had renal abnormalities
Madrigal 1988	Comparison of short versus standard duration antibiotic for lower UTI - included in Michalek 2003
Marild 2009	Included children were required to have fever
McCracken 1981	> 20% of children had fever, abdominal/flank pain and contralateral tenderness indicating pyelonephritis
Moe 1977	Not all included children had bacteriologically proven UTI
Montini 2003	Children had pyelonephritis
Montini 2007	Children had pyelonephritis
Nolan 1989	Half of the included children had fever, loin pain and/or back pain
Noorbaksh 2004	Included children had pyelonephritis
Olking 1971	Some children had renal abnormalities; although the results refer to patients with and without abnormalities, no numbers are included so data cannot be extracted
Paloux 1986	Half of included children had known renal abnormalities
Pitt 1982	More than half of the included children had abdominal pain and/or fever
Pylkkanen 1981	Compared 10-day treatment with 42-day treatment in children
Repeno 1984	Children with fever were not analysed separately from children without fever
Rodriguez 1983	Included children had fever

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15

(Continued)

Rubens 1984	52% of children had fever and were not reported separately from those without
Russo 1989	Majority of included children had fever
Schach 1972	Most children received concomitant surgical therapy
Sembei 1985	Some patients had fever and/or lumbar pain and were not reported separately from patients without
Stanfeld 1975	Symptoms not reported. Approximately half of included children had reflux, but grade of reflux was not reported
Stigmann 1985	Most included children had fever
Sullivan 1980	No symptoms of UTI reported. Bacteriological definition of UTI only
Tambic 1992	As per Michalek 2003. Study was excluded because significantly more patients (32/59) with pyelonephritis 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
Tapanya 1999	Included children were required to have fever
Tong 2005	Some children had pyelonephritis but were not reported separately from those without
Toporovskii 1988	Included children presented with fever
Varese 1987	One third of included children had known renal abnormalities and are not presented separately from those without
Vlaskovic 1972	Included children had pyelonephritis
Vlaskovic 1974	Included children had pyelonephritis
Weber 1982	More than half of the included children had fever and were not reported separately from those without
Wiemers 1979	Comparison of short versus standard duration antibiotic for lower UTI - included in Michalek 2003
Zaki 1986	Comparison of short versus standard duration antibiotic for lower UTI - included in Michalek 2003

randomised controlled trial- RCT; UTI - urinary tract infection

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16

## 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
1 Persistent bacteriuria	6	238	Risk Ratio (M-H, Random, 95% CI)	2.01 [1.06, 3.80]
1.1 Amoxicillin	4	131	Risk Ratio (M-H, Random, 95% CI)	1.97 [0.96, 4.33]
1.2 Other antibiotics	2	97	Risk Ratio (M-H, Random, 95% CI)	2.09 [0.71, 6.18]
2 Persistent symptoms	1		Risk Ratio (M-H, Random, 95% CI)	Totals not selected
3 Recurrence	2	79	Risk Ratio (M-H, Random, 95% CI)	1.38 [0.55, 3.50]
4 Persistent bacteriuria 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
1 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 [0.45, 5.26]
3 Re-infection	1		Risk Ratio (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]

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17

### 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
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

### Comparison 6. Single-dose fosfomycin versus single-dose nitroimidazole

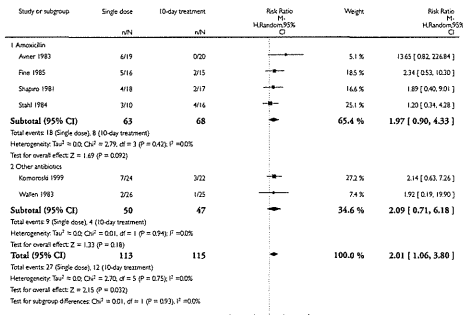
Outcome or subgroup title	No. of studies	No. of participants	Statistical method	Effect size
1 Persistent bacteriuria	1		Risk Ratio (M-H, Random, 95% CI)	Totals not selected
2 Recurrence	1		Risk Ratio (M-H, Random, 95% CI)	Totals not selected

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18

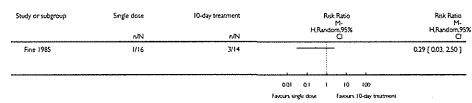
**Analysis 1.1. Comparison 1 Single-dose versus conventional 10-day treatment, Outcome 1 Persistent bacteriuria.**

Review: Antibiotics for treating lower urinary tract infection in children  
 Comparison: 1 Single-dose versus conventional 10-day treatment  
 Outcome: 1 Persistent bacteriuria



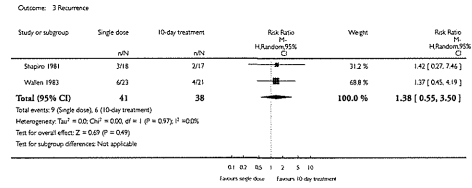
**Analysis 1.2. Comparison 1 Single-dose versus conventional 10-day treatment, Outcome 2 Persistent symptoms.**

Review: Antibiotics for treating lower urinary tract infection in children  
 Comparison: 1 Single-dose versus conventional 10-day treatment  
 Outcome: 2 Persistent symptoms



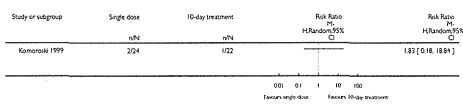
**Analysis 1.3. Comparison 1 Single-dose versus conventional 10-day treatment, Outcome 3 Recurrence.**

Review: Antibiotics for treating lower urinary tract infection in children  
 Comparison: 1 Single-dose versus conventional 10-day treatment  
 Outcome: 3 Recurrence



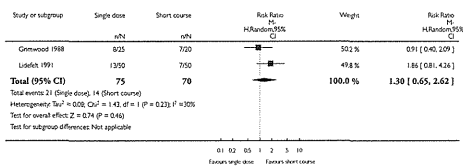
**Analysis 1.4. Comparison 1 Single-dose versus conventional 10-day treatment, Outcome 4 Persistent bacteriuria and symptoms.**

Review: Antibiotics for treating lower urinary tract infection in children  
 Comparison: 1 Single-dose versus conventional 10-day treatment  
 Outcome: 4 Persistent bacteriuria and symptoms



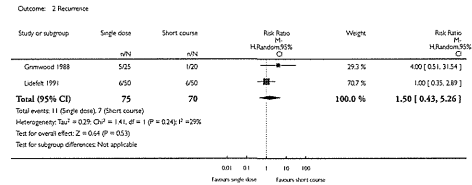
**Analysis 2.1. Comparison 2 Single-dose versus short-course (3-7 days) treatment, Outcome 1 Persistent bacteriuria.**

Review: Antibiotics for treating lower urinary tract infection in children  
 Comparison: 2 Single-dose versus short-course (3-7 days) treatment  
 Outcome: 1 Persistent bacteriuria



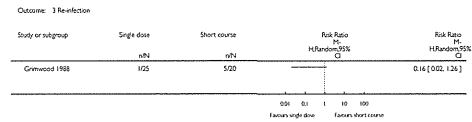
**Analysis 2.2. Comparison 2 Single-dose versus short-course (3-7 days) treatment, Outcome 2 Recurrence.**

Review: Antibiotics for treating lower urinary tract infection in children  
 Comparison: 2 Single-dose versus short-course (3-7 days) treatment  
 Outcome: 2 Recurrence



**Analysis 2.3. Comparison 2 Single-dose versus short-course (3-7 days) treatment, Outcome 3 Re-infection.**

Review: Antibiotics for treating lower urinary tract infection in children  
 Comparison: 2 Single-dose versus short-course (3-7 days) treatment  
 Outcome: 3 Re-infection

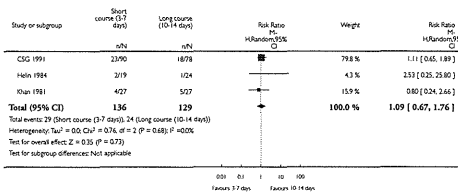


**Analysis 3.1. Comparison 3 Short-course (3-7 days) versus long-course (10-14 days) treatment, Outcome 1 Persistent bacteriuria.**

Review: Antibiotics for treating lower urinary tract infection in children

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

Outcome: 1 Persistent bacteriuria

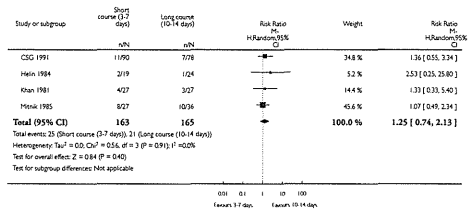


**Analysis 3.2. Comparison 3 Short-course (3-7 days) versus long-course (10-14 days) treatment, Outcome 2 Recurrence.**

Review: Antibiotics for treating lower urinary tract infection in children

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

Outcome: 2 Recurrence

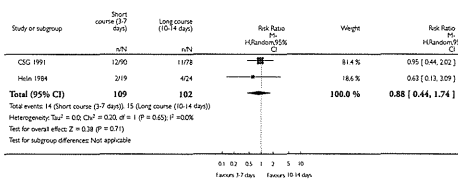


**Analysis 3.3. Comparison 3 Short-course (3-7 days) versus long-course (10-14 days) treatment, Outcome 3 Re-infection.**

Review: Antibiotics for treating lower urinary tract infection in children

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

Outcome: 3 Re-infection

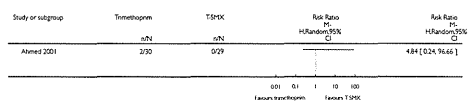


**Analysis 4.2. Comparison 4 Trimethoprim (10 days) versus trimethoprim+sulfamethoxazole (10 days), Outcome 2 Persistent symptoms.**

Review: Antibiotics for treating lower urinary tract infection in children

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

Outcome: 2 Persistent symptoms

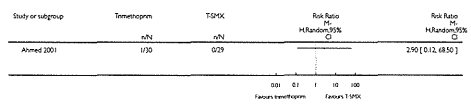


**Analysis 4.3. Comparison 4 Trimethoprim (10 days) versus trimethoprim+sulfamethoxazole (10 days), Outcome 3 Recurrence.**

Review: Antibiotics for treating lower urinary tract infection in children

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

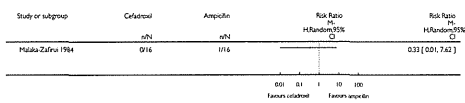
Outcome: 3 Recurrence





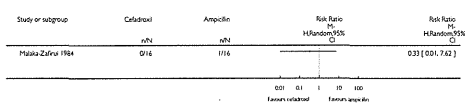
**Analysis 5.1. Comparison 5 Cefadroxil (10 days) versus ampicillin (10 days), Outcome 1 Persistent bacteriuria.**

Review: Antibiotics for treating lower urinary tract infection in children  
 Comparison: 5 Cefadroxil (10 days) versus ampicillin (10 days)  
 Outcome: 1 Persistent bacteriuria



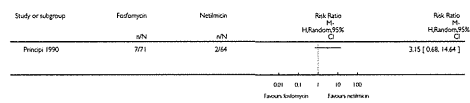
**Analysis 5.2. Comparison 5 Cefadroxil (10 days) versus ampicillin (10 days), Outcome 2 Persistent symptoms.**

Review: Antibiotics for treating lower urinary tract infection in children  
 Comparison: 5 Cefadroxil (10 days) versus ampicillin (10 days)  
 Outcome: 2 Persistent symptoms



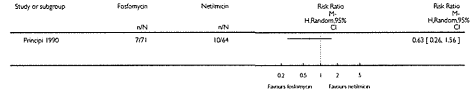
**Analysis 6.1. Comparison 6 Single-dose fosfomycin versus single-dose netilmicin, Outcome 1 Persistent bacteriuria.**

Review: Antibiotics for treating lower urinary tract infection in children  
 Comparison: 6 Single-dose fosfomycin versus single-dose netilmicin  
 Outcome: 1 Persistent bacteriuria



**Analysis 6.2. Comparison 6 Single-dose fosfomycin versus single-dose netilmicin, Outcome 2 Recurrence.**

Review: Antibiotics for treating lower urinary tract infection in children  
 Comparison: 6 Single-dose fosfomycin versus single-dose netilmicin  
 Outcome: 2 Recurrence



**APPENDICES**

**Appendix 1. Electronic search strategy**

Database	Search terms used
CENTRAL	1. child*.ti,ab,kw 2. (infant* or babies or neonat* or newborn* or toddler*).ti,ab,kw 3. (adolescent* or pubert* or pubesc* or prepubert* or prepubes* or juvenile* or youth* or teen*).ti,ab,kw 4. (pediatr* or paediatr*).ti,ab,kw 5. (boys or girls).ti,ab,kw 6. (#1 OR #2 OR #3 OR #4 OR #5) 7. MeSH descriptor Urinary Tract Infections explode all trees 8. MeSH descriptor Cystitis explode all trees 9. MeSH descriptor Pyelonephritis, this term only 10. urinary tract mucosa infection*.kw 11. cystitis*.ti,ab,kw 12. pyeloneph*.ti,ab,kw 13. bacteriuri*.ti,ab,kw 14. (pyuria or pyuria* or pyuria).ti,ab,kw 15. (uti or uti).ti,ab,kw 16. ((bladder* or genitourin* or renal or ureter* or ureth* or urin* or urinal* or urogen*) near5 (infect* or bacteri* or microbio*))ti,ab 17. (bladder* near5 (acute* or ulcra)).ti,ab 18. (#7 OR #8 OR #9 OR #10 OR #11 OR #12 OR #13 OR #14 OR #15 OR #16 OR #17) 19. #6 AND #18 20. SR-RENAL 21. #19 AND NOT 20 22. Anti-infective next agentickw 23. Antiinfective next agentickw 24. MeSH descriptor Anti-Infective Agents, Utiary explode all trees 25. antibiotic next agentickw 26. antibiotic*.ti,ab 27. bacteriocid*.ti,ab 28. ((antimycobacteri* or antibacteri* or bacteriocid*) near5 agent*).ti,ab 29. ((antiseptic* or anti-infective* or antimicrobic*) near5 unia*).ti,ab 30. penicillin*.ti,ab,kw 31. amoxic*.ti,ab,kw 32. amoxicil*.ti,ab,kw 33. augmentin*.ti,ab,kw 34. ampicillin*.ti,ab,kw 35. penbrtin*.ti,ab,kw 36. ceftazid*.ti,ab,kw 37. ceftioxi*.ti,ab,kw 38. cefadrox*.ti,ab,kw 39. cefproz*.ti,ab,kw 40. kefzol*.ti,ab,kw 41. ceftim*.ti,ab,kw

(Continued)

- 42. supra\*.ti,ab,kw
- 43. ceftazim\*.ti,ab,kw
- 44. ceforan\*.ti,ab,kw
- 45. klaforan\*.ti,ab,kw
- 46. cephalosporin\*.ti,ab,kw
- 47. cefiprone\*.ti,ab,kw
- 48. ceftriaxon\*.ti,ab,kw
- 49. cefepime\*.ti,ab,kw
- 50. erodax\*.ti,ab,kw
- 51. ceftin\*.ti,ab,kw
- 52. vintox\*.ti,ab,kw
- 53. cefixime\*.ti,ab,kw
- 54. forum\*.ti,ab,kw
- 55. ceftriaxon\*.ti,ab,kw
- 56. neoptilim\*.ti,ab,kw
- 57. cefprozim\*.ti,ab,kw
- 58. sinacep\*.ti,ab,kw
- 59. sinacep\*.ti,ab,kw
- 60. gentamicin\*.ti,ab,kw
- 61. cidebmycin\*.ti,ab,kw
- 62. genticin\*.ti,ab,kw
- 63. methemycin\*.ti,ab,kw
- 64. hexamin\*.ti,ab,kw
- 65. hiprez\*.ti,ab,kw
- 66. nitrofurantoin\*.ti,ab,kw
- 67. furandiazol\*.ti,ab,kw
- 68. macrodantin\*.ti,ab,kw
- 69. nitrothoprim\*.ti,ab,kw
- 70. trimethoprim\*.ti,ab,kw
- 71. monocerin\*.ti,ab,kw
- 72. andimocillin\*.ti,ab,kw
- 73. mefloquine\*.ti,ab,kw
- 74. selaxid\*.ti,ab,kw
- 75. amikacin\*.ti,ab,kw
- 76. aminoglycosid\*.ti,ab,kw
- 77. aminoglycosid\*.ti,ab,kw
- 78. tobramycin\*.ti,ab,kw
- 79. robecin\*.ti,ab,kw
- 80. robecin\*.ti,ab,kw
- 81. quinolone\*.ti,ab,kw
- 82. \*4-quinolone\*.kw
- 83. \*4-Quinolone Derivative\*.kw
- 84. netilmicin\*.ti,ab,kw
- 85. neillin\*.ti,ab,kw
- 86. (#22 OR #23 OR #24 OR #25 OR #26 OR #27 OR #28 OR #29 OR #30 OR #31 OR #32 OR #33 OR #34 OR #35 OR #36 OR #37 OR #38 OR #39 OR #40 OR #41 OR #42 OR #43 OR #44 OR #45 OR #46 OR #47 OR #48 OR #49 OR #50 OR #51 OR #52 OR #53 OR #54 OR #55 OR #56 OR #57 OR #58 OR #59 OR #60 OR #61 OR #62 OR #63 OR #64 OR #65 OR #66 OR #67 OR #68

(Continued)

68 OR #69 OR #70 OR #71 OR #72 OR #73 OR #74 OR #75 OR #76 OR #77 OR #78 OR #79 OR #80 OR #81 OR #82 OR #83 OR #84 OR #85  
87. (#21 AND #80)

MEDLINE (OVID SP)

1. exp Child/
2. exp Infant/
3. Adolescent/
4. Puberty/
5. child\$.rw
6. (pediatric or paediatric).rw
7. (boys or girls).rw
8. (infant\$ or babies or neonat\$ or newborn\$ or toddler\$).rw
9. (adolescent\$ or pubert\$ or pubesc\$ or prepubert\$ or juvenile\$ or youth\$ or teen\$).rw
10. or/1-9
11. exp Urinary Tract Infection/
12. exp Cystitis/
13. Pyelonephritis/
14. (uti or urti).rw
15. bacteriuria\$.rw
16. (pyuria or pyuric or pyuria).rw
17. cystitis.rw
18. (bladder\$ adj\$ (acute\$ or chronic)).rw
19. ((bladder\$ or genitourin\$ or renal or ureter\$ or ureth\$ or urin\$ or urolog\$ or urogen\$) adj\$ (infect\$ or bacteri\$ or microbiol\$)).rw
20. pyelonephri\$.rw
21. pyelocysti\$.rw
22. or/11-21
23. and/10,22
24. Anti-Infective Agent/
25. Anti-Bacterial Agent/
26. exp Anti-Infective Agents, Urinary/
27. antibiotic\$.rw
28. bacteriocid\$.rw
29. antibacteri\$.rw
30. antimicrobacteri\$.rw
31. antiinfect\$.rw
32. antiinfective\$.rw
33. anti-infective.rw
34. Penicillin/
35. penicillin\$.rw
36. Amoxicillin/
37. amoxicilli\$.rw
38. amoxicilli\$.rw
39. amoxicil\$.rw
40. Amoxicillin-Clavulanic Acid Combination/
41. augmentin\$.rw
42. Ampicillin/
43. ampicillin\$.rw

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71

(Continued)

44. penicillin\$.rw
45. Cefadroxil/
46. cefadroxilli\$.rw
47. cefadroxilli\$.rw
48. cephalos\$.rw
49. Cephalosin/
50. cephalosin\$.rw
51. cefaclor\$.rw
52. cefaclor\$.rw
53. ceporex\$.rw
54. ketec\$.rw
55. Cefixime/
56. cefiximi\$.rw
57. cephalim\$.rw
58. suprax\$.rw
59. Cefprozime/
60. cefprozimi\$.rw
61. cephoxim\$.rw
62. claforan\$.rw
63. klaforan\$.rw
64. Cephaloplatin/
65. cephaloplati\$.rw
66. Ceftriaxime/
67. ceftriaximi\$.rw
68. cepodoxim\$.rw
69. ceftriax\$.rw
70. Cephradine/
71. cefradini\$.rw
72. cefradini\$.rw
73. Cefadime/
74. cefadime\$.rw
75. formim\$.rw
76. Ceftriaxone/
77. ceftriaxon\$.rw
78. rocephin\$.rw
79. Cefuroxime/
80. cefuroxim\$.rw
81. sinactin\$.rw
82. sinactin\$.rw
83. Gentamicin/
84. gentamicin\$.rw
85. edomycin\$.rw
86. genticin\$.rw
87. Methenamine/
88. methenamin\$.rw
89. heamim\$.rw
90. hiprex.rw
91. Nitrofurantoin/

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71

(Continued)

92. nitrofurantoi\$.rw
93. furadantin\$.rw
94. macrodantin\$.rw
95. Trimethoprim/
96. Trimethoprim-Sulfamethoxazole Combination/
97. trimethoprim\$.rw
98. monosulfim\$.rw
99. Amdinocillin/
100. amdinocilli\$.rw
101. mactinam\$.rw
102. adexic\$.rw
103. Amikacin/
104. amikacin\$.rw
105. Aminoglycoside/
106. aminoglycoside\$.rw
107. aminoglycoside\$.rw
108. Tobramycin/
109. tobramycin\$.rw
110. nebecin\$.rw
111. tobi.rw
112. Quinolones/
113. 4-Quinolones/
114. quinolone\$.rw
115. Neflomicin/
116. neflomicin\$.rw
117. neclini\$.rw
118. or/24-117
119. and/23,118

EMBASE (OVID SP)

1. exp Child/
2. exp Newborn/
3. Adolescent/
4. exp Adolescence/
5. exp Childhood/
6. child\$.rw
7. (pediat\$ or paediat\$).rw
8. (boys or girls).rw
9. (infant\$ or babies or neonat\$ or newborn\$ or toddler\$).rw
10. (adolescent\$ or pubert\$ or pubesc\$ or prepubert\$ or juvenile\$ or youth\$ or teen\$).rw
11. or/1-10
12. exp Urinary Tract Infection/
13. exp Cystitis/
14. exp Pyelonephritis/
15. Bacteriuria/
16. Pyuria/
17. (uti or urti).rw
18. bacteriuria\$.rw
19. (pyuria or pyuric or pyuria).rw
20. cystitis.rw

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71

(Continued)

21. (bladder\$ adj\$ (acute\$ or chronic)).rw
22. ((bladder\$ or genitourin\$ or renal or ureter\$ or ureth\$ or urin\$ or urolog\$ or urogen\$) adj\$ (infect\$ or bacteri\$ or microbiol\$)).rw
23. pyelonephri\$.rw
24. pyelocysti\$.rw
25. or/12-24
26. and/11,25
27. Anti-Infective Agent/
28. Antibiotic Agent/
29. antibiotic\$.rw
30. bacteriocid\$.rw
31. ((antimicrobacteri\$ or antibacteri\$ or bacteriocid\$) adj\$ agent\$).rw
32. Penicillin Derivative/ or Penicillin G/
33. penicilli\$.rw
34. exp Urinary Tract Anti-Infective Agent/
35. ((antiseptic\$ or antiinfective\$) adj\$ urin\$).rw
36. Amoxicillin/
37. amoxicilli\$.rw
38. amoxicilli\$.rw
39. amoxicil\$.rw
40. Amoxicillin Plus Clavulanic Acid/
41. augmentin\$.rw
42. Ampicillin/
43. ampicillin\$.rw
44. penicillin\$.rw
45. Cefadroxil/
46. cefadroxilli\$.rw
47. cephalos\$.rw
48. Cefalexin/
49. cephalosin\$.rw
50. cefalexin\$.rw
51. cefaclor\$.rw
52. ceporex\$.rw
53. ketec\$.rw
54. Cefixime/
55. cefiximi\$.rw
56. cephalim\$.rw
57. suprax\$.rw
58. Cefprozime/
59. cefprozimi\$.rw
60. cephoxim\$.rw
61. claforan\$.rw
62. klaforan\$.rw
63. Cephaloplatin Derivative/
64. cephaloplati\$.rw
65. Cefuroxime/
66. cefuroxim\$.rw
67. cepodoxim\$.rw

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71

(Continued)

- 68. orcloxS.rw
- 69. CefradinS.rw
- 70. cefradinS.rw
- 71. cephradins.rw
- 72. vcloxS.rw
- 73. CefadimS.rw
- 74. cefadimS.rw
- 75. forumS.rw
- 76. CeftriaxolS.rw
- 77. ceftriaxolS.rw
- 78. rocephins.rw
- 79. CefuroximS.rw
- 80. cefuroximS.rw
- 81. zinatS.rw
- 82. zinatS.rw
- 83. GentamicinS.rw
- 84. gentamicinS.rw
- 85. cidomycinS.rw
- 86. genticinS.rw
- 87. MethenaminS.rw
- 88. methenaminS.rw
- 89. hexaminS.rw
- 90. lipexS.rw
- 91. NitrofurantoinS.rw
- 92. nitrofurantoinS.rw
- 93. furandantS.rw
- 94. macrodantS.rw
- 95. TrimethoprimS.rw
- 96. CotrimoxazoleS.rw
- 97. nitrothoprimS.rw
- 98. monorinS.rw
- 99. MecillinamS.rw
- 100. meclimamS.rw
- 101. andimocillinS.rw
- 102. selaxS.rw
- 103. AmikacinS.rw
- 104. amikacinS.rw
- 105. AminoglycosideS.rw
- 106. aminoglycosideS.rw
- 107. TobramycinS.rw
- 108. TobramycinS.rw
- 109. Tobramycin SulfateS.rw
- 110. tobramycinS.rw
- 111. neobiS.rw
- 112. ubiS.rw
- 113. QuinoloneS.rw
- 114. 4-Quinolone DerivativeS.rw
- 115. quinoloneS.rw

(Continued)

- 116. NeohimicilS.rw
- 117. neohimicilS.rw
- 118. neohimicilS.rw
- 119. orf27-118
- 120. and26,119

### Appendix 2. Risk of bias assessment tool

Potential source of bias	Assessment criteria
<b>Random sequence generation</b> Selection bias (biased allocation to interventions) due to inadequate generation of a randomised sequence	<b>Low risk of bias:</b> Random number table; computer random number generator; coin tossing; shuffling cards or envelopes; throwing dice; drawing of lots; minimisation (minimisation may be implemented without a random element, and this is considered to be equivalent to being random)  <b>High risk of bias:</b> Sequence generated by odd or even date of birth; date (or day) of admission; sequence generated by hospital or clinic record number; allocation by judgement of the clinician; by preference of the participant; based on the results of a laboratory test or a series of tests by availability of the intervention  <b>Unclear:</b> Insufficient information about the sequence generation process to permit judgement
<b>Allocation concealment</b> Selection bias (biased allocation to interventions) due to inadequate concealment of allocations prior to assignment	<b>Low risk of bias:</b> Randomisation method described that would not allow investigator/participant to know or influence intervention group before eligible participant entered in the study (e.g. central allocation, including telephone, web-based, and pharmacy-controlled, randomisation; sequentially numbered drug containers of identical appearance; sequentially numbered, opaque, sealed envelopes)  <b>High risk of bias:</b> Using an open random allocation schedule (e.g. a list of random numbers); assignment envelopes were used without appropriate safeguards (e.g. if envelopes were unsealed or non-opaque or not sequentially numbered); alternation or rotation; date of birth case record number; any other explicitly uncontrolled procedure  <b>Unclear:</b> Randomisation stated but no information on method used is available
<b>Blinding of participants and personnel</b> Performance bias due to knowledge of the allocated interventions by participants and personnel during the study	<b>Low risk of bias:</b> No blinding or incomplete blinding, but the review authors judge that the outcome is not likely to be influenced by lack of blinding; blinding of participants and key study personnel ensured, and unlikely that the blinding could have been broken  <b>High risk of bias:</b> No blinding or incomplete blinding, and the outcome is likely to be influenced by lack of blinding; blinding of key study participants and personnel attempted, but likely that the blinding could have been broken, and the outcome is likely to be influenced by lack of blinding  <b>Unclear:</b> Insufficient information to permit judgement

(Continued)

<b>Blinding of outcome assessment</b> Detection bias due to knowledge of the allocated interventions by outcome assessors	<b>Low risk of bias:</b> No blinding of outcome assessment, but the review authors judge that the outcome measurement is not likely to be influenced by lack of blinding; blinding of outcome assessment ensured, and unlikely that the blinding could have been broken  <b>High risk of bias:</b> No blinding of outcome assessment, and the outcome measurement is likely to be influenced by lack of blinding; blinding of outcome assessment, but likely that the blinding could have been broken, and the outcome measurement is likely to be influenced by lack of blinding  <b>Unclear:</b> Insufficient information to permit judgement
<b>Incomplete outcome data</b> Attrition bias due to amount, nature or handling of incomplete outcome data	<b>Low risk of bias:</b> No missing outcome data; reasons for missing outcome data unlikely to be related to true outcome (for survival data, censoring unlikely to be introducing bias); missing outcome data balanced in numbers across intervention groups, with similar reasons for missing data across groups, for dichotomous outcome data, the proportion of missing outcomes compared with observed events risk not enough to have a clinically relevant impact on the intervention effect estimate; for continuous outcome data, plausible effect size (difference in means or standardised difference in means) among missing outcomes not enough to have a clinically relevant impact on observed effect size; missing data have been imputed using appropriate methods  <b>High risk of bias:</b> Reasons for missing outcome data likely to be related to true outcome, with either imbalance in numbers or reasons for missing data across intervention groups; for dichotomous outcome data, the proportion of missing outcomes compared with observed events risk enough to induce clinically relevant bias in intervention effect estimate; for continuous outcome data, plausible effect size (difference in means or standardised difference in means) among missing outcomes enough to induce clinically relevant bias in observed effect size; 'as-treated' analysis done with substantial departure of the intervention received from that assigned at randomisation; potentially inappropriate application of

(Continued)

<b>Selective reporting</b> Reporting bias due to selective outcome reporting	<b>Simple imputation</b> <b>Unclear:</b> Insufficient information to permit judgement  <b>Low risk of bias:</b> The study protocol is available and all of the study's pre-specified (primary and secondary) outcomes that are of interest in the review have been reported in the pre-specified way; the study protocol is not available but it is clear that the published reports include all expected outcomes, including those that were pre-specified (convincing text of this nature may be uncommon)  <b>High risk of bias:</b> Not all of the study's pre-specified primary outcomes have been reported; one or more primary outcomes are reported using measurements, analysis methods or subsets of the data (e.g. subgroups) that were not pre-specified; one or more reported primary outcomes were not pre-specified (unless clear justification for their reporting is provided, such as an unexpected adverse effect); one or more outcomes of interest in the review are reported incompletely so that they cannot be entered in a meta-analysis; the study report fails to include results for a key outcome that would be expected to have been reported for such a study  <b>Unclear:</b> Insufficient information to permit judgement
<b>Other bias</b> Bias due to problems not covered elsewhere in the table	<b>Low risk of bias:</b> The study appears to be free of other sources of bias  <b>High risk of bias:</b> Had a potential source of bias related to the specific study design; stopped early due to some data-dependent process (including a formal stopping rule); had extreme baseline imbalance; has been claimed to have been fraudulent; had some other problem  <b>Unclear:</b> Insufficient information to assess whether an important risk of bias exists; insufficient rationale or evidence that an identified problem will introduce bias

### HISTORY

Protocol first published: Issue 4, 2007  
Review first published: Issue 8, 2012

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
- Assessment for inclusion: AF, RM
- Quality assessment: AF, RM
- Data extraction: AF, RM
- Data entry into RevMan: AF
- Data analysis: AF, RM
- Disagreements resolution: ML, KT

#### DECLARATIONS OF INTEREST

- Antia Fitzgerald: Some of this work was undertaken 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 those of the authors and not necessarily those of NICE.
- Monica Lakhanpaul: I was the Clinical Director at the National Collaborating Centre for Women's Health and led the development of the NICE Urinary Tract Infection Guideline. I am no longer the Clinical Director but remain on the NCC-WCH board and I am a NICE Fellow and member of the NHS evidence advisory team.

#### 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-infection (growth of new bacteria)
- In some studies urine samples were collected using non-invasive methods (clean-catch, urine collection bag or pad) but if urine was unobtainable, several studies included the option of a supra-pubic aspiration or catheter samples. We included studies that collected urine using supra-pubic aspiration or catheters, as the difficulties in collecting urine from children, particularly infants can be problematic.
- We initially defined recurrence as at least three episodes of cystitis/lower 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.

#### INDEX TERMS Medical Subject Headings (MeSH)

Anti-Bacterial Agents [administration & dosage; \*therapeutic use]; Anti-Infective Agents, Urinary [administration & dosage; \*therapeutic use]; Bacteriuria [\*drug therapy]; Drug Administration Schedule; Infants, Newborn; Randomized Controlled Trials as Topic; Urinary Tract Infections [\*drug therapy]

#### MeSH check words

Adolescent; Child; Child, Preschool; Humans; Infant