National Institute for Health and Care Excellence

Draft

Urinary tract infection (recurrent): antimicrobial prescribing

[A] Evidence review NICE guideline NG112

October 2018 (amended 2024)

Update information

In 2024, we amended the recommendations on referral and seeking specialist advice, and on oestrogen. We did not review the evidence on these areas but made these changes based on committee expertise. See the <u>section in this evidence review about the 2024 guideline update</u> and the update information section in the <u>2024 version of this guideline</u> for more information.

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ISBN: xxx-x-xxxx-xxxx-x

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1 **1** Context

2 1.1 Background

Urinary tract infection (UTI) is a non-specific term that refers to infection anywhere in the
 urinary tract. This evidence review covers the prevention of UTI in women (including
 pregnant women), men and children with recurrent UTI, who do not have a catheter. Lower
 UTI, acute pyelonephritis, and catheter-associated UTI are covered in separate evidence
 reviews.

8 Recurrent UTI includes recurrence of lower UTIs (cystitis) and/or upper UTIs (acute

9 pyelonephritis), but repeated pyelonephritis should prompt further investigation. See NICE
 10 antimicrobial prescribing guidelines on <u>lower UTI</u> and <u>acute pyelonephritis</u> for background
 11 information.

- 12 Recurrent UTIs are repeated UTIs with a frequency of at least 3 UTIs in the last year or 2
- 13 UTIs in the last 6 months (European Association of Urology (EAU) guidelines on urological
- 14 <u>infections</u> [2017]). This may be due to relapse or reinfection:
- Relapse is recurrent UTI with the same strain of organism. Relapse is the likely cause if
 UTI recurs within a short period (for example within 2 weeks) after treatment.
- Reinfection is recurrent UTI with a different strain or species of organism. Reinfection is
 the likely cause if UTI recurs more than 2 weeks after treatment.

The number of recurrences that is regarded as clinically significant depends on the risks of
infection and the impact of UTI on the person (EAU guideline [2017]). Lower UTI (cystitis)
recurs within a year in 25 to 50% of women, usually as reinfections (rather than relapses)
(NICE clinical knowledge summary – UTI (lower) - women).

Recurrent UTIs are common in women. Risk factors in young and pre-menopausal women
include sexual intercourse, new sexual partner, mother with a history of UTI and history of
UTI as a child. In post-menopausal and elderly women, risk factors include history of UTI
before menopause, urinary incontinence, atrophic vaginitis due to oestrogen deficiency,

- 27 increased post-void urine volume, and urine catheterisation and functional status
- 28 deterioration in elderly institutionalised women (EAU guideline [2017]).
- Some people (mainly women) may be able to identify 1 or more triggers that often brings on a UTI. These triggers may vary for different people, and include sexual intercourse, going for long walks and wearing occlusive underwear.
- Risk factors that may predispose men to recurrent UTIs include abnormalities of urinary tract function or structure, incomplete bladder emptying and immunosuppression (NICE clinical knowledge summary, UTI (lower), men)
- 34 knowledge summary <u>UTI (lower) men</u>).
- 35 Risk factors for recurrent UTI in children include abnormalities of urinary tract function or
- 36 structure, for example vesicoureteric reflux, spinal abnormalities and constipation;
- dysfunctional elimination syndrome; and infection or irritation of the genital area that prevents
 regular voiding (NICE clinical knowledge summary <u>UTI children</u>).
- 39 The diagnosis of recurrent UTI should be confirmed by urine culture. Extensive routine
- 40 investigations such as cystoscopy and imaging are not routinely recommended, but may be
- 41 performed in some circumstances such as when renal calculi or outflow obstruction is
- 42 suspected (EAU guideline [2017]).
- The management of suspected community-acquired bacterial urinary tract infection in adults
 aged 16 years and over is covered in the NICE quality standard on <u>urinary tract infection in</u>

- adults (2015). This includes women who are pregnant, people with indwelling catheters and people with other diseases or medical conditions such as diabetes. The quality standard was developed to contribute to a reduction in emergency admissions for acute conditions that should not usually require hospital admission, and improvements in health-related quality of life. It includes a <u>placeholder statement</u> on the treatment of recurrent UTI, which is an area of error that has been prioritized by the Quality Standard Advisory Committee but for which is an
- 6 care that has been prioritised by the Quality Standards Advisory Committee but for which no
 7 source guidance was currently available. A placeholder statement indicates the need for
- 8 evidence-based guidance to be developed in this area.
- 9 The NICE guideline on <u>urinary tract infection in under 16s</u> (2007) defines recurrent UTI in 10 children as:
- 11 2 or more episodes of UTI with acute pyelonephritis/upper UTI, or
- 1 episode of UTI with acute pyelonephritis plus 1 or more episode of UTI with cystitis/lower UTI, or
- 3 or more episodes of UTI with cystitis/lower UTI.

15 The NICE guideline on urinary tract infection in under 16s (2007) makes recommendations 16 on the diagnosis of UTI in infants and children. All infants younger than 3 months with 17 suspected UTI should be referred to paediatric specialist care and a urine sample should be 18 sent for urgent microscopy and culture. These infants should be managed in accordance with 19 the recommendations for this age group in the NICE guideline on fever in under 5s (2013). 20 Infants and children who have had recurrent UTIs should undergo ultrasound (within 6 weeks) (see the NICE guideline on urinary tract infection in under 16s (2007) for more 21 22 information).

UTIs are usually caused by bacteria from the gastrointestinal tract entering the urethra and ascending into the bladder. The most common causative pathogen in uncomplicated UTIs, in 70 to 95% of cases, is *Escherichia coli* (*E. coli*). *Staphylococcus saprophyticus* accounts for 5 to 10% of cases and occasionally other Enterobacteriaceae, such as *Proteus mirabilis* and Klebsiella species are isolated.

1.2 Managing infections that require antibiotics

In most cases, managing a UTI will require antibiotic treatment, but antibiotics should only be started when there is clear evidence of infection. Antibiotic prophylaxis may also be an option in people with recurrent UTI, to reduce the risk of recurrent infections. The NICE guideline on urinary tract infection in under 16s (2017) recommends that antibiotic prophylaxis may be considered in infants and children with recurrent UTI.

34 1.2.1 Self-care

35 The NICE guideline on antimicrobial stewardship: changing risk-related behaviours in the

36 <u>general population</u> (2017) recommends that people should be given verbal advice and
 37 written information that they can take away about how to manage their infection themselves
 38 at home with self-care if it is safe to do so.

- 39 Self-care options that have been used to relieve symptoms in UTI include paracetamol or 40 non-steroidal anti-inflammatory drugs, cranberry products and urine alkalinising agents.
- 41 Other strategies have also been used to reduce the risk of recurrent infections. These
- 42 include avoiding known risk factors, behavioural changes (for example, reducing fluid intake,
- 43 habitual and post-coital delayed urination and wearing occlusive underwear), probiotics,
- 44 cranberry products and D-mannose (see <u>Clinical effectiveness</u>).

1 **1.2.2** Back-up antibiotic prescribing strategies

2 The NICE guideline on antimicrobial stewardship: changing risk-related behaviours in the

general population (2017) recommends that if the person has been given a <u>back-up antibiotic</u>
 prescription, they should be told:

- 4 prescription, they should be told:
- 5 How to self-care to manage their symptoms.
- What the antimicrobials would be used for, if needed.
- How to recognise whether they need to use the antimicrobials, and if so:
- 8 o how to get them 9 o when to start tak
 - \circ when to start taking or using them
- 10 \circ how to take them.

11 1.2.3 Antibiotic prescribing strategies

- 12 The NICE guideline on antimicrobial stewardship: systems and processes for effective
- antimicrobial medicine use (2015) recommends that when antimicrobials are prescribed,
 prescribers should:
- Consider supplying antimicrobials in pack sizes that correspond to local (where available)
 and national guidelines on course lengths.
- Follow local (where available) or national guidelines on prescribing the shortest effective course, the most appropriate dose, and route of administration.
- Undertake a clinical assessment and document the clinical diagnosis (including symptoms) in the patient's record and clinical management plan.
- Document in the patient's records (electronically wherever possible):
- 22 o the reason for prescribing an antimicrobial
- o the plan of care as discussed with the patient, their family member or carer (as appropriate), including the planned duration of any treatment.
- Take into account the benefits and harms for an individual patient associated with the particular antimicrobial, including:
- 27 o possible interactions with other medicines or any food and drink
- the patient's other illnesses, for example, the need for dose adjustment in a patient with
 renal impairment
- 30 o any drug allergies (these should be documented in the patient's record)
- the risk of selection for organisms causing healthcare associated infections, for
 example, *C. difficile*.

Document in the patient's records the reasons for any decision to prescribe outside local (where available) or national guidelines.

- The NICE guideline on antimicrobial stewardship: changing risk-related behaviours in the general population (2017) recommends that resources and advice should be available for
- 37 people who are prescribed antimicrobials to ensure they are taken as instructed at the
- 38 correct dose, via the correct route, for the time specified. Verbal advice and written
- information that people can take away about how to use antimicrobials correctly should begiven, including:
- not sharing prescription-only antimicrobials with anyone other than the person they were
 prescribed or supplied for
- not keeping them for use another time
- returning unused antimicrobials to the pharmacy for safe disposal and not flushing them
 down toilets or sinks.

1 1.3 Safety netting advice

2 The NICE guideline on antimicrobial stewardship: changing risk-related behaviours in the

3 general population (2017) recommends that safety netting advice should be shared with

everyone who has an infection (regardless of whether or not they are prescribed or suppliedwith antimicrobials). This should include:

- how long symptoms are likely to last with and without antimicrobials
- 7 what to do if symptoms get worse
- what to do if they experience adverse effects from the treatment
- when they should ask again for medical advice.

The NICE clinical knowledge summary on UTI (lower) - women suggests advising all women
 with recurrent UTI to seek medical attention if they:

- 12 develop fever or loin pain, because of suspected acute pyelonephritis, or
- do not respond to treatment with the first-choice antibiotic, because this may be due to a resistant organism.

15 For men with recurrent UTI, the NICE clinical knowledge summary on UTI (lower) – men

16 suggests that men are advised about measures that may reduce the risk of recurrent UTIs,

17 such as to maintain sufficient fluid intake (at least 2 litres per day) to avoid dehydration. If

18 hospital admission is not needed and empirical antibiotics are started, follow up should be

arranged, for example after 48 hours, to check the response to treatment and the urine

culture results. If symptoms persist after antibiotic treatment referral for specialist urologicalassessment may be needed.

The NICE guideline on urinary tract infection in under 16s (2007) recommends that all infants
 younger than 3 months with suspected UTI should be referred immediately to paediatric
 specialist care. All infants and children 3 months or older with recurrent UTI should be

25 assessed by a paediatric specialist.

1.4 Symptoms and signs of a more serious illness or condition (red flags)

Complications of lower UTI include ascending infection leading to pyelonephritis, renalfailure, and sepsis.

The NICE clinical knowledge summary on UTI (lower) - women suggests routinely referring
 the following women with recurrent UTIs:

- who have a risk factor for an abnormality of the urinary tract
- 33 who are immunocompromised or who have diabetes
- who have a known abnormality of their renal tract who might benefit from surgical correction
- who have not responded to preventive treatments.
- 37 In pregnancy, asymptomatic bacteriuria can lead to pyelonephritis; and symptomatic UTI has

been associated with developmental delay or cerebral palsy in the infant, and foetal death.

For women with visible or non-visible haematuria an urgent 2-week wait referral should be arranged if a urological cancer is suspected (NICE clinical knowledge summary on UTI

- 41 (lower) women).
- 42 For men with recurrent UTI, the NICE clinical knowledge summary on UTI (lower) men
- 43 suggests that alternative conditions such as urethritis are considered. At least 50% of men

1 with recurrent UTI will have prostate involvement, which may lead to complications such as

2 prostatic abscess or chronic bacterial prostatitis. Urinary stones are also a possibility, more

3 likely with *Proteus mirabilis* infection which is associated with stone formation in the renal

4 collecting ducts. Emergency admission to hospital is recommended if a man with a

5 suspected lower UTI is severely unwell with symptoms or signs suggestive of urosepsis (for 6 example nausea and vomiting, confusion, tachypnoea, tachycardia, or hypotension). If

example nausea and vomiting, confusion, tachypnoea, tachycardia, or hypotension). If
hospital admission is not needed and empirical antibiotics are started, follow up should be

arranged, for example after 48 hours, to check the response to treatment and the urine

9 culture results. If symptoms persist after antibiotic treatment referral for specialist urological

10 assessment may be needed.

11 Treatment failure (due to relapse or reinfection) is more likely in men with risk factors for

12 complications (see NICE antimicrobial prescribing guideline on UTI: acute pyelonephritis).

13 Prognosis partly depends on whether any underlying cause can be treated or removed, such

14 as urinary stone extraction. For men with suspected urological cancer an urgent 2-week

15 referral should be arranged. A non-urgent referral for bladder cancer should be considered in

16 men aged 60 years and over with recurrent or persistent unexplained UTI (NICE clinical

17 knowledge summary – UTI (lower) – men).

18 In children, UTIs can lead to renal scarring, but more often this is preceded by acute

19 pyelonephritis rather than cystitis, and it is more common in children with vesicoureteral

20 reflux. UTIs in childhood have also been associated with hypertension (if there is severe or

bilateral renal scarring) and renal insufficiency or failure (if febrile UTIs are treated late; NICE

22 clinical knowledge summary on UTI - children).

23 1.5 2024 Guideline update

An update of this guideline was conducted in 2024 which made new recommendations, or updated existing recommendations, in the below areas.

26 **1.5.1 Methenamine hippurate**

Methenamine hippurate is an antiseptic drug and was within the scope of the 2018 guideline
 as an antimicrobial pharmacological intervention. However, methenamine hippurate was
 classified as an antibiotic so it was grouped with antibiotics in the <u>Clinical effectiveness</u> and
 <u>Safety and tolerability sections</u> of this evidence review. No recommendations were made on
 the use of methenamine hippurate in the 2018 guideline.

As a result of <u>exceptional surveillance in 2022</u>, a new evidence review was conducted to examine the evidence on the effectiveness of methenamine hippurate prophylaxis as an alternative to antibiotic prophylaxis for treating recurrent UTI. In the 2024 evidence review, methenamine hippurate was classified as an antiseptic to clarify the distinction from antibiotics. Evidence supporting the 2024 recommendations on methenamine hippurate is described in the <u>2024 evidence review</u> and does not draw on the evidence on methenamine hippurate described in this evidence review.

39 1.5.2 Oestrogen

After publication of the 2018 guideline, stakeholder feedback was received regarding the
recommendation on vaginal oestrogen and what was meant by the 'lowest effective dose'.
The wording of the existing recommendation was reviewed by the committee for the 2024
update to improve clarity and implementation. They agreed that the phrase 'lowest effective
dose' was confusing and were aware that the BNF contains information about recommended
doses, so they removed this from the recommendation. They also removed the example of
estriol cream from the recommendation and agreed that preferences for different types of

1 vaginal oestrogen preparations should be discussed with the person because they did not

2 want to imply that estriol cream was the preferred preparation; the 2018 committee agreed

3 they could not make firm conclusions from the evidence or their experience about different

4 vaginal oestrogen products.

5 The 2024 guideline committee discussed all the 2018 recommendations on oestrogen and agreed that these should be extended to cover the perimenopausal stage. The committee 6 7 agreed, based on their knowledge and experience, that recurrent UTI often first occurs when 8 oestrogen starts to decline in the perimenopausal period, which can last many years. The 9 committee were concerned that limiting the use of vaginal oestrogen to after menopause would mean either unnecessarily delaying a potentially beneficial treatment, perhaps for 10 years, or that more people may receive antibiotic prophylaxis than necessary. Therefore, 11 12 although the evidence on oestrogen was not reviewed as part of the 2024 update, the committee agreed updating the recommendations to be more inclusive was essential from 13 14 both an equality perspective and to support antimicrobial stewardship aims. 15 The 2018 guideline recommended that possible adverse effects and the uncertainty of

16 endometrial safety with long-term or repeated use of oestrogens was discussed to ensure 17 shared decision-making. However, there were concerns that this recommendation may have 18 been informed from evidence on high dose vaginal oestrogens that are not used in the UK. 19 Therefore, the committee agreed that the point about endometrial safety should be removed 20 from the recommendation. The committee were aware of the Medicines and Healthcare 21 products Regulatory Agency [MHRA] drug safety update on hormone replacement therapy 22 (2019) and they agreed it was important for prescribers to discuss this with people and that 23 they should be reassured that serious side effects are very rare when using vaginal 24 oestrogen. They also included a cross-reference to the NICE guideline on menopause for 25 information about the use of vaginal oestrogen for people with a history of breast cancer.

The committee also agreed that the recommendation on not offering oral oestrogens should be amended to include all systemic hormone therapy because there are high dose oestrogen products available in non-oral preparations and, in the committee's experience, these should not be used to for recurrent UTI due to risks associated with higher doses of oestrogen.

301.5.3Use of gender-inclusive language and the explicit inclusion of trans and
non-binary people in the guideline recommendations

The committee for the 2024 update reviewed the language of the recommendations to bring them up to date and address equality issues. The committee agreed that recommendations for women in the 2018 guideline should also apply to trans men and non-binary people with a female urinary system. Similarly, recommendations for men should also apply to trans women and non-binary people with a male genitourinary system.

37 The committee discussed that there was no evidence to inform how recommendations 38 should apply to people who have had gender reassignment surgery that involved structural 39 alteration of the urethra. They did not want to explicitly exclude people who have had gender 40 reassignment surgery from the recommendations as, in their experience, some of the 41 recommendations may be beneficial for these groups. However, they agreed that when 42 considering different management options in this population specialist advice should always 43 be sought. Therefore, they agreed to include people who have had gender reassignment 44 surgery that involved structural alteration of the urethra in the recommendation on referral 45 and seeking specialist advice.

Evidence selection 2 1

2 A range of evidence sources are used to develop antimicrobial prescribing guidelines. These

- fall into 2 broad categories: 3
- 4 Evidence identified from the literature search (see section 2.1 below)
- · Evidence identified from other information sources. Examples of other information sources 5 6 used are shown in the interim process guide (2017).
- 7 See appendix A: evidence sources for full details of evidence sources used.

2.1 Literature search 8

9 A literature search was developed to identify evidence for the effectiveness and safety of 10 interventions for managing all urinary tract infections (UTIs) (see appendix C: literature search strategy for full details). The literature search identified 6,695 references. These 11 references were screened using their titles and abstracts and 133 full text references were 12 obtained and assessed for relevance. Thirty-eight full text references of systematic reviews 13 14 and randomised controlled trials (RCTs) were assessed as relevant to the guideline review 15 question (see appendix B: review protocol). Ten percent of studies were screened to establish inter-rater reliability, and this was within the required threshold of 90%. 16 17 Thirteen of the 38 references were prioritised by the committee as the best available

evidence and were included in this evidence review (see appendix F: included studies). The 18

19 methods for identifying, selecting and prioritising the best available evidence are described in the interim process guide. 20

21 The 25 references that were not prioritised for inclusion are listed in appendix I: studies not prioritised. Also see appendix E: evidence prioritisation for more information on study 22 23 selection.

24 The remaining 95 references were excluded. These are listed in appendix J: excluded 25 studies with reasons for their exclusion.

26 Four further systematic reviews were identified following stakeholder consultation and an 27 updated search (May 2018). Luis et al. (2017) is a systematic review and Ledda et al. (2017) 28 is an RCT both covering cranberry products, however, both studies were deprioritised as another systematic review also identified following stakeholder consultation on the same 29 intervention was prioritised (Fu et al. [2017)]; see appendix I: studies not prioritised). Fu et al. 30 31 conducted a meta-analysis comparing cranberry products with placebo or no treatment in 32 non-pregnant women. A third systematic review (Roshdibonab et al. [2017)]) conducted the same comparison in children and was also included in the guideline. The remaining 15 33 references identified in the updated search were excluded. These are listed in appendix J: 34 excluded studies with reasons for their exclusion. 35

36 See also appendix D: study flow diagram.

Summary of included studies 37 **2.2**

A summary of the included studies is shown in tables 1 to 3. Details of the study citation can 38 be found in appendix F: included studies. An overview of the quality assessment of each 39 40 included study is shown in appendix G: quality assessment of included studies.

41

1 Table 1: Summary of included studies: non-pharmacological interventions

Study	Number of participants	Population	Intervention	Comparison	Primary outcome		
Probiotics (lactobacillus)							
Grin et al. 2013 Systematic review. Multiple countries. Follow-up up to 12 months	n=294 (5 RCTs)	Premenopausal women with history of UTI, defined as one or more UTIs within the last 12 months	Lactobacillus (pessaries or oral; in 3 studies lactobacillus given after a course of antibiotics), for prophylaxis	Placebo	Incidence of recurrent urinary tract infections		
Schwenger et al. 2015 Systematic review. Multiple countries. Follow-up up to 28 days	n=735 (9 RCTs and quasi- RCTs)	Adults and children with history of at least 1 UTI or current UTI, 1 study in healthy women (some studies included children with VUR)	Probiotics in any formulation for prophylaxis	Placebo Antibiotics	Symptomatic bacterial urinary tract infection		
D-Mannose							
Kranjcec et al. 2014 RCT Croatia Follow-up 6 months	n=308	Non-pregnant women with history of UTI, defined as at least 2 UTIs in the last 6 months and/or 3 UTIs in the last year	Oral d-mannose for prophylaxis	Antibiotic (nitrofurantoin) No treatment	Number of women experiencing a urinary tract infection		
Cranberry products							
Fu et al. 2017 Systematic review Multiple countries 6 to 12 months follow up	n=1498 (7 RCTs)	Generally healthy non- pregnant women with a history of UTI	Cranberry products (juice, tablets and powder capsules) for prophylaxis	Placebo or no treatment	Number of women experiencing a urinary tract infection		
Jepson et al. 2012 Systematic review. Multiple countries. Follow-up up to 12 months	n=4,473 (24 RCTs)	Adults susceptible to UTI including: people with a history of recurrent lower UTI (defined as more than	Cranberry products (juice, concentrate, capsules, or tablets) for prophylaxis	Placebo, no treatment, water, methenamine hippurate, antibiotics or lactobacillus	Number (incidence) of confirmed urinary tract infection		

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	Number of				
Study	participants	Population	Intervention	Comparison	Primary outcome
		2 episodes in the last year); pregnant women; older people, people with cancer or spinal injury/neuropathic bladder and children with first or subsequent UTI			
Roshdibonab et al. 2017 Multiple countries 3 to 17 months follow up	n=794 (10 RCTs)	Children with history of UTI	Cranberry juice or capsules for prophylaxis	Placebo or antibiotics	Number of urinary tract infections
Beerepoot et al. 2011 RCT Netherlands Follow-up up to 15 months	n=221	Premenopausal women with a history of recurrent UTI, defined as at least 3 self-reported UTIs in the last year	Cranberry capsules for prophylaxis	Antibiotic (co-trimoxazole)	Number of symptomatic urinary tract infections over 12 months Proportion of patients with at least 1 symptomatic urinary tract infection during 12 months of prophylaxis use Median time to the first symptomatic urinary tract infection
Uberos et al. 2012 RCT Spain Follow-up up to 12 months	n=192	Children 1 month to 13 years, with a history of recurrent UTI (defined as at least 2 episodes in the last 6 months), VUR of any degree or renal pelvic dilation associated with UTI	Cranberry syrup for prophylaxis	Antibiotic (trimethoprim)	Number of urinary tract infection and safety
Abbreviations: RCT, Ran	idomised controlled trial; \	/UR, Vesicoureteral reflux			

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1 Table 2: Summary of included studies: non-antimicrobial pharmacological interventions

Study	Number of participants	Population	Intervention	Comparison	Primary outcome			
Oestrogens	Oestrogens							
Perrotta et al. 2008 Systematic review. Multiple countries Follow-up up to 4 years	n=3,345 (9 RCTs)	Post-menopausal women	Oral oestrogens, with or without progestogens; or vaginal oestrogens, delivered by vaginal ring, vaginal pessaries, vaginal tablets	Placebo or antibiotics	Women with recurrent urinary tract infections Urinary tract infections Time until recurrence Number of urinary infections/person/year			

2 Table 3: Summary of included studies: antimicrobials

Study	Number of participants	Population	Intervention	Comparison	Primary outcome			
Antibiotics versus places	Antibiotics versus placebo or no treatment							
Albert et al. 2004 Systematic review Multiple countries. Follow-up not clearly reported	n=1,120 (19 RCTs)	Non-pregnant women (both pre- and post- menopausal women) with at least 2 UTIs in the last year	Antibiotics of various classes administered for at least 6 months	Placebo, antibiotics or another pharmacological non- antibiotic treatment	Number of recurrences per patient-year using 1) microbiological criteria and 2) clinical criteria Proportion of patients who had severe side effects Proportion of patients who had mild side effects			
Dai et al. 2010 Systematic review Multiple countries Follow-up varied according to study	n=1,093 (7 RCTS)	Children with or without VUR	Antibiotics of various classes	Placebo	Deterioration of renal scars			
Muller et al. 2017 Systematic review	n=3,052 (26 RCTs)	Adults and children (authors conducted a mixed analysis of studies in adults,	Nitrofurantoin	Placebo	Occurrence of urinary tract infection			

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Study	Number of participants	Population	Intervention	Comparison	Primary outcome
Multiple countries. Follow-up varied according to study		children or both); the ages of participants involved were not reported consistently, if at all.			Mild adverse effects Emergence of resistance
Schneeberger al. 2015 Systematic review US Follow-up until delivery	n=200 (1 RCT)	Pregnant women with history of 1 or more UTIs before or during pregnancy	Nitrofurantoin and close monitoring	Close monitoring alone	Recurrent urinary tract infection before birth (recurrent pyelonephritis, recurrent cystitis) Preterm birth (less than 37 weeks) Small for gestational age
Williams and Craig 2011 Systematic review Multiple countries. Follow-up varied according to study	n=1,557 (12 RCTs)	Children (without VUR), however studies in which less than 50% of the population had VUR (any grade) were included.	Antibiotics of various classes	Placebo	Recurrence of urinary tract infections Microbial resistance to prophylactic drug Adverse events Withdrawals due to adverse events
Antibiotics versus other	antibiotics				
Muller et al. 2017 Systematic review Multiple countries. Follow-up varied according to study	n=3,052 (26 RCTs)	Adults and children (authors conducted a mixed analysis of studies in adults, children or both); the ages of participants involved were not reported consistently, if at all.	Nitrofurantoin	Different antibiotic classes: Beta-lactams Quinolones Co-trimoxazole Trimethoprim Methenamine hippurate	Occurrence of urinary tract infection Mild adverse effects
Albert et al. 2004 Systematic review	n=1,120 (19 RCTs)	Non-pregnant women (both pre- and post- menopausal women)	Antibiotics of various cla	asses	Number of recurrences per patient-year using 1) microbiological

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Study	Number of participants	Population	Intervention	Comparison	Primary outcome
Multiple countries. Follow-up not clearly reported		with at least 2 UTIs in the last year			criteria and 2) clinical criteria Proportion of patients who had severe side effects Proportion of patients who had mild side effects
Williams and Craig 2011 Systematic review Multiple countries. Follow-up varied according to study	n=1,557 (12 RCTs)	Children (without VUR), however studies in which less than 50% of the population had VUR (any grade) were included.	Antibiotics of various classes		Recurrence of urinary tract infections Microbial resistance to prophylactic drug Adverse events Withdrawals due to adverse events
Duration of antibiotic trea	atment (adults)				
Zhong et al. 2011 RCT China Follow-up12 months	n=68	Postmenopausal women	Antibiotic (continuous low-dose daily)	Antibiotic (intermittent patient-initiated single- dose)	Occurrence of urinary tract infection Conditions predisposing to antibiotic use Adverse events
Abbreviations: RCT, Rar	ndomised controlled trial; \	/UR, vesicoureteral reflux			

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Draft for consultation

1

3 Clinical effectiveness

Full details of clinical effectiveness are shown in <u>appendix H: GRADE profiles</u>. The
 main results are summarised below.

4 3.1 Non-pharmacological interventions

5 3.1.1 Lactobacillus (probiotic) in non-pregnant women

6 The evidence review for lactobacillus is based on 2 meta-analyses (Grin et al. 2013

7 and <u>Schwenger et al. 2015</u>). The studies cover lactobacillus compared with placebo,

8 and lactobacillus compared with antibiotics.

9 Lactobacillus versus placebo

10 The evidence review for lactobacillus versus placebo is based on Grin et al. 2013 (5 11 RCTs, n=294), which included studies in premenopausal women with a history of 12 prior urinary tract infection (UTI) (defined as 1 or more UTIs within the last 12 months 13 prior to entry to the study). In 2 studies included in the meta-analysis, participants 14 received a course of lactobacillus following a UTI treated with antimicrobials until the 15 infection cleared. Four studies treated the women with vaginal pessaries containing 16 lactobacillus, the remaining study used a lactobacillus drink preparation. The strains of Lactobacillus spp. included across the studies were: L. rhamnosus GR-1, L. 17 18 fermentum B-54, L. casei v rhamnosus LCR35, L. rhamnosus GG, and L. crispastus 19 CTV-05. The composition of the different preparations varied among the different 20 studies. The pessaries were administered daily, 5 days a week or twice a week. The 21 length of treatment ranged from 5 days to 12 months. Length of follow-up was also 22 inconsistent between studies, ranging from 4 weeks to 12 months. 23 The populations included in the studies were mostly premenopausal adult women.

25 Only 1 study reported the age range of included in the meta-analysis defined UTI with 26 microbiological criteria that ranged from 10³ colony forming units per millilitre

(CFU/mL) to $10^5 CFU/mL$. In some studies, women were already receiving antibiotic

treatment for their UTI and, in 1 study the women were healthy and had no infection.

29 Lactobacillus spp. did not significantly reduce the risk of recurrent UTIs in

premenopausal women when compared with placebo (5 RCTs, n=194: 29.9% versus
34.7%; risk ratio [RR] 0.85 95% CI 0.58 to 1.25; low quality evidence). When authors
restricted the analysis to studies that only used 'effective strains' of lactobacillus (as
defined by the authors), results were statistically significant (2 RCTs, n=127, 16.1%
versus 32.3%: RR 0.51, 95% CI 0.26 to 0.99; NNT 7 [95% CI 4 to 64]; moderate
quality evidence).

36 Lactobacillus versus antibiotics

37 The evidence review for lactobacillus versus antibiotics is based on a single RCT 38 (NAPRUTI Study II 2006) reported within a systematic review (Schwenger et al. 39 2015). The 'Non-antibiotic versus Antibiotic Prophylaxis for Recurrent Urinary Tract 40 Infections' (NAPRUTI) study compared Lactobacillus spp. (L. rhamnosus GR-1 and L. reuteri RC-14) with co-trimoxazole as prophylaxis for the prevention of UTIs in 41 42 postmenopausal women with recurrent UTIs. Patients randomised to receive 43 lactobacillus took 1 capsule containing at least 10⁹ CFUs of L. rhamnosus GR-1 and L. reuteri RC-14 twice a day and 1 placebo capsule at night for 12 months. Patients 44

- randomised to receive co-trimoxazole took a 480 mg tablet at night, and 1 placebo
 capsule twice a day for 12 months.
- Schwenger et al. (2015) defined the rate of UTIs in each treatment group as the
 number of patients experiencing at least 1 UTI, not the number of UTIs in a treatment
 group.
- 6 There was no significant difference in the number of symptomatic infections between 7 women treated with lactobacillus and those treated with antibiotics (1 RCT, n=223: 74.8% versus 66.7% DD 1.12, 05% CL 0.05 to 1.22 low guality suidenes)
- 8 74.8% versus 66.7%; RR 1.12, 95% CI 0.95 to 1.33; low quality evidence).

9 Sensitivity analysis was conducted to determine the effect of imputing data

10 (participants with missing data were assumed to have negative outcomes, known as

11 worst case scenario), or ignoring missing data on the reported outcome. When a 12 worst case scenario was applied for those randomised to the lactobacillus treatment 13 group, there was a significant increase in the number of symptomatic bacterial UTIs 14 seen in this group compared with those receiving antibiotics (1 RCT, n=223: 79.1% 15 versus 66.7%; RR 1.19 95% CI 1.01 to 1.4; NNT 8 [95% CI 5 to 114]; moderate 16 quality evidence). However, when a worst case scenario was applied for antibiotics, 17 there was a significant increase in the number of symptomatic bacterial UTIs seen in 18 this group compared with those receiving lactobacillus (1 RCT, n=223: 74.8% versus

- 19 89.8%; RR 0.83 95% CI 0.74 to 0.94; NNT 7 [95% CI 4.0 to 19.0]; moderate quality
- 20 evidence).

21 3.1.2 D-Mannose in non-pregnant women

22 The evidence review for D-mannose is based on 1 RCT (Kranjcec et al. 2014, n=308) 23 comparing D-mannose (200 ml of 1% solution once daily in the evening) with no 24 treatment, or an antibiotic (nitrofurantoin 50 mg once daily in the evening). Kranjcec 25 et al. (2014) included non-pregnant women who presented with current UTI and a 26 history of recurrent UTI. The authors defined the latter as 2 episodes in the last 6 27 months or 3 episodes in the last year. Authors based the diagnosis of UTI on a 28 microbiological assessment (≥10³ CFUs per ml) as well as lower urinary tract 29 symptoms such as dysuria, frequency and urgency. All women in the study took 30 antibiotics (ciprofloxacin 500 mg twice a day) for 1 week for their current UTI. The 31 median age was between 48 and 52 years, and 47.4% of participants were 32 postmenopausal. The authors assessed effectiveness as the number of participants 33 presenting with 1 recurrent UTI during the study period.

34 **D-mannose versus no treatment**

D-mannose was significantly more effective in preventing recurrent UTI in nonpregnant women compared with no treatment over the 6-month study period
(Kranjcec et al. 2014, n=205: 14.6% versus 60.8%; RR 0.24, 95% CI 0.15 to 0.39;

38 NNT 3 [95% Cl 2 to 3]; high quality evidence).

39 **D-mannose versus antibiotic**

40 D-mannose did not show a significant benefit in reducing recurrent UTIs in non-

41 pregnant women when compared with antibiotics (nitrofurantoin 50 mg a day) over

42 the 6-month study period (Kranjcec et al. 2014, n=206: 14.6% versus 20.4%; RR

- 43 0.71, 95% CI not stated, calculated by NICE as 95% CI 0.39 to 1.31; low quality
- 44 evidence).

1 3.1.3 Cranberry products

2 The evidence review for cranberry products is based on 1 systematic review (Jepson 3 et al. 2012,) and 2 RCTs (Beerepoot et al. 2011 and Uberos et al. 2012). The 2 RCTs provided evidence on antimicrobial resistance (see section 5). Across all publications 4 5 included, authors defined recurrent UTI as 3 episodes of infection in the last 12 6 months or 2 episodes of infection in the last 6 months. Participants received 7 cranberry products either in liquid form (juice or syrup) or solid form (capsules or 8 tablets). Cranberry products were compared with placebo, no treatment or antibiotics. Two further systematic reviews were identified following stakeholder consultation and 9 10 an updated search. Fu et al. (2017) conducted a meta-analysis comparing cranberry

products with placebo or no treatment in non-pregnant women and <u>Roshdibonab et</u>
 al. (2017) conducted the same comparison in children.

13 Cranberry products in women

14 Two systematic reviews (Jepson et al. 2012 and Fu et al. 2017) and 1 RCT

15 (Beerepoot et al. 2011) assessed the efficacy of cranberry products for preventing

16 UTIs in women. The studies included women with recurrent or previous UTI. Age

17 groups varied across the studies from young women to elderly women and not all

18 studies specified whether pregnant women were excluded. The main outcome of

19 interest was reduction of recurrent UTIs, defined as participants with 1 or more UTI,

20 or repeat symptomatic UTI.

21 Cranberry products versus placebo or no treatment

22 <u>Jepson et al. (2012)</u> identified 4 RCTs that compared cranberry products (juice, syrup 23 or tablets) with matched placebo or no treatment. The concentration of cranberry 24 products as well as the frequency of administration varied across the studies. The 25 age of women also varied across the studies from 21 to 72 years. Across the studies, 26 authors used microbiological criteria and symptoms to assess UTIs. Some studies 27 required >10⁴ CFUs/ml in a sample, and others ≥10⁵ CFUs/ml in a sample.

Jepson et al. 2012 found that prophylactic cranberry products for 3, 6 or 12 months did not show a significant benefit in the number of women who had one or more UTI

30 during follow up (4 RCTs, n=594: 19.9% versus 22.8%; RR 0.74, 95% CI 0.42 to

31 1.31; very low quality evidence) when compared with placebo or no treatment.

32 **Evidence identified following stakeholder consultation**

33 Fu et al. (2017) compared cranberry in either juice or capsule form, for preventing 34 UTIs in non-pregnant women, with a follow up of 6 to 12 months. Age of participants 35 varied from 21 to 72 years old. The included studies differed in their definition of UTI, 36 with most trials defining UTI through the presence of symptoms, and 4 requiring 37 confirmed bacteriuria of varying thresholds. This data adds an additional 4 unique 38 RCTs to the Jepson et al. (2012) analysis, including a total of 501 additional 39 participants. Furthermore, 3 of the RCTs included in Fu et al. (2017), are also 40 included in Jepson et al. (2012), while 1 RCT included in Jepson et al. (2012) is not 41 included in Fu et al. (2017).

42 Cranberry juice or capsules significantly reduced the incidence of UTI in non-

- 43 pregnant women, diagnosed either by symptom presence or culture confirmation,
- 44 compared with placebo or no treatment (7 RCTs, n=1498: 20.7% versus 26.5%; RR
- 45 0.74, 95% CI 0.55 to 0.98; very low quality evidence). When restricted to UTIs

confirmed by culture, this difference was not significantly significant (5 RCTs, n=912: 1 2 19.8% versus 24.0%; RR 0.71, 95% CI 0.45 to 1.12; very low quality evidence).

3 Cranberry juice did not significantly reduce the incidence of UTI, diagnosed either by 4 symptom presence or culture confirmation, compared with placebo or no treatment (6 5 RCTs, n= 1272: 22.0% versus 26.6%; RR 0.79, 95% CI 0.59 1.06, very low quality 6 evidence). However, cranberry tablets did significantly reduce incidence of UTI 7 compared with placebo (2 RCTs, n= 276: 13.5% versus 28.0%; RR 0.48, 95% CI 8 0.29 to 0.79; low quality evidence).

9 Cranberry products versus antibiotics

10 Jepson et al. 2012 identified 2 RCTs that compared cranberry products (tablets 11 500 mg) with antibiotics (trimethoprim 100 mg or co-trimoxazole 480 mg). The 12 frequency of administration varied across the studies. The age of women varied 13 across the studies, with 1 study recruiting women aged 45 years and older, and the 14 other study including premenopausal women who were older than 18 years. It was 15 unclear whether pregnant women were excluded. Both RCTs used microbiologic criteria to confirm UTIs. One study required >10⁴ CFUs/ml in a urine sample while the 16 17 other required $\geq 10^5$ CFUs/ml. The duration of the studies was 6 or 12 months. 18 Prophylactic cranberry products did not show a significant benefit in reducing

- 19 recurrent UTIs in women (2 RCTs, n=344: 51.1% versus 40.4%; RR 1.31, 95% CI
- 0.85 to 2.02; moderate quality evidence) when compared with antibiotics 20
- 21 (trimethoprim or co-trimoxazole).

22 Cranberry products versus placebo or no treatment in pregnant women

23 One systematic review (Jepson et al. 2012) assessed the efficacy of cranberry 24 products for preventing UTIs pregnant women, which included 2 studies of cranberry 25 products (juice) compared with matched placebo or water. No data were identified for 26 comparisons with antibiotics. The authors did not provide details on whether 27 pregnant women had previous or current UTIs. One study was available as abstract 28 only and did not include information on diagnosis of UTI or treatment length. The 29 authors of the second study confirmed UTI using microbiological criteria (>10⁸ CFUs 30 per ml of single organism) and symptoms such as dysuria, frequency or urgency. The 31 length of this study was 5 to 7 months. The main outcome reported was the reduction 32 of recurrent UTIs, defined as participants with 1 or more UTI, or repeat symptomatic 33 UTI.

- 34 Prophylactic cranberry products did not show a significant benefit in reducing
- 35 recurrent UTIs in pregnant women (Jepson et al. 2012, 2 RCTs, n=674: 56.6%
- 36 versus 55.6%; RR 1.04, 95% CI 0.93 to 1.17; moderate quality evidence) when
- 37 compared with placebo or no treatment. No other relevant outcomes were reported.

38 Cranberry products versus placebo or no treatment in elderly men and women

39 One systematic review (Jepson et al. 2012) assessed the efficacy of cranberry 40 products for preventing UTIs in older people (men and women), which included 2 41 RCTs. These RCTs covered whether cranberry products (juice or capsules) were more effective than matched placebo or no treatment in adults aged 60 years and 42 43 over. In 1 study, patients took 300 ml cranberry juice or matched placebo juice. It was 44 unclear whether this was taken once a day or more frequently. In the other study 45 patients took a 650 mg cranberry capsule once or twice a day. The studies included 46 people who were either admitted to acute medicine for the elderly assessment. 47 rehabilitation units for elderly people, or lived in care facilities. One study only

- 1 included elderly people with dementia. Both RCTs used microbiologic criteria and
- 2 symptoms to confirm UTI. One study required >10⁴ CFUs/ml in a urine sample while
- 3 the other required $\geq 10^8$ CFUs/ml. No data were identified for comparisons with
- 4 antibiotics. The main outcome reported was participants with 1 or more UTI at follow
- 5 up, measured using urine culture.
- 6 Prophylactic cranberry products did not show a significant benefit in reducing
- 7 recurrent UTIs in older people (men and women) when compared with placebo or no
- 8 treatment during a 6-month treatment period (2 RCTs, n=413: 9.7% versus 12.6%;
- 9 RR 0.75, 95% ČI 0.39 to 1.44; very low quality evidence).

10 Cranberry products in children

11 Two systematic reviews (Jepson et al. 2012 and Roshdibonab et al. 2017) assessed

12 the efficacy of cranberry products for preventing UTIs in children. The included

13 studies covered whether cranberry products were more effective than placebo or no

treatment, or antibiotics. The main outcome reported was reduction of recurrent UTI

15 defined as participants with 1 or more UTI or repeated symptomatic UTI.

16 Cranberry products versus placebo or no treatment

Jepson et al. (2012) identified 2 RCTs comparing cranberry products (concentrate or
 juice) with matched placebo or no treatment. One publication included only girls aged

19 3 to 14 years with an average age of 7 years and 6 months. The other publication did

20 not specify the sex or ages of the children. The authors of 1 publication used

symptoms and microbiological criteria (> 10⁸ CFUs per ml) to diagnose UTI, whereas
 the other publication did not specify diagnostic criteria.

23 Prophylactic cranberry products did not show a significant benefit in reducing

recurrent UTIs in children over the 6-month study period (2 RCTs, n=309: 16.3%

versus 29.5%; RR 0.48, 95% CI 0.19 to 1.22; low quality evidence) when compared
with placebo or no treatment.

27 Evidence identified following stakeholder consultation

28 Roshdibonab et al. (2017) included 8 RCTs comparing cranberry taken daily in juice 29 or capsule form, with placebo for recurrent UTI in children, with 2 to 12 month follow 30 up. Children were aged between 1 to 13 years, with UTI diagnosed by positive urine 31 culture in all studies. Two of the RCTs included in the meta-analysis included 32 children with catheters. This data includes an additional 6 RCTs to the Jepson et al. 33 (2012) analysis, including a total of 262 additional participants. The 2 RCTs included 34 in Jepson et al. (2012) for this population are also included in Roshdibonab et al. (2017). 35

Children using cranberry juice or capsules showed a significant reduction in
incidence of culture confirmed UTI compared with children taking placebo (8 RCTs,
n= 571: OR 0.31, 95% CI 0.21 to 0.46; very low quality evidence).

39 Cranberry products versus antibiotics

40 Jepson et al. (2012) identified 1 RCT comparing cranberry products (syrup) with

41 antibiotics (trimethoprim 8 mg/kg). The authors included children between 1 month

- 42 and 13 years, and mean ages ranged from 28.3 to 30.7 months. Children either
- 43 presented with recurrent UTI (2 or more infections in 6 months), vesicoureteric reflux
- 44 of any degree, pyelic ectasia or hydronephrosis, or anatomical kidney disorder.

- 1 Jepson et al. 2012 found that prophylactic cranberry products did not show a
- 2 significant benefit in reducing recurrent UTIs in children (1 RCT, n=192: 10.7%
- 3 versus 15.4%; RR 0.69, 95% CI 0.32 to 1.51; low quality evidence) when compared
- 4 with antibiotics (trimethoprim) over the 6-month study period.

5 3.2 Non-antimicrobial pharmacological interventions

6 3.2.1 Oestrogens in post-menopausal women

7 The evidence review for oestrogens (with or without progestogens) is based on 1 8 systematic review of 9 RCTs (Perrotta et al. 2008, n=3,345). The author's objective 9 was to examine the efficacy of oestrogen in decreasing the rate of recurrent urinary 10 tract infection (UTI) in postmenopausal women and its safety. All studies within the 11 systematic review included post-menopausal women with recurrent UTI (defined as 3 12 episodes of infection in the last 12 months or 2 episodes of infection in the last 6 13 months). The systematic review included comparisons of oral oestrogen versus 14 placebo, vaginal oestrogen versus placebo, and vaginal oestrogen versus oral antibiotics. The main efficacy outcome was reduction in recurrent UTI. 15

16 Oral oestrogens compared with placebo

17 Perrotta et al. (2008) identified 4 RCTs that reported on the efficacy of oral 18 oestrogens compared with placebo in post-menopausal women. These included 1 19 large study (n=2,654) with a duration of up to 4 years, and 3 smaller studies (fewer 20 than 100 participants each) with durations of 12 weeks or 6 months. The age of 21 women varied across the studies, with the large study recruiting participants less 22 than 80 years of age, while another study reported mean age of 88 years. In the 23 large study the oestrogen preparation also contained a progestogen. There was no 24 significant reduction in recurrent UTI when oral oestrogen was compared with 25 placebo (4 RCTs, n=2,798: 11.3% versus 10.4%; RR 1.08, 95% CI 0.88 to 1.33; 26 moderate quality evidence).

27 Vaginal oestrogens compared with placebo or no treatment

28 Perrotta et al. (2008) identified 2 small RCTs that reported on the efficacy of vaginal 29 oestrogens compared with placebo or no treatment. The trials differed in the 30 administration method of oestrogens and comparator used. One RCT compared an 31 oestrogen-releasing vaginal ring with no treatment while the other compared topically 32 applied vaginal oestrogen cream with placebo cream. The age of the participants 33 was not reported, and the results were presented separately for each study, not 34 pooled in a meta-analysis. Oestrogen administered via a vaginal ring (Estring) 35 showed a statistically significant benefit for reducing recurrent UTI compared with no 36 treatment during the 36 week study period (1 RCT, n=108: 50.9% versus 80%; RR 37 0.64, 95% CI 0.47 to 0.86; NNT 4 [95% CI 3 to 9]; moderate quality evidence). 38 Similarly, oestrogen administered topically (oestriol cream) showed a significant 39 reduction in recurrent UTI when compared with placebo during an 8-month study 40 period (1 RCT, n=93: 16% versus 62.8%; RR 0.25 95% CI 0.13 to 0.5; NNT 3 [95%

41 CI 2 to 4]; high quality evidence).

42 Vaginal oestrogens versus antibiotics

- 43 Perrotta et al. (2008) identified 2 RCTs that reported on the efficacy of vaginal
- 44 oestrogens (pessary or cream) compared with oral antibiotics (nitrofurantoin or
- 45 ofloxacin). Both studies included post-menopausal women. However, ages or

- 1 diagnostic criteria for UTI were not specified. Perrotta et al. (2008) presented the
- 2 results of the studies separately as the authors felt that results could not be pooled
- 3 due to high heterogeneity. There were significantly more UTIs at the end of the 9-
- 4 month study period with vaginal oestrogens delivered via pessary compared with
- 5 nitrofurantoin 100 mg a day (1 RCT, n=171; 67.4% versus 51.8%; RR 1.3, 95% CI
- 6 1.01 to 1.68; low quality evidence). In contrast, vaginal oestrogen cream (premarin
- cream) was significantly more effective than of loxacin 600 mg a day in reducing
 recurrent UTI at the end of the 3-month study period (1 RCT, n=42; 7.4% versus)
- recurrent UTI at the end of the 3-month study period (1 RCT, n=42; 7.4% versus
 80%; RR 0.09 95% CI 0.02 to 0.36; NNT 2 [95% CI 2 to 2]; low quality evidence).
- 10 This benefit only lasted as long as participants were on prophylaxis, with no benefit
- 11 seen 2 months after stopping (1 RCT, n=42; 7.4% versus 13.3%; RR 0.56 95% CI
- 12 0.09 to 3.55; very low quality evidence).

3.3 Antimicrobials in non-pregnant women

14 The evidence review for antimicrobials in non-pregnant women is based on 1

- 15 systematic review (Albert et al. 2004), and 1 RCT (Zhong et al. 2011). The included
- 16 studies assessed antibiotics compared with placebo, and the duration of antibiotic
- 17 treatment.

18 3.3.1 Antibiotics compared with placebo

Albert et al. (2004) included 10 RCTs comparing antibiotics with placebo (n=1,120), assessing the efficacy and safety of antibiotic prophylaxis to prevent recurrent urinary tract infection (UTI) in adult non-pregnant women. Participants were included if they had experienced at least 2 episodes of uncomplicated UTI in the previous year, and were aged over 14 years old. The authors performed sensitivity analysis, excluding trials that had different inclusion criteria or tested different prophylaxis schedules.

25 In 8 RCTs, antibiotic prophylaxis was given for 6 months, and in 2 RCTs it was given 26 for 12 months. The antibiotic dose regimens used in the studies included: 27 ciprofloxacin 125 mg post-coital (women were instructed to take ciprofloxacin as a 28 single dose after sexual intercourse), co-trimoxazole 40/200 mg daily, cephalexin 29 125 mg daily, nitrofurantoin 50 mg daily, nitrofurantoin 100 mg daily, norfloxacin 30 200 mg daily and cinoxacin 250 mg daily). In all studies, prophylaxis was stopped in 31 each case of recurrent infection. Recurrence was defined as the presence of 32 bacteriuria and the clinical symptoms of UTI.

33 Antibiotic prophylaxis, when compared with placebo, significantly reduced the 34 recurrence of UTI during the prophylactic period of 6 to 12 months, when using 35 microbiological criteria (10 RCTs, n=372: 12.3% versus 65.5%; RR 0.21 95% CI 0.13 to 0.34; NNT 2 [95% CI 2 to 3]; high quality evidence) and clinical criteria (7 RCTs, 36 37 n=257: 7.4% versus 51.2%; RR 0.15 95% CI 0.08 to 0.28; NNT 3 [95% CI 2 to 3]; 38 high guality evidence). However, this effect was diminished when recurrence was 39 reported after the prophylactic period (2 RCTs, n=70: 52.3% versus 57.7%; RR 0.82 40 95% CI 0.44 to 1.53; very low quality evidence).

41 **3.3.2 Choice of antibiotic**

42 Although Albert et al. (2004) reported outcomes for studies which compared different 43 antibiotic choices, these studies were included in a larger meta-analysis (Muller et al.

44 2017), which is described in <u>section 3.5.2</u> of this evidence review.

1 3.3.3 Antibiotic dosing and course length

2 Zhong et al. (2011) (n=83) compared the efficacy and safety of intermittent single-3 dose antibiotic prophylaxis versus continuous antibiotic prophylaxis over 12 months. 4 The study included postmenopausal women who had experienced 3 or more UTIs 5 within a 12-month period. The average number of UTIs prior to entry was 6 approximately 5 infections in the previous year, in both treatment groups. Participants 7 took antibiotics either continuously over the study period or used single-dose 8 antibiotics whenever they were exposed to conditions that might trigger UTI. These 9 conditions were determined from the women's experience and included working or 10 walking for a long time, sexual intercourse, travelling, or micturition delay. It was 11 unclear whether women took their intermittent antibiotics before or after exposure to 12 triggers for UTI. The choice of antibiotic (nitrofurantoin, norfloxacin, ciprofloxacin, 13 amoxicillin, co-trimoxazole, cefaclor or cefuroxime) in both groups was done on a 14 case by case basis and depended on the woman's previous use of antibiotics and 15 the outcome of an antimicrobial susceptibility test. Dose varied by antibiotic but was 16 the same for an individual antibiotic. Diagnosis of UTI was based on microscopic 17 pyuria in a urine test.

18 The authors reported the number of episodes of UTI per year, the number of

episodes per year per patient as well as the number of patients having 1, 2, 3, and up
to 12 episodes per year. There was no statistically significant difference between the
intermittent single-dose and continuous treatment regimens (Zhong et al. 2011, n=68:
80.6% versus 70.3%; RR and 95% CI not stated; calculated by NICE as RR 1.15
95% CI 0.87 to 1.51; moderate quality evidence).

24 One study in Albert et al. 2004 (Melekos et al. 1997), compared ciprofloxacin 125 mg 25 taken as a single dose immediately after sexual intercourse, and ciprofloxacin taken 26 as a single dose at night. The study was conducted in pre-menopausal women aged 27 18 to 45, who were sexually active and had ≥3 documented lower UTIs in the last 12 28 months. They found no significant difference in the number of women experiencing at 29 least one microbiological recurrence whilst on prophylaxis (1 RCT, n=135: 2.9% 30 versus 3.1%; RR 0.93 95% CI 0.13 to 6.4; low quality evidence), or the number of 31 women experiencing at least one clinical recurrence whilst on prophylaxis (1 RCT, 32 n=135: 5.7% versus 4.6%; RR 1.24 95% CI 0.29 to 5.32; low quality evidence). 33 Authors noted no significant difference between groups, in the microbiological recurrence after the prophylactic period (low quality evidence). 34

35 **3.4** Antimicrobials in pregnant women

36 The evidence review for antimicrobials in pregnant women is based on 1 systematic review (Schneeberger et al. 2015). This review covers whether antibiotics are more 37 effective than clinical surveillance alone (no treatment) in preventing recurrent urinary 38 39 tract infection (UTI). Schneeberger et al. (2015) planned to assess the effectiveness 40 of pharmacological and non-pharmacological interventions for the prevention of 41 recurrent UTI in pregnant women. However, only a single RCT was identified as 42 meeting the inclusion criteria, which compared a continuous course of nitrofurantoin 43 and close monitoring until delivery, with close monitoring alone.

44 **3.4.1** Nitrofurantoin compared with no treatment (monitoring alone)

45 Pregnant women who were admitted to hospital with a clinical diagnosis of acute

46 pyelonephritis were included into the study. Clinical diagnosis included the presence

- 47 of costovertebral angle and 2 of the following symptoms: temperature $\geq 101^{\circ}$ F, pyuria,
- 48 or bacteriuria (>10³ gram-negative organisms per ml). Women randomised to receive

- antibiotics were given nitrofurantoin 50 mg three times a day for the remainder of the 1
- 2 pregnancy in conjunction with close monitoring. Monitoring was defined as fortnightly
- visits to the clinic until the 36th week of pregnancy, after which time they were seen 3
- 4 weekly until delivery. Urine tests were also conducted at each visit.

5 Nitrofurantoin significantly reduced the incidence of asymptomatic bacteriuria in 6 pregnant women when compared with monitoring alone (1 RCT, n=102: 32.6% 7 versus 59.3%; RR 0.55 0.95% CI 0.34 to 0.89; NNT 4 [95% CI 3 to 13]; moderate 8 guality evidence). However, nitrofurantoin did not significantly reduce recurrent 9 pyelonephritis (n=167: 7.3% versus 8.2%; RR 0.89, 95% CI 0.31 to 2.53; low quality evidence) or recurrent UTI (n=167: 2.4% versus 8.2%; RR 0.3, 95% CI 0.06 to 1.38; 10 11 low quality evidence) in pregnant women. Furthermore, nitrofurantoin did not show any additional benefit compared with monitoring alone for the following outcomes: 12 number of preterm births <37 weeks, birthweight, 5 minute Apgar score <7, and 13 14 miscarriage (very low to low quality evidence).

15 3.4.2 Choice of antibiotic

16 No evidence from systematic reviews or RCTs was identified.

17 3.4.3 Antibiotic dosing and course length

18 No evidence from systematic reviews or RCTs was identified.

Antimicrobials in adults and children (mixed 3.5 19 population analysis) 20

21 The evidence review for antimicrobials in men, women and children is based on 1 22 systematic review (Muller et al. 2017). This study did not stratify analysis by gender or age, but reported overall outcomes. Most studies included had a mixed gender 23 24 population in either adults or children. The included studies cover antibiotics versus 25 placebo and antibiotics versus other antibiotics.

26 3.5.1 Antibiotics compared with placebo

27 Nitrofurantoin versus placebo

28 Muller et al. (2017), which included 26 RCTs (n=3.052), assessed the effectiveness 29 of nitrofurantoin (various doses: 100 mg a day, 100 mg twice a day, 100 three times a 30 day, 75 mg a day, 50 mg a day or 50 mg twice a day, 1mg/kg (children aged 2 to 18 31 years), 1.5 mg/kg (children, age not reported), 2 mg/kg (children aged 2 to 12 32 years)), given as long-term prophylaxis (defined as greater than 14 days), for the 33 primary or secondary prevention of urinary tract infection (UTI) in men, non-pregnant 34 women (pre- or post-menopausal) and children (predominantly female children). The 35 authors did not define primary or secondary prophylaxis. Most included studies recruited people with recurrent UTI; however, the study specific definition of recurrent 36 37 UTI was not reported. A few studies conducted in children included children with 38 neurogenic bladder requiring catheterisation. The ages of children included in the 39 individual studies was not reported in all studies, or reported in a consistent manner. 40 The duration of antibiotic prophylaxis varied among studies, and ranged from 5 41 weeks to 24 months. Muller et al. (2017) also assessed short-term prophylaxis 42 (defined as 3 to 14 days). However, the studies included looked at surgical

prophylaxis which is not relevant to this evidence review. 43

- 1 Nitrofurantoin when given as primary or secondary long-term prophylaxis (for 5
- 2 weeks to 24 months) significantly reduced the occurrence of UTI in adults and
- children compared with placebo or no treatment (8 RCTs, n=491: 22.5% versus 59%; 3
- 4 RR 0.38, 95% CI 0.28 to 0.50; NNT 3 [95% CI 3 to 4]; low guality evidence).
- 5 One controlled trial included in Muller et al. (2017) which could not be included in the
- 6 meta-analysis (due to lack of randomisation) compared nitrofurantoin, methenamine
- 7 hippurate and no treatment in older men and women. Those who were allocated to
- 8 receive no treatment received almost twice as many antibiotic courses than any other
- 9 groups (no results were reported, only described narratively).

3.5.2 Choice of antibiotic 10

11 Muller et al. (2017) assessed the effectiveness of nitrofurantoin compared with a

12 range of other antibiotics (amoxicillin, penicillin, pivmecillinam, cefaclor, cefixime,

13 cinoxacin, norfloxacin, co-trimoxazole, trimethoprim, methenamine hippurate) and

14 stratified the analysis according to antibiotic class. The duration of antibiotic

15 prophylaxis varied among studies, and ranged from 3 months to 24 months.

16 Nitrofurantoin compared with other antibiotics (overall)

17 There was no significant difference between nitrofurantoin and other antibiotics in

18 reducing the incidence of recurrent UTI in adults and children (22 RCTs, n=1,319:

19 23.3% versus 26.1%; RR 0.93, 95% CI 0.69 to 1.26; very low quality evidence).

20 Nitrofurantoin versus methenamine hippurate

21 Using nitrofurantoin as prophylaxis for the prevention of recurrent UTI significantly

22 reduced the incidence of UTI in adults and children compared with methenamine

23 hippurate (2 RCTs, n=196: 35.8% versus 51.2%; RR 0.60, 95% CI 0.43 to 0.85; NNT

24 7 [95% CI 4 to 102]; low quality evidence).

25 Nitrofurantoin versus trimethoprim

26 There was no significant difference between nitrofurantoin and trimethoprim in

27 reducing the incidence of UTI in adults or children (5 RCTs, n=350: 22.5% versus 28 29.3%; RR 0.81, 95% CI 0.38 to 1.71; very low quality evidence).

29 Nitrofurantoin versus co-trimoxazole

- 30 There was no significant difference between nitrofurantoin and co-trimoxazole in
- 31 reducing the incidence of UTI in adults or children (4 RCTs, n=81: 12% versus 8.9%;
- 32 RR 1.42, 95% CI 0.17 to 12.0; very low quality evidence).

33 Nitrofurantoin versus beta-lactam antibiotics

- 34 There was no significant difference between nitrofurantoin and or beta-lactam
- 35 antibiotics in reducing the incidence of recurrent UTI in adults and children (5 RCTs,
- 36 n=249: 16.5% versus 22.4%; RR 0.84, 95% CI 0.49 to 1.44; very low quality 37
- evidence).

38 Nitrofurantoin versus quinolones

- 39 There was no significant difference between nitrofurantoin and quinolones in
- 40 reducing the incidence of recurrent UTI in adults and children (3 RCTs, n=186: 29.8%
- 41 versus 14.7%; RR 2.26, 95% CI 0.73 to 7; very low guality evidence).

1 3.5.3 Antibiotic dosing and course length

2 Muller et al. (2017) conducted a meta-analysis to assess the effect of different

3 nitrofurantoin dosing regimens for long-term prophylaxis in adult participants (100 mg

4 daily, 75 mg daily, 50 mg daily and 50 mg twice daily). The studies used to calculate

5 the effect of dose on the incidence of urinary tract infections were not reported by

6 Muller et al. (2017), neither were they identifiable from the supplementary material.

7 They reported no significant differences between the different regimens (absolute

8 figures not reported; p=0.08, $l^2=53\%$; unable to give GRADE quality rating).

9 3.6 Antimicrobials in children

10 The evidence review for antimicrobials in children is based on 2 systematic reviews 11 (Dai et al. 2010, and Williams and Craig 2011). The included studies cover antibiotics 12 versus placebo and antibiotics versus other antibiotics. Some studies included a 13 small proportion of children diagnosed with vesicoureteric reflux, but most excluded 14 children with grades 4 and 5, or recruited only those with milder/less symptomatic 15 grades (1-3), which typically resolved in most children without intervention.

16 3.6.1 Antibiotics compared with placebo

Williams and Craig (2011), which included 5 RCTs (n=1,069), assessed the efficacy 17 18 of antibiotic prophylaxis compared with placebo in children with recurrent urinary tract 19 infection (UTI). Not all the included studies had clear inclusion and exclusion criteria, 20 and the authors pointed out that it is likely that children were misclassified in the 21 individual studies due to the poor inclusion criteria, and this may impact upon the 22 generalisability of the overall findings. The ages of children included in the studies 23 varied, with 1 study including children from birth to 18 years, and in other studies no 24 age range was reported. The definition of recurrent UTI was not consistent across 25 the studies. However, 1 of the studies included in the review excluded children with a 26 history of urinary tract infection. The length of prophylaxis also differed between 27 studies, with the majority of children receiving antibiotics for at least 6 months. In 2 28 studies, the length of prophylaxis was not reported. The antibiotics used were 29 nitrofurantoin (50 mg daily [children weighing >20 kg], 25 mg daily [children weighing 30 <20 kg], and co-trimoxazole [trimethoprim 2 mg/kg/daily and sulfamethoxazole 31 10 mg/kg/daily]. Studies which had a population of children in which more than 50% 32 were diagnosed with any grade of vesicoureteral reflux were excluded from the 33 systematic review.

34 Antibiotic prophylaxis did not significantly reduce the recurrence of symptomatic UTI 35 compared with placebo or no treatment (4 RCTs, n=1,024: 10.5% versus 17.2%; RR 36 0.75, 95% CI 0.36 to 1.53; very low quality evidence). This did not change when the 37 analysis was restricted to studies that only included children without vesicoureteral 38 reflux (3 RCTs, n=491: 7.3% versus 13.8%; RR 0.56 95% CI 0.15 to 2.12; very low 39 guality evidence). There was no significant difference in the rate of antimicrobial 40 resistance to the prophylactic antibiotic in children who received antibiotics compared 41 with placebo (Williams and Craig 2011, 2 RCTs, n=118: 35.3% versus 16.4%; RR 42 2.4, 95% CI 0.62 to 9.26; very low quality evidence). Similarly, antibiotics offered no 43 significant benefit over the use of placebo or no treatment in the number of repeat 44 positive cultures obtained in children (very low quality evidence).

45 Another systematic review (<u>Dai et al. 2010</u>) also assessed the effect of long-term

46 antibiotic prophylaxis in children (aged less than 18 years old) for the prevention of

47 recurrent UTI. Long-term prophylaxis was defined by the authors as antibiotics given

- 1 were included in the studies. Six out of 7 studies compared co-trimoxazole with
- 2 placebo for a duration of 3 to 24 months.
- 3 Antibiotics did not significantly reduce the rate of deteriorated renal scars in children
- 4 when compared with placebo or no treatment (Dai et al. 2010, 7 RCTs, n=1,093:
- 5 2.9% versus 3.5%; RR 0.95 95% CI 0.51 to 1.78; very low quality evidence).

6 3.6.2 Choice of antibiotic

7 Williams and Craig (2010) assessed the choice of antibiotics for prophylactic use in8 the prevention of recurrent UTI in children.

9 Nitrofurantoin versus trimethoprim

- 10 Nitrofurantoin (1 to 1.5 mg/kg daily) significantly reduced the risk of obtaining a 11 repeat positive culture at the end of prophylaxis (6 months) compared with
- trimethoprim (2–3 mg/kg daily) in children being treated to prevent recurrent UTI (1
- 13 RCT, n=60: 20% versus 61.7%; RR 0.3, 95% CI 0.2 to 0.6; NNT 3 [95% CI 2 to 8];
- 14 moderate quality evidence).

15 Nitrofurantoin versus co-trimoxazole

- 16 Nitrofurantoin (1 to 2 mg/kg daily) significantly reduced the recurrence of
- 17 symptomatic UTI at 6 months compared with co-trimoxazole (2 mg/kg daily) (1 RCT,
- n=132: 25.8% versus 45.5%; RR 0.57, 95% CI 0.35 to 0.92; NNT 6 [95% CI 3 to 27];
 very low guality evidence).

20 Nitrofurantoin versus cefixime

- 21 Nitrofurantoin (1 mg/kg daily) did not reduce the risk of obtaining a repeat positive
- 22 culture at the end of prophylaxis (6 to 12 months) compared with cefixime (2 mg/kg
- 23 daily; 1 RCT, n=57: 10% versus 7.4%; risk difference 0.03 95% CI -0.12 to 0.17;
- 24 moderate quality evidence).

25 3.6.3 Antibiotic dosing and course length

26 No evidence from systematic reviews or RCTs was identified.

4 Safety and tolerability

2 Details of safety and tolerability outcomes from studies included in the evidence

3 review are shown in <u>appendix H: GRADE profiles</u>. The main results are summarised
4 below.

See the <u>summaries of product characteristics</u>, <u>British National Formulary</u> (BNF) and
 <u>BNF for children</u> (BNF-C) for information on contraindications, cautions and adverse
 effects of individual medicines, and for appropriate use and dosing in specific
 populations, for example, hepatic impairment, renal impairment, pregnancy and
 breastfeeding.

10 4.1 Non-pharmacological interventions

11 4.1.1 Probiotics (lactobacillus)

12 No safety data were reported for lactobacillus compared with placebo. Schwenger et al. (2015) assessed the effect of probiotic prophylaxis for the prevention of recurrent 13 14 urinary tract infection (UTI) in adults (men and non-pregnant women) and children 15 compared with antibiotics. Safety data were described in 4 studies included in the 16 review, however they were not pooled in the analysis (justification not provided). A 17 single study (NAPRUTI Study II 2006) compared probiotics with antibiotics, and 18 showed there is no significant difference in the number of adverse events 19 experienced by those who receive antibiotics compared with those who receive 20 probiotics (1 RCT, n=152: 5.6% versus 11.8%; RR and 95% CI not stated; calculated 21 by NICE as RR 0.47, 95% CI 0.20 to 1.12; low quality evidence). In the same study, 22 there is no significant difference between the proportions of participants who 23 experienced at least 1 adverse event having received probiotics compared with those 24 who received antibiotics (1 RCT, n= 152: 52.8% versus 58.3%; RR and 95% CI not 25 stated; calculated by NICE as RR 0.91 95% CI 0.73 to 1.13; low guality evidence). 26 Another study included in the review (Stapleton et al. 2011), reported that a single 27 participant withdrew from treatment in the lactobacillus group due to a lack of 28 efficacy.

29 4.1.2 D-Mannose

<u>Kranjcec et al. (2014)</u> assessed the safety of D-mannose compared with an antibiotic
 (nitrofurantoin) in non-pregnant women who presented with current UTI and a history
 of recurrent UTI. While Kranjcec et al. (2014) included a no treatment study arm, no
 adverse events were reported for these participants.

34 D-mannose versus placebo or no treatment

35 No relevant evidence was identified.

36 D-mannose versus antibiotic

- 37 D-mannose significantly reduced adverse events, such as diarrhoea, nausea, and
- 38 vaginal burning, in non-pregnant women when compared with nitrofurantoin (n=206:
- 39 7.8% versus 28.2%; RR 0.28, 95% CI 0.13 to 0.57; NNH 5 [95% CI 4 to 10]; high
- 40 quality evidence).

4.1.3 Cranberry 1

2 Jepson et al. 2012 assessed the safety of prophylactic cranberry products (24 RCTs,

3 n=4,473) comparing cranberry products with placebo or no treatment, or antibiotics.

4 The authors pooled safety data (any gastrointestinal effect) across several adult

5 subgroups including women, and elderly women and men. Data on children were not 6 available.

7 Cranberry products versus placebo or no treatment

8 Prophylactic cranberry products in comparison with placebo or no treatment did not

9 significantly affect the incidence of any gastrointestinal adverse events (4 RCTs,

10 n=597: 3% versus 3.3%; RR 0.83, 95% CI 0.31 to 2.27; low quality evidence).

11 Cranberry products versus antibiotics

12 Prophylactic cranberry products in comparison with antibiotics did not significantly

13 affect the incidence of gastrointestinal adverse events (2 RCTs, n=344: 9.6% versus

14 12.0%; RR 0.78, 95% CI 0.42 to 1.42; low quality evidence).

15 **4.2** Non-antimicrobial pharmacological interventions

4.2.1 16 Oestrogens

17 Hormone replacement therapy (HRT) increases the risk of venous thromboembolism,

18 stroke, endometrial cancer (reduced by a progestogen), breast cancer, and ovarian

19 cancer; there is an increased risk of coronary heart disease in women who start

20 combined HRT more than 10 years after menopause (MHRA Drug Safety Update,

21 November 2015; British National Formulary [BNF], December 2017). Before

22 prescribing HRT, health professionals should consider carefully the potential benefits

23 and risks for every woman. The minimum effective dose of HRT should be used for 24

the shortest duration (MHRA Drug Safety Update, November 2015). The endometrial

25 safety of long-term or repeated use of topical vaginal oestrogens is uncertain; 26 treatment should be reviewed at least annually, with special consideration given to

27 any symptoms of endometrial hyperplasia or carcinoma (BNF August 2018).

28 Perrotta et al. (2008) identified 2 small RCTs that reported on the safety of oral

29 oestrogens compared with placebo. Adverse events reported in these RCTs were

30 breast tenderness or discomfort, or vaginal bleeding or spotting. There were

31 significantly more adverse events with oral oestrogen compared with placebo

32 (Perrotta et al. 2008, 2 RCTs, n=104; 23.5% versus 3.8%; RR 5.11, 95% CI 1.39 to

33 18.76; NNH 5 [95% CI 3 to 14]; high quality evidence).

34 Perrotta et al. (2008) also identified 2 RCTs that reported on the safety of vaginal 35 oestrogens compared with placebo. Safety results were reported in 2 ways, as 36 pooled analysis and RCT-based results. Overall, results suggested that vaginal

37 oestrogen was associated with more adverse events (vaginal bleeding,

38 nonphysiologic discharge, vaginal irritation, burning, or itching) when compared with

39 placebo (2 RCTs, n=201: 23.3% versus 5.1%; RR 4.57, 95% CI 1.81 to 11.5; NNH 5

40 [95% CI 3 to 11]; low quality evidence). Furthermore, there were significantly more

41 adverse events (burning, itching, or vaginal bleeding) with vaginal oestrogen

42 compared with oral antibiotics (Perrotta et al. 2008, 2 RCTs, n=216: 16.4% versus

43 0%; RR 12.86, 95% CI 1.75 to 94.29; NNH 6 [95% CI 4 to 10]; moderate quality 44 evidence).

Antimicrobials 4.3 1

2 Antibiotic-associated diarrhoea is estimated to occur in 2 to 25% of people taking

antibiotics, depending on the antibiotic used (NICE clinical knowledge summary 3 4 [CKS]: diarrhoea - antibiotic associated).

5 About 10% of the general population claim to have a penicillin allergy; this has often been because of a skin rash that occurred during a course of penicillin in childhood. 6 Fewer than 10% of people who think they are allergic to penicillin are truly allergic. 7 8 Therefore, penicillin allergy can potentially be excluded in 9% of the population. 9 People with a history of immediate hypersensitivity to penicillins may also react to cephalosporins and other beta-lactam antibiotics. The most common side effect with 10 11 penicillins is diarrhoea, which can also cause antibiotic-associated colitis. Diarrhoea 12 is most common with broad-spectrum penicillins (such as amoxicillin and co-13 amoxiclav) (BNF August 2018).

14 Quinolones, including ciprofloxacin, cause arthropathy in the weight-bearing joints of 15 immature animals and are generally not recommended in children or young people

16 who are growing (BNF August 2018).

17 Nitrofurantoin should be used with caution in those with renal impairment. Adults 18 (especially the elderly) and children on long-term therapy should be monitored for

19 liver function and pulmonary symptoms, with nitrofurantoin discontinued if there is a 20 deterioration in lung function (BNF August 2018).

21 Trimethoprim has a teratogenic risk in the first trimester of pregnancy (folate 22 antagonist), and manufacturers advise avoidance during pregnancy (BNF August 23 2018).

24 Co-trimoxazole is currently under restriction for use in the UK. It is advised that it

25 should only be used in UTI where there is bacteriological evidence of sensitivity to

26 co-trimoxazole. Co-trimoxazole should be used with caution in those with asthma, or

27 people with blood disorders, GP6D deficiency or infants under 6 weeks (except for

28 treatment or prophylaxis of pneumocystis pneumonia) (BNF August 2018).

4.3.1 29 Antibiotics in non-pregnant women

A systematic review (Albert et al. 2004) assessed the safety of antibiotic prophylaxis 30 31 for the prevention of recurrent UTI in non-pregnant women.

32 Antibiotic prophylaxis did not significantly increase the incidence of severe side

33 effects compared with placebo (10 RCTs, n=420: 4% versus 2.1%; RR 1.58, 95% CI

34 0.47 to 5.28; low quality evidence). However, antibiotics did increase the incidence of

35 'other side effects' (defined as non-serious side effects such vagina itching and

36 nausea) compared with placebo (10 RCTs, n=420: 15.1% versus 7.7%; RR 1.78,

95% CI 1.06 to 3.00; NNH 13 [95% CI 7 to 70]; low quality evidence). 37

38 One RCT included in the systematic review (Melekos et al. 1997) found no significant 39 difference in the number of non-serious side effects, between premenopausal women

40 who took ciprofloxacin (125 mg) as a single dose immediately after sexual

41 intercourse, or once daily at night (1 RCT, n=135: 5.7% versus 13.8%; RR 0.41 95%

42 CI 0.13 to 1.28; low quality evidence).

43 Zhong et al. (2011) (n=83) found that intermittent single-dose antibiotics significantly

44 reduced the incidence of adverse events compared with continuous antibiotics (n=73:

45 63.6% versus 92.5%; RR and 95% CI not stated; calculated by NICE as RR 0.69

46 95% CI 0.52 to 0.9; NNH 3 [95% CI 2 to 9]; moderate quality evidence).

1 4.3.2 Antibiotics in pregnant women

No evidence was identified regarding the safety of antibiotic prophylaxis in pregnant
 women.

4 4.3.3 Antibiotics in adults and children

5 <u>Muller et al. (2017)</u> assessed the safety of nitrofurantoin, given as long-term

6 prophylaxis (defined as greater than 14 days) for the primary or secondary

7 prevention of UTI in men, non-pregnant women (pre- or post-menopausal) and

8 children (predominantly female children).

9 Overall, the use of nitrofurantoin as prophylaxis (for at least 3 months) for recurrent

10 UTI, significantly increased the risk of experiencing mild (not defined) adverse effects

11 compared with other antibiotics (amoxicillin, penicillin, pivmecillinam, cefaclor,

12 cefixime, cinoxacin, norfloxacin, co-trimoxazole, trimethoprim, or methenamine

13 hippurate) (22 RCTs n=1,205: 30.6% versus 11.7%; RR 2.24 95% CI 1.77 to 2.83;

14 NNH 5 [95% CI 4 to 6]; low quality evidence).

When specific antibiotics were compared, there were significantly more mild adverse
effects with nitrofurantoin compared with beta-lactams (5 RCTs, n=275: 25% versus
12.2%; RR 1.99, 95% CI 1.19 to 3.32; NNH 7 [95% CI 4 to 28]; very low quality
evidence); trimethoprim (4 RCTs, n=330: 42% versus 14.6%; RR 2.20 95% CI 1.51
to 3.20; NNH 3 [95% CI 2 to 4]; moderate quality evidence); and methenamine

20 hippurate (2 RCTs, n=196: 35.8% versus 7%; RR 4.22, 95% CI 2.06 to 8.67; NNH 3

21 [95% Cl 2 to 6]; moderate quality evidence).

However, when nitrofurantoin was compared with quinolones or co-trimoxazole, there
were no significant differences in the number of mild adverse effects (very low quality
evidence).

25 4.3.4 Antibiotics in children

Williams and Craig (2011) assessed the safety of antibiotic prophylaxis in
comparison with placebo or no treatment in children with recurrent UTI. Antibiotics
did not significantly affect the incidence of adverse events reported (2 RCTs, n=914:
3.8% versus 2.4%; RR 2.31, 95% CI 0.03 to 170.67; very low quality evidence) or the
number of withdrawals due to adverse events (2 RCTs, n=576: 1.4% versus 3.5%;
RR 0.40, 95% CI 0.13 to 1.26; very low quality evidence).

Nitrofurantoin significantly reduced the incidence of adverse events compared with
trimethoprim (1 RCT, n=60: 25.8% versus 62.1%; RR 0.42, 95% CI 0.21 to 0.81;
NNH 2 [95% CI 1 to 8]; low quality evidence).

Nitrofurantoin significantly increased the incidence of adverse events compared with cefixime (1 RCT, n=120: 61.7% versus 28.3%; risk difference 2.18, 95% CI 1.39 to

37 3.41; NNH 3 [95% CI 2 to 6]; moderate guality evidence).

38

5 Antimicrobial resistance

2 The consumption of antimicrobials is a major driver for the development of antibiotic 3 resistance in bacteria, and the 3 major goals of antimicrobial stewardship are to:

- 5 resistance in bacteria, and the 5 major goals of antimicrobial steward
- optimise therapy for individual patients
- 5 prevent overuse, misuse and abuse, and
- minimise development of resistance at patient and community levels.

7 The NICE guideline on antimicrobial stewardship: systems and processes for

8 <u>effective antimicrobial medicine use</u> (2015) recommends that the risk of antimicrobial

9 resistance for individual patients and the population as a whole should be taken into

10 account when deciding whether or not to prescribe an antimicrobial.

- When antimicrobials are necessary to treat an infection that is not life-threatening, a
 narrow-spectrum antibiotic should generally be first choice. Indiscriminate use of
- 13 broad-spectrum antibiotics creates a selective advantage for bacteria resistant even
- 14 to these 'last-line' broad-spectrum agents, and also kills normal commensal flora
- 15 leaving people susceptible to antibiotic-resistant harmful bacteria such as *C. difficile*.
- 16 For infections that are not life-threatening, broad-spectrum antibiotics (for example,

17 co-amoxiclav, quinolones and cephalosporins) need to be reserved for second-

18 choice treatment when narrow-spectrum antibiotics are ineffective (<u>CMO report</u>

19 <u>2011</u>).

20 The ESPAUR report 2016 reported that antimicrobial consumption declined

significantly between 2014 and 2015, with community prescribing from general and

22 dental practice decreasing by more than 6%. Antibiotic prescribing in primary care in

23 2015 is at the lowest level since 2011, with broad-spectrum antibiotic use (antibiotics

that are effective against a wide range of bacteria) continuing to decrease in primarycare.

26 5.1 Antimicrobial resistance in the included studies

27 5.1.1 Cranberry products

28 Beerepoot et al. (2011) (n=221) reported that *E. coli* isolates from women receiving co-trimoxazole showed antibiotic resistance for amoxicillin, trimethoprim, and co-29 30 trimoxazole at 1 month prophylaxis (70% resistance). This reduced at 1 and 31 3 months after stopping prophylaxis, returning to baseline at 12 months. E. coli 32 isolates from women receiving cranberry products did not show antibiotic resistance. However, prophylactic cranberry products did reduce the development of antibiotic 33 34 resistance in premenopausal women compared with prophylaxis with co-trimoxazole 35 (moderate quality evidence).

Uberos et al. 2016 (n=192) found that cranberry products did not show a significant
 benefit in reducing the development of antibiotic resistance in children (n=192;

- 38 narrative results reported; moderate quality evidence). This study included an
- 39 unknown proportion of children with vesicoureteral reflux.

40 5.1.2 Antibiotic prophylaxis

- 41 Muller et al. (2017) reported resistance data from 1 RCT (n=15) comparing
- 42 nitrofurantoin prophylaxis with placebo in children. Weekly urine cultures showed *E*.
- 43 *coli* cultures were replaced over time by resistant strains including *Klebsiella* and
- 44 *Pseudomonas* spp., in children receiving nitrofurantoin prophylaxis, and this change

- 1 was not seen in children receiving placebo. However, there were no reports of
- 2 infection from resistant strains (low quality evidence).
- 3 Another RCT included in the systematic review (n= 130) compared nitrofurantoin and
- 4 trimethoprim prophylaxis in children. At baseline, 9% (6/67) of children randomised to
- 5 nitrofurantoin carried nitrofurantoin resistant strains, which decreased to 7% (4/60)
- 6 during prophylaxis. 8% (5/63) of children randomised to trimethoprim carried
- 7 trimethoprim resistant strains at baseline, which increased to 47% (28/60) throughout
- 8 prophylaxis (very low quality evidence).

6 Other considerations

2 6.1 Resource impact

3 6.1.1 Antibiotic prophylaxis

4 Recommended antibiotics (nitrofurantoin, trimethoprim, amoxicillin and cefalexin) are

5 available as generic formulations, but there is currently no generic formulation of 6 pivmecillinam, see <u>Drug Tariff</u> for costs.

7 Nitrofurantoin 25mg/5ml oral suspension is more expensive than other oral

8 suspensions, such as trimethoprim 50mg/5ml. The cost of a 300 ml bottle of

9 nitrofurantoin is £446.95 compared with £4.87 for a 100 ml bottle of trimethoprim

10 (<u>Drug Tariff</u>, September 2018).

11 6.2 Medicines adherence

12 Medicines adherence may be a problem for some people with medicines that require

13 frequent dosing (for example, some antibiotics) or longer treatment duration (for

14 example, with antibiotic prophylaxis). See the NICE guideline on <u>medicines</u>

15 <u>adherence</u>).

16 6.3 Regulatory status

17 6.3.1 Oestrogens

18 A range of oral and vaginal oestrogens (for example, estradiol), with or without

19 progestogens, are available for use in managing menopausal symptoms and

20 prevention of osteoporosis. See the <u>summaries of product characteristics</u> for

21 information on licensed indications of individual medicines. None are specifically

22 licensed for preventing recurrent urinary tract infections, so use for this indication

would be <u>off label</u>. The prescriber should follow relevant professional guidance,
 taking full responsibility for the decision. Informed consent should be obtained and

documented. See the General Medical Council's <u>Good practice in prescribing and</u>

26 <u>managing medicines and devices for further information</u>.

27 6.3.2 Antibiotics

Amoxicillin is not licensed for preventing UTIs, so use for this indication would be <u>off</u> <u>label</u>. The prescriber should follow relevant professional guidance, taking full

30 responsibility for the decision. Informed consent should be obtained and

documented. See the <u>General Medical Council's Good practice in prescribing and</u>

32 <u>managing medicines and devices</u> for further information.

33

7 Terms used in the guideline

2 7.1.1 Vesicoureteric reflux

- 3 Vesicoureteric reflux occurs when there is damage to the valve between the bladder
- 4 and the ureters (tubes which carry urine away from the kidney into the bladder),
- 5 causing it to no longer working properly. This means that urine may flow backwards,
- 6 and sometimes reach as far back as the kidneys. This is problematic when the urine
- 7 is infected with bacteria, as the infection can reach the kidneys, and result in a very
- 8 severe urinary tract infection otherwise known as acute pyelonephritis, or worse. This
- 9 is common in children (1 in 100), and can lead to multiple urinary tract infections.
- 10 Most children with the condition, find that it resolves as they get older without
- 11 intervention.

1 Appendices

2

Appendix A: Evidence Sources

Key area	Key question(s)	Evidence sources
Background	 What is the natural history of the infection? What is the expected duration and severity of symptoms with or without antimicrobial treatment? What are the most likely causative organisms? What are the usual symptoms and signs of the infection? What are the known complication rates of the infection, with and without antimicrobial treatment? Are there any diagnostic or prognostic factors to identify people who may or may not benefit from an antimicrobial? 	 NICE guideline CG160: Fever in under 5s: assessment and initial management (2017) NICE guideline NG15: Antimicrobial stewardship: systems and processes for effective antimicrobial medicine use (2015) NICE guideline NG63: Antimicrobial stewardship: changing risk-related behaviours in the general population (2017) NICE guideline CG54: Urinary tract infection in under 16s: diagnosis and management (updated 2017) NICE Quality standard QS90: Urinary tract infections in adults (2015) NICE Clinical knowledge summary on UTI (lower) – women NICE Clinical knowledge summary on UTI (lower) – men
Safety netting	What safety netting advice is needed for managing the infection?	 NICE clinical knowledge summary on <u>UTI</u> (lower) - women NICE clinical knowledge summary on <u>UTI</u> (lower) - men

Key area	Key question(s)	Evidence sources
		NICE guideline CG54: <u>Urinary tract infection</u> <u>in under 16s: diagnosis and management</u> (updated 2017)
Red flags	What symptoms and signs suggest a more serious illness or condition (red flags)?	 NICE clinical knowledge summary on <u>UTI</u> (lower) - women NICE clinical knowledge summary on <u>UTI</u> (lower) - men NICE clinical knowledge summary on <u>UTI - children</u>
Non-pharmacological interventions	What is the clinical effectiveness and safety of non- pharmacological interventions for managing the infection or symptoms?	Evidence review - see appendix F for included studies
Non-antimicrobial pharmacological interventions	What is the clinical effectiveness and safety of non- antimicrobial pharmacological interventions for managing the infection or symptoms?	 Evidence review - see appendix F for included studies <u>MHRA Drug Safety Update (November 2015)</u> <u>British National Formulary (BNF)</u> (August 2018)
Antimicrobials	What is the clinical effectiveness and safety of antimicrobials for managing the infection or symptoms?	 Evidence review - see appendix F for included studies NICE guideline CG160: Fever in under 5s: assessment and initial management (2017) NICE clinical knowledge summary on diarrhoea – antibiotic associated BNF (August 2018)
	Which people are most likely to benefit from an antimicrobial?	 Evidence review - see appendix F for included studies
	Which antimicrobial should be prescribed if one is indicated (first, second and third line treatment, including people with drug allergy)?	Evidence review - see appendix F for included studies

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Key area	Key question(s)	Evidence sources
	What is the optimal dose, duration and route of administration of antimicrobials?	 Evidence review - see appendix F for included studies <u>BNF</u> (August 2018) <u>BNF for children</u> (BNF-C) (August 2018) <u>Summary of product characteristics</u>
Antimicrobial resistance	What resistance patterns, trends and levels of resistance exist both locally and nationally for the causative organisms of the infectionWhat is the need for broad or narrow spectrum antimicrobials?What is the impact of specific antimicrobials on the development of future resistance to that and other antimicrobials?	 Evidence review - see appendix F for included studies NICE guideline NG15: <u>Antimicrobial</u> <u>stewardship: systems and processes for</u> <u>effective antimicrobial medicine use</u> (2015)<u>European surveillance programme for</u> <u>antimicrobial utilisation and resistance</u> (ESPAUR) report (2016) <u>Chief medical officer (CMO) report (2011)</u>
Resource impact	What is the resource impact of interventions (such as escalation or de-escalation of treatment)?	 Evidence review - see appendix F for included studies <u>Drug Tariff</u> (September 2018)
Medicines adherence	What are the problems with medicines adherence (such as when longer courses of treatment are used)?	 Evidence review - see appendix F for included studies NICE guideline NG76: <u>Medicines adherence:</u> <u>involving patients in decisions about</u> <u>prescribed medicines and supporting</u> <u>adherence</u> (2009)
Regulatory status	What is the regulatory status of interventions for managing the infection or symptoms?	 <u>Summary of product characteristics</u> General Medical Council's <u>Good practice in prescribing and managing medicines and devices</u> (2013)

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² Appendix B: Review protocol

Review	protocol for recur	rent urinary tract infections	Notes
I	Review question	What pharmacological (antimicrobial and non-antimicrobial) and non-pharmacological interventions are effective in managing recurrent urinary tract infections (UTIs)?	 antimicrobial includes antibiotics (treatment and prophylaxis) non-antimicrobial includes analgesia and cranberry products search will include terms for recurrent urinary tract infections
II	Types of review question	Intervention questions will primarily be addressed through the search.	These will, for example, also identify natural history in placebo groups and causative organisms in studies that use laboratory diagnosis, and relative risks of differing management options.
III	Objective of the review	 To determine the effectiveness of prescribing interventions in managing recurrent UTIs to address antimicrobial resistance. In line with the major goals of antimicrobial stewardship. This includes interventions that lead prescribers to: optimise therapy for individuals reduce overuse, misuse or abuse of antimicrobials. All of the above will be considered in the context of national antimicrobial resistance patterns where available, if not available committee expertise will be used to guide decision-making. 	 The secondary objectives of the review of studies will include: indications for prescribing an antimicrobial (for example 'red flags' and illness severity, thresholds for treatment and individual patient factors affecting choice of antimicrobial indications for no or delayed antimicrobial indications for non-antimicrobial interventions

Review	protocol for recur	rent urinary tract infections	Notes
			 antimicrobial choice, optimal dose, duration (specifically length of treatment) and route for specified antimicrobial(s) the natural history of the infection
IV	Eligibility criteria – population/ disease/ condition/ issue/domain	 Population: Adults and children (aged 72 hours and older) with recurrent UTIs (lower or upper) of any severity. The definition of 'recurrence' of UTI varies: 2 UTIs in 6 months or ≥3 UTIs in 1 year in non-pregnant women (Source: PHE guidance: definition has also been applied to all women). 2 or more UTIs in a 3-month period in men aged 16 years and over (Source: CKS) In children (NICE CG54) 2 or more episodes of UTI with acute pyelonephritis/upper UTI or 1 episode of UTI with acute pyelonephritis/upper UTI plus 1 or more episode of UTI with cystitis/lower UTI, or 3 or more episodes of UTI with cystitis/lower UTI This review protocol includes recurrent UTI (defined by any of the above criteria) in 	 Subgroups of interest, those: with protected characteristics under the Equality Act 2010. with true allergy pregnant women men children (possible age groups) older people (frailty, care home resident, dementia) people with 'complicated¹' lower UTI people with upper UTI people with risk factors² for increased resistance
		non-pregnant and pregnant women, men and children. Consideration will be given to	

¹ Complicated UTI: UTI with one or more factors that predispose to persistent infection, recurrent infection or treatment failure, such as abnormal urinary tract, virulent organism, impaired host defences (diabetes mellitus, immunocompromised) or impaired renal function (Source: CKS)

² Risk factors for increased resistance include: care home resident, recurrent UTI, previous hospitalisation, unresolving urinary symptoms, recent travel to country with increased resistance, previous UTI resistant to antibiotics (previous antibiotic use [trimethoprim]) (Source PHE management of infection guidance)

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Review	protocol for recur	rent urinary tract infections	Notes
		differing management in subgroups based on age, gender, pregnancy, complicating factors and risk of resistance. Studies that use for example symptoms or signs (prognosis), clinical diagnosis or microbiological methods for diagnosing the condition.	
V	Eligibility criteria – intervention(s)/ exposure(s)/ prognostic factor(s)	 The review will include studies which include: Non-pharmacological interventions³. Non-antimicrobial pharmacological interventions⁴. Antimicrobial pharmacological interventions⁵. For the treatment of recurrent UTI in primary, secondary or other care settings (for example walk-in-centres, urgent care, and minor ailment schemes) either by prescription or by any other legal means of supply of medicine (for example patient group direction). 	Limited to those interventions commonly in use (as agreed by the committee)
VI	Eligibility criteria – comparator(s)/ control or reference (gold) standard	 Any other plausible strategy or comparator, including: Placebo or no treatment Non-pharmacological interventions. Non-antimicrobial pharmacological interventions. Other antimicrobial pharmacological interventions. 	

³Non-pharmacological interventions include: no intervention, watchful waiting, delayed prescribing, self-care prevention (avoiding bubble bath, appropriate wiping etc.)

⁴ Non-antimicrobial pharmacological interventions include: analgesics, NSAIDs, cranberry products, barley, D-mannose

⁵ Antimicrobial pharmacological interventions include: delayed (back-up) prescribing, standby or rescue therapy, prophylaxis (including post-coital and rotation of antibiotics) narrow or broad spectrum, single, dual or triple therapy, escalation or de-escalation of treatment. Antibiotics included in the search include those named in current guidance (plus the class to which they belong) plus other antibiotics agreed by the committee

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Review	v protocol for recur	rent urinary tract infections	Notes
VII	Outcomes and	a) Clinical outcomes such as:mortality	The committee have agreed that the following outcomes are critical:
	prioritisation	 infection cure rates (number or proportion of people with resolution of symptoms at a given time point, incidence of escalation of treatment) 	 reduction in number of recurrent⁶ episodes
		 time to clinical cure (mean or median time to resolution of illness) 	reduction in symptoms
		reduction in symptoms (duration or severity)	(duration or severity) for example difference in time
		rate of complications with or without treatment	to substantial improvement
		• safety, tolerability, and adverse effects.	• time to clinical cure (mean
		 b) Thresholds or indications for antimicrobial treatment (which people are most, or least likely to benefit from antimicrobials) 	or median time to resolution of illness)
		 c) Changes in antimicrobial resistance patterns, trends and levels as a result of treatment. 	 rate of complications⁷ (including mortality and
		d) Patient-reported outcomes, such as medicines adherence, patient experience and patient satisfaction.	deterioration in renal function) with or without
		e) Ability to carry out activities of daily living.	treatment, including escalation of treatment
		f) Service user experience.	health and social care
		g) Health and social care related quality of life, including long-term harm or disability.	utilisation (including length
		 h) Health and social care utilisation (including length of stay, planned and unplanned contacts). 	of stay, ITU stays, planned and unplanned contacts).
		The Committee considered which outcomes should be prioritised when multiple outcomes are reported (critical and important outcomes). Additionally, the Committee were asked to consider what clinically important features of study design may be important for this condition (for example length of study follow-up, treatment	 thresholds or indications for antimicrobial treatment (which people are most, or least likely to benefit from antimicrobials)

⁶ Recurrence may be due to underlying causes which require further investigation (for example stones, less usual pathogens etc)
⁷ Ascending infection leading to pyelonephritis, renal failure or sepsis, and in pregnancy, pre-term labour developmental delay or cerebral palsy in the infant and foetal death. In men, prostate involvement. Also urinary stones, risk of blood infections (bacteraemia), renal abscess, renal scarring in children, neonatal sepsis.

Review	protocol for recur	rent urinary tract infections	Notes
		failure/recurrence, important outcomes of interest such as sequela or progression to more severe illness).	 an individual's risk factors for resistance and choice of antibiotic The committee have agreed that the following outcomes are important: patient-reported outcomes, such as medicines adherence, patient experience changes in antimicrobial resistance patterns, trends and levels as a result of treatment
VIII	Eligibility criteria – study design	 The search will look for: Systematic review of randomised controlled trials (RCTs) RCTs If insufficient evidence is available progress to: Controlled trials Systematic reviews of non-randomised controlled trials Non-randomised controlled trials Observational and cohort studies Pre and post intervention studies (before and after) Time series studies 	Committee to advise the NICE project team on the inclusion of information from other condition specific guidance and on whether to progress due to insufficient evidence.
IX	Other inclusion exclusion criteria	 The scope sets out what the guidelines will and will not include (exclusions). Further exclusions specific to this guideline include: non-English language papers, studies that are only available as abstracts in relation to antimicrobial resistance, non-UK papers. 	

Review	protocol for recur	rent urinary tract infections	Notes
Х	Proposed sensitivity/ sub- group analysis, or meta- regression	The search may identify studies in population subgroups (for example adults, older adults, children (those aged under 18 years of age), and people with co-morbidities or characteristics that are protected under the Equality Act 2010 or in the NICE equality impact assessment). These will be analysed within these categories to enable the production of management recommendations.	
XI	Selection process – duplicate screening/ selection/ analysis	 All references from the database searches will be downloaded, de-duplicated and screened on title and abstract against the criteria above. A randomly selected initial sample of 10% of records will be screened by two reviewers independently. The rate of agreement for this sample will be recorded, and if it is over 90% then remaining references will screened by one reviewer only. Disagreement will be resolved through discussion. Where abstracts meet all the criteria, or if it is unclear from the study abstract whether it does, the full text will be retrieved. If large numbers of papers are identified and included at full text, the Committee may consider prioritising the evidence for example, evidence of higher quality in terms of study type or evidence with critical or highly important outcomes. 	
XII	Data management (software)	Data management will be undertaken using EPPI-reviewer software. Any pairwise meta-analyses will be performed using Cochrane Review Manager (RevMan5). 'GRADEpro' will be used to assess the quality of evidence for each outcome.	
XIII	Information sources – databases and dates	 Medline; Medline in Process; Embase; Cochrane database of systematic reviews (CDSR); Database of abstracts of effectiveness (DARE) (legacy); Cochrane Central Register of Controlled Trials (CENTRAL); Health Technology Assessment (HTA) database; Clinicaltrials.gov All the above to be searched from 2006 to present day. Filters for systematic reviews, RCTs and comparative studies to be applied, unless numbers without filters are low Searches to be limited to studies reported in English. 	

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Review	Review protocol for recurrent urinary tract infections Notes			
		Animal studies and conference abstracts to be excluded		
		Medicines and Healthcare products Regulatory Agency (MHRA) website; European Medicines Agency (EMA) website; U.S. Food and Drug Administration (FDA) website; Drug Tariff; MIMs		
		 The above to be searched for advice on precautions, warnings, undesirable effects of named antimicrobials. 		
XIV	Identify if an update	Not applicable at this time.		
XV	Author contacts	Web: <u>https://www.nice.org.uk/guidance/indevelopment/gid-apg10002</u>		
		Email: <u>infections@nice.org.uk</u>		
XVI	Highlight if amendment to previous protocol	For details please see the interim process guide (2017).		
XVII	Search strategy – for one database	For details please see appendix C.		
XVIII	Data collection process – forms/duplicate	GRADE profiles will be used, for details see appendix H.		
XIX	Data items – define all variables to be collected	GRADE profiles will be used, for details see appendix H.		
хх	Methods for assessing bias at	Standard study checklists will be used to critically appraise individual studies. For details please see section 6.2 of <u>Developing NICE guidelines: the manual</u> . The risk of		

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Review	protocol for recurr	rent urinary tract infections	Notes
	outcome/ study level	bias across all available evidence will be evaluated for each outcome using an adaptation of the 'Grading of Recommendations Assessment, Development and Evaluation (GRADE) toolbox' developed by the international GRADE working group http://www.gradeworkinggroup.org/	
XXI	Criteria for quantitative synthesis (where suitable)	For details please see section 6.4 of <u>Developing NICE guidelines: the manual</u>	
XXII	Methods for analysis – combining studies and exploring (in)consistency	For details please see the interim process guide (2017)	
XXIII	Meta-bias assessment – publication bias, selective reporting bias	For details please see the interim process guide (2017)	
XXIV	Assessment of confidence in cumulative evidence	For details please see the interim process guide (2017)	
XXV	Rationale/ context – Current management	For details please see the introduction to the evidence review in the guideline.	
XXVI	Describe contributions of	A multidisciplinary committee developed the guideline. The committee was convened by NICE and chaired by Dr Tessa Lewis in line with section 3 of <u>Developing NICE</u> guidelines: the manual.	

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Review	protocol for recur	Notes	
	authors and guarantor	Staff from NICE undertook systematic literature searches, appraised the evidence, conducted meta-analysis and cost-effectiveness analysis where appropriate, and drafted the guideline in collaboration with the committee. For details please see the methods chapter of the full guideline.	
XXVII	Sources of funding/support	Developed and funded by NICE.	
XXVIII	Name of sponsor	Developed and funded by NICE.	
XXIX	Roles of sponsor	NICE funds and develops guidelines for those working in the NHS, public health, and social care in England.	

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Appendix C: Literature search strategy

2

3 <u>1 Search format</u>

- 4 The search strategy has been designed to cover four UTI protocols and it takes the following format:
- 5 Urinary Tract Infections
- 6 AND (Named Antibiotics OR Classes of Antibiotics OR Pain Relief OR NSAIDs OR Cranberry
- 7 Products OR Alkalinising agents OR Bladder instillations OR Drinking Fluids OR Prescribing
- 8 Strategies OR Self Care OR Catheter Removal)
- 9 AND (Systematic Reviews OR Randomised Controlled Trials OR Observational Studies)
- 10 AND Limits
- 11 Note there is an additional search in this format:
 - Named Antibiotics AND Drug Resistance AND Limits
- 12 13

14 <u>2 Overview of search results</u>

	No. of hits in	Position in the
	MEDLINE	strategy
Search without any limits	65,619	Line 178
Search with limits	14,263	Line 184
Search with limits and Systematic Reviews	2,428	Line 200
Search with limits and RCTs (not SRs)	2,230	Line 217
Search with limits and Observational Studies (not SRs or RCTs)	3,795	Line 240
Search with limits (without SRs, RCTs, Observational)	5,810	Line 241
Named Antibiotics AND Drug Resistance	48,201	Line 257
Named Antibiotics AND Drug Resistance with Limits	20,072	Line 262

15

16 <u>3 Contents of the search strategy</u>

Main concepts	Coverage	Position in strategy
Urinary Tract Infections	Urinary tract infections Cystitis Vesico-ureteral reflux Pyelonephritis Catheter-Related Infections Bacteriuria Urosepsis	Lines 1-20
Named Antibiotics	Urethritis Trimethoprim Nitrofurantoin Fosfomycin Methenamine hippurate Gentamicin Amikacin Tobramycin	Lines 21-84

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	Amoxicillin	
	Ampicillin	
	Co-amoxiclav	
	Pivmecillinam	
	Cefalexin	
	Cefotaxime	
	Cefixime	
	Ceftriaxone	
	Ciprofloxacin	
	Ofloxacin	
	Colistin	
	Ertapenem	
	Doxycycline	
	Septrin	
	Chloramphenicol	
	Tazocin	
	Aztreonam	
	Temocillin	
	Tigecycline	
	Vancomycin	
	Teicoplanin	
	Linezolid	
	Cefuroxime	
	Cefradine	
	Ceftazidime	
	Levofloxacin	
Classes of Antibiotics	Aminoglycosides	Lines 86-93
	Penicillins	LINE3 00-30
	Cephalosporins	
	Quinolones	
	Carbapenems	
	Tetracyclines	
Pain Relief	Paracetamol	
	Ibuprofen	Lines 96-111
	Naproxen	
	Codeine	
	Diclofenac	
	Analgesics	
	Non-steroidal anti-inflammatory drugs	
Non-pharmaceutical products	Cranberry products	Lines 113-119
	Barley products	
	Barley products D-Mannose	
Alkalinising agents		Lines 121-127
Alkalinising agents	D-Mannose Potassium citrate	
Alkalinising agents	D-Mannose Potassium citrate Sodium citrate	
	D-Mannose Potassium citrate Sodium citrate Sodium bicarbonate	Lines 121-127
Alkalinising agents Bladder instillations	D-Mannose Potassium citrate Sodium citrate Sodium bicarbonate Chlorhexidine solution	
Bladder instillations	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solution	Lines 121-127 Lines 129-133
	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solutionFluid therapy	Lines 121-127
Bladder instillations	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solutionFluid therapyDrinking water, beverages, fluids or	Lines 121-127 Lines 129-133
Bladder instillations Drinking Fluids	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solutionFluid therapyDrinking water, beverages, fluids orliquids	Lines 121-127 Lines 129-133 Lines 135-139
Bladder instillations	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solutionFluid therapyDrinking water, beverages, fluids orliquidsWatchful waiting	Lines 121-127 Lines 129-133
Bladder instillations Drinking Fluids	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solutionFluid therapyDrinking water, beverages, fluids orliquidsWatchful waitingNo intervention	Lines 121-127 Lines 129-133 Lines 135-139
Bladder instillations Drinking Fluids	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solutionFluid therapyDrinking water, beverages, fluids orliquidsWatchful waiting	Lines 121-127 Lines 129-133 Lines 135-139
Bladder instillations Drinking Fluids	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solutionFluid therapyDrinking water, beverages, fluids orliquidsWatchful waitingNo interventionActive surveillance	Lines 121-127 Lines 129-133 Lines 135-139
Bladder instillations Drinking Fluids	D-Mannose Potassium citrate Sodium citrate Sodium bicarbonate Chlorhexidine solution Sodium chloride solution Fluid therapy Drinking water, beverages, fluids or liquids Watchful waiting No intervention Active surveillance Delayed treatment	Lines 121-127 Lines 129-133 Lines 135-139
Bladder instillations Drinking Fluids	D-Mannose Potassium citrate Sodium citrate Sodium bicarbonate Chlorhexidine solution Sodium chloride solution Fluid therapy Drinking water, beverages, fluids or liquids Watchful waiting No intervention Active surveillance Delayed treatment Prescribing times	Lines 121-127 Lines 129-133 Lines 135-139
Bladder instillations Drinking Fluids Prescribing Strategies	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solutionFluid therapyDrinking water, beverages, fluids orliquidsWatchful waitingNo interventionActive surveillanceDelayed treatmentPrescribing timesAntibiotic prophylaxis	Lines 121-127 Lines 129-133 Lines 135-139 Lines 141-160
Bladder instillations Drinking Fluids	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solutionFluid therapyDrinking water, beverages, fluids orliquidsWatchful waitingNo interventionActive surveillanceDelayed treatmentPrescribing timesAntibiotic prophylaxisSelf management	Lines 121-127 Lines 129-133 Lines 135-139
Bladder instillations Drinking Fluids Prescribing Strategies	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solutionFluid therapyDrinking water, beverages, fluids orliquidsWatchful waitingNo interventionActive surveillanceDelayed treatmentPrescribing timesAntibiotic prophylaxisSelf managementSelf care secondary prevention	Lines 121-127 Lines 129-133 Lines 135-139 Lines 141-160
Bladder instillations Drinking Fluids Prescribing Strategies	D-MannosePotassium citrateSodium citrateSodium bicarbonateChlorhexidine solutionSodium chloride solutionFluid therapyDrinking water, beverages, fluids orliquidsWatchful waitingNo interventionActive surveillanceDelayed treatmentPrescribing timesAntibiotic prophylaxisSelf management	Lines 121-127 Lines 129-133 Lines 135-139 Lines 141-160

	Systematic Reviews	
	Reviews	
Randomised Controlled Trials	RCTs	Lines 201-215
	Controlled Clinical Trials	
	Cross over studies	
Observational Studies	Observational Study	Lines 218-238
	Epidemiologic Studies	
	Case-Control Studies	
	Cohort Studies	
	Cross-Sectional Studies	
	Controlled Before-After Studies	
Limits	2006-Current	Lines 179-184
	Exclude Animal studies	
	Exclude letters, editorials and letters	
Additional search	Drug resistance	Lines 242-262

2 <u>4 Key to search operators</u>

1	Medical Subject Heading (MeSH) term
Ехр	Explodes the MeSH terms to retrieve narrower terms in the hierarchy
.ti	Searches the title field
.ab	Searches the abstract field
*	Truncation symbol (searches all word endings after the stem)
adj <i>n</i>	Adjacency operator to retrieve records containing the terms within a specified number (<i>n</i>) of words of each other

3

4 <u>5 Search strategy for MEDLINE</u>

5 Database(s): Ovid MEDLINE(R) Epub Ahead of Print, In-Process & Other Non-Indexed Citations, Ovid

6 MEDLINE(R) Daily and Ovid MEDLINE(R) 1946 to Present

7 Search Strategy:

#	Searches	Results
1	exp urinary tract/	406398
2	exp urinary tract infections/	42175
3	exp cystitis/	8814
4	vesico-ureteral reflux/	7753
5	exp pyelonephritis/	14154
6	exp Urinary Calculi/	32650
7	Urethritis/	4483
8	Catheters, Indwelling/	17219
9	Urinary Catheters/	530
10	Urinary Catheterization/	13329
11	Catheter-Related Infections/	3344
12	Catheter Obstruction/	139

	13	(UTI or CAUTI or RUTI or cystitis* or bacteriuria* or pyelonephriti* or pyonephrosi* or pyelocystiti* or pyuri* or VUR or urosepsis* or uroseptic* or urosepses* or urethritis*).ti,ab.	38919
	14	((urin* or renal* or kidney*) adj1 (system* or tract* or calculus or calculi* or stone* or sepsis*)).ti,ab.	82884
	15	((bladder* or genitourin* or genito urin* or kidney* or pyelo* or renal* or ureter* or ureth* or urin* or urolog* or urogen*) adj3 (infect* or bacteria* or microbial* or block* or obstruct* or catheter* or inflamm*)).ti,ab.	87091
	16	((upper or lower) adj3 urin*).ti,ab.	21980
	17	(bladder* adj3 (ulcer* or ulcus)).ti,ab.	151
	18	(schistosomiasis adj3 (haematobia or hematobia or urin*)).ti,ab.	966
	19	((vesicorenal* or vesicoureteral* or vesicoureteric* or vesico renal* or vesico ureteral* or vesico ureteric* or bladder* or cystoureteral* or ureter* or urether* or nephropathy*) adj3 (backflow* or reflux*)).ti,ab.	7989
2	20	or/1-19	576113
2	21	Trimethoprim/	6280
2	22	(Trimethoprim* or Monotrim*).ti,ab.	14565
2	23	Nitrofurantoin/	2517
2	24	(Nitrofurantoin* or Genfura* or Macrobid*).ti,ab.	2980
2	25	Fosfomycin/	1685
2	26	(Fosfomycin* or Phosphomycin* or Fosfocina* or Monuril* or Monurol* or Fomicyt*).ti,ab.	2378
2	27	Methenamine/	1045
2	28	(Methenamine* or hexamine* or hippurate* or Hiprex*).ti,ab.	2411
2	29	Gentamicins/	17268
3	30	(Gentamicin* or Cidomycin*).ti,ab.	21976
3	31	Amikacin/	3751
3	32	(amikacin* or Amikin*).ti,ab.	8118
3	33	Tobramycin/	3973
3	34	(tobramycin* or Nebcin*).ti,ab.	6203
3	35	Amoxicillin/	8654
3	36	(Amoxicillin* or Amoxil*).ti,ab.	12541
3	37	Ampicillin/	12932
3	38	ampicillin*.ti,ab.	20478
3	39	Amoxicillin-Potassium Clavulanate Combination/	2301

(co-amoxiclav* or Coamoxiclav* or Amox-clav* or Amoxicillin-Clavulanic Acid* or Amoxicillin-40 Potassium Clavulanate Combination* or Amoxi-Clavulanate* or Clavulanate Potentiated Amoxycillin Potassium* or Clavulanate-Amoxicillin Combination* or Augmentin*).ti,ab. 41 Amdinocillin Pivoxil/ (pivmecillinam* or Pivamdinocillin* or Selexid*).ti,ab. Cefalexin/ (Cefalexin* or Cephalexin* or Keflex*).ti,ab. Cefotaxime/ cefotaxime*.ti,ab. Cefixime/ (cefixime* or Suprax*).ti,ab. Ceftriaxone/ (ceftriaxone* or Rocephin*).ti,ab. Ciprofloxacin/ (Ciprofloxacin* or Ciproxin*).ti,ab. Ofloxacin/ (ofloxacin* or Tarivid*).ti,ab. Colistin/ (Colistin* or Colistimethate* or Colimycin* or Coly-Mycin* or Colymycin* or Colomycin* or Promixin*).ti,ab. (Ertapenem* or Invanz*).ti,ab. Doxycycline/ (Doxycycline* or Efracea* or Periostat* or Vibramycin*).ti,ab. Trimethoprim, Sulfamethoxazole Drug Combination/ (Septrin* or Co-trimoxazole* or Cotrimoxazole* or Sulfamethoxazole Trimethoprim Comb* or Trimethoprim Sulfamethoxazole Comb*).ti,ab. Chloramphenicol/ (Chloramphenicol* or Cloranfenicol* or Kemicetine* or Kloramfenikol*).ti,ab. Piperacillin/ (Tazocin* or Piperacillin* or Tazobactam*).ti,ab. Aztreonam/ (Aztreonam* or Azactam*).ti,ab. (Temocillin* or Negaban*).ti,ab. (Tigecycline* or Tygacil*).ti,ab.

70	Vancomycin/	11836
71	(Vancomycin* or Vancocin*).ti,ab.	22446
72	Teicoplanin/	2067
73	(Teicoplanin* or Targocid*).ti,ab.	3233
74	Linezolid/	2421
75	(Linezolid* or Zyvox*).ti,ab.	4568
76	Cefuroxime/	2037
77	(Cefuroxime* or Cephuroxime* or Zinacef* or Zinnat* or Aprokam*).ti,ab.	3919
78	Cefradine/	540
79	(Cefradine* or Cephradine* or Nicef*).ti,ab.	699
80	Ceftazidime/	3461
81	(Ceftazidime* or Fortum* or Tazidime*).ti,ab.	7727
82	Levofloxacin/	2708
83	(Levofloxacin* or Evoxil* or Tavanic*).ti,ab.	6119
84	or/21-83	214218
85	20 and 84	18255
86	exp aminoglycosides/	142346
87	exp penicillins/	76761
88	exp cephalosporins/	39233
89	exp quinolones/	41144
90	exp Carbapenems/	8711
91	exp Tetracyclines/	44511
92	(Aminoglycoside* or Penicillin* or Cephalosporin* or Quinolone* or Carbapenem* or Tetracycline*).ti,ab.	120900
93	or/86-92	359234
94	20 and 93	22544
95	Anti-Infective Agents, Urinary/	2557
96	Acetaminophen/	15854
97	(paracetamol* or acetaminophen* or Panadol* or perfalgan* or calpol*).ti,ab.	20775
98	lbuprofen/	7581
99	(ibuprofen* or arthrofen* or ebufac* or rimafen* or brufen* or calprofen* or feverfen* or nurofen* or orbifen*).ti,ab.	11191
100	Naproxen/	3730
101	(Naproxen* or Naprosyn* or Stirlescent*).ti,ab.	5450

102 Codeine/	4237
103 (codeine* or Galcodine*).ti,ab.	4407
104 Diclofenac/	6823
(Diclofenac* or Voltarol* or Dicloflex* or Econac* or Fenactol* or Volsaid* or Enstar* or Diclomax* 105 or Motifene* or Rhumalgan* or Pennsaid*).ti,ab.	9698
106 (nsaid* or analgesic*).ti,ab.	87160
107 ((nonsteroid* or non steroid*) adj3 (anti inflammator* or antiinflammator*)).ti,ab.	34162
108 analgesics/	43460
109 exp analgesics, non-narcotic/	299959
110 analgesics, short-acting/	8
111 or/96-110	400073
112 20 and 111	10492
113 Vaccinium macrocarpon/	645
114 (cranberry* or cranberries* or vaccinium macrocarpon*).ti,ab.	1247
115 Hordeum/	8153
116 (barley* or hordeum*).ti,ab.	15407
117 Mannose/	8489
118 (mannose* or d-mannose* or dmannose*).ti,ab.	24493
119 or/113-118	45484
120 20 and 119	1500
121 potassium citrate/	245
122 (potassium citrate* or Effercitrate*).ti,ab.	546
123 (sodium citrate* or Cymalon* or Cystocalm* or Micolette* or Micralax*).ti,ab.	2644
124 sodium bicarbonate/	4205
125 (sodium bicarbonate* or S-Bicarb* or SodiBic* or Thamicarb* or Polyfusor*).ti,ab.	5477
((alkalizer* or alkalinisation* or alkalinization* or alkalinising or alkalinizing) adj3 (drug* or agent* or 126 therap*)).ti,ab.	191
127 or/121-126	10890
128 20 and 127	1049
129 Chlorhexidine/	7123
130 ((chlorhexidine or sodium chloride*) adj3 (solution* or diluent* or instillation* or intravesical*)).ti,ab.	3327
131 Administration, Intravesical/	3418
132 (bladder* adj3 (instillat* or drug admin*)).ti,ab.	540
133 or/129-132	13618

134	20 and 133	1976
135	Drinking/ or Drinking Behavior/	19308
136	Fluid therapy/	17515
137	exp Beverages/	114331
138	((water* or fluid* or liquid* or beverage* or drinks) adj3 (consumption* or consume* or consuming* or intake* or drink* or hydrat* or rehydrat*)).ti,ab.	80871
139	or/135-138	210996
140	20 and 139	6845
141	watchful waiting/	2278
142	Antibiotic Prophylaxis/	11779
143	"no intervention*".ti,ab.	6125
144	(watchful* adj2 wait*).ti,ab.	2077
145	(wait adj2 see).ti,ab.	1225
146	(active* adj2 surveillance*).ti,ab.	5705
147	(expectant* adj2 manage*).ti,ab.	2738
148	((prescription* or prescrib*) adj4 ("red flag" or strateg* or appropriat* or inappropriat* or unnecessary or defer* or delay* or no or non or behaviour* or behavior* or optimal or optimi* or reduc* or decreas* or declin* or rate* or improv* or postcoital* or postcoitus* or postsex* or postintercourse* or post coital* or post coitus* or post sex* or post intercourse* or night* or nocturnal* or prophylaxis* or prophylactic* or prevent* or preoperative* or pre operative* or perioperative* or peri operative* or postoperative* or post operative*)).ti,ab.	25168
149	((misuse* or "mis-use*" or overuse* or "over-use*" or "over-prescri*" or abuse*) adj4 (bacter* or antibacter* or anti-bacter* or "anti bacter*" or antimicrobial or anti-microbial or "anti microbial" or antibiot* or anti-biot* or "anti biot*")).ti,ab.	1761
150	((delay* or defer*) adj3 (treat* or therap* or interven*)).ti,ab.	26341
151	or/141-150	82704
152	anti-infective agents/ or exp anti-bacterial agents/ or exp anti-infective agents, local/	844581
153	(antibacter* or anti-bacter* or antibiot* or anti-biot* or antimicrobial* or anti-microbial*).ti,ab.	401551
154	152 or 153	1017858
155	(postcoital* or postcoitus* or postsex* or postintercourse* or post coital* or post coitus* or post sex* or post intercourse* or night* or nocturnal* or delay* or defer* or back-up* or backup* or immediate* or rapid* or short* or long* or standby or "stand by" or rescue or escalat* or "de- escalat*" or (prescribing adj strateg*) or "red flag*" or prevent* or prophylaxis* or	4758691
450	prophylactic*).ti,ab.	0000
156	Coitus/	6880

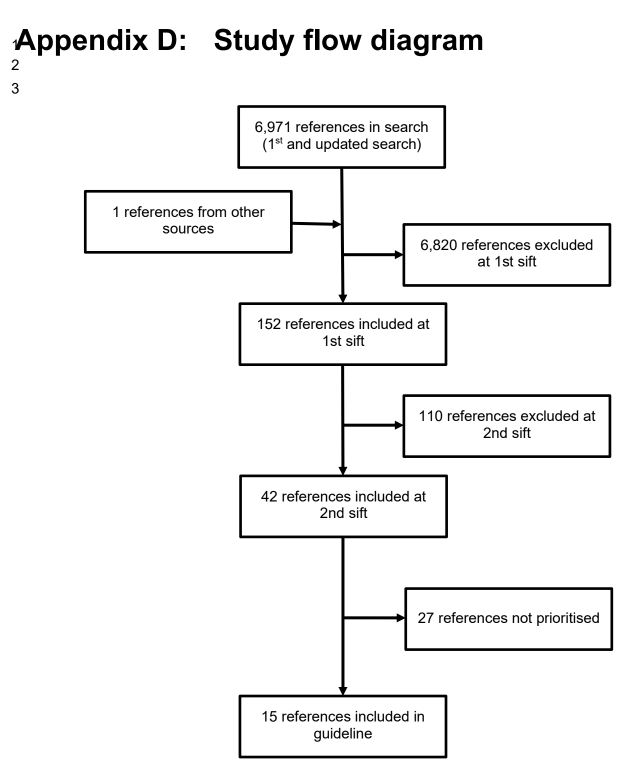
156 Coitus/

157 Inappropriate prescribing/	1695
158 or/155-157	4764914
159 154 and 158	221871
160 151 or 159	292655
161 20 and 160	15345
162 Self Care/ or self medication/	32883
163 ((self or selves or themsel*) adj4 (care or manag*)).ti,ab.	33223
164 Secondary Prevention/	17180
165 Hygiene/	14900
166 Baths/	4966
167 Soaps/	2343
((postcoital* or postcoitus* or postsex* or postintercourse* or post coital* or post coitus* or post	
sex* or post intercourse* or postmicturit* or micturit* or postmicturat* or micturat* or urinat* or	
168 defecat* or toilet* or lavatory or lavatories or perineal* or perineum*) adj3 (prophylaxis* or	1611
prophylactic* or treatment* or wipe* or wiping or hygiene* or hygienic* or clean* or douche* or	
douching* or bath* or soap* or wash* or shower*)).ti,ab.	
169 (second* adj3 prevent*).ti,ab.	21506
170 or/162-169	112930
171 20 and 170	1919
172 or/8-10	29047
173 Device Removal/	10427
174 172 and 173	753
(Catheter* adj3 (care* or removal* or removing* or remove* or "take* out" or "taking out" or 175 change* or changing* or clean* or wash* or bath* or hygiene* or hygienic*)).ti,ab.	10138
176 174 or 175	10561
177 20 and 176	5423
178 85 or 94 or 95 or 112 or 120 or 128 or 134 or 140 or 161 or 171 or 177	65619
179 limit 178 to yr="2006 -Current"	21429
180 limit 179 to english language	19392
181 Animals/ not (Animals/ and Humans/)	4291504
182 180 not 181	15047
183 limit 182 to (letter or historical article or comment or editorial or news)	784
184 182 not 183	14263
185 Meta-Analysis.pt.	74747

186 Meta-Analysis as Topic/	15461
187 Network Meta-Analysis/	34
188 Review.pt.	2230816
189 exp Review Literature as Topic/	9193
190 (metaanaly* or metanaly* or (meta adj3 analy*)).ti,ab.	109466
191 (review* or overview*).ti.	389897
192 (systematic* adj5 (review* or overview*)).ti,ab.	109630
193 ((quantitative* or qualitative*) adj5 (review* or overview*)).ti,ab.	7343
194 ((studies or trial*) adj2 (review* or overview*)).ti,ab.	36022
195 (integrat* adj3 (research or review* or literature)).ti,ab.	8769
196 (pool* adj2 (analy* or data)).ti,ab.	22123
197 (handsearch* or (hand adj3 search*)).ti,ab.	7550
198 (manual* adj3 search*).ti,ab.	4715
199 or/185-198	2487695
200 184 and 199	2428
201 Randomized Controlled Trial.pt.	448607
202 Controlled Clinical Trial.pt.	91938
203 Clinical Trial.pt.	508233
204 exp Clinical Trials as Topic/	304614
205 Placebos/	34193
206 Random Allocation/	89847
207 Double-Blind Method/	143336
208 Single-Blind Method/	23779
209 Cross-Over Studies/	40867
210 ((random* or control* or clinical*) adj3 (trial* or stud*)).ti,ab.	1003782
211 (random* adj3 allocat*).ti,ab.	28603
212 placebo*.ti,ab.	189958
213 ((singl* or doubl* or trebl* or tripl*) adj (blind* or mask*)).ti,ab.	153095
214 (crossover* or (cross adj over*)).ti,ab.	74298
215 or/201-214	1721840
216 184 and 215	2933
217 216 not 200	2230
218 Observational Studies as Topic/	1959
219 Observational Study/	31517
60	

220 Epidemiologic Studies/	7369
221 exp Case-Control Studies/	834068
222 exp Cohort Studies/	1623327
223 Cross-Sectional Studies/	234990
224 Controlled Before-After Studies/	218
225 Historically Controlled Study/	97
226 Interrupted Time Series Analysis/	243
227 Comparative Study.pt.	1770190
228 case control*.ti,ab.	102767
229 case series.ti,ab.	52479
230 (cohort adj (study or studies)).ti,ab.	133481
231 cohort analy*.ti,ab.	5462
232 (follow up adj (study or studies)).ti,ab.	43245
233 (observational adj (study or studies)).ti,ab.	70390
234 longitudinal.ti,ab.	186074
235 prospective.ti,ab.	454707
236 retrospective.ti,ab.	381342
237 cross sectional.ti,ab.	245513
238 or/218-237	3929955
239 184 and 238	5469
240 239 not (200 or 216)	3795
241 184 not (200 or 216 or 240)	5810
242 exp Drug Resistance, Bacterial/	72249
243 exp Drug Resistance, Multiple/	28752
244 ((bacter* or antibacter* or anti-bacter* or "anti bacter*") adj4 (resist* or tolera*)).ti,ab.	34156
245 ((antibiot* or anti-biot* or "anti biot*") adj4 (resist* or tolera*)).ti,ab.	42316
246 (multi* adj4 drug* adj4 (resist* or tolera*)).ti,ab.	12134
247 (multidrug* adj4 (resist* or tolera*)).ti,ab.	38335
248 (multiresist* or multi-resist* or "multi resist*").ti,ab.	6214
249 ((microb* or antimicrob* or anti-microb* or "anti microb*") adj4 (resist* or tolera*)).ti,ab.	22368
250 (superbug* or super-bug* or "super bug*").ti,ab.	448
251 Superinfection/	1644
(superinvasion* or super-invasion* or "super invasion*" or superinfection* or super-infection* or 252 "super infection*").ti,ab.	5185

253 R Factors/	4157
254 "r factor*".ti,ab.	3648
255 (resist* factor* or "r plasmid*" or resist* plasmid*).ti,ab.	5218
256 or/242-255	180317
257 84 and 256	48201
258 limit 257 to yr="2006 -Current"	25203
259 limit 258 to english language	23256
260 259 not 181	20939
261 limit 260 to (letter or historical article or comment or editorial or news)	867
262 260 not 261	20072



Appendix E: Evidence prioritisation

Key questions	Included	studies ¹	Studies no	t prioritised ²
	Systematic reviews	RCTs	Systematic reviews	RCTs
Which non-pharmacological intervention	ons are effective?			
Lactobacillus	<u>Grin et al. 2013</u> Schwenger et al. 2015	-	Beerepoot et al. 2013	Stapleton et al. 2011
D-Mannose	Kranjcec et al. 2014	-	-	Porru et al. 2014
Cranberry products	<u>Fu et al. 2017</u> <u>Jepson et al. 2012</u> <u>Roshdibonab et al. 2017</u>	Beerepoot et al. 2011 Uberos et al. 2012	Beerepoot et al. 2013 Luis et al. 2017 Wang et al. 2012	Afshar et al. 2012 Bailey et al. 2007 Barbosa-Cesnik et al. 2011 Bianco et al. 2012 Bosmans et al. 2014 Caljouw et al. 2014 Ferrara et al. 2009 Ledda et al. 2015 Maki et al. 2016 McMurdo et al. 2010 Salo et al. 2012 Sengupta et al. 2011 Singh et al. 2016 Stapleton et al. 2012 Takahashi et al. 2013 van den Hout et al. 2014
Which non-antimicrobial pharmacologi	ical interventions are effective	?		
Oestrogens	Perrotta et al. 2008	-	Beerepoot et al. 2013	-
Is antibiotic prophylaxis effective?				
Antibiotic prophylaxis versus placebo	<u>Albert et al. 2004</u> <u>Muller et al. 2017</u> <u>Williams and Craig 2011</u>	-	Mathew 2010 Mori et al. 2009 Price et al. 2016	Norinder et al. 2006

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Key questions	Included studies ¹		Studies not prioritised ²	
	Systematic reviews	RCTs	Systematic reviews	RCTs
	Schneeberger et al. 2012			
Which antibiotic prophylaxis is most effe	ective?			
Antibiotic prophylaxis versus different antibiotic prophylaxis	<u>Dai et al. 2010</u> <u>Williams and Craig 2011</u>	-	Albert et al. 2004	Antachopoulos et al. 2016
What is the optimal dosage, duration and route of administration of antibiotic prophylaxis?				
Dosage	-	-	-	-
Course length	Albert et al. 2004	Zhong et al. 2011	-	-
Route of administration	-	-	-	-
¹ See <u>appendix F</u> for full references of included studies ² See <u>appendix I</u> for full references of not-prioritised studies, with reasons for not prioritising these studies				

² Appendix F: Included studies

Albert X, Huertas I, Pereiro II, Sanfelix J, Gosalbes V, and Perrota C (2004) Antibiotics for
 preventing recurrent urinary tract infection in non-pregnant women. The Cochrane database
 of systematic reviews (3), CD001209

- 6 Beerepoot Marielle A. J, ter Riet, Gerben, Nys Sita, van der Wal, Willem M, de Borgie,
- 7 Corianne A J. M, de Reijke, Theo M, Prins Jan M, Koeijers Jeanne, Verbon Annelies,
- 8 Stobberingh Ellen, and Geerlings Suzanne E (2011) Cranberries vs antibiotics to prevent
- 9 urinary tract infections: a randomized double-blind noninferiority trial in premenopausal
- 10 women. Archives of internal medicine 171(14), 1270-8
- Dai B, Liu Y, Jia J, and Mei C (2010) Long-term antibiotics for the prevention of recurrent
 urinary tract infection in children: a systematic review and meta-analysis. Archives of disease
 in childhood 95(7), 499-508
- Fu Z, Liska D, Talan D., Chung M. Cranberry Reduces the Risk of Urinary Tract Infection
 Recurrence in Otherwise Healthy Women: A Systematic Review and Meta-Analysis. Journal
 of Nutrition. 2017; 147(12):2282-2288
- 17 Grin Peter M, Kowalewska Paulina M, Alhazzan Waleed, and Fox-Robichaud Alison E
- (2013) Lactobacillus for preventing recurrent urinary tract infections in women: meta analysis. The Canadian journal of urology 20(1), 6607-14
- Jepson RG, Williams G, and Craig JC (2012) Cranberries for preventing urinary tract
 infections. The Cochrane database of systematic reviews 10, CD001321
- Kranjcec Bojana, Papes Dino, and Altarac Silvio (2014) D-mannose powder for prophylaxis
 of recurrent urinary tract infections in women: a randomized clinical trial. World journal of
 urology 32(1), 79-84
- Muller A E, Verhaegh E M, Harbarth S, Mouton J W, and Huttner A (2017) Nitrofurantoin's
 efficacy and safety as prophylaxis for urinary tract infections: a systematic review of the
 literature and meta-analysis of controlled trials. Clinical microbiology and infection: the official
- 28 publication of the European Society of Clinical Microbiology and Infectious Diseases,
- Perrotta C, Aznar M, Mejia R, Albert X, and Ng C W (2008) Oestrogens for preventing
 recurrent urinary tract infection in postmenopausal women. The Cochrane database of
 systematic reviews (2), CD005131
- 32 Roshdibonab F, Mohammadbager FazlJoo S, Torbati M, Mohammadi Gh, Asadloo M,
- 33 Noshad H. The Role of Cranberry in Preventing Urinary Tract Infection in Children; a
- 34 Systematic Review and Meta-Analysis. Int J Pediatr 2017; 5(12): 6457-68. DOI:
- 35 10.22038/ijp.2017.27041.2327
- Schneeberger Caroline, Geerlings Suzanne E, Middleton Philippa, and Crowther Caroline A
 (2015) Interventions for preventing recurrent urinary tract infection during pregnancy. The
 Cochrane database of systematic reviews 11, CD009279
- 39 Schwenger Erin M, Tejani Aaron M, and Loewen Peter S (2015) Probiotics for preventing
- 40 urinary tract infections in adults and children. The Cochrane database of systematic reviews
 41 (12), CD008772
- 42 Uberos J, Nogueras-Ocana M, Fernandez-Puentes V, Rodriguez-Belmonte R, Narbona-
- 43 Lopez E, Molina-Carballo A, and Munoz-Hoyos A (2012) Cranberry syrup vs trimethoprim in

- the prophylaxis of recurrent urinary tract infections among children: A controlled trial. Open
 Access Journal of Clinical Trials 4, 31-38
- Williams G, and Craig JC (2011) Long-term antibiotics for preventing recurrent urinary tract
 infection in children. The Cochrane database of systematic reviews (3), CD001534

5 Zhong Y H, Fang Y, Zhou J Z, Tang Y, Gong S M, and Ding X Q (2011) Effectiveness and

6 safety of patient initiated single-dose versus continuous low-dose antibiotic prophylaxis for

7 recurrent urinary tract infections in postmenopausal women: a randomized controlled study.

8 The Journal of international medical research 39(6), 2335-43

Appendix G: Quality assessment of included studies

G.1 Lactobacillus

3 Table 4: Overall risk of bias/quality assessment – systematic reviews (<u>SR checklist</u>)

Study reference	Grin et al. 2013
Did the review address a clearly focused question?	Yes
Did the authors look for the right type of papers?	Yes
Do you think all the important, relevant studies were included?	Yes
Did the review's authors do enough to assess the quality of the included studies?	Yes
If the results of the review have been combined, was it reasonable to do so?	Yes
What are the overall results of the review?	See GRADE profiles
How precise are the results?	See GRADE profiles
Can the results be applied to the local population?	Yes – lactobacillus preparations are available in the UK
Were all important outcomes considered?	No – only a single outcome was reported
Are the benefits worth the harms and costs?	See GRADE profiles

4 Table 5: Overall risk of bias/quality assessment – systematic reviews (SR checklist)

Study reference	Schwenger et al. 2015
Did the review address a clearly focused question?	Yes
Did the authors look for the right type of papers?	Yes
Do you think all the important, relevant studies were included?	Yes
Did the review's authors do enough to assess the quality of the included studies?	Yes
If the results of the review have been combined, was it reasonable to do so?	Yes
What are the overall results of the review?	See GRADE profiles
How precise are the results?	See GRADE profiles
Can the results be applied to the local population?	Yes
Were all important outcomes considered?	No - only a single outcome was reported
Are the benefits worth the harms and costs?	See GRADE profiles

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G.2 D-Mannose

2 Table 6: Overall risk of bias/quality assessment – randomised controlled trials (RCT checklist)

Study reference	Kranjcec et al. 2014
Did the trial address a clearly focused issue?	Yes
Was the assignment of patients to treatments randomised?	Yes
Were patients, health workers and study personnel blinded?	Unclear ^a
Were the groups similar at the start of the trial?	Yes
Aside from the experimental intervention, were the groups treated equally?	Yes
Were all of the patients who entered the trial properly accounted for at its conclusion?	Yes
How large was the treatment effect?	See GRADE profiles
How precise was the estimate of the treatment effect?	See GRADE profiles
Can the results be applied in your context? (or to the local population)	Yes
^a Not specified	

G.3 Cranberry products

4 Table 7: Overall risk of bias/quality assessment – systematic reviews (SR checklist)

Study reference	Jepson et al. 2012
Did the review address a clearly focused question?	Yes
Did the authors look for the right type of papers?	Yes
Do you think all the important, relevant studies were included?	Yes
Did the review's authors do enough to assess the quality of the included studies?	Yes
If the results of the review have been combined, was it reasonable to do so?	Yes
What are the overall results of the review?	See GRADE profiles
How precise are the results?	See GRADE profiles
Can the results be applied to the local population?	Yes
Were all important outcomes considered?	Yes
Are the benefits worth the harms and costs?	See GRADE profiles

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2 Table 8: Overall risk of bias/quality assessment – systematic reviews (<u>SR checklist</u>)

Study reference	Fu et al. 2018
Did the review address a clearly focused question?	Yes
Did the authors look for the right type of papers?	Yes
Do you think all the important, relevant studies were included?	Yes
Did the review's authors do enough to assess the quality of the included studies?	Yes
If the results of the review have been combined, was it reasonable to do so?	Yes
What are the overall results of the review?	See GRADE profiles
How precise are the results?	See GRADE profiles
Can the results be applied to the local population?	Yes
Were all important outcomes considered?	Yes
Are the benefits worth the harms and costs?	See GRADE profiles

3 Table 9: Overall risk of bias/quality assessment – systematic reviews (SR checklist)

Study reference	Roshidibonab et al. 2018
Did the review address a clearly focused question?	Yes
Did the authors look for the right type of papers?	Yes
Do you think all the important, relevant studies were included?	Yes
Did the review's authors do enough to assess the quality of the included studies?	Yes
If the results of the review have been combined, was it reasonable to do so?	Yes
What are the overall results of the review?	See GRADE profiles
How precise are the results?	See GRADE profiles
Can the results be applied to the local population?	Yes
Were all important outcomes considered?	Yes
Are the benefits worth the harms and costs?	See GRADE profiles

⁴

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1 Table 10: Overall risk of bias/quality assessment – randomised controlled trials (<u>RCT checklist</u>)

Beerepoot et al. 2011	Uberos et al. 2012	
Yes	Yes	
See	See GRADE profiles	
See GRADE profiles		
Yes	Yes ^a	
	2011YesYesYesYesYesYesYesSeeSee	

^a Patient population included children with vesicoureteral reflux (VUR)

G.4 Oestrogens

3 Table 11: Overall risk of bias/quality assessment – systematic reviews (SR checklist)

Study reference	Perrotta et al. 2008
Did the review address a clearly focused question?	Yes
Did the authors look for the right type of papers?	Yes
Do you think all the important, relevant studies were included?	Yes
Did the review's authors do enough to assess the quality of the included studies?	Yes
If the results of the review have been combined, was it reasonable to do so?	Yes
What are the overall results of the review?	See GRADE profiles
How precise are the results?	See GRADE profiles
Can the results be applied to the local population?	Yes
Were all important outcomes considered?	Yes
Are the benefits worth the harms and costs?	See GRADE profiles

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G.5 Antimicrobials in non-pregnant women

2 Table 12: Overall risk of bias/quality assessment – systematic reviews (<u>SR checklist</u>)

Study reference	Albert et al. 2004	Muller et al. 2017
Did the review address a clearly focused question?	Yes	Yes
Did the authors look for the right type of papers?	Yes	Yes
Do you think all the important, relevant studies were included?	No	No
Did the review's authors do enough to assess the quality of the included studies?	Yes	Yes
If the results of the review have been combined, was it reasonable to do so?	Yes ^a	Yes ^a
What are the overall results of the review?	See GRADE profiles	
How precise are the results?	See GRADE profiles	
Can the results be applied to the local population?	Unclear ^b	Unclear ^b
Were all important outcomes considered?	Yes °	Yes ^c
Are the benefits worth the harms and costs?	See GRADE profiles	
^a 9 studies could not be pooled in a meta-analysis due to uncommon features in the individual studies ^b Not all the antibiotics reviewed are available for use in the UK		
° The review planned to assess a number of outcomes, but there was no evidence available for all outcomes		

3 Table 13: Overall risk of bias/quality assessment – randomised controlled trials (<u>RCT checklist</u>)

Study reference	Zhong et al. 2011 ^a	
Did the trial address a clearly focused issue?	Yes	
Was the assignment of patients to treatments randomised?	Yes	
Were patients, health workers and study personnel blinded?	Unclear	
Were the groups similar at the start of the trial?	Yes	
Aside from the experimental intervention, were the groups treated equally?	Yes	
Were all of the patients who entered the trial properly accounted for at its conclusion?	Yes	
How large was the treatment effect?	See GRADE profiles	
How precise was the estimate of the treatment effect?	See GRADE profiles	
Can the results be applied in your context? (or to the local population)	Yes	
^a Summary statistics, risk ratio, and 95% confidence interval (CI) not reported; calculated by NIC	E	

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G.6 Antimicrobials in pregnant women

2 Table 14: Overall risk of bias/quality assessment – systematic reviews (SR checklist)

Study reference	Schneeberger et al. 2015
Did the review address a clearly focused question?	Yes
Did the authors look for the right type of papers?	Yes
Do you think all the important, relevant studies were included?	Yes
Did the review's authors do enough to assess the quality of the included studies?	Yes
If the results of the review have been combined, was it reasonable to do so?	N/A
What are the overall results of the review?	See GRADE profiles
How precise are the results?	See GRADE profiles
Can the results be applied to the local population?	Yes
Were all important outcomes considered?	Yes
Are the benefits worth the harms and costs?	See GRADE profiles

G.7 Antimicrobials in a mixed population of adults and children

4 Table 15: Overall risk of bias/quality assessment – systematic reviews (SR checklist)

Study reference	Muller et al. 2017
Did the review address a clearly focused question?	Yes
Did the authors look for the right type of papers?	Yes
Do you think all the important, relevant studies were included?	Unclear ^a
Did the review's authors do enough to assess the quality of the included studies?	Yes
If the results of the review have been combined, was it reasonable to do so?	Yes
What are the overall results of the review?	See GRADE profiles
How precise are the results?	See GRADE profiles
Can the results be applied to the local population?	Yes
Were all important outcomes considered?	No ^b
Are the benefits worth the harms and costs?	See GRADE profiles

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Study reference

Muller et al. 2017

^a Studies were included if they were controlled trials, evaluating oral doses of nitrofurantoin. The majority of studies included were randomised (81%), with a small proportion double-blinded (27%).

^b The study did not report all the secondary outcomes they planned *a priori*.

G.8 Antimicrobials in children

2 Table 16: Overall risk of bias/quality assessment – systematic reviews (SR checklist)

Study reference	Dai et al. 2010	Williams and Craig 2011
Did the review address a clearly focused question?	Yes	Yes
Did the authors look for the right type of papers?	Yes	Yes
Do you think all the important, relevant studies were included?	Yes	Yes
Did the review's authors do enough to assess the quality of the included studies?	Yes	Yes
If the results of the review have been combined, was it reasonable to do so?	Yes	Yes
What are the overall results of the review?	See GRA	DE profiles
How precise are the results?	See GRA	DE profiles
Can the results be applied to the local population?	Unclearª	Unclear ^a
Were all important outcomes considered?	Yes	No ^b
Are the benefits worth the harms and costs?	See GRA	DE profiles

^a Most studies did not report a clear inclusion and exclusion criteria for participants entry into the study; it was possible for patients to be misclassified, there was also significant heterogeneity despite the use of a random effects model.

^b Not all planned outcomes were reported; and in some studies 'repeat positive urine culture' was reported instead of the recurrence of urinary tract infection

3

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Appendix H: GRADE profiles

Lactobacillus **H.**2

Table 17: GRADE profile – lactobacillus versus placebo in premenopausal women 3

			Quality asses	sment			No of pati	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Lactobacillus	Placebo	Relative (95% CI)	Absolute		
Risk of re	current urinal	ry tract infect	ion (follow-up 1-12	2 months)								
5 ¹					very serious²	none	44/147 (29.9%)	51/147 (34.7%)		52 fewer per 1000 (from 146 fewer to 87 more)	⊕⊕OO LOW	CRITICAL
Risk of re	current urina	ry tract infect	ion - sensitivity ar	alysis of only el	fective strain	ns of lactobacillus	³ (follow-up 1	-12 mont	ths)			
2 ¹				no serious indirectness	serious ⁴	none	10/62 (16.1%)	21/65 (32.3%)	RR 0.51 (0.26 to 0.99)		⊕⊕⊕O MODERATE	CRITICAL
Abbreviati	ons: CI, confid	ence interval;	RR, risk ratio									

¹ Grin et al. 2013

4567

² Downgraded 1 level – at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable harm or appreciable benefit
 ³ Effective strains of lactobacillus as defined by study authors
 ⁴ Downgraded 1 level – at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with placebo

8 Table 18: GRADE profile – lactobacillus versus antibiotics in non-pregnant women

			Quality asse	ssment			No of p	atients	E	ffect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Lactobacillus	Co- trimoxazole	Relative (95% Cl)	Absolute		
Symptom	atic bacteria	urinary tra	ct infection									
	randomised trials	serious ²	N/A	no serious indirectness	serious ³	none	86/115 (74.8%)	72/108 (66.7%)	RR 1.12 (0.95 to 1.33)	80 more per 1000 (from 33 fewer to 220 more)	⊕⊕OO LOW	CRITICAL
Symptom	atic bacteria	urinary tra	ct infection - wors	st case scenario	probiotics							
		no serious risk of bias	N/A	no serious indirectness	serious ³	none	91/115 (79.1%)	72/108 (66.7%)		127 more per 1000 (from 7 more to 267 more)	⊕⊕⊕O MODERATE	CRITICAL
Symptom	atic bacteria	urinary tra	ct infection - wors	st case scenario	antibiotics		-	•				•

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			Quality asse	ssment			No of p	atients	E	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Lactobacillus	Co- trimoxazole	Relative (95% CI)	Absolute		
1 ¹		no serious risk of bias	N/A	no serious indirectness	serious ⁴	none	86/115 (74.8%)	97/108 (89.8%)	RR 0.83 (0.74 to 0.94)	153 fewer per 1000 (from 234 fewer to 54 fewer)	⊕⊕⊕O MODERATE	CRITICAL
No. of pe	ople experier	ncing at leas	t 1 adverse even	t								
1 ¹	randomised trials		no serious inconsistency	no serious indirectness	serious ⁴	none	66/125 (52.8%)	74/127 (58.3%)	NICE analysis: RR 0.91 (0.73 to 1.13) ⁵	52 fewer per 1000 (from 157 fewer to 76 more)	⊕⊕OO LOW	CRITICAL
Number o	of adverse ev	ents	•	•	•	•					•	
1 ¹	randomised trials		no serious inconsistency	no serious indirectness	serious ⁴	none	7/125 (5.6%)	15/127 (11.8%)	NICE analysis: RR 0.47 (0.2 to 1.12) ⁵	63 fewer per 1000 (from 94 fewer to 14 more)	⊕⊕OO LOW	CRITICAL
Abbreviati	ons: N/A, not	applicable; C	CI, confidence inter	val; RR, risk ratio	2							

¹ Schwenger et al. 2015 (NAPRUTI Study II 2006)

² Downgraded 1 level - high risk of attrition bias

³ Downgraded 1 level – at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with lactobacillus
 ⁴ Downgraded 1 level – at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with co-trimoxazole
 ⁵ RR and 95% CI not reported, calculated by NICE assuming an intention-to-treat analysis was done

H.2 D-mannose in non-pregnant women

Table 19: GRADE profile – D-mannose versus no treatment 7

			Quality as	sessment			No of p	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	D-mannose	No treatment	Relative (95% CI)	Absolute		
Participar	nts with recuri	ent urinary tr	act infection			•						
1 ¹		no serious risk of bias	-		no serious imprecision	none	15/103 (14.6%)	62/102 (60.8%)	RR 0.24 (0.15 to 0.39) ²	462 fewer per 1000 (from 517 fewer to 371 fewer)		CRITICAL
Abbreviatio	ons: CI, Confid	ence interval;	N/A, Not applica	able; RR, Relative	risk			-			•	

¹ Kranjcec et al. 2014

1

2345

8 9

² 95% confidence interval not stated; intervals calculated by NICE

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Table 20: GRADE profile – D-mannose versus antibiotics 1

Quality assessment										No of p	atients	Effect		Quality	Importance
N	lo of st	tudies		Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	D-mannose	Antibiotics	Relative (95% Cl)	Absolute		
articipants with r	ecurre	nt urinary tra	act infection	n											
1					no serious risk of bias	N/A		very serious ²	none	15/103 (14.6%)	21/103 (20.4%)	(0.39 to	59 fewer per 1000 (from 124 fewer to 63 more)		CRITICAL
dverse events				r					I						
¹ randomised no trials serio risk c bias	us	no serious indirectness	no serious imprecision			8/103 (7.8%)		29/103 28.2%)	RR 0.28 (0.13 0.57) ³		fewer per 10 245 fewer to fewer)		⊕⊕⊕ HIGH	C	CRITICAL

¹ Kranjcec et al. 2014 ² Downgraded 2 levels - at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm ³ 95% confidence interval not stated; calculated by NICE

2 3 4

Cranberry products H.\$

Table 21: GRADE profile - cranberry products versus placebo or no treatment in women 6

			Quality asse	essment			No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cranberry products	Placebo or no treatment	Relative (95% Cl)	Absolute		
Participar	nts with one o	or more UTI a	t follow-up									
	randomised trials	no serious risk of bias		no serious indirectness	very serious ³	none	64/322 (19.9%)	62/272 (22.8%)		59 fewer per 1000 (from 132 fewer to 71 more)		CRITICAL
Abbreviatio	ons: UTI, urina	ary tract infect	on; CI, Confider	nce interval; N/A,	Not applicable	e; RR, Relative risk		•	•	•		•

¹ Jepson et al. 2012

7 8 9

² Downgraded 1 level – heterogeneity >50% ³ Downgraded 2 levels - at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm

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1

Data from study identified following consultation

Table 22: GRADE profile – cranberry products versus placebo or no treatment in women

			Quality as	sessment	_		No o	f patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cranberry	Placebo or no treatment	Relative (95% Cl)	Absolute		
Incidence	of urinary tra	ct infectio	on (symptom o	r culture confirm	ned) (follow-u	p 6 to 12 months)						
7 ¹	randomised trials	serious ²	serious ³	no serious indirectness	serious ⁴	none	165/796 (20.7%)	186/702 (26.5%)	RR 0.74 (0.55 to 0.98)	69 fewer per 1000 (from 5 fewer to 119 fewer)	⊕000 VERY LOW	CRITICAL
Incidence	of urinary tra	ct infectio	on (culture con	firmed) (follow-u	ip 6 to 12 mo	nths)						
5 ¹	randomised trials	serious ²	serious ³	no serious indirectness	serious ⁴	none	100/504 (19.8%)	98/408 (24.0%)	RR 0.71 (0.45 to 1.12)	70 fewer per 1000 (from 132 fewer to 29 more)	⊕000 VERY LOW	CRITICAL
Abbreviati	ons: Cl, confide	ence interv	al; RR, relative	risk								
Ū						t with no meaningfu bo or no treat				0		
			Quality as	sessment				f patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Quality as: Inconsistency	sessment Indirectness	Imprecision	Other considerations			Relative (95% Cl)	Effect Absolute	Quality	Importance
studies		bias	Inconsistency	Indirectness	•		No o Cranberry	f patients Placebo or no			Quality	Importance
studies	of urinary tra	bias ct infectio	Inconsistency on (symptom o serious ³	Indirectness	•	considerations	No o Cranberry	f patients Placebo or no		Absolute 56 fewer per 1000	Quality ⊕000 VERY LOW	Importance CRITICAL
studies Incidence 6 ¹	e of urinary tra randomised trials	bias ct infection serious ²	Inconsistency on (symptom o serious ³	Indirectness r culture confirm no serious indirectness	ied) (follow-ι	considerations up 6 to 12 months)	No o Cranberry juice 146/663	f patients Placebo or no treatment 162/609	(95% CI) RR 0.79	Absolute 56 fewer per 1000 (from 109 fewer to 16	⊕000 VERY	

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			Quality ass	essment			No of patie	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cranberry tablets or capsules	Placebo	Relative (95% CI)	Absolute		p
ncidence	of urinary tr	act infecti	on (symptom or o	ulture confirmed	d) (follow-up	6 to 12 months)						
2 ¹	randomised trials	serious ²	no serious inconsistency	no serious indirectness	serious ³	none	18/133 (13.5%)	40/143 (28.0%)	RR 0.48 (0.29 to 0.79)	145 fewer per 1000 (from 199 fewer to 59 fewer)	⊕⊕OO LOW	CRITICAL

Table 22: GRADE profile – cranberry products versus antibiotics in women 1

trials serious risk of bias inconsistency indirectness indirectness (51.1%) (40.4%) 2.02) fewer to 412 more) MODERATE Development of antibiotic resistance - premenopausal women				Quality ass	essment			No of p			Effect	Quality	Importance
21 randomised trials no serious inconsistency no serious indirectness serious ² none 91/178 (51.1%) 67/166 (40.4%) RR 1.31 (0.85 to 2.02) 125 more per 1000 (from 61 fewer to 412 more)		Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cranberry products	Antibiotics	Relative (95% Cl)	Absolute		
trials serious risk of bias inconsistency indirectness indirectness (51.1%) (40.4%) 2.02) fewer to 412 more) MODERATE Development of antibiotic resistance - premenopausal women Image: Comparison of trials no serious risk of bias N/A no serious indirectness serious ⁴ none N=21 E. coli isolates from women receiving co-trimoxazole showed antibiotic resistance for amoxicillin, trimethoprim, and trimethoprim-sulfamethoxazole at 1 month prophylaxis (70% resistance). This reduced at 1 month and 3 months after stopping of prophylaxis, returning to baseline at 12 months. E. coli isolates from women receiving cranberry Image: Comparison of the top in the top i	Repeat s	ymptomatic	urinary t	tract infection									
1 ³ randomised trials no N/A no serious indirectness serious ⁴ none N=221 E. coli isolates from women receiving co-trimoxazole showed antibiotic resistance for amoxicillin, trimethoprim-sulfamethoxazole at 1 month prophylaxis (70% resistance). This reduced at 1 month and 3 months after stopping of prophylaxis, returning to baseline at 12 months. E. coli isolates from women receiving cranberry Image: CRITICA women receiving co-trimoxazole showed antibiotic resistance for amoxicillin, trimethoprim-sulfamethoxazole at 1 month prophylaxis (70% resistance). This reduced at 1 month and 3 months after stopping of prophylaxis, returning to baseline at 12 months. E. coli isolates from women receiving cranberry Image: CRITICA women receiving co-trimoxazole showed antibiotic resistance for amoxicillin, trimethoprim-sulfamethoxazole at 1 month prophylaxis (70% resistance). This reduced at 1 month and 3 months after stopping of prophylaxis, returning to baseline at 12 months. Image: CRITICA women receiving co-trimoxazole showed antibiotic resistance). This reduced at 1 month and 3 months after stopping of prophylaxis, returning to baseline at 12 months. Image: CRITICA women receiving co-trimoxazole at 12 months. <td< td=""><td></td><td>trials</td><td>serious risk of</td><td></td><td></td><td>serious²</td><td>none</td><td></td><td></td><td></td><td></td><td></td><td>CRITICAL</td></td<>		trials	serious risk of			serious ²	none						CRITICAL
trialsserious risk of biasindirectnessMODERATEvisk of biasindirectnessindirectnessMODERATEvisk of biasvisk of biasvisk of biasvisk of visk of biasMODERATEvisk of biasvisk of biasvisk of visk of biasMODERATEvisk of biasvisk of visk of biasvisk of visk of visk of biasMODERATEvisk of biasvisk of visk of visk of biasvisk of visk of 	Develop	ment of anti	biotic res	istance - preme	nopausal won	nen							
		trials	serious risk of	N/A		serious ⁴	none	N=2	221	showed antibio trimethoprim, and ti 1 month prophylaxis at 1 month and prophylaxis, retur E. coli isolates fro	tic resistance for amoxicillin, rimethoprim-sulfamethoxazole at s (70% resistance). This reduced d 3 months after stopping of rning to baseline at 12 months. om women receiving cranberry		CRITICAL

¹Jepson et al. 2012

234 5

²Downgraded 1 level - at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm ³Beerepoot et al. 2011

⁴ Downgraded 1 level – not assessable

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Table 23: GRADE profile – cranberry products versus placebo or no treatment in pregnant women 1

			Quality as	sessment			No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cranberry products	Placebo or no treatment	Relative (95% Cl)	Absolute		
Participar	nts with one	or more U	TI at follow-up									
	randomised trials	serious ²	no serious inconsistency	no serious indirectness	no serious imprecision	none	184/325 (56.6%)	194/349 (55.6%)	RR 1.04 (0.93 to 1.17)	22 more per 1000 (from 39 fewer to 94 more)	⊕⊕⊕O MODERATE	CRITICAL
Abbreviati	ons: CI, Confi	dence inte	rval; N/A, Not appl	icable; RR, Relat	ive risk	-				•		

¹Jepson et al. 2012

2 3 ² Downgraded by 1 level - high drop-out rate across the studies

Table 24: GRADE profile – cranberry products versus placebo or no treatment in elderly women and men 4

			Quality asse	essment			No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cranberry products	Placebo or no treatment	Relative (95% CI)	Absolute		
Participan	nts with one o	r more UT	l at follow-up									
	randomised trials				very serious ³	none	20/207 (9.7%)	26/206 (12.6%)	RR 0.75 (0.39 to 1.44)	32 fewer per 1000 (from 77 fewer to 56 more)	⊕OOO VERY LOW	CRITICAL
Abbreviatio	ons: CI, Confic	lence inter	val; N/A, Not applic	able; RR, Relative	e risk			•			•	

5 6 7

¹ Jepson et al. 2012

² Downgraded by 1 level - high drop-out rate across the studies ³ Downgraded 2 levels - at a 95% confidence interval, data are consistent with no meaningful difference, appreciable benefit or appreciable harm

8 Table 25: GRADE profile – cranberry products versus placebo or no treatment in adults

			Quality asses	ssment			No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cranberry products	Placebo or no treatment	o Relative (95% CI) Absolute			
Adverse e	events - any g	astrointestin	al effect					·				
4 ¹					very serious²	none	10/328 (3%)	9/269 (3.3%)		6 fewer per 1000 (from 23 fewer to 42 more)		CRITICAL
	ons: CI, Confid	dence interval	N/A, Not applicabl	e; RR, Relative ri	sk					•		

9 ¹ Jepson et al. 2012

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1 ² Downgraded 2 levels - at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm

Table 26: GRADE profile – cranberry products versus antibiotics in adults 2

			Quality asses	sment			No of pa	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cranberry products	Antibiotics	Relative (95% Cl)	Absolute		
Adverse e	vents – gastr	ointestinal	•	•			·		•	•	• • •	
_				no serious indirectness	very serious ⁱⁱ	none	17/178 (9.6%)	20/166 (12.0%)(RR 0.78 (0.42 to 1.42)	27 fewer per 1000 (from 70 fewer to 51 more)	⊕⊕OO LOW	CRITICAL
Abbreviati	ons: CI, Confid	ence interval;	N/A, Not applicable	; RR, Relative ris	k							

¹ Jepson et al. 2012

3 4 5 ² Downgraded 1 level – heterogeneity >50%
 ³ Downgraded 1 level - at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with placebo or no treatment

6 Table 27: GRADE profile – cranberry products versus placebo or no treatment in children

			Quality asse	essment			No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cranberry products	Placebo or no treatment	Relative (95% Cl)	Absolute		
Participan	ts with one o	r more UTI at	follow-up		•			•	• • •		•	
		no serious risk of bias		no serious indirectness	serious ³	none	25/153 (16.3%)	46/156 (29.5%)	RR 0.48 (0.19 to 1.22)	153 fewer per 1000 (from 239 fewer to 65 more)	⊕⊕OO LOW	CRITICAL
Abbreviatio	 ons: Cl, Confid	ence interval;	N/A, Not applic	able; RR, Relative	e risk					more)		

7 ¹Jepson et al. 2012

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1

² Downgraded 2 levels - at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm

Data from study identified following consultation Table 31: GRADE profile – cranberry products versus placebo in children Quality assessment No of patients Effect Importance Quality Relative Risk of bias Inconsistency Indirectness Imprecision Other considerations No of studies Design Cranberry Placebo (95% CI) Incidence of urinary tract infection (culture confirmed) (follow-up 2 to 12 months) randomised trials serious² serious³ serious⁴ serious⁵ n =293 n =278 OR 0.31 (0.21 to 0.46)6 CRITICAL none ⊕000 VERY LOW Abbreviations: CI, confidence interval; OR, odds ratio ¹ Roshdibonab et al. 2017 ² Downgraded 1 level - 5 of the included studies have high or unclear risk of bias in at least 1 quality assessment criteria domain, as reported by study authors ³ Downgraded 1 level - heterogeneity ≥50% ⁴ Downgraded 1 level - 2 of the included studies included people with catheters ⁵ Downgraded 1 level - not assessable

2 Table 28: GRADE profile – cranberry products versus antibiotics in children

			Quality ass	essment			No of p			Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Cranberry products	Antibiotics	Relative (95% Cl)	Absolute		
Repeat s	symptomatic	urinary t	ract infection									
1 ¹	randomised trials		no serious inconsistency		very serious²	none	8/75 18/117 (10.7%) (15.4%)		RR 0.69 (0.32 to 1.51)	48 fewer per 1000 (from 105 fewer to 78 more)	⊕⊕OO LOW	CRITICAL
Develop	ment of antil	biotic resi	stance	•								
1 ³	randomised trials		no serious inconsistency	no serious indirectness	serious ⁴	none	N=	192	were observed resistance to amo 2.7; P-value not si	tween the treatment branches in the rate of percentage of oxicillin or co-trimoxazole ($\chi 2$ = gnificant and $\chi 2$ = 0.3; P-value ificant, respectively).	⊕⊕⊕O MODERATE	CRITICAL

¹ Jepson et al. 2012

3 4

² Downgraded 2 levels - at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable harm or appreciable benefit

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1 2 ³ Uberos et al. 2012

⁴ Downgraded 1 level - not assessable

Oestrogens in post-menopausal women H.4

Table 29: GRADE profile - oral oestrogen versus placebo or no treatment 4

			Quality ass	essment			No of pa	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Oral oestrogen	Placebo	Relative (95% CI)	Absolute		
Urinary tr	act infection	at the end of	f the treatment per	iod								
4 ¹	randomised trials		no serious inconsistency	no serious indirectness	serious ²	none	157/1389 (11.3%)	147/1409 (10.4%)		8 more per 1000 (from 13 fewer to 34 more)	⊕⊕⊕O MODERATE	CRITICAL
All advers	se events	•	-	•	•	•	•		•		•	
2 ¹	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	12/51 (23.5%)	2/53 (3.8%)	RR 5.11 (1.39 to 18.76)	155 more per 1000 (from 15 more to 670 more)	⊕⊕⊕⊕ HIGH	CRITICAL
	ons: Cl, Confi et al. 2010	dence interva	I; N/A, Not applicat	le; RR, Relative r	isk	l				,		I

5 6

² Downgraded 1 level - at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with oral oestrogen

Table 30: GRADE profile – vaginal oestrogen versus placebo or no treatment 7

			Quality as	sessment			No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Vaginal oestrogen	Placebo/no treatment	Relative (95% Cl)	Absolute		
Urinary tr	act infection	at the end o	of the treatmen	t period (estrad	iol-releasing sili	cone vaginal ring	[Estring] vs	no treatment)				
1 ¹		no serious risk of bias	N/A	no serious indirectness	serious ²	none	27/53 (50.9%)	44/55 (80%)	RR 0.64 (0.47 to 0.86)	288 fewer per 1000 (from 424 fewer to 112 fewer)	⊕⊕⊕O MODERATE	CRITICAL
Urinary tr	act infection	at the end o	f the treatmen	t period (topical	ly applied intra-	aginal oestriol cr	eam vs place	bo)				
		no serious risk of bias	N/A	no serious indirectness	no serious imprecision	none	8/50 (16%)	27/43 (62.8%)	RR 0.25 (0.13 to 0.5)	471 fewer per 1000 (from 546 fewer to 314 fewer)	⊕⊕⊕⊕ HIGH	CRITICAL
Any adve	rse event				<u>.</u>			•				
		no serious risk of bias	serious ³	no serious indirectness	serious ²	none	24/103 (23.3%)	5/98 (5.1%)	RR 4.57(1.81 to 11.5)	190 more per 1000 (from 17 fewer to 1000 more)	⊕⊕OO LOW	CRITICAL
Abbreviati	ons: CI, Confi	dence interva	al; N/A, Not app	licable; RR, Rela	ative risk							

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¹ Perrotta et al. 2010

1 2 3

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² Downgraded 1 level - at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with no treatment ³ Downgraded 1 level - heterogeneity > 50%

Table 31: GRADE profile – vaginal oestrogen versus oral antibiotics 4

			Quality ass	essment			No of p	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Vaginal oestrogen	Oral antibiotics	Relative (95% Cl)	Absolute		
Urinary tr	act infection	at the end o	f the treatment pe	eriod (oestriol-c	ontaining vagin	al pessary vs ora	antibiotics)					
	randomised trials	serious ²		no serious indirectness	serious ³	none	58/86 (67.4%)	44/85 (51.8%)	RR 1.3 (1.01 to 1.68)	155 more per 1000 (from 5 more to 352 more)	⊕⊕OO LOW	CRITICAL
Urinary tr	act infection	at the end o	f the treatment pe	eriod (Vaginal o	estrogens [intra	vaginal premarin	cream] vs or	al antibiotics)			
	randomised trials	very serious ⁴		no serious indirectness	no serious imprecision	none	2/27 (7.4%)	12/15 (80%)	RR 0.09 (0.02 to 0.36)	728 fewer per 1000 (from 784 fewer to 512 fewer)	⊕⊕OO LOW	CRITICAL
Urinary tr	act infection	2 months af	ter treatment				•	•				
	randomised trials	very serious ⁴	N/A	no serious indirectness	very serious ⁵	none	2/27 (7.4%)	2/15 (13.3%)	RR 0.56 (0.09 to 3.55)	59 fewer per 1000 (from 121 fewer to 340 more)	⊕OOO VERY LOW	CRITICAL
Adverse e	events											
	randomised trials	no serious risk of bias		no serious indirectness	serious ⁶	none	19/116 (16.4%)	0/100 (0%)	RR 12.86 (1.75 to 94.29)	-	⊕⊕⊕O MODERATE	CRITICAL
Abbreviati	ons: CI, Confi	dence interva	al; N/A, Not applica	ble; RR, Relative	e risk		•	•				

¹ Perrotta et al. 2010

² Downgraded 1 level - large drop-out rate (29%)
 ³ Downgraded 1 level - at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with vaginal oestrogen

⁴ Downgrade 2 levels - small study, 2:1 randomisation, relative short treatment duration compared to other studies (3 months), unclear why antibiotic treatment would result in 80% recurrent UTI

⁵ Downgraded 2 levels - at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm

⁶ Downgraded 1 level - very wide CI interval

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Antimicrobials in non-pregnant women H.5

2 Table 32: GRADE profile – antibiotics versus placebo

			Quality asso	essment			No of p	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Antibiotic	Placebo	Relative (95% Cl)	Absolute		
Patients w	ith at least or	ne microbiolo	gical recurrence d	luring prophylaxi	s			•				•
10 ¹	randomised trials				no serious imprecision	none	24/195 (12.3%)	116/177 (65.5%)		518 fewer per 1000 (from 570 fewer to 433 fewer)	⊕⊕⊕⊕ HIGH	CRITICAL
Patients w	ith at least or	ne clinical rec	urrence during pr	ophylaxis								
7 ¹	randomised trials			no serious indirectness	no serious imprecision	none	10/136 (7.4%)	62/121 (51.2%)	RR 0.15 (0.08 to 0.28)	436 fewer per 1000 (from 471 fewer to 369 fewer)	⊕⊕⊕⊕ HIGH	CRITICAL
Patients w	ith at least or	ne microbiolo	gical recurrence a	fter prophylaxis	•	•	•		•			
2 ¹	randomised trials	no serious risk of bias	serious ²	no serious indirectness	very serious ³	none	23/44 (52.3%)	15/26 (57.7%)	RR 0.82 (0.44 to 1.53)	104 fewer per 1000 (from 323 fewer to 306 more)	⊕OOO VERY LOW	CRITICAL
Severe sid	le effects	•	Ρ	1	1			,	I			
10 ¹	randomised trials	no serious risk of bias		no serious indirectness	very serious ³	none	9/225 (4%)	4/195 (2.1%)	RR 1.58 (0.47 to 5.28)	12 more per 1000 (from 11 fewer to 88 more)	⊕⊕OO LOW	CRITICAL
Other side	effects (non	serious side	effects)	•	•	•	•		•			
10 ¹	randomised trials	no serious risk of bias	serious ²	no serious indirectness	serious ⁴	none	34/225 (15.1%)	15/195 (7.7%)	RR 1.78 (1.06 to 3.00)	60 more per 1000 (from 5 more to 154 more)	⊕⊕OO LOW	CRITICAL
Abbreviatio	ons: CI, Confid	ence interval;	RR, Relative risk				•	•	•			•

¹ Albert et al. 2004

34 56

² Downgraded 1 level – heterogeneity > 50%
 ³ Downgraded 2 levels – at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm
 ⁴ Downgraded 1 level – at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with antibiotics

Table 33: GRADE profile – single-dose versus continuous antibiotic prophylaxis in postmenopausal women 7

			Quality ass	essment			No of patie	ents	Ef	ffect	Quality	Importance		
No of studies	studies Design bias inconsistency indirectness imprecision conside						Intermittent patient- initiated single dose antibiotics	Continuous antibiotics	Relative (95% CI)	Absolute				
Patients														
	atients with at least 1 recurrent urinary tract infection randomised no serious N/A no serious serious ² none trials risk of bias						25/31 (80.6%) ³	26/37 (70.3%) ³	No summary statistic reported	105 more per 1000 (from 91	⊕⊕⊕O MODERATE	CRITICAL		

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			Quality ass	essment			No of patie	ents	E	ffect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Intermittent patient- initiated single dose antibiotics	Continuous antibiotics	Relative (95% Cl)	Absolute		
									NICE analysis RR 1.15 (0.87 to 1.51) ⁴	fewer to 358 more)		
Any adve	erse events											
1 ¹	trials	no serious risk of bias	N/A	no serious indirectness	serious ⁵	none	21/33 (63.6%) ³	37/40 (92.5%) ³	No summary statistic reported NICE analysis RR 0.69 (0.52 to 0.9) ⁴	287 fewer per 1000 (from 444 fewer to 93 fewer)	⊕⊕⊕O MODERATE	CRITICAL

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2345

¹ Zhong et al. 2011 ² Downgraded 1 level - at a minimal important difference of 25%, data are consistent with no meaningful difference appreciable harm with intermittent patient-initiated single dose antibiotics ³ Summary statistics not stated; calculated by NICE

⁴ Risk ratio and 95% confidence interval not stated; calculated by NICE
 ⁵ Downgraded 1 level - at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with continuous antibiotics

Table 34: GRADE profile – single-dose versus continuous antibiotic prophylaxis in pre-menopausal women 6

			Quality asso	essment			No of p	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Post coital ciprofloxacin	Continuous ciprofloxacin	Relative (95% CI)	Absolute		
Patients v	vith at least o	ne microbio	logical recurre	nce during prop	hylaxis							
1 ¹		no serious risk of bias		no serious indirectness	very serious²	none	2/70 (2.9%)	2/65 (3.1%)	RR 0.93 (0.13 to 6.4)	2 fewer per 1000 (from 27 fewer to 166 more)	⊕⊕OO LOW	CRITICAL
Patients v	vith at least o	ne clinical r	ecurrence duri	ng prophylaxis					•			
1 ¹		no serious risk of bias		no serious indirectness	very serious²	none	4/70 (5.7%)	3/65 (4.6%)	RR 1.24 (0.29 to 5.32)	11 more per 1000 (from 33 fewer to 199 more)	⊕⊕OO LOW	CRITICAL
Other sid	e effects (non	-serious sid	le effects)						•	· · · · · · ·		
1 ¹		no serious risk of bias		no serious indirectness	very serious²	none	4/70 (5.7%)	9/65 (13.8%)	RR 0.41 (0.13 to 1.28)	82 fewer per 1000 (from 120 fewer to 39 more)	⊕⊕OO LOW	CRITICAL
Patients v	vith at least o	ne microbio	logical recurre	nce after proph	ylaxis			•	· · · ·	••		

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			Quality asso	essment			No of p	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Post coital ciprofloxacin	Continuous ciprofloxacin	Relative (95% Cl)	Absolute		
		no serious risk of bias			very serious²	none	25/70 (35.7%)	21/65 (32.3%)	RR 1.11 (0.69 to 1.77)	36 more per 1000 (from 100 fewer to 249 more)		CRITICAL

Abbreviations: CI, Confidence interval; N/A, Not applicable; RR, Relative risk

¹ Albert et al. 2004 (Melekos et al. 1998) ² Downgraded 2 levels - at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm 1 2

H.6 Antimicrobials in pregnant women

Table 35: GRADE profile – nitrofurantoin and close monitoring versus close monitoring 4

			Quality ass	essment			No of pat	ients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin and close surveillance	Close surveillance alone	Relative (95% CI)	Absolute		
Recurren	t pyelonephr	itis										
-		no serious risk of bias	N/A	no serious indirectness	very serious²	none	6/82 (7.3%)	7/85 (8.2%)	RR 0.89 (0.31 to 2.53)	9 fewer per 1000 (from 57 fewer to 126 more)	⊕⊕OO LOW	CRITICAL
Recurren	t urinary trac	t infection	(cystitis)									
-		no serious risk of bias	N/A	no serious indirectness	very serious²	none	2/82 (2.4%)	7/85 (8.2%)	RR 0.3 (0.06 to 1.38)	58 fewer per 1000 (from 77 fewer to 31 more)	⊕⊕OO LOW	CRITICAL
Asympto	matic bacteri	iuria in won	nen with 90% c	linical attendar	ice		•			•		
		no serious risk of bias	N/A	no serious indirectness	serious ³	none	14/43 (32.6%)	35/59 (59.3%)	RR 0.55 (0.34 to 0.89)	267 fewer per 1000 (from 392 fewer to 65 fewer)		CRITICAL
Preterm b	oirth (<37 we	eks)										
		no serious risk of bias	N/A	no serious indirectness	very serious²	none	7/73 (9.6%)	6/74 (8.1%)	RR 1.18 (0.42 to 3.35)	15 more per 1000 (from 47 fewer to 191 more)	⊕⊕OO LOW	CRITICAL
Birthweig	ht (g) (Bette	r indicated	by higher value	es)								
	randomised trials	serious ⁴	N/A	no serious indirectness	serious⁵	none	71	76		wer (327.2 lower to 1.2 higher)	⊕⊕OO LOW	CRITICAL
5-min Ap	gar score <7											

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			Quality ass	essment			No of pati	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin and close surveillance	Close surveillance alone	Relative (95% CI)	Absolute		
1 ¹	randomised trials	serious ⁴		no serious indirectness	very serious²	none	2/73 (2.7%)	1/74 (1.4%)	RR 2.03 (0.19 to 21.87)	14 more per 1000 (from 11 fewer to 282 more)	⊕OOO VERY LOW	CRITICAL
Miscarria	iges											
1 ¹	randomised trials	serious ⁴		no serious indirectness	very serious²	none	3/82 (3.7%)	1/85 (1.2%)	RR 3.11 (0.33 to 29.29)	25 more per 1000 (from 8 fewer to 333 more)	⊕000 VERY LOW	CRITICAL
Abbreviat	ions: N/A ,not	applicable;	CI, confidence i	nterval: RR, risk	ratio							

¹ Schneeberger et al. 2015

² Downgraded 2 levels - at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm

³ Downgraded 1 level - at a 95% confidence interval, data are consistent with no meaningful difference, appreciable benefit with nitrofurantoin

⁴ Downgraded 1 level -it is unclear how the lack of blinding would have led to a under or over estimation of effect

⁵ Downgraded 1 level – at a minimal important difference of 0.5 standard deviation of the close surveillance arm, data are consistent with no meaningful difference or appreciable benefit with close surveillance alone

H.7 Antimicrobials in a mixed population of adults and children

2 Table 36: GRADE profile – nitrofurantoin versus placebo in adults and children

			Quality asse	ssment			No of pati	ents		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin	Placebo	Relative (95% Cl)	Absolute		
Occurrenc	e of urinary tr	act infection	on									
-	randomised trials		no serious inconsistency		no serious imprecision	none	38/169 (22.5%)	190/322 (59%)	RR 0.38 (0.28 to 0.5)	366 fewer per 1000 (from 425 fewer to 295 fewer)	⊕⊕OO LOW	CRITICAL
Abbreviatio	ons: N/A ,not ap	plicable; C	I, confidence interva	al: RR, risk rati	0							

¹ Muller et al. 2017

² Downgraded 1 level - high risk of bias associated with the lack of randomisation in 3 studies; randomisation was unclear in 3 studies.

³ Downgraded by 1 level - one study included patients with spinal cord injury, another study included children with neurogenic bladder

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1 ⁵ Downgraded by 1 level – not assessable

2 Table 37: GRADE profile - nitrofurantoin versus antibiotics in adults and children

			Quality asse	essment			No of pa	tients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin	Antibiotics	Relative (95% CI)	Absolute		
Occurren	ce of urinary t	ract infect	ion									
22 ¹		,	no serious inconsistency		very serious ³	none	119/511 (23.3%)	211/808 (26.1%)	RR 0.93 (0.68 to 1.26)	18 fewer per 1000 (from 84 fewer to 68 more)	⊕000 VERY LOW	CRITICAL
Mild adve	rse effects		·									
22 ¹		,	no serious inconsistency	no serious indirectness	no serious	none	154/503 (30.6%)	82/702 (11.7%)	RR 2.24 (1.77 to 2.83)	145 more per 1000 (from 90 more to 214 more)	⊕⊕OO LOW	CRITICAL
Abbreviatio	ons: CI, confide	ence interv	al: RR, risk ratio									

3 4 5

7 8 9

² Downgraded 2 levels - majority of evidence was to be of high risk of bias, which is likely to affect the measurement of the outcome ³ Downgraded 2 levels – at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm

Table 38: GRADE profile – nitrofurantoin versus methenamine hippurate in adults and children 6

			Quality as	sessment			No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin	Methenamine hippurate	Relative (95% CI)	Absolute		
Occurren	ce of urinary	tract infe	ection									
	randomised trials		no serious inconsistency	no serious indirectness	serious ³	none	24/67 (35.8%)	66/129 (51.2%)	RR 0.60 (0.43 to 0.85)	205 fewer per 1000 (from 292 fewer to 77 fewer)	⊕⊕OO LOW	CRITICAL
Mild side	effects			<u>.</u>	-					<u>.</u>		
21	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	24/67 (35.8%)	9/129 (7%)	RR 4.22 (2.06 to 8.67)	225 more per 1000 (from 74 more to 535 more)	⊕⊕⊕O MODERATE	CRITICAL

¹ Muller et al. 2017

¹ Muller et al. 2017

² Downgraded by 1 level as majority of evidence has high risk of bias, which is likely to affect the measurement of the outcome
 ³ Downgraded 1 level – at a 95% confidence interval, data are consistent with no meaningful difference or appreciable harm with methenamine hippurate

Table 39: GRADE profile – nitrofurantoin versus trimethoprim in adults and children 10

Quality assessment No of patients Effect Quality Importance

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No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin	Trimethoprim	Relative (95% Cl)	Absolute		
Occurren	ce of urinary	tract infe	ction									
5 ¹	randomised trials	serious ²		no serious indirectness	very serious ⁴	none	32/142 (22.5%)	61/208 (29.3%)	RR 0.81 (0.38 to 1.71)	56 fewer per 1000 (from 182 fewer to 208 more)	⊕000 VERY LOW	IMPORTANT
Mild adve	erse effects											
4 ¹	randomised trials				no serious imprecision	none	58/138 (42%)	28/192 (14.6%)	RR 2.20 (1.51 to 3.2)	175 more per 1000 (from 74 more to 321 more)	⊕⊕⊕O MODERATE	CRITICAL
Abbreviati	ons: CI, confid	lence inter	rval: RR, risk ratio				•					

¹ Muller et al. 2017

² Downgraded by 1 level as majority of evidence has high risk of bias

³ Downgraded 1 level – heterogeneity > 50% ⁴ Downgraded 2 levels – at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm

5 Table 40: GRADE profile - nitrofurantoin versus co-trimoxazole in adults and children

			Quality asso	essment			No of pa	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin	Co- trimoxazole	Relative (95% Cl)	Absolute		
Occurren	ce of urinary	tract infec	tion	-	•		•					
4 ¹	randomised trials	very serious²	no serious inconsistency	serious ³	very serious ⁴	none	3/25 (12%)	5/56 (8.9%)	RR 1.42 (0.17 to 12)	37 more per 1000 (from 74 fewer to 982 more)	⊕OOO VERY LOW	CRITICAL
Mild adve	rse effects	•	•	-	•		•					•
1 ¹	randomised trials	serious ²	N/A	no serious indirectness	very serious ⁴	none	1/6 (16.7%)	1/13 (7.7%)	RR 2.17 (0.16 to 29.1)	90 more per 1000 (from 65 fewer to 1000 more)	⊕OOO VERY LOW	CRITICAL
Abbreviatio	ons: CI, confid	ence interv	al: RR, risk ratio				•					

¹ Muller et al. 2017

² Downgraded by 1 level as majority of evidence has high risk of bias
 ³ Downgraded by 1 level as one study included children with vesicoureteral reflux
 ⁴ Downgraded 2 levels – at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm

10 Table 41: GRADE profile – nitrofurantoin versus beta-lactams in adults and children

Quality assessment No of patients Effect Quality I	Importance	nce
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studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin	Beta- lactams	Relative (95% Cl)	Absolute		
Occurrence	e of urinary tr	ract infecti	ion									
	rials	,		no serious indirectness	very serious ³	none	19/115 (16.5%)	30/134 (22.4%)	RR 0.84 (0.49 to 1.44)	36 fewer per 1000 (from 114 fewer to 99 more)	⊕000 VERY LOW	CRITICAL
tri	rials	serious ²	no serious inconsistency al: RR, risk ratio	no serious indirectness	serious ⁴	none	32/128 (25%)	18/147 (12.2%)	RR 1.99 (1.19 to 3.32)	121 more per 1000 (from 23 more to 284 more)	⊕000 VERY LOW	CRITICAL

¹ Muller et al. 2017

² Downgraded 1 level - majority of evidence has very high risk of bias

³ Downgraded 2 levels – at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm ⁴ Downgraded 1 level – at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with nitrofurantoin

5 Table 42: GRADE profile - nitrofurantoin versus guinolones in adults and children

Quality assessment				No of patients		Effect		Quality	Importance			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin	Quinolones	Relative (95% Cl)	Absolute		
Occurrenc	Decurrence of urinary tract infection											
3 ¹	randomised trials	very serious²	serious ³		very serious⁵	none	25/84 (29.8%)	15/102 (14.7%)	RR 2.26 (0.73 to 7)	185 more per 1000 (from 40 fewer to 882 more)	⊕000 VERY LOW	CRITICAL
Mild adver	se effects											
3 ¹	randomised trials	very serious²	serious ³		very serious⁵	none	24/112 (21.4%)	19/118 (16.1%)	RR 1.37 (0.79 to 2.36)	60 more per 1000 (from 34 fewer to 219 more)	⊕000 VERY LOW	CRITICAL
Abbreviatio	ns: Cl, confide	nce interval	: RR, risk ratio	•		•	•					

¹ Muller et al. 2017

² Downgraded by 1 level as majority of evidence has high risk of attrition bias, which is likely to affect the measurement of the outcome

³ Downgraded 1 level – heterogeneity > 50%

⁴ Downgraded 1 level as nitrofurantoin was compared to cinoxacin (not available in the UK), in 2 of the studies

⁵ Downgraded 2 levels – at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm

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Draft GRADE profiles

H.8 Antimicrobials in children

2 Table 43: GRADE profile – antibiotic versus placebo or no treatment

			Quality ass	essment			No of	f patients		Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Antibiotic	Placebo or no treatment	Relative (95% Cl)	Absolute		
Recurrer	nce of sympto	omatic uri	nary tract infecti	on - no vesicou	ireteral reflux	x						
3 ¹	randomised trials	serious ²	serious ³	no serious indirectness	very serious⁵	none	20/273 (7.3%)	30/218 (13.8%)	RR 0.56 (0.15 to 2.12)	61 fewer per 1000 (from 117 fewer to 154 more)	⊕000 VERY LOW	CRITICAL
Recurrer	nce of sympto	omatic uri	nary tract infecti	on								
4 ¹	randomised trials	serious ²	serious ³	serious ⁴	very serious⁵	none	58/553 (10.5%)	81/471 (17.2%)	RR 0.75 (0.36 to 1.53)	43 fewer per 1000 (from 110 fewer to 91 more)	⊕000 VERY LOW	CRITICAL
Repeat p	ositive cultu	re		•								
4 ¹	randomised trials	serious ²	very serious ³	serious ⁴	serious ⁶	none	43/270 (15.9%)	76/197 (38.6%)	RR 0.31 (0.08 to 1.18)	266 fewer per 1000 (from 355 fewer to 69 more)	⊕000 VERY LOW	CRITICAL
Microbia	l resistance t	o prophy	lactic drug									
2 ¹	randomised trials		no serious inconsistency	no serious indirectness	very serious⁵	none	18/51 (35.3%)	11/67 (16.4%)	RR 2.4 (0.62 to 9.26)	230 more per 1000 (from 62 fewer to 1000 more)	⊕000 VERY LOW	CRITICAL
All adve	se events			•								
2 ¹	randomised trials	serious ²	very serious ³	serious ⁴	very serious⁵	none	19/499 (3.8%)	10/415 (2.4%)	RR 2.31 (0.03 to 170.67)	32 more per 1000 (from 23 fewer to 1000 more)	⊕000 VERY LOW	CRITICAL
Withdrav	val due to ad	verse eve	nts		1							L
2 ¹	randomised trials	serious ²	no serious inconsistency	serious ⁴	very serious⁵	none	4/288 (1.4%)	10/288 (3.5%)	RR 0.40 (0.13 to 1.26)	21 fewer per 1000 (from 30 fewer to 9 more)	⊕000 VERY LOW	CRITICAL
Rate of r	new or deterio	orated ren	al scars	•			•				•	•
7 ⁷	randomised trials		no serious inconsistency	serious ⁴	very serious⁵	none	17/578 (2.9%)	18/515 (3.5%)	RR 0.95 (0.51 to 1.78)	2 fewer per 1000 (from 17 fewer to 27 more)	⊕000 VERY LOW	CRITICAL
Emerger	ice of resista	nce		·	• 	•	•					·
1 ⁸	randomised trials	serious ⁹	N/A	no serious indirectness	serious ¹⁰	none	1	ד=15	resistant stra	ere replaced over treatment by ains in children receiving but not in children receiving placebo.	⊕⊕OO LOW	CRITICAL

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- ¹ Williams and Craig 2011
- ² Downgraded by 2 levels due to a very high risk of bias lack of randomisation, lack of blinding, selective reporting of outcomes
- ³ Downgraded 1 level heterogeneity > 50%

⁴ Downgraded by 1 level as most studies did not report a clear inclusion and exclusion criteria for participants entry into the study; it was possible for patients to be misclassified

- ⁵ Downgraded 2 levels at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm
- ⁶ Downgraded 1 level at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with placebo
- ⁷ Dai et al. 2010

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⁸ Muller et al. 2017

⁹ Downgraded 1 level - high risk of bias associated with the lack of randomisation in 3 studies and randomisation was unclear in 3 studies included in systematic review; unclear which studies this is relevant to

¹⁰ Downgraded 1 level – not assessable

12 **Table 44: GRADE profile – Nitrofurantoin versus trimethoprim**

			Quality as	sessment			No of p	atients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin		Relative (95% Cl)	Absolute	Quanty	importance
Repeat p	ositive cult	ure		-								
		no serious risk of bias	N/A		no serious imprecision	none	12/60 (20%)	37/60 (61.7%)	RR 0.3 (0.2 to 0.6)	432 fewer per 1000 (from 493 fewer to 247 fewer)	⊕⊕⊕O MODERATE	CRITICAL
Adverse	events											
		no serious risk of bias	N/A	serious ²	serious ³	none	8/31 (25.8%)	18/29 (62.1%)	RR 0.42 (0.21 to 0.81)	360 fewer per 1000 (from 490 fewer to 118 fewer)	⊕⊕OO LOW	CRITICAL
Emerger	nce of resista	ance		•				•				
	randomised trials	serious⁵	N/A	serious ⁶	serious ⁷	none	n=67	n=63	prophylax accessibl	nce rates linked to nitrofurantoin is reduced (9% to 7%; quality not e) whereas rates associated with rim prophylaxis increased (8% to 47%)	⊕⊕OO VERY LOW	CRITICAL
Abbrevia	tions: N/A, no	ot applica	able; CI, confider	nce interval: RF	R, risk ratio							

¹ Williams and Craig 2011

² Downgraded by 1 level as 30 children had vesicoureteral reflux or significant structural abnormalities

³ Downgraded 1 level – at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with trimethoprim

⁴ Muller et al. 2017

⁵ Downgraded 1 level - high risk of bias associated with the lack of randomisation in 3 studies and randomisation was unclear in 3 studies included in systematic review; unclear which studies this is relevant to

⁶ Downgraded by 1 level - study included children with neurogenic bladder

⁷ Downgraded 1 level – not assessable

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Draft for consultation

Table 45: GRADE profile – Nitrofurantoin versus co-trimoxazole 1

Quality assessme				sment			No of patients		Effect		Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin	Co- trimoxazole	Relative (95% Cl)	Absolute		
Recurrence	ecurrence of symptomatic urinary tract infection											
	randomised trials	serious ²	N/A	serious ³	serious ⁴	none	17/66 (25.8%)	30/66 (45.5%)	RR 0.57 (0.35 to 0.92)	195 fewer per 1000 (from 295 fewer to 36 fewer)	⊕000 VERY LOW	CRITICAL
Microbial	resistance to	prophylac	tic drugs									
	randomised trials		no serious inconsistency	serious ³	serious ⁴	none	10/29 (34.5%)	45/67 (67.2%)	RR 0.54 (0.31 to 0.92)	309 fewer per 1000 (from 463 fewer to 54 fewer)	⊕000 VERY LOW	CRITICAL
Abbreviatio	ons: N/A, not a	oplicable; C	CI, confidence interv	al: RR, risk ra	tio		•	-				

¹ Williams and Craig 2011

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² Downgraded 2 levels - a very high risk of bias - lack of randomisation, lack of blinding, selective reporting of outcomes
 ³ Downgraded 1 level – classification of children was unclear
 ⁴ Downgraded 1 level – at a minimal important difference of 25%, data are consistent with no meaningful difference or appreciable harm with co-trimoxazole

6 Table 46: GRADE profile – Nitrofurantoin versus cefixime

Quality assessment						No of patients		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Nitrofurantoin	Cefixime	(95% CI)	Quanty	importance
Repeat pos	peat positive culture										
1 ¹	randomised trials	serious ²	N/A		no serious imprecision	none	3/30 (10%)	2/27 (7.4%)	Risk difference 0.03 (-0.12 to 0.17)	⊕⊕⊕O MODERATE	CRITICAL
Adverse ev	vents				·						
1 ¹	randomised trials	serious ²	N/A		no serious imprecision	none	37/60 (61.7%)	17/60 (28.3%)	Risk difference 2.18 (1.39 to 3.41)	⊕⊕⊕O MODERATE	CRITICAL
Abbreviatio	ns: N/A, not appl	icable; CI, c	onfidence interv	al: RR, risk ratio	•	•	•				

¹ Williams et al. 2011

² Downgraded 1 level - most studies did not report a clear inclusion and exclusion criteria for participants entry into the study; it was possible for patients to be misclassified, not all planned outcomes were reported

¹Jepson et al. 2012

² Downgraded 2 levels - at a minimal important difference of 25%, data are consistent with no meaningful difference, appreciable benefit or appreciable harm

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Appendix I: Studies not-prioritised

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Study reference		Reason
Afshar K, Stothers L, Scott H, and MacNeily A Cranberry juice for the prevention of pediatric infection: a randomized controlled trial. The Jo 188(4 Suppl), 1584-7	urinary tract	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Antachopoulos Charalampos, Ioannidou Maria Athanasios, Iosifidis Elias, Katragkou Aspasia Paschalis, Kollios Konstantinos, and Roilides Comparison of cotrimoxazole vs. second-gene cephalosporins for prevention of urinary tract children. Pediatric nephrology (Berlin, and Ge 2271-2276	n, Kadiltzoglou Emmanuel (2016) eration infections in	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Bailey David T, Dalton Carol, Joseph Daughe Tempesta Michael S (2007) Can a concentrat extract prevent recurrent urinary tract infection pilot study. Phytomedicine : international journ and phytopharmacology 14(4), 237-41	ted cranberry ns in women? A	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Barbosa-Cesnik Cibele, Brown Morton B, Bux Lixin, DeBusscher Joan, and Foxman Betsy (2 juice fails to prevent recurrent urinary tract info a randomized placebo-controlled trial. Clinical diseases : an official publication of the Infection Society of America 52(1), 23-30	2011) Cranberry ection: results from l infectious	RCT included in a systematic review that has been prioritised
Beerepoot M A. J, Geerlings S E, van Haarst, N Mensing, ter Riet, and G (2013) Nonantibio recurrent urinary tract infections: a systematic analysis of randomized controlled trials. The J 190(6), 1981-9	tic prophylaxis for review and meta-	A higher quality systematic review has been prioritised (Perrotta et al. 2011)
Beerepoot Maj, Ter Riet G, Nys S, Wal Wm, E Tm, Prins Jm, Koeijers J, Verbon A, Stobberir Geerlings Se (2013) Lactobacilli versus antibi- urinary tract infections: A randomized, double- trial in postmenopausal women. [Dutch]. Nede voor geneeskunde 157(10),	ngh Ee, and otics to prevent -blind, noninferiority	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Bianco L, Perrelli E, Towle V, Ness Ph, and Ju (2012) Pilot randomized controlled dosing stu- capsules for reduction of bacteriuria plus pyur home residents. Journal of the American Geri 60(6), 1180-1	dy of cranberry ia in female nursing	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Bosmans JE, Beerepoot MA, Prins JM, ter Rie SE (2014) Cost-effectiveness of cranberries v prevent urinary tract infections in premenopau randomized clinical trial. PloS one 9(4), e9193	rs antibiotics to usal women: a	No or fewer critical outcomes reported
Caljouw Monique A. A, van den Hout, Wilbert Achterberg Wilco P, Cools Herman J. M, and (2014) Effectiveness of cranberry capsules to tract infections in vulnerable older persons: a randomized placebo-controlled trial in long-ter Journal of the American Geriatrics Society 62	Gussekloo Jacobijn prevent urinary double-blind rm care facilities.	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Ferrara Pietro, Romaniello Luciana, Vitelli Ott Serva Martina, and Cataldi Luigi (2009) Crant prevention of recurrent urinary tract infections controlled trial in children. Scandinavian journ	perry juice for the a randomized:	RCT included in a systematic review that has been prioritised

nephrology 43(5), 369-72

Study reference	Reason
Ledda A, Bottari A, Luzzi R, Belcaro G, Hu S, Dugall M, Hosoi M, Ippolito E, Corsi M, Gizzi G, Morazzoni P, Riva A, Giacomelli L, and Togni S (2015) Cranberry supplementation in the prevention of non-severe lower urinary tract infections: a pilot study. European review for medical and pharmacological sciences 19(1), 77-80	A higher quality systematic review has been prioritised (Fu et al. 2017)
Luís, Â, Domingues F and Pereira L. Can Cranberries Contribute to Reduce the Incidence of Urinary Tract Infections? A Systematic Review with Meta-Analysis and Trial Sequential Analysis of Clinical Trials. The Journal of Urology. Sept 2017;198(3):614–621	A higher quality systematic review has been prioritised (Fu et al. 2017)
Maki Kevin C, Kaspar Kerrie L, Khoo Christina, Derrig Linda H, Schild Arianne L, and Gupta Kalpana (2016) Consumption of a cranberry juice beverage lowered the number of clinical urinary tract infection episodes in women with a recent history of urinary tract infection. The American journal of clinical nutrition 103(6), 1434-42	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Mathew JL. Antibiotic prophylaxis following urinary tract infection in children: a systematic review of randomized controlled trials. Indian pediatrics. 2010 Jul 1;47(7):599-605.	A higher quality systematic review has been prioritised (Williams and Craig 2011)
McMurdo Marion E. T, Argo Ishbel, Phillips Gabby, Daly Fergus, and Davey Peter (2009) Cranberry or trimethoprim for the prevention of recurrent urinary tract infections? A randomized controlled trial in older women. The Journal of antimicrobial chemotherapy 63(2), 389-95	RCT included in a systematic review that has been prioritised
Mori et al. 2009, Antibiotic prophylaxis for children at risk of developing urinary tract infection: a systematic review. Acta paediatrica (Oslo, and Norway : 1992) 98(11), 1781-6	A higher quality systematic review has been prioritised (Williams and Craig 2011)
Norinder Birgit Stattin, Norrby Ragnar, Palmgren Ann-Chatrin, Hollenberg Sofia, Eriksson Ulla, and Nord Carl Erik (2006) Microflora changes with norfloxacin and pivmecillinam in women with recurrent urinary tract infection. Antimicrobial agents and chemotherapy 50(4), 1528-30	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Porru D, Parmigiani A, Tinelli C, Barletta D, Choussos D, Di Franco C, Bobbi V, Bassi S, Miller O, Gardella B, Nappi R E, Spinillo A, and Rovereto B (2014) Oral D-mannose in recurrent urinary tract infections in women: A pilot study. Journal of Clinical Urology 7(3), 208-213	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Price Jameca Renee, Guran Larissa A, Gregory W Thomas, and McDonagh Marian S (2016) Nitrofurantoin vs other prophylactic agents in reducing recurrent urinary tract infections in adult women: a systematic review and meta-analysis. American journal of obstetrics and gynecology 215(5), 548-560	A higher quality systematic review has been prioritised (Muller et al. 2017)
Salo Jarmo, Uhari Matti, Helminen Merja, Korppi Matti, Nieminen Tea, Pokka Tytti, and Kontiokari Tero (2012) Cranberry juice for the prevention of recurrences of urinary tract infections in children: a randomized placebo-controlled trial. Clinical infectious diseases : an official publication of the Infectious Diseases Society of America 54(3), 340-6	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Sengupta K, Alluri K V, Golakoti T, Gottumukkala G V, Raavi J, Kotchrlakota L, Sigalan S C, Dey D, Ghosh S, and Chatterjee A (2011) A randomized, double blind, controlled, dose dependent clinical trial to evaluate the efficacy of a proanthocyanidin standardized whole cranberry (Vaccinium macrocarpon) powder on infections of the urinary tract. Current Bioactive Compounds 7(1), 39-46	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review

Study reference	Reason
Singh Iqbal, Gautam Lokesh Kumar, and Kaur Iqbal R (2016) Effect of oral cranberry extract (standardized proanthocyanidin- A) in patients with recurrent UTI by pathogenic E. coli: a randomized placebo-controlled clinical research study. International urology and nephrology 48(9), 1379-86	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Stapleton Ann E, Au-Yeung Melissa, Hooton Thomas M, Fredricks David N, Roberts Pacita L, Czaja Christopher A, Yarova-Yarovaya Yuliya, Fiedler Tina, Cox Marsha, and Stamm Walter E (2011) Randomized, placebo-controlled phase 2 trial of a Lactobacillus crispatus probiotic given intravaginally for prevention of recurrent urinary tract infection. Clinical infectious diseases : an official publication of the Infectious Diseases Society of America 52(10), 1212-7	RCT included in a systematic review that has been prioritised
Stapleton Ann E, Dziura James, Hooton Thomas M, Cox Marsha E, Yarova-Yarovaya Yuliya, Chen Shu, and Gupta Kalpana (2012) Recurrent urinary tract infection and urinary Escherichia coli in women ingesting cranberry juice daily: a randomized controlled trial. Mayo Clinic proceedings 87(2), 143-50	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
Takahashi Satoshi, Hamasuna Ryoichi, Yasuda Mitsuru, Arakawa Soichi, Tanaka Kazushi, Ishikawa Kiyohito, Kiyota Hiroshi, Hayami Hiroshi, Yamamoto Shingo, Kubo Tatsuhiko, and Matsumoto Tetsuro (2013) A randomized clinical trial to evaluate the preventive effect of cranberry juice (UR65) for patients with recurrent urinary tract infection. Journal of infection and chemotherapy : official journal of the Japan Society of Chemotherapy 19(1), 112-7	This RCT does not provide additional evidence that adds to the evidence from a prioritised systematic review
van den Hout WB, Caljouw MA, Putter H, Cools HJ, and Gussekloo J (2014) Cost-effectiveness of cranberry capsules to prevent urinary tract infection in long-term care facilities: economic evaluation with a randomized controlled trial. Journal of the American Geriatrics Society 62(1), 111-6	No or fewer critical outcomes reported
Wang Chih-Hung, Fang Cheng-Chung, Chen Nai-Chuan, Liu Sot Shih-Hung, Yu Ping-Hsun, Wu Tao-Yu, Chen Wei-Ting, Lee Chien-Chang, and Chen Shyr-Chyr (2012) Cranberry-containing products for prevention of urinary tract infections in susceptible populations: a systematic review and meta-analysis of randomized controlled trials. Archives of internal medicine 172(13), 988-96	A higher quality systematic review has been prioritised (Jepson et al. 2012)

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Appendix J: Excluded studies

Study reference	Reason for exclusion
Altarac Silvio, and Papes Dino (2014) Use of D-mannose in prophylaxis of recurrent urinary tract infections (UTIs) in women. BJU international 113(1), 9-10	Publication/study type (literature review)
Aydin A, Ahmed K, Zaman I, Khan M S, and Dasgupta P (2015) Recurrent urinary tract infections in women. Obstetrical and Gynecological Survey 70(10), 621-622q2	Publication/study type (literature review)
Beerepoot Maj, Ter Riet G, Nys S, Wal Wm, Borgie Cajm, Reijke Tm, Prins Jm, Koeijers J, Verbon A, Stobberingh E E, and Geerlings S E (2012) Predictive value of Escherichia coli susceptibility in strains causing asymptomatic bacteriuria for women with recurrent symptomatic urinary tract infections receiving prophylaxis. Clinical Microbiology and Infection. 1;18(4).	Publication/study type
Beversdorf D Q, Galloway H S, Foster Sr, R T, and Tatum P E (2011) Preventing recurrent urinary tract infections in a woman with dementia. Clinical Geriatrics 19(11), 33-35	Unable to source study
Wiese Birgitt, and Gagyor Ildiko (2016) Recurrent urinary tract	Publication/study type (retrospective long-term follow- up analysis)
Bonetta A, Derelli R, and Pierro F (2011) Cranberry extracts reduce urinary tract infections during radiotherapy for prostate adenocarcinoma. Anticancer research 31(5), 1849-1850	Unable to source study
Braga Luis H, Pemberton Julia, Heaman Jessie, DeMaria Jorge, and Lorenzo Armando J (2014) Pilot randomized, placebo controlled trial to investigate the effect of antibiotic prophylaxis on the rate of urinary tract infection in infants with prenatal hydronephrosis. The Journal of urology 191(5 Suppl), 1501-7	Not a relevant study
Brandstrom P (2011) The swedish reflux trial. Pediatric nephrology (Berlin, and Germany) 26(9), 1733	Not relevant population
Brandström P, Jodal U, Sillén U, and Hansson S (2011) The Swedish reflux trial: review of a randomized, controlled trial in children with dilating vesicoureteral reflux. Journal of pediatric urology 7(6), 594-600	Not relevant population
Brandstrom P, and Hansson S (2013) Growth in children with dilating VUR-a follow up of the swedish reflux trial. Pediatric nephrology (Berlin, and Germany) 28(8), 1391	Not relevant population
Canning D A (2010) Antibiotic prophylaxis and recurrent urinary tract infection in children. Journal of Urology 184(5), 2135	Unable to source study
Cayley Jr, and W E (2013) Are cranberry products effective for the prevention of urinary tract infections?. American Family Physician 88(11), 745-746	Publication/study type (commentary)
Cote J, Caillet S, Doyon G, Sylvain JF, and Lacroix M (2010) Bioactive compounds in cranberries and their biological properties. Critical reviews in food science and nutrition 50(7), 666-79	Not a relevant study
	Poor relevance against search terms (intervention)

Study reference	Reason for exclusion
Damiano R, Quarto G, Bava I, Ucciero G, De Domenico, R, Palumbo M I, and Autorino R (2011) Erratum: Prevention of recurrent urinary tract infections by intravesical administration of hyaluronic acid and chondroitin sulphate: A placebo-controlled randomised trial (European Urology (2011) 59 (645-651)). European Urology 60(1), 193	Poor relevance against search terms (intervention)
Dessi A, Atzei A, and Fanos V (2011) Cranberry in children: Prevention of recurrent urinary tract infections and review of the literature. Brazilian Journal of Pharmacognosy 21(5), 807-813	Publication/study type (literature review)
De Vita, Davide, and Giordano Salvatore (2012) Effectiveness of intravesical hyaluronic acid/chondroitin sulfate in recurrent bacterial cystitis: a randomized study. International urogynecology journal 23(12), 1707-13	Poor relevance against search terms (intervention)
De Vita, Davide, Antell Henrik, and Giordano Salvatore (2013) Effectiveness of intravesical hyaluronic acid with or without chondroitin sulfate for recurrent bacterial cystitis in adult women: a meta-analysis. International urogynecology journal 24(4), 545- 52	Poor relevance against search terms (intervention)
Dieter A A (2015) Cranberry capsules (2 taken twice daily for an average 38 days) reduce the risk of postoperative urinary tract infection in women undergoing benign gynaecological surgery involving intraoperative catheterisation. Evidence-Based Medicine 20(4), 137	Publication/study type (commentary)
Donabedian H (2006) Nutritional therapy and infectious diseases: a two-edged sword. Nutrition journal 5, 21	Not a relevant study
Dotis J, Printza N, Stabouli S, Pavlaki A, Samara S, and Papachristou F (2014) Efficasy of cranberry capsules to prevent recurences of urinary tract infections. Pediatric nephrology (Berlin, and Germany) 29(9), 1793-4	Unable to source study
Duenas-Garcia O F, Sullivan G, Hall C D, Flynn M K, and O'Dell K (2016) Pharmacological agents to decrease new episodes of recurrent lower urinary tract infections in postmenopausal women. A systematic review. Female Pelvic Medicine and Reconstructive Surgery 22(2), 63-69	Not a relevant study
Durham Spencer H, Stamm Pamela L, and Eiland Lea S (2015) Cranberry Products for the Prophylaxis of Urinary Tract Infections in Pediatric Patients. The Annals of pharmacotherapy 49(12), 1349-56	Publication/study type (literature review)
Edmonson M Bruce, and Eickhoff Jens C (2017) Weight Gain and Obesity in Infants and Young Children Exposed to Prolonged Antibiotic Prophylaxis. JAMA pediatrics 171(2), 150- 156	Not relevant population
Eells Samantha J, McKinnell James A, and Miller Loren G (2011) Daily cranberry prophylaxis to prevent recurrent urinary tract infections may be beneficial in some populations of women. Clinical infectious diseases : an official publication of the Infectious Diseases Society of America 52(11), 1393-5	Publication/study type (commentary)
Epp Annette, Larochelle Annick, Lovatsis Danny, Walter Jens- Erik, Easton William, Farrell Scott A, Girouard Lise, Gupta Chander, Harvey Marie-Andree, Robert Magali, Ross Sue, Schachter Joyce, Schulz Jane A, Wilkie David, Ehman William, Domb Sharon, Gagnon Andree, Hughes Owen, Konkin Jill, Lynch Joanna, Marshall Cindy, Society of, Obstetricians, Gynaecologists of, and Canada (2010) Recurrent urinary tract infection. Journal of obstetrics and gynaecology Canada : JOGC	Publication/study type (literature review)

Study reference	Reason for exclusion
= Journal d'obstetrique et gynecologie du Canada : JOGC 32(11), 1082-101	
Espino M, Areses R, Meseguer Cg, Pena A, Melgosa M, Ruperez M, Mitjavilla M, and Albillos Jc (2012) Antibiotic prophylaxis inhighdegree vesicoureteral reflux. Prospective, randomized and multicentric study. Preliminary results. Pediatric nephrology (Berlin, and Germany) 27(9), 1648-9	Publication/study type (commentary)
Falakaflaki B, Fallah R, Jamshidi Mr, Moezi F, and Torabi Z (2007) Comparison of nitrofurantoin and trimethoprim- sulphamethoxazole for long-term prophylaxis in children with recurrent urinary tract infections. International Journal of Pharmacology 3(2), 179-82	Not relevant population
Fanos V, Atzei A, Zaffanello M, Piras A, and Cataldi L (2006) Cranberry and prevention of urinary tract infections in children. Journal of chemotherapy (Florence, and Italy) 18 Spec no 3, 21- 4	Publication/study type (literature review)
Fernández-Puentes V, Uberos J, Rodríguez-Belmonte R, Nogueras-Ocaña M, Blanca-Jover E, and Narbona-López E (2015) Efficacy and safety profile of cranberry in infants and children with recurrent urinary tract infection. Anales de pediatria (barcelona, and spain : 2003) 82(6), 397-403	Unable to source
Flower Andrew, Wang Li-Qiong, Lewith George, Liu Jian Ping, and Li Qing (2015) Chinese herbal medicine for treating recurrent urinary tract infections in women. The Cochrane database of systematic reviews (6), CD010446	Does not reflect usual UK practice
Fonseca Fernando F, Tanno Fabio Y, and Nguyen Hiep T (2012) Current options in the management of primary vesicoureteral reflux in children. Pediatric clinics of North America 59(4), 819-34	Not relevant population
Foxman B, Cronenwett AE, Spino C, Berger MB, and Morgan DM (2015) Cranberry juice capsules and urinary tract infection after surgery: results of a randomized trial. American journal of obstetrics and gynecology 213(2), 194.e1-8	Not relevant population
Foxman Betsy, Cronenwett Anna E. W, Spino Cathie, Berger Mitchell B, and Morgan Daniel M (2015) Cranberry juice capsules and urinary tract infection after surgery: results of a randomized trial. American journal of obstetrics and gynecology 213(2), 194.e1-8	Duplicate
Fromentin E, Vostalova J, Vidlar A, Galandakova A, Vrbkova J, Ulrichova J, Student V, and Simanek V (2014) A randomized, double-blind, placebo-controlled clinical trial to investigate the efficacy of cranberry fruit powder (Pacran) in the prevention of recurrent urinary tract infection in women. FASEB journal 28(1 suppl. 1),	Abstract only
Gallien P, and Reymann Jm (2008) Cranberry for prevention of urinary tract infections in multiple sclerosis patients. ClinicalTrials gov (www clinicaltrials gov) (accessed 4 November 2010),	Publication/study type (study registration)
Gallien Philippe, Amarenco Gerard, Benoit Nicolas, Bonniaud Veronique, Donze Cecile, Kerdraon Jacques, de Seze, Marianne, Denys Pierre, Renault Alain, Naudet Florian, and Reymann Jean Michel (2014) Cranberry versus placebo in the prevention of urinary infections in multiple sclerosis: a multicenter, randomized, placebo-controlled, double-blind trial. Multiple sclerosis (Houndmills, Basingstoke, and England) 20(9), 1252-9	Not relevant population
Garin Eduardo H, Olavarria Fernando, Garcia Nieto, Victor , Valenciano Blanca, Campos Alfonso, and Young Linda (2006)	Not relevant population

Study reference	Person for evolution
Study reference Clinical significance of primary vesicoureteral reflux and urinary	Reason for exclusion
antibiotic prophylaxis after acute pyelonephritis: a multicenter, randomized, controlled study. Pediatrics 117(3), 626-32	
Gautam L, Singh I, Gautam Lk, Kaur Ir, Rai S, and Joshi Mk (2014) Effect of oral cranberry extract (standardised proanthocyanidin-a) on the uropathogenic bacteria in urine of patients with subclinical/recurrent uti: A randomised placebo controlled clinical study. Indian journal of urology 30, S152	Abstract only
Gupta A (2007) Cranberry and Prevention of UTI - A Comprehensive Approach. http://www.clinicaltrials.gov,	Publication/study type (study registration)
Gucuk Adnan, Burgu Berk, Gokce Ilker, Mermerkaya Murat, and Soygur Tarkan (2013) Do antibiotic prophylaxis and/or circumcision change periurethral uropathogen colonization and urinary tract infection rates in boys with VUR?. Journal of pediatric urology 9(6 Pt B), 1131-6	Not a relevant study
Gupta K, and Trautner B W (2013) Diagnosis and management of recurrent urinary tract infections in non-pregnant women. BMJ (Online) 346(7910), f3140	Publication/study type (literature review)
Handeland Maria, Grude Nils, Torp Torfinn, and Slimestad Rune (2014) Black chokeberry juice (Aronia melanocarpa) reduces incidences of urinary tract infection among nursing home residents in the long terma pilot study. Nutrition research (New York, and N.Y.) 34(6), 518-25	Not a relevant study
Hari P, Sarin Y K, and Mathew J L (2014) Antimicrobial prophylaxis for children with vesicoureteral reflux. Indian Pediatrics 51(7), 571-574	Not relevant population
Hari Pankaj, Hari Smriti, Sinha Aditi, Kumar Rakesh, Kapil Arti, Pandey Ravindra Mohan, and Bagga Arvind (2015) Antibiotic prophylaxis in the management of vesicoureteric reflux: a randomized double-blind placebo-controlled trial. Pediatric nephrology (Berlin, and Germany) 30(3), 479-86	Not relevant population
Higgs R (2010) Pediatrics: Modest effect of prophylactic antibiotics on UTI in children. Nature Reviews Urology 7(1), 5	Publication/study type (commentary)
Hodson E M, Wheeler D M, Vimalchandra D, Smith G H, and Craig J C (2007) Interventions for primary vesicoureteric reflux. The Cochrane database of systematic reviews (3), CD001532	Not relevant population
Jepson RG, Mihaljevic L, and Craig J (2000) Cranberries for preventing urinary tract infections. The Cochrane database of systematic reviews (2), CD001321	Updated systematic review available
Jepson RG, Mihaljevic L, and Craig J (2001) Cranberries for preventing urinary tract infections. The Cochrane database of systematic reviews (3), CD001321	Updated systematic review available
Jepson RG, Mihaljevic L, and Craig J (2004) Cranberries for preventing urinary tract infections. The Cochrane database of systematic reviews (2), CD001321	Updated systematic review available
Jepson Ruth G, and Craig Jonathan C (2007) A systematic review of the evidence for cranberries and blueberries in UTI prevention. Molecular nutrition & food research 51(6), 738-45	Updated systematic review available
Jepson R G, and Craig J C (2008) Cranberries for preventing urinary tract infections. The Cochrane database of systematic reviews (1), CD001321	Updated systematic review available
Jodal Ulf, Smellie Jean M, Lax Hildegard, and Hoyer Peter F (2006) Ten-year results of randomized treatment of children with severe vesicoureteral reflux. Final report of the International	Not relevant population

Study reference	Reason for exclusion
Reflux Study in Children. Pediatric nephrology (Berlin, and	
Germany) 21(6), 785-92	
Juthani-Mehta Manisha, Van Ness, Peter H, Bianco Luann, Rink Andrea, Rubeck Sabina, Ginter Sandra, Argraves Stephanie, Charpentier Peter, Acampora Denise, Trentalange Mark, Quagliarello Vincent, and Peduzzi Peter (2016) Effect of Cranberry Capsules on Bacteriuria Plus Pyuria Among Older Women in Nursing Homes: A Randomized Clinical Trial. JAMA 316(18), 1879-1887	Not relevant population
LaPlante K L, Gill C M, and Rowley D (2017) Cranberry capsules for bacteriuria plus pyuria in nursing home residents. JAMA - Journal of the American Medical Association 317(10), 1078	Publication/study type (commentary)
Larcombe James (2015) Urinary tract infection in children: recurrent infections. BMJ clinical evidence 2015,	Publication/study type (Review of systematic reviews/RCTs)
Lee B B, Simpson J M, Craig J C, and Bhuta T (2007) Methenamine hippurate for preventing urinary tract infections. The Cochrane database of systematic reviews (4), CD003265	Not relevant population
Lee Linda C, Lorenzo Armando J, and Koyle Martin A (2016) The role of voiding cystourethrography in the investigation of children with urinary tract infections. Canadian Urological Association journal = Journal de l'Association des urologues du Canada 10(5-6), 210-214	Not a relevant study
Lee Seung Joo, Shim Yoon Hee, Cho Su Jin, and Lee Jung Won (2007) Probiotics prophylaxis in children with persistent primary vesicoureteral reflux. Pediatric nephrology (Berlin, and Germany) 22(9), 1315-20	Not relevant population
Lee Seung Joo, and Lee Jung Won (2015) Probiotics prophylaxis in infants with primary vesicoureteral reflux. Pediatric nephrology (Berlin, and Germany) 30(4), 609-13	Not relevant population
Ledda A, Bottari A, Luzzi R, Belcaro G, Hu S, Dugall M, Hosoi M, Ippolito E, Corsi M, Gizzi G, Morazzoni P, Riva A, Giacomelli L, and Togni S (2015) Cranberry supplementation in the prevention of non-severe lower urinary tract infections: a pilot study. European review for medical and pharmacological sciences 19(1), 77-80	Publication/study type (observational study)
Leo V, Cappelli V, Massaro Mg, Tosti C, and Morgante G (2017) Evaluation of the effects of a natural dietary supplement with cranberry, Noxamicina® and D-mannose in recurrent urinary infections in perimenopausal women. Minerva ginecologica 69(4), 336-341	Non-English language
Lo V, Wah Y, and Maggio L (2011) Antibiotic prophylaxis to prevent recurrent UTI in children. American Family Physician 84(2), 3-4	Publication/study type (commentary)
Long Elliot, Colquhoun Samantha, and Carapetis Jonathan R (2006) Antibiotic prophylaxis for the prevention of recurrent urinary tract infections in children. Advances in experimental medicine and biology 582, 243-9	Publication/study type (book article)
Lorenzo A J, and Braga L H. P (2013) Use of cranberry products does not appear to be associated with a significant reduction in incidence of recurrent urinary tract infections. Evidence-Based Medicine 18(5), 181-182	Publication/study type (commentary)
Mattoo Tej K (2007) Medical management of vesicoureteral refluxquiz within the article. Don't overlook placebos. Pediatric nephrology (Berlin, and Germany) 22(8), 1113-20	Not a relevant study

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Mattoo Tej K, Chesney Russell W, Greenfield Saul P, Hoberman	Not relevant population
Alejandro, Keren Ron, Mathews Ranjiv, Gravens-Mueller Lisa, Ivanova Anastasia, Carpenter Myra A, Moxey-Mims Marva, Majd Massoud, Ziessman Harvey A, and Investigators Rivur Trial (2016) Renal Scarring in the Randomized Intervention for Children with Vesicoureteral Reflux (RIVUR) Trial. Clinical journal of the American Society of Nephrology : CJASN 11(1), 54-61	Not relevant population
Mazokopakis Elias E, Karefilakis Christos M, and Starakis Ioannis K (2009) Efficacy of cranberry capsules in prevention of urinary tract infections in postmenopausal women. Journal of alternative and complementary medicine (New York, and N.Y.) 15(11), 1155	Publication/study type (commentary)
Mohseni Mohammad-Javad, Aryan Zahra, Emamzadeh-Fard Sahra, Paydary Koosha, Mofid Vahid, Joudaki Hasan, and Kajbafzadeh Abdol-Mohammad (2013) Combination of probiotics and antibiotics in the prevention of recurrent urinary tract infection in children. Iranian journal of pediatrics 23(4), 430-8	Not relevant population
Mutlu Hatice, and Ekinci Zelal (2012) Urinary tract infection prophylaxis in children with neurogenic bladder with cranberry capsules: randomized controlled trial. ISRN pediatrics 2012, 317280	Not relevant population
Naber Kurt G, Cho Yong-Hyun, Matsumoto Tetsuro, and Schaeffer Anthony J (2009) Immunoactive prophylaxis of recurrent urinary tract infections: a meta-analysis. International journal of antimicrobial agents 33(2), 111-9	Does not reflect usual UK practice
Nachum Z, Braverman M, Letova Ygz, Salim R, and Chazan B (2015) The effect of preventive antibiotic treatment in the postpartum period on urinary tract infection (UTI) rate in women treated during pregnancy for recurrent UTI e a prospective randomized controlled study. American journal of obstetrics and gynecology 212(1 suppl. 1), S399-s400	Abstract only
Nagler Evi Vt, Williams Gabrielle, Hodson Elisabeth M, and Craig Jonathan C (2011) Interventions for primary vesicoureteric reflux. The Cochrane database of systematic reviews (6), CD001532	Not a relevant population
Nct (2008) Prospective, randomized, double-blind, placebo- controlled study on parallel groups evaluating the efficacy and safety of cranberry (Vaccinium Macrocarpon) in prevention of urinary tract infections in multiple sclerosis patients. clinicaltrials.gov/ct2/show/NCT00280592,	Publication/study type (trial registration)
Nct (2008) Cranberry for UTI prevention in residents of long term care facilities (PACS). clinicaltrials.gov/ct2/show/NCT00596635,	Publication/study type (trial registration)
Nct, and Sumit D (2014) A Clinical Trial to Determine the Extent to Which Probiotic Therapy Reduces Side Effects of Antibiotic Prophylaxis in Pediatric Neurogenic Bladder Patients With a History of Recurrent Urinary Tract Infections. Http://clinicaltrials.gov/show/NCT02044965,	Publication/study type (trial registration)
Nelson Caleb P, Hoberman Alejandro, Shaikh Nader, Keren Ron, Mathews Ranjiv, Greenfield Saul P, Mattoo Tej K, Gotman Nathan, Ivanova Anastasia, Moxey-Mims Marva, Carpenter Myra A, and Chesney Russell W (2016) Antimicrobial Resistance and Urinary Tract Infection Recurrence. Pediatrics 137(4),	Not relevant population
Neveus Tryggve, Brandstrom Per, Linner Tina, Jodal Ulf, and Hansson Sverker (2012) Parental experiences and preferences regarding the treatment of vesicoureteral reflux. Scandinavian journal of urology and nephrology 46(1), 26-30	Not a relevant study

Study reference	Reason for exclusion
Nordenstrom Josefin, Holmdahl Gundela, Brandstrom Per, Sixt Rune, Stokland Eira, Sillen Ulla, and Sjostrom Sofia (2016) The Swedish infant high-grade reflux trial: Study presentation and vesicoureteral reflux outcome. Journal of pediatric urology,	Not relevant population
Nordenstrom J, Sillen U, Holmdahl G, Linner T, Stokland E, and Sjostrom S (2016) The Swedish Infant High-grade Reflux Trial - Bladder function. Journal of pediatric urology ,	Not a relevant study
Opperman E A (2010) Cranberry is not effective for the prevention or treatment of urinary tract infections in individuals with spinal cord injury. Spinal cord 48(6), 451-6	Not relevant population
Ostrovsky D A (2017) Cranberry Capsules do not Appear to Reduce Bacteriuria and Pyuria in Elderly Women Residing in Nursing Homes. Explore 13(3), 226-227	Publication/study type (literature review)
Perez-Gaxiola G (2011) Review: Antibiotic prophylaxis may not prevent recurrent symptomatic urinary tract infection in children. Archives of Disease in Childhood: Education and Practice Edition 96(5), 198	Abstract only
Pouwels Koen B, Visser Sipke T, and Hak Eelko (2013) Effect of pravastatin and fosinopril on recurrent urinary tract infections. The Journal of antimicrobial chemotherapy 68(3), 708-14	Poor relevance against search terms (interventions)
British Medical Journal Publishing Group (2013) Prevention of recurrent urinary tract infections in women. Drug and therapeutics bulletin 51(6), 69-72	Publication/study type (literature review)
Rego L L, Glazer C S, and Zimmern P E (2016) Risks of long- term use of nitrofurantoin for urinary tract prophylaxis in the older patient. Urological Science 27(4), 193-198	Publication/study type (literature review)
Salo J, Kontiokari T, Helminen M, Korppi M, Nieminen T, Pokka T, and Uhari M (2010) Randomized trial of cranberry juice for the prevention of recurrences of urinary tract infections in children. Clinical microbiology and infection 16(Suppl 2), S385-s386	Unable to source
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Sen Ayan (2006) Recurrent cystitis in non-pregnant women. Clinical evidence (15), 2558-64	Publication/study type (review of systematic reviews and RCTs)
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Shmuely H, Ofek I, Weiss EI, Rones Z, and Houri-Haddad Y (2012) Cranberry components for the therapy of infectious disease. Current opinion in biotechnology 23(2), 148-52	Not a clinical study
Stepanova N, Kruglikov V, Lebid L, and Kolesnyk M (2013) Oral lactobacilli vs antibiotic prophylaxis for recurrent urinary tract infections in premenopausal women. European Urology, and Supplements 12(1), e892	Publication/study type (literature review)

Study reference	Reason for exclusion
Sumukadas D, Davey P, and McMurdo M E. T (2009) Recurrent urinary tract infections in older people: The role of cranberry products. Age and Ageing 38(3), 255-257	Publication/study type (commentary)
Sung Jennifer, and Skoog Steven (2012) Surgical management of vesicoureteral reflux in children. Pediatric nephrology (Berlin, and Germany) 27(4), 551-61	Not relevant population
Takahashi S (2012) Prevention of acute uncomplicated cystitis by cranberry juice. International journal of urology 19, 410	Abstract only
Takvani A, Gokani C, and Malaviya P (2015) Vesicoureteric reflux-a prospective study of 11 years. European Urology, and Supplements 14(2), e505-e505a	Not a relevant study
Thomas J (2011) Cranberry juice fails to prevent recurring urinary tract infections. Australian Journal of Pharmacy 92(1092), 81	Abstract only
Uberos J, Rodrguez-Belmonte R, Fernndez-Puentes V, Narbona-Lpez E, Molina-Carballo A, and Munoz-Hoyos A (2010) Cranberry syrup vs. trimethoprim in the prophylaxis of recurrent urinary infection: A double-blind randomized clinical trial. Acta paediatrica 99(Suppl 462), 48	Abstract only
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Uehara Shinya, Monden Koichi, Nomoto Koji, Seno Yuko, Kariyama Reiko, and Kumon Hiromi (2006) A pilot study evaluating the safety and effectiveness of Lactobacillus vaginal suppositories in patients with recurrent urinary tract infection. International journal of antimicrobial agents 28 Suppl 1, S30-4	Abstract only
Vasileiou I, Katsargyris A, Theocharis S, and Giaginis C (2013) Current clinical status on the preventive effects of cranberry consumption against urinary tract infections. Nutrition research (New York, and N.Y.) 33(8), 595-607	Publication/study type (literature review)
Vicariotto Franco (2014) Effectiveness of an association of a cranberry dry extract, D-mannose, and the two microorganisms Lactobacillus plantarum LP01 and Lactobacillus paracasei LPC09 in women affected by cystitis: a pilot study. Journal of clinical gastroenterology 48 Suppl 1, S96-101	Publication/study type (observational study)
Vidlar A, Vostalova J, Vacek J, Kosina P, Vrbkova J, Ulrichova J, Student V, and Simanek V (2011) The effect of cranberry (Vaccini um macrocarpon) on the recurrence urinary tract infection in women. European Urology, and Supplements 10(9), 622	Abstract only
Vostalova Jitka, Vidlar Ales, Simanek Vilim, Galandakova Adela, Kosina Pavel, Vacek Jan, Vrbkova Jana, Zimmermann Benno F, Ulrichova Jitka, and Student Vladimir (2015) Are High Proanthocyanidins Key to Cranberry Efficacy in the Prevention of Recurrent Urinary Tract Infection?. Phytotherapy research : PTR 29(10), 1559-67	Publication/study type (literature review)
Wald E (2010) Antibiotic prophylaxis can prevent recurrent infection in children with urinary tract infections. Journal of Pediatrics 156(5), 856-857	Abstract only
Wan KS, Liu CK, Lee WK, Ko MC, and Huang CS (2016) Cranberries for Preventing Recurrent Urinary Tract Infections in Uncircumcised Boys. Alternative therapies in health and medicine 22(6), 20-23	Not relevant intervention

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Williams GJ, Lee A, and Craig JC (2001) Long-term antibiotics for preventing recurrent urinary tract infection in children. The Cochrane database of systematic reviews (4), CD001534	Updated systematic review available
Williams GJ, Wei L, Lee A, and Craig JC (2006) Long-term antibiotics for preventing recurrent urinary tract infection in children. The Cochrane database of systematic reviews (3), CD001534	Updated systematic review available
Williams GJ, Craig JC, and Carapetis JR (2013) Preventing urinary tract infections in early childhood. Advances in experimental medicine and biology 764, 211-8	Publication/study type (literature review)

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