National Institute for Health and Care Excellence

Final

Renal and ureteric stones: assessment and management

Imaging for diagnosis

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Final

This evidence review was developed by the National Guideline Centre



FINAL

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Contents

1	Imag	jing for	diagnosis of renal and ureteric stones	5
	1.1	and ur MRI to	v question: In people with suspected (or under investigation for) renal eteric stones, how accurate is ultrasound, plain abdominal radiograph or identify whether a renal or ureteric stone is present, as indicated by the nee standard, non-contrast CT?	5
	1.2	Introdu	uction	5
	1.3	PICO	able	5
	1.4	Clinica	I evidence	6
		1.4.1	Included studies	6
		1.4.2	Excluded studies	6
		1.4.3	Summary of clinical studies included in the evidence review	7
		1.4.4	Quality assessment of clinical studies included in the evidence review \ldots	. 10
	1.5	Econo	mic evidence	. 13
		1.5.1	Included studies	. 13
		1.5.2	Excluded studies	. 13
		1.5.3	Unit costs	. 13
	1.6	Resou	rce costs	. 14
	1.7	Evider	ice statements	. 14
		1.7.1	Clinical evidence statements	. 14
		1.7.2	Health economic evidence statements	. 14
		1.8.1	Interpreting the evidence	. 14
Ref	ferenc	:es		. 18
Арј	pendi	ces		. 29
	Appe	endix A:	Review protocols	. 29
	Арре	endix B:	Literature search strategies	. 32
	Арре	endix C:	Clinical evidence selection	. 42
	Арре	endix D:	Clinical evidence tables	. 43
	Appe	endix E:	Coupled sensitivity and specificity forest plots and sROC curves	. 65
	Арре	endix F:	Health economic evidence selection	. 68
	Арре	endix G:	Health economic evidence tables	. 69
	Арре	endix H:	Excluded studies	. 70

1 Imaging for diagnosis of renal and ureteric stones

1.1 Review question: In people with suspected (or under investigation for) renal and ureteric stones, how accurate is ultrasound, plain abdominal radiograph or MRI to identify whether a renal or ureteric stone is present, as indicated by the reference standard, non-contrast CT?

1.2 Introduction

Imaging which provides an accurate and timely diagnosis of a stone in a patient presenting with acute renal colic is essential to manage the patient in the most appropriate way. An accurate diagnosis is essential as the site and size of the stone and anatomical features of the patient are important in defining the most appropriate treatment options. There are a variety of imaging modalities used to assess patients with suspected renal colic including ultra sound, CT scanning with radiation and MRI scanning. CT is more expensive than ultrasound but the extra cost may be outweighed by avoiding additional investigations if the first test misses the diagnosis. There is uncertainty about which imaging modality should be the first line investigation in the acute setting of suspected renal colic. Similarly there are concerns regarding radiation doses in certain groups, children and pregnant women and the question will address the most suitable imaging test in these groups.

1.3 PICO table

For full details see the review protocol in appendix A.

Population	People (adults, children and young people) with suspected (or under investigation for) renal and ureteric stones				
Target condition	Renal and ureteric stones				
Index tests	 Plain abdominal radiograph (conventional, KUB) Ultrasound Magnetic resonance imaging 				
Reference standard	Non contrast computed tomography				
Statistical measures	 Specificity Sensitivity Positive Predictive Value Negative Predictive Value 				
Study design	Cross-sectional studies, cohort studies Case–control and case series studies should be included only if there is no other evidence				

Table 1:	PICO	characteristics	of review	question
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1.4 Clinical evidence

1.4.1 Included studies

A search was conducted for prospective and retrospective cohort studies assessing the diagnostic test accuracy of ultrasound, MRI or plain abdominal radiograph to identify whether the condition is present (as indicated by the reference standard) in people under investigation for renal and ureteric stones.

Thirteen studies were included in the review;^{16, 18, 20, 42, 52-54, 57, 63, 95, 105, 119, 124} these are summarised in Table 2 below. Evidence from these studies is summarised in the clinical evidence summary below (Table 3).

See also the study selection flow chart in appendix C, sensitivity and specificity forest plots and receiver operating characteristics (ROC) curves in appendix E, and study evidence tables in appendix D.

1.4.2 Excluded studies

See the excluded studies list in appendix H.

4.3 Summary of clinical studies included in the evidence review

Table 2: Summary of studies included in the evidence review

Study	Population	Target condition	Index test	Reference standard	Comments
Chan 2008 ¹⁶	n=100 People being investigated for suspected urolithiasis Ireland	Urolithiasis	Plain abdominal radiograph (KUB)	Unenhanced CT	Diagnosing number of patients with stones
Cifci 2016 ¹⁸	n=159 People who were admitted to the urology department with suspected acute urinary calculi (flank pain, hematuria or patients with a history of urinary calculi) Turkey	Ureteral calculi	MRI (MRU with a B- TFE sequence)	Unenhanced CT	Diagnosing number of patients with stones Calculi located in the kidneys and the bladder were not documented. Only calculi located in ureters were documented. Reports results from two independent observers
de Souza 2007 ²⁰	n=52 People referred for evaluation of acute renal colic Brazil	Ureterolithiasis	US	Non-contrast helical CT	Diagnosing number of patients with stones

Study	Population	Target condition	Index test	Reference standard	Comments
Haroun 2010 ⁴²	n=156 People who underwent UHCT scan and US for suspicion of urolithiasis Jordan	Renal stones Ureteral stones	US (B-mode)	Unenhanced CT	Diagnosing number of patients with stones Study reports sensitivity and specificity for all stones (including urinary bladder) or for renal and ureteric stones separately. Currently extracted all stones as no raw data is reported for separate types of stone.
Kanno 2017 ⁵²	n=822 People with acute flank pain, hematuria, or a history of urinary stones who had KUB, US and NCCT on the same day Japan	Renal stones	US (greyscale) Plain abdominal radiograph (KUB)	Non-contrast CT	Diagnosing number of kidneys with stones
Kanno 2014a ⁵⁴	n=428 People with acute flank pain, hematuria, or a history of urinary stones who had US and NCCT on the same day Japan	Renal stones	US	Non-contrast CT	Diagnosing number of kidneys with stones Study reports results for 'individual stone' and for 'specific stone'. Currently extracted 'individual stone' data. The study includes 856 kidneys, but in the results there are only 853.

Study	Population	Target condition	Index test	Reference standard	Comments
Kanno 2014b ⁵³	n=428 People with symptoms such as acute flank pain or hematuria Japan	Ureteral stones	US	Non-contrast CT	Diagnosing number of ureters with stones
Kielar 2012 ⁵⁷	n=51 People with flank pain Canada	Urolithiasis	US (greyscale)	Unenhanced CT	Diagnosing number of stones
Levine 1997 ⁶³	n=152 People with acute flank pain who had a CT within 4 hours of plain radiography USA	Ureteral stones	Plain abdominal radiography	Unenhanced helical CT	Diagnosing number of patients with stones
Passerotti 2009 ⁹⁵	n=50 Children who had signs, symptoms or a history suggestive of urolithiasis who had US and CT within 0.5-8 hours of each other	Nephrolithiasis/urinary stones	US	Non-contrast CT	Diagnosing number of patients with stones 24% of participants wer asymptomatic at presentation but were evaluated because of a history of urolithiasis

Study	Population	Target condition	Index test	Reference standard	Comments
	USA				
Resorlu 2015 ¹⁰⁵	n=500 People with acute flank pain who had CT and US within 10 days Turkey	Urinary stones (kidney and ureter)	US (grey scale)	Non-contrast CT	Diagnosing number of patients with stones
Semins 2013 ¹¹⁹	n=22 People with suspected acute ureteral calculus USA	Obstructing stones	MRI (MRU (non- contrast HASTE [Half- Fourier single shot turbo spin-echo]))	Non-contrast CT	Diagnosing number of patients with stones
Sternberg 2016 ¹²⁴	n=155 People with suspected renal colic who had US and CT within 1 day Lebanon	Urinary calculi	Renal US	Non-contrast CT	Diagnosing number of patients with stones

See appendix D for full evidence tables.

1.4.4 Quality assessment of clinical studies included in the evidence review

Table 3: Clinical evidence summary: diagnostic test accuracy for Ultrasound, plain abdominal radiograph and MRI in adults						
Index Test (Threshold)	Number of studies	n	Quality	Sensitivity % (95% CI)	Specificity % (95% CI)	
<u>Ultrasound</u>						
Ultrasound	7	4189	VERY LOW ^{a,b,d} due to risk of bias, very serious inconsistency and very serious imprecision	Pooled ^e : 0.60 (0.38, 0.79)	Pooled ^e : 0.90 (0.79, 0.97)	
Plain abdominal radiogram	<u>oh</u>					
Plain abdominal radiograph	3	1895	VERY LOW ^{a,d} due to risk of bias, very serious imprecision	Pooled ^e : 0.58 (0.29, 0.58)	Pooled ^e : 0.90 (0.41, 1.00)	
MRI						
MRI	3	159	LOW ^{a,d} due to risk of bias, serious imprecision	0.66 (0.55, 0.76) ^f	0.96 (0.89, 0.99) ^f	
		159	MODERATE ^a due to risk of bias	0.72 (0.61, 0.81) ^f	1.00 (0.95, 1.00) ^f	
		22	LOW ^{a,d} due to risk of bias, serious imprecision	0.84 (0.60, 0.97) ^f	1.00 (0.29, 1.00) ^f	

Table 3: Clinical evidence summary: diagnostic test accuracy for Ultrasound, plain abdominal radiograph and MRI in adults

The assessment of the evidence quality was conducted with emphasis on sensitivity as this was identified by the committee as the primary measure in guiding decisionmaking. The committee set the sensitivity threshold at 95% as the acceptable level to recommend a test.

(a) Risk of bias was assessed using the QUADAS-2 checklist. The evidence was downgraded by 1 increment if the majority of studies were rated at high risk of bias, and downgraded by 2 increments if the majority of studies were rated at very high risk of bias.

(b) Inconsistency was assessed by inspection of the sensitivity and specificity plots

(c) Indirectness was assessed using the QUADAS-2 checklist items referring to applicability. The evidence was downgraded by 1 increment if the majority of studies were seriously indirect, and downgraded by 2 increments if the majority of studies are very seriously indirect

(d) Imprecision was assessed based on inspection of the confidence region of sensitivity in the diagnostic meta-analysis or, where diagnostic meta-analysis has not been conducted, assessed according to the range of confidence intervals in the individual studies. The evidence was downgraded by 1 increment when the range of the confidence interval around the point estimate was 20–40%, and downgraded by 2 increments when there was a range of >40%

(e) Pooled sensitivity/specificity from diagnostic meta-analysis

(f) Could not be pooled due to insufficient data. Individual sensitivity values and their coupled specificity is presented.

Table 4: Clinical evidence summary: diagnostic test accuracy for Ultrasound, plain abdominal radiograph and MRI in children

Index Test (Threshold)	Number of studies	n	Quality	Specificity % (95% Cl)	Sensitivity % (95% CI)
Ultrasound					
Ultrasound	1	52	LOW ^{a,d} due to risk of bias, serious imprecision	1.00 (0.79, 1.00)	0.76 (0.59, 0.89)

The assessment of the evidence quality was conducted with emphasis on sensitivity as this was identified by the committee as the primary measure in guiding decisionmaking. The committee set the sensitivity threshold at 95% as the acceptable level to recommend a test.

(a) Risk of bias was assessed using the QUADAS-2 checklist. The evidence was downgraded by 1 increment if the majority of studies were rated at high risk of bias, and downgraded by 2 increments if the majority of studies were rated at very high risk of bias.

(b) Inconsistency was assessed by inspection of the sensitivity and specificity plots

(c) Indirectness was assessed using the QUADAS-2 checklist items referring to applicability. The evidence was downgraded by 1 increment if the majority of studies were seriously indirect, and downgraded by 2 increments if the majority of studies are very seriously indirect

(d) Imprecision was assessed based on inspection of the confidence region of sensitivity in the diagnostic meta-analysis or, where diagnostic meta-analysis has not been conducted, assessed according to the range of confidence intervals in the individual studies. The evidence was downgraded by 1 increment when the range of the confidence interval around the point estimate was 20–40%, and downgraded by 2 increments when there was a range of >40%

5

1.5 Economic evidence

1.5.1 Included studies

No relevant health economic studies were identified.

1.5.2 Excluded studies

No health economic studies that were relevant to this question were excluded due to assessment of limited applicability or methodological limitations.

See also the health economic study selection flow chart in appendix F.

1.5.3 Unit costs

Relevant unit costs are provided below to aid consideration of cost effectiveness.

Diagnostic imaging	Detail	Unit cost
Plain abdominal radiograph	Direct access plain film Currency code: DAPF	£29.78
Ultrasound	Ultrasound Scan with duration of less than 20 minutes, without Contrast Currency code: RD40Z	£51.59
Computerised Tomography (CT)	<u>Adults:</u> Computerised Tomography Scan of One Area, without Contrast, 19 years and over Currency code: RD20A	£85.56
	<u>Children:</u> Computerised Tomography Scan of One Area, without Contrast, between 6 and 18 years	£91.67
	Currency code: RD20B Computerised Tomography Scan of One Area, without Contrast, 5 years and under Currency code: RD20C	£94.72
Magnetic Resonance Imaging (MRI)	<u>Adults:</u> Magnetic Resonance Imaging Scan of One Area, without Contrast, 19 years and over Currency code: RD01A	£138.24
	<u>Children:</u> Magnetic Resonance Imaging Scan of One Area, without Contrast, between 6 and 18 years	£135.88
	Currency code: RD01B Magnetic Resonance Imaging Scan of One Area, without Contrast, 5 years and under Currency code: RD01C	£160.59

Table 5: UK costs of diagnostic imaging techniques

Source: NHS reference costs 2016/17⁸⁷.

1.6 Resource costs

The recommendations made by the committee based on this review (see section **Error!** eference source not found.) are not expected to have a substantial impact on resources.

1.7 Evidence statements

1.7.1 Clinical evidence statements

Thirteen studies that evaluated 3 diagnostic tests were included in the review. Of these, the committee noted that all tests demonstrated poor sensitivity for identifying renal and ureteric stones. The evidence was of Moderate to Very Low quality. Evidence was identified for the following diagnostic tests: ultrasound, plain abdominal radiograph, MRI.

- **Ultrasound**: Low quality evidence from 1 study with 52 children with suspected stones showed that ultrasound has a sensitivity of 76% and a specificity of 100%. Very Low quality evidence from 7 studies with 4189 adults with suspected stones showed that ultrasound has a sensitivity of 60% and a specificity of 90%.
- **Plain abdominal radiograph**: Very Low quality from 3 studies with 1895 adults with suspected stones demonstrated that plain abdominal radiograph had a sensitivity of 58% and a specificity of 90.
- **MRI**: Three studies investigated the diagnostic accuracy of MRI, however they were unable to be pooled due to insufficient data. Moderate quality evidence from one study of 159 adults showed that the MRI has a sensitivity and specificity respectively of 72% and 100%. Low quality evidence from 1 study with 159 adults with suspected stones showed that the MRI has a sensitivity and specificity respectively of 66% and 96%. Low quality evidence from 1 study with suspected stones showed that the MRI has a sensitivity and specificity respectively of 66% and 96%. Low quality evidence from 1 study with suspected stones showed that the MRI has a sensitivity and specificity respectively of 84% and 100%.

1.7.2 Health economic evidence statements

• No relevant economic evaluations were identified.

1.8 The committee's discussion of the evidence

1.8.1 Interpreting the evidence

1.8.1.1 The diagnostic measures that matter most

Diagnostic accuracy for renal and ureteric stones was the outcome prioritised for this review. Sensitivity was considered the most important measure by the committee for this review question because it was considered that false positives are rare and not hugely problematic. The consequences of missing a patient with a renal or ureteric stone could include being sent for further imaging or investigations and a delay in treatment, potentially resulting in damage to the kidney.

1.8.1.2 The quality of the evidence

The evidence for ultrasound in adults was very low quality, due to very serious imprecision and very serious inconsistency, and in children the evidence was low quality due to imprecision. There was also a risk of bias for both populations, due to uncertainty regarding whether the results of the index or reference standard tests were interpreted without knowledge of the other test, and uncertainty regarding participants excluded from the analysis. The evidence for plain abdominal radiograph was low quality, due to very serious imprecision and a high risk of bias. The evidence for MRI ranged from low to moderate quality, due to risk of bias and imprecision. The committee noted a number of other factors that varied between the studies, such as the amount of time between the index and reference standard test, and the expertise of the clinician interpreting the test results.

1.8.1.3 Benefits and harms

Evidence for adults and children and young people, and for those with symptomatic and asymptomatic suspected stones was searched for, however no evidence was identified where the majority of the population was people with a suspected asymptomatic stone. The committee agreed that the recommendations should only apply to those who were symptomatic of a renal or ureteric stone.

The committee noted that no mode of imaging met the pre-specified threshold for sensitivity, which was set at 95%, and therefore concluded that none of the imaging modalities were as effective as the reference standard; non-contrast CT. The committee considered the consequences of a low sensitivity, and noted that this could include being sent for other imaging or investigations, and potentially a delay to treatment. The committee noted that there are risks associated with CT such as the exposure to radiation, however considered that this did not outweigh the risks associated with not diagnosing a stone.

On the other hand it was discussed how there may be some groups where the radiation risk is a concern; such as women who may be pregnant, in which case ultrasound is current practice in place of CT. The committee discussed the timing of CT imaging for diagnosis, and noted that access to CT is not currently the same across the country. It was also noted that there are harms associated with not carrying out imaging urgently, such as delay to diagnosis, and delay to treatment which may increase the risk of deterioration in renal function. Therefore, the committee agreed that based on clinical expertise and opinion of the committee, CT imaging should be done within 24 hours. The committee highlighted that CT should only be offered for those with suspected renal colic, rather than any abdominal pain without additional indicators or reasons for suspicion of renal colic. Other imaging modalities may be more appropriate where renal colic is not the suspected cause of abdominal pain.

The committee considered that for the paediatric population, there was no evidence for MRI or plain abdominal radiograph, and the evidence for ultrasound suggested that it did not meet the pre-specified threshold for sensitivity. The committee noted that in current UK practice, ultrasound is often used as the first line method of diagnosis. They considered the benefits of ultrasound include no dose of ionising radiation; however it is not as sensitive as CT and is open to wide operator variation. The committee discussed the risks associated with using CT in young children concerning the radiation, such as the increased lifetime malignancy risk, and considered that this was a very serious and potentially severe harm. The committee also discussed that renal and ureteral colic is often lower on the differential diagnosis list in children compared to adults, and presentation is much more commonly atypical. Therefore, they agreed that it was important not to increase the amount of unnecessary CT's being carried out, given that they are associated with harms. Based on this the committee agreed that US should be offered as first line imaging, and that low-dose non-contrast CT should be considered only when there is diagnostic uncertainty following US.

1.8.1.4 Cost effectiveness and resource use

No economic evidence was identified for this question.

The committee were presented with the costs of the different imaging techniques. MRI is the most costly, followed by CT, ultrasound, and plain abdominal radiograph.

A test with a low sensitivity will miss people and create a lot of false negatives. Poor specificity will result in more people being diagnosed as having stones when they do not (false positives). The implications of low sensitivity would be that people's condition could get worse as they have not been diagnosed as having renal stones, which could result in more

emergency care or higher risks, and require further investigation. The implication of low specificity would be unnecessary management that the patient doesn't need, and delays identification of the true underlying condition.

The reference standard is a CT. This is one of the more expensive tests. MRI is the most expensive test, and as it is less effective than CT (because CT is the reference standard and therefore assumed to be 100% accurate); this makes MRI a dominated alternative.

Plain abdominal radiograph or ultrasound are also assumed to be by default less accurate than CT. This has to be traded off against their lower cost.

In adults, the sensitivity of ultrasound was 0.6 and specificity was 0.9. Assuming a prevalence of renal stones of 60% in a population being imaged because of pain, then if 1000 people are imaged using ultrasound, this will mean out of 600 that have a stone, only 360 are correctly identified as having a stone, and 240 will be false negatives (i.e. missed people with renal stones). There will also be 40 false positives.

Plain abdominal radiograph had similar sensitivity of 0.58 and specificity of 0.9. Using the same assumptions as above, this would lead to only 348 people identified as correctly having a stone, 252 false negatives and 40 false positives.

The cost of the different types of imaging over a cohort of 1000 people would be around £86,000 for CT, £52,000 for ultrasound, and £30,000 for plain abdominal radiograph. Spreading the cost difference between CT and ultrasound, and CT and plain abdominal radiograph, over the individuals correctly identified as having stones, would lead to a cost per correct stone diagnosis of £143 for both CT and ultrasound, and £86 per correct diagnosis for plain abdominal radiograph. Those who were missed with the less accurate techniques of ultrasound and plain abdominal radiograph will consume further resources, as they

will be diagnosed at a later point, which means more imaging/tests, and any GP attendances/hospital admissions because they have been misdiagnosed and still have a stone, or potential adverse events because of the delay in diagnosis. Therefore, it could cost more in the long run to use a lower cost technique initially, because of the lower accuracy. These calculations didn't consider the false positives who would also face delay in achieving their real diagnosis and consume unnecessary resources.

There is also the impact on quality of life to consider from being misdiagnosed.

Additionally, more accurate techniques such as CT can have other benefits unrelated to the condition in question. For example, it was committee opinion that there are around 10% incidental findings such as abdominal aneurysm, which is a life threatening condition if missed.

The committee felt that the evidence had not shown that any other technique was as good as CT, based on their pre-specified threshold for sensitivity of 95%. CT is generally already current practice for diagnosing renal stones in adults, and therefore wasn't considered likely to have a large resource impact. The committee felt that the costs of misdiagnosing people were likely to outweigh the additional cost of undertaking CT over other techniques, and there were also other benefits like incidental findings or other diagnoses with similar presentations to acute renal colic (for example a leaking abdominal aortic aneurysm). Therefore a recommendation was made to offer CT in adults with suspected renal colic. The urgency with which the CT should take place was debated, as although immediately would be ideal, this may not always be feasible in all locations particularly out of hours, and some hospitals would ask someone to come back the following day. So although CT is generally the gold standard for diagnosis of renal stones and this is established practice, how quickly this takes place can be variable. The committee felt that by specifying 'urgent' and outlining that this means within 24 hours, would allow some flexibility in hospitals where this cannot happen within a few hours.

It was discussed whether the recommendation would lead to an increase in referrals from primary care/the community if CT needs to be offered within 24 hours to diagnose renal colic. The committee felt strongly that CT should be the gold standard method for diagnosing renal colic, in which case it may be that those previously presenting in the community with renal colic still had imaging, but on an outpatient/community basis. In which case it may be the

same number of patients being imaged, but the imaging is being brought forward to within a shorter time frame. However, there may be an increase in referrals if some people were not being imaged, but managed in the community. The additional impact is uncertain.

It is important to make clear that the recommendation is specifying a population with suspected renal colic, which means this suspicion (based on history and clinical examination) has to be in place clinically, and this would be a subset of people presenting to hospital with abdominal/flank pain in general.

For children, there was only evidence for ultrasound, which showed a sensitivity of 0.76 compared to CT. There is more caution around imaging with a higher radiation burden in children because of their age and the cumulative effect of imaging over their lifetime. Because of these concerns, a recommendation was made for ultrasound as first line imaging and low dose non-contrast CT, if there is still uncertainty around the diagnosis and there is a high degree of suspicion for renal colic. This is in keeping with current practice where ultrasound is more likely to be first line imaging. The paediatric population suspected with renal colic is likely to be small as renal stones in this population itself is very small.

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Appendices

Appendix A: Review protocols

Table 6:	Review protocol: What is the clinical and cost effectiveness of performing
	imaging for diagnosing renal and ureteric stones?

agnosing renal and ureteric stones?
Content
What is the clinical and cost effectiveness of performing imaging for diagnosing renal and ureteric stones?
Diagnostic review A review of health economic evidence related to the same review question was conducted in parallel with this review. For details see the health economic review protocol for this NICE guideline.
To evaluate the diagnostic test accuracy of imaging techniques in diagnosing renal and ureteric stones.
People (adults, children and young people) with suspected (or under investigation for) renal and ureteric stones
 Plain abdominal radiograph (conventional, KUB) Ultrasound Magnetic resonance imaging
Non contrast computed tomography
 Specificity Sensitivity Positive Predictive Value Negative Predictive Value
Cross-sectional studies, cohort studies, Case–control and case series studies should be included only if there is no other evidence (as they are biased).
Bladder stones Open surgery for renal (kidney and ureteric) stones Laparoscopic nephrolithotomy and pyelolithotomy Non-English language studies
 Strata: Adults (≥16 years) Children and young people (<16 years) Pregnant women
Studies are sifted by title and abstract. Potentially significant publications obtained in full text are then assessed against the inclusion criteria specified in this protocol.
 Sensitivity and specificity are calculated using Cochrane Review Manager (RevMan5). Diagnostic meta-analyses are conducted using WinBUGS14 and graphically presented using RevMan5. Endnote for bibliography, citations, sifting and reference management

Information sources – databases and dates	Clinical search databases to be used: Medline, Embase, Cochrane Library Date: all years Health economics search databases to be used: Medline, Embase, NHSEED, HTA Date: Medline, Embase from 2014 NHSEED, HTA – all years Language: Restrict to English only Supplementary search techniques: backward citation searching Key papers: Not known
Identify if an update	Not applicable
Author contacts	https://www.nice.org.uk/guidance/indevelopment/gid-ng10033
Highlight if amendment to previous protocol	For details please see section 4.5 of Developing NICE guidelines: the manual.
Search strategy – for one database	For details please see appendix B
Data collection process – forms / duplicate	A standardised evidence table format will be used, and published as appendix D of the evidence report.
Data items – define all variables to be collected	For details please see evidence tables in Appendix D (clinical evidence tables) or H (health economic evidence tables).
Methods for assessing bias at outcome / study level	Standard study checklists are used to critically appraise individual studies. For details please see section 6.2 of Developing NICE guidelines: the manual. The risk of bias is evaluated for each outcome on a study using the QUADAS-2 checklist.
Criteria for quantitative synthesis	For details please see section 6.4 of Developing NICE guidelines: the manual.
Methods for quantitative analysis – combining studies and exploring (in)consistency	For details please see the separate Methods report for this guideline.
Meta-bias assessment – publication bias, selective reporting bias	For details please see section 6.2 of Developing NICE guidelines: the manual. [Consider exploring publication bias for review questions where it may be more common, such as pharmacological questions, certain disease areas, etc. Describe any steps taken to mitigate against publication bias, such as examining trial registries.]
Confidence in cumulative evidence	For details please see sections 6.4 and 9.1 of Developing NICE guidelines: the manual.
Rationale / context – what is known	For details please see the introduction to the evidence review.
Describe contributions of authors and guarantor	A multidisciplinary committee developed the evidence review. The committee was convened by the National Guideline Centre (NGC) and chaired by Andrew Dickinson in line with section 3 of Developing NICE guidelines: the manual. Staff from NGC undertook systematic literature searches, appraised the evidence, conducted meta-analysis and cost-effectiveness analysis where appropriate, and drafted the evidence review in collaboration with the committee. For details please see Developing NICE guidelines: the manual.

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Sources of funding / support	NGC is funded by NICE and hosted by the Royal College of Physicians.
Name of sponsor	NGC is funded by NICE and hosted by the Royal College of Physicians.
Roles of sponsor	NICE funds NGC to develop guidelines for those working in the NHS, public health and social care in England.
PROSPERO registration number	Not registered

Table 7: Health economic review protocol

Review question	All questions – health economic evidence
Objective s	To identify economic studies relevant to any of the review questions.
Search criteria	 Populations, interventions and comparators must be as specified in the individual review protocol above. Studies must be of a relevant economic study design (cost-utility analysis, cost-effectiveness analysis, cost-benefit analysis, cost-consequences analysis, comparative cost analysis). Studies must not be a letter, editorial or commentary, or a review of economic evaluations. (Recent reviews will be ordered although not reviewed. The bibliographies will be checked for relevant studies, which will then be ordered.) Unpublished reports will not be considered unless submitted as part of a call for evidence. Studies must be in English.
Search strategy	An economic study search will be undertaken using population-specific terms and an economic study filter – see Appendix G [in the Full guideline].
Review strategy	 Studies not meeting any of the search criteria above will be excluded. Studies published before 2002, abstract-only studies and studies from non-OECD countries or the USA will also be excluded. Each remaining study will be assessed for applicability and methodological limitations using the NICE economic evaluation checklist which can be found in Appendix G of the 2014 NICE guidelines manual.⁸⁶ Inclusion and exclusion criteria If a study is rated as both 'Directly applicable' and with 'Minor limitations' then it will be included in the guideline. An economic evidence table will be completed and it will be included in the economic evidence profile. If a study is rated as either 'Not applicable' or with 'Very serious limitations' then it will usually be excluded from the guideline. If it is excluded then an economic evidence table will not be completed and it will not be included in the economic evidence profile. If a study is rated as 'Partially applicable', with 'Potentially serious limitations' or both then there is discretion The health economist will make a decision based on the relative applicability and quality of the available evidence for that question, in discussion with the Committee if required. The ultimate aim is to include economic studies that are helpful for decision-making in the context of the guideline and the current NHS setting. If several studies are considered of sufficiently high applicability and methodological quality that they could all be included, then the health economist, in discussion with the Committee if required, may decide to include only the most applicability and to selectively

methodological limitations will be listed with explanation as excluded economic studies in Appendix M.

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

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

Economic study type:

- Cost-utility analysis (most applicable).
- Other type of full economic evaluation (cost-benefit analysis, cost-effectiveness analysis, cost-consequences analysis).
- Comparative cost analysis.
- Non-comparative cost analyses including cost-of-illness studies will have been excluded before being assessed for applicability and methodological limitations. *Year of analysis:*
- The more recent the study, the more applicable it will be.
- Studies published in 2002 or later but that depend on unit costs and resource data entirely or predominantly from before 2002 will be rated as 'Not applicable'.
- Studies published before 2002 will have been excluded before being assessed for applicability and methodological limitations.

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

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

Appendix B: Literature search strategies

The literature searches for this review are detailed below and complied with the methodology outlined in Developing NICE guidelines: the manual 2014, updated 2017 https://www.nice.org.uk/guidance/pmg20/resources/developing-nice-guidelines-the-manual-pdf-72286708700869

For more detailed information, please see the Methodology Review. [Add cross reference]

B.1 Clinical search literature search strategy

Searches were constructed using a PICO framework where population (P) terms were combined with Intervention (I) and in some cases Comparison (C) terms. Outcomes (O) are rarely used in search strategies for interventions as these concepts may not be well described in title, abstract or indexes and therefore difficult to retrieve. Search filters were applied to the search where appropriate.

Database	Dates searched	Search filter used
Medline (OVID)	1946 – 29 November 2017	Exclusions Randomised controlled trials Systematic review studies Observational studies

Table 8: Database date parameters and filters used

Database	Dates searched	Search filter used
		Diagnostic tests studies
Embase (OVID)	1974 – 29 November 2017	Exclusions Randomised controlled trials Systematic review studies Observational studies Diagnostic tests studies
The Cochrane Library (Wiley)	Cochrane Reviews to 2017 Issue 11 of 12 CENTRAL to 2017 Issue 10 of 12 DARE, and NHSEED to 2015 Issue 2 of 4 HTA to 2016 Issue 4 of 4	None

Medline (Ovid) search terms

1.	exp urolithiasis/
2.	(nephrolitiasis or nephrolith or nephroliths or urolithias?s or ureterolithias?s).ti,ab.
3.	((renal or kidney* or urinary or ureter* or urethra*) adj3 (stone* or calculi or calculus or calculosis or lithiasis or c?olic*)).ti,ab.
4.	stone disease*.ti,ab.
5.	((calculi or calculus or calcium oxalate or cystine) adj3 (crystal* or stone* or lithiasis)).ti,ab.
6.	or/1-5
7.	letter/
8.	editorial/
9.	news/
10.	exp historical article/
11.	Anecdotes as Topic/
12.	comment/
13.	case report/
14.	(letter or comment*).ti.
15.	or/7-14
16.	randomized controlled trial/ or random*.ti,ab.
17.	15 not 16
18.	animals/ not humans/
19.	exp Animals, Laboratory/
20.	exp Animal Experimentation/
21.	exp Models, Animal/
22.	exp Rodentia/
23.	(rat or rats or mouse or mice).ti.
24.	or/17-23
25.	6 not 24
26.	limit 25 to English language
27.	exp Tomography/
28.	tomograph*.ti,ab.
29.	(NCCT or CT or UHCT).ti,ab.
30.	((CAT or body) adj2 scan*).ti,ab.

31.	or/27-30
32.	Radiography/
33.	Radiography, Abdominal/
34.	Urography/
35.	(radiograph* or x ray* or xray* KUB or urograph*).ti,ab.
36.	or/32-35
37.	Ultrasonography/
38.	(ultrasonograph* or ultrasound or ultrasonic or sonograph* or echograph* or echotomograph*).ti,ab.
39.	(ultra adj2 (sound or sonic)).ti,ab.
40.	(sound* adj2 (wave* or frequenc*)).ti,ab.
41.	(US adj3 imag*).ti,ab.
42.	or/37-41
43.	Magnetic Resonance Imaging/
44.	((magnetic or nuclear) adj2 resonance adj3 imag*).ti,ab.
45.	(MRI or NMR or NMRI or fMRI or MR).ti,ab.
46.	or/43-45
47.	31 or 36 or 42 or 46
48.	26 and 47
49.	exp "sensitivity and specificity"/
50.	(sensitivity or specificity).ti,ab.
51.	((pre test or pretest or post test) adj probability).ti,ab.
52.	(predictive value* or PPV or NPV).ti,ab.
53.	likelihood ratio*.ti,ab.
54.	likelihood function/
55.	((area under adj4 curve) or AUC).ti,ab.
56.	(receive* operat* characteristic* or receive* operat* curve* or ROC curve*).ti,ab.
57.	(diagnos* adj3 (performance* or accurac* or utilit* or value* or efficien* or effectiveness)).ti,ab.
58.	gold standard.ab.
59.	or/49-58
60.	randomized controlled trial.pt.
61.	controlled clinical trial.pt.
62.	randomi#ed.ti,ab.
63.	placebo.ab.
64.	randomly.ti,ab.
65.	Clinical Trials as topic.sh.
66.	trial.ti.
67.	or/60-66
68.	Meta-Analysis/
69.	exp Meta-Analysis as Topic/
70.	(meta analy* or metanaly* or metaanaly* or meta regression).ti,ab.
71.	((systematic* or evidence*) adj3 (review* or overview*)).ti,ab.
72.	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.

73.	(search strategy or search criteria or systematic search or study selection or data extraction).ab.
74.	(search* adj4 literature).ab.
75.	(medline or pubmed or cochrane or embase or psychlit or psyclit or psychinfo or psycinfo or cinahl or science citation index or bids or cancerlit).ab.
76.	cochrane.jw.
77.	((multiple treatment* or indirect or mixed) adj2 comparison*).ti,ab.
78.	or/68-77
79.	Epidemiologic studies/
80.	Observational study/
81.	exp Cohort studies/
82.	(cohort adj (study or studies or analys* or data)).ti,ab.
83.	((follow up or observational or uncontrolled or non randomi#ed or epidemiologic*) adj (study or studies or data)).ti,ab.
84.	((longitudinal or retrospective or prospective or cross sectional) and (study or studies or review or analys* or cohort* or data)).ti,ab.
85.	Controlled Before-After Studies/
86.	Historically Controlled Study/
87.	Interrupted Time Series Analysis/
88.	(before adj2 after adj2 (study or studies or data)).ti,ab.
89.	or/79-88
90.	exp case control study/
91.	case control*.ti,ab.
92.	or/90-91
93.	89 or 92
94.	Cross-sectional studies/
95.	(cross sectional and (study or studies or review or analys* or cohort* or data)).ti,ab.
96.	or/94-95
97.	89 or 96
98.	89 or 92 or 96
99.	59 or 67 or 78 or 98
100.	48 and 99

Embase (Ovid) search terms

1.	exp urolithiasis/
2.	(nephrolitiasis or nephrolith or nephroliths or urolithias?s or ureterolithias?s).ti,ab.
3.	((renal or kidney* or urinary or ureter* or urethra*) adj3 (stone* or calculi or calculus or calculosis or lithiasis or c?olic*)).ti,ab.
4.	stone disease*.ti,ab.
5.	((calculi or calculus or calcium oxalate or cystine) adj3 (crystal* or stone* or lithiasis)).ti,ab.
6.	or/1-5
7.	letter.pt. or letter/
8.	note.pt.
9.	editorial.pt.
10.	case report/ or case study/
11.	(letter or comment*).ti.

13. randomized controlled trial/ or random*.ti,ab. 14. 12 not 13 15. animal/ not human/ 16. nonhuman/ 17. exp Animal Experiment/ 18. exp Experimental Animal/ 19. animal model/ 20. exp Rodent/ 21. (rat or rats or mouse or mice).ti. 22. or/14-21 23. 6 not 22 24. limit 23 to English language 25. exp 'tomography/ 26. tomograph'.ti,ab. 27. (NCCT or CT or UHCT).ti,ab. 28. ((CAT or body) adj2 scan*).ti,ab. 29. or/25-28 30. *radiography/ 31. *abdominal radiography/ 32. *urography/ 33. (radiograph or x ray* or xray* KUB or urograph*).ti,ab. 34. or/30-33 35. *echography 36. (Ultrasonograph* or x ray* or xray* KUB or urograph* or echograph* or echograph* or echotomograph* or echoto	12.	or/7-11
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effectiveness)).ti,ab.	53.	(receive* operat* characteristic* or receive* operat* curve* or ROC curve*).ti,ab.
	54.	
55. diagnostic accuracy/	55.	diagnostic accuracy/

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56.	diagnostic test accuracy study/								
57.	gold standard.ab.								
58.	or/47-57								
59.	random*.ti,ab.								
60.	factorial*.ti,ab.								
61.	(crossover* or cross over*).ti,ab.								
62.	((doubl* or singl*) adj blind*).ti,ab.								
63.	(assign* or allocat* or volunteer* or placebo*).ti,ab.								
64.	(assign or allocat or volunteer or placedo").tl,ab.								
65.									
	single blind procedure/ randomized controlled trial/								
66.									
67.	double blind procedure/								
68.	or/59-67								
69.	systematic review/								
70.	meta-analysis/								
71.	(meta analy* or metanaly* or metanaly* or meta regression).ti,ab.								
72.	((systematic* or evidence*) adj3 (review* or overview*)).ti,ab.								
73.	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.								
74.	(search strategy or search criteria or systematic search or study selection or data extraction).ab.								
75.	(search* adj4 literature).ab.								
76.	(medline or pubmed or cochrane or embase or psychilt or psyclit or psychinfo or psycinfo or cinahl or science citation index or bids or cancerlit).ab.								
77.	cochrane.jw.								
78.	((multiple treatment* or indirect or mixed) adj2 comparison*).ti,ab.								
79.	or/69-78								
80.	Clinical study/								
81.	Observational study/								
82.	family study/								
83.	longitudinal study/								
84.	retrospective study/								
85.	prospective study/								
86.	cohort analysis/								
87.	follow-up/								
88.	cohort*.ti,ab.								
89.	87 and 88								
90.	(cohort adj (study or studies or analys* or data)).ti,ab.								
91.	 ((follow up or observational or uncontrolled or non randomi#ed or epidemiologic*) adj (study or studies or data)).ti,ab. 								
92.	((longitudinal or retrospective or prospective or cross sectional) and (study or studies or review or analys* or cohort* or data)).ti,ab.								
93.	(before adj2 after adj2 (study or studies or data)).ti,ab.								
94.	or/80-86,89-93								
95.	exp case control study/								
96.	case control*.ti,ab.								
	or/95-96								

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98.	94 or 97
99.	cross-sectional study/
100.	(cross sectional and (study or studies or review or analys* or cohort* or data)).ti,ab.
101.	or/99-100
102.	94 or 101
103.	94 or 97 or 101
104.	58 or 68 or 79 or 103
105.	46 and 104

Cochrane Library (Wiley) search terms

#1.	MeSH descriptor: [Urolithiasis] explode all trees						
#1.	(nephrolitiasis or nephrolith or nephroliths or urolithias?s or ureterolithias?s):ti,ab						
#3.	((renal or kidney* or urinary or ureter* or urethra*) near/3 (stone* or calculi or calculus or calculosis or lithiasis or c?olic*)):ti,ab						
#4.	stone disease*:ti,ab						
#5.	((calculi or calculus or calcium oxalate or cystine) near/3 (crystal* or stone* or lithiasis)):ti,ab						
#6.	(or #1-#5)						
#7.	MeSH descriptor: [Tomography] explode all trees						
#8.	tomograph*:ti,ab						
#9.	(NCCT or CT or UHCT):ti,ab						
#10.	((CAT or body) near/2 scan*):ti,ab						
#11.	(or #7-#10)						
#12.	MeSH descriptor: [Radiography] this term only						
#13.	MeSH descriptor: [Radiography, Abdominal] this term only						
#14.	MeSH descriptor: [Urography] explode all trees						
#15.	(radiograph* or x ray* or xray or KUB or urograph*):ti,ab						
#16.	(or #12-#15)						
#17.	MeSH descriptor: [Ultrasonography] this term only						
#18.	(ultrasonograph* or ultrasound or ultrasonic or sonograph* or echograph* or echotomograph*):ti,ab						
#19.	(ultra near/2 (sound or sonic)):ti,ab						
#20.	(sound* near/2 (wave* or frequenc*)):ti,ab						
#21.	(US near/3 imag*):ti,ab						
#22.	(or #17-#21)						
#23.	MeSH descriptor: [Magnetic Resonance Imaging] this term only						
#24.	((magnetic or nuclear) near/2 resonance near/3 imag*):ti,ab						
#25.	(MRI or NMR or NMRI or fMRI or MR):ti,ab						
#26.	(or #23-#25)						
#27.	#11 or #16 or #22 or #26						
#28.	#6 and #27						

B.2 Health Economics literature search strategy

Health economic evidence was identified by conducting a broad search relating to renal and ureteric stones population in NHS Economic Evaluation Database (NHS EED – this ceased to be updated after March 2015) and the Health Technology Assessment database (HTA) with no date restrictions. NHS EED and HTA databases are hosted by the Centre for

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Research and Dissemination (CRD). Additional searches were run on Medline and Embase for health economics studies.

Database	Dates searched	Search filter used
Medline	2014 – 9 March 2018	Exclusions Health economics studies
Embase	2014 – 9 March 2018	Exclusions Health economics studies
Centre for Research and Dissemination (CRD)	HTA - Inception – 9 March 2018 NHSEED - Inception to March 2015	None

Table 9: Database date parameters and filters used

Medline (Ovid) search terms

1.	exp urolithiasis/						
2.	(nephrolitiasis or nephrolith or nephroliths or urolithias?s or ureterolithias?s).ti,ab.						
3.	((renal or kidney* or urinary or ureter* or urethra*) adj3 (stone* or calculi or calculus or calculosis or lithiasis or c?olic*)).ti,ab.						
4.	stone disease*.ti,ab.						
5.	((calculi or calculus or calcium oxalate or cystine) adj3 (crystal* or stone* or lithiasis)).ti,ab.						
6.	or/1-5						
7.	letter/						
8.	editorial/						
9.	news/						
10.	exp historical article/						
11.	Anecdotes as Topic/						
12.	comment/						
13.	case report/						
14.	(letter or comment*).ti.						
15.	or/7-14						
16.	randomized controlled trial/ or random*.ti,ab.						
17.	15 not 16						
18.	animals/ not humans/						
19.	exp Animals, Laboratory/						
20.	exp Animal Experimentation/						
21.	exp Models, Animal/						
22.	exp Rodentia/						
23.	(rat or rats or mouse or mice).ti.						
24.	or/17-23						
25.	6 not 24						
26.	limit 25 to English language						
27.	Economics/						
28.	Value of life/						
29.	exp "Costs and Cost Analysis"/						
30.	exp Economics, Hospital/						

31.	exp Economics, Medical/						
32.	Economics, Nursing/						
33.	Economics, Pharmaceutical/						
34.	exp "Fees and Charges"/						
35.	exp Budgets/						
36.	budget*.ti,ab.						
37.	cost*.ti.						
38.	(economic* or pharmaco?economic*).ti.						
39.	(price* or pricing*).ti,ab.						
40.	(cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab.						
41.	(financ* or fee or fees).ti,ab.						
42.	(value adj2 (money or monetary)).ti,ab.						
43.	or/27-42						
44.	26 and 43						

Embase (Ovid) search terms

1.	exp urolithiasis/							
2.	(nephrolitiasis or nephrolith or nephroliths or urolithias?s or ureterolithias?s).ti,ab.							
3.	((renal or kidney* or urinary or ureter* or urethra*) adj3 (stone* or calculi or calculus or calculosis or lithiasis or c?olic*)).ti,ab.							
4.	stone disease*.ti,ab.							
5.	((calculi or calculus or calcium oxalate or cystine) adj3 (crystal* or stone* or lithiasis)).ti,ab.							
6.	or/1-5							
7.	letter.pt. or letter/							
8.	note.pt.							
9.	editorial.pt.							
10.	case report/ or case study/							
11.	(letter or comment*).ti.							
12.	or/7-11							
13.	randomized controlled trial/ or random*.ti,ab.							
14.	12 not 13							
15.	animal/ not human/							
16.	nonhuman/							
17.	exp Animal Experiment/							
18.	exp Experimental Animal/							
19.	animal model/							
20.	exp Rodent/							
21.	(rat or rats or mouse or mice).ti.							
22.	or/14-21							
23.	6 not 22							
24.	limit 23 to English language							
25.	health economics/							

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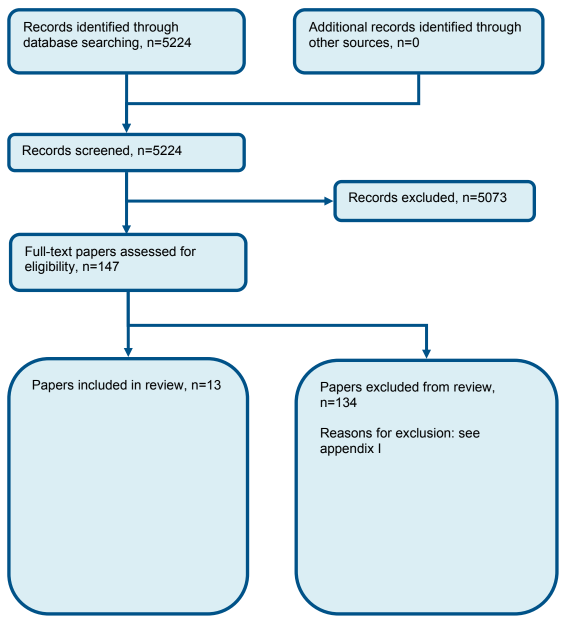
26. exp economic evaluation/ 27. exp health care cost/ 28. exp fee/ 29. budget/ 30. funding/							
28. exp fee/ 29. budget/							
29. budget/	exp health care cost/						
	exp fee/						
20 funding/	budget/						
31. budget*.ti,ab.							
32. cost*.ti.	cost*.ti.						
33. (economic* or pharmaco?economic*).ti.	(economic* or pharmaco?economic*).ti.						
34. (price* or pricing*).ti,ab.							
35. (cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab.							
36. (financ* or fee or fees).ti,ab.							
37. (value adj2 (money or monetary)).ti,ab.	(value adj2 (money or monetary)).ti,ab.						
38. or/25-37	or/25-37						
39. 24 and 38							

NHS EED and HTA (CRD) search terms

#1.	MeSH DESCRIPTOR urolithiasis EXPLODE ALL TREES					
#2.	(((nephrolitiasis or nephrolith or urolithiasis)))					
#3.	((((renal or kidney or urinary or ureteric or ureteral or ureter or urethra*) adj2 (stone* or calculi or calculus or calculosis or lithiasis or colic))))					
#4.	((stone disease*))					
#5.	((((calculi or calculus) adj2 (stone* or lithiasis))))					
#6.	(#1 OR #2 OR #3 OR #4 OR #5)					

Appendix C: Clinical evidence selection

Figure 1: Flow chart of clinical study selection for the review of What is the clinical and cost effectiveness of performing imaging for diagnosing renal and ureteric stones?



Appendix D: Clinical evidence tables

Chan 2008
Retrospective cohort study
Data source: retrospective review of imaging
Recruitment: consecutive patients being investigated for suspected urolithiasis
n = 100
Age, mean (SD): 45 (SD not reported)
Gender (male to female ratio): 63:37
Ethnicity: Not reported
Setting: Not reported
Country: Ireland
Inclusion criteria: Patients presenting with acute renal colic who had both a KUB and UHCT within a 3 hour period.
Exclusion criteria: Not reported
Urolithiasis
Index test KUB using a standard KUB protocol.
Reference standard
CT. All CT imaging was obtained with a spiral CT scanner using a standard low dose protocol (100 mAs, 120 kVp, 1.4 pitch, single breath-hold) extending from the top of the kidneys to the base of the bladder.
UHCT and KUB pairs were divided into two separate groups and read by two radiologists in consensus who were experienced in
genitourological radiology. The UHCT and KUB subsets were reviewed at separate time intervals to prevent internal bias. All revisions were done on a picture archiving and communications system (PACS).
Time between measurement of index test and reference standard: 3 hours

Reference	Chan 2008							
Reference								
2×2 table		Reference standard +	Reference standard -	Total				
	Index test +	39	2	41				
	Index test -	20	39	59				
	Total	59	41	100				
Statistical	Index text: KUB							
measures		% (52.61, 77.92)						
	Specificity: 95.12% (83.47, 99.40)							
	PPV: 95.12% (8							
	NPV: 66.1% (57							
	PLR: 13.55 (3.4							
	NLR: 0.36 (0.25	9, 0.51)						
Source of	Not reported							
funding								
Limitations	Risk of bias: None							
Linitationio	Indirectness: None							
Comments								
Reference	Cifci 2016							
Study type	Prospective cohort study							
Study	Data source: patients admitted to the urology department							
methodology								
Number of	450							
patients	n = 159							
Patient	Age, mean (SC): 41.5 years (13 years)							
characteristics								
	Gender (male to female ratio): 120:39							
	Ethnicity: Not reported							
	Setting: Not reported							

Reference	Cifci 2016							
	Country: Turkey							
		Inclusion criteria: patients who were admitted to the urology department with suspected acute urinary calculi (flank pain, hematuria or						
	patients with a history of urinary calculi)							
	Exclusion criteria: Images that did not meet the required conditions (inadequate anatomy and image quality) were excluded from the study. Patients with MRU images performed more than 3 hours after CT examination were also excluded from the study, due to a potential change in the stone localization. Additionally, patients who were not able to hold their breath, yielding images with respirator							
Townst		e artifacts, were exclude	d from the study.					
Target condition(s)	Ureteral calculi							
Index test(s)	Index test		1 E Taala (T) wait (latara	Currencen Dhiline	Medical Cystema, The Netherlands) and a			
and reference standard					Medical Systems, The Netherlands) and a bic symphysis Avial coronal and sadittal B-TEE			
	four-channel phased-array body coil was used, from the top of the kidneys to the pubic symphysis. Axial, coronal and sagittal B-TFE (TR: 3.5–5, TE: 1.5–2.5, slice thickness: +4 mm and gap: -2.5 mm) images were taken while breath was held from the top of the							
	kidneys to the public symphysis.							
	Reference standard CT. Abdominal non-enhanced CT (Philips, MX 8000, The Netherlands) was performed from the top of the kidneys to the pubic							
	symphysis, while breath was held, at 80–120 kV, 100–120 mAs, and 5a -mm slice thickness with 3-mm reconstruction. All patients w hydrated a minimum of 1 hour before CT and MRU imaging. No contrast media was used.							
	l ime between n	neasurement of index tes	st and reference standar	d: 1-3 hours				
2×2 table		Reference standard +	Reference standard -	Total				
observer 1	Index test +	56	3	59				
	Index test -	29	71	100				
	Total	85	74	159				
Statistical	Index text: KUB							
measures	Sensitivity: 65.88% (54.80, 75.82)							
	Specificity: 95.95% (88.61, 99.16)							
	PPV: 94.92% (85.91, 98.28) NPV: 71.00% (64.48, 76.75)							
	PLR: 16.25 (5.31, 49.75)							
	NLR: 0.36 (0.26, 0.48)							

45

Reference	Cifci 2016				
2×2 table		Reference standard +	Reference standard -	Total	
observer 2	Index test +	61	0	61	
	Index test -	24	74	98	
	Total	85	74	159	
Statistical measures		76% (60.96, 81.00) .00% (95.14, 100.00) 68.72, 81.23)			
Source of funding	No funding				
Limitations	Risk of bias: ref Indirectness: hi	ference standard story of stones			
Comments					

Reference	De Souza 2007
Study type	Prospective cohort study
Study methodology	Data source: patients referred from the emergency department
	Recruitment: consecutive patients being evaluated for acute flank pain
Number of patients	n = 52

Reference	De Souza 2007					
Patient characteristics	Age, mean (SD): Not reported					
	Gender (male to female ratio): Not reported					
	Ethnicity: Not reported					
	Setting: Not reported					
	Country: Brazil					
	Inclusion criteria: Not reported					
	Exclusion criteria: known renal diseases or imaging signs of pyelonephritis, chronic renal insufficiency, nephrocalcinosis and staghorn calculus.					
Target condition(s)	Ureteral stone					
Index test(s)	Index test					
and reference standard	US performed transabdominally, after ingestion of water. Sonography was performed by senior radiology residents and immediately checked by attending radiologists, using a Philips SD800 scanner (Philips Medical Systems, Eindhoven, Netherlands) with a convex (curved phased array) transducer (2-5 MHz) and transducer frequencies selected to optimize the imaging of the kidneys, ureters and bladder. The US diagnosis of ureteral calculi required the demonstration of an intraluminal hyperechoic structure causing acoustic shadowing. The presence of collecting system dilatation was also evaluated. No patient underwent transvaginal or transrectal sonographic examination.					
	Reference standard CT. NCT scans were acquired after US examination, on a Tomoscan EV-EV1 (Philips Medical Systems, Eindhoven, Netherlands) usir Secura Release 1.3 software. The scan parameters included helical data acquisition, with section thickness 3-5 mm, using 120 kV and 200 mAs and a pitch of 1-1.5. Images were obtained during apnea, from the top of the kidneys to the base of the bladder, and no contrast medium was used. The NCT images were interpreted by a senior resident, using an electronic workstation (Philips), and subsequently reviewed by three experienced abdominal radiologists in a blinded manner.					
	Time between measurement of index test and reference standard: both were performed within 8 hours of the onset of colic (average of 4 hours between the two tests)					
2×2 table	Reference standard + Reference standard - Total					
	Index test + 9 0 9					
	Index test - 31 12 43					

Reference	De Souza 2	2007						
	Total	40	12	52				
Statistical measures	Specificity: PPV: 100.00	22.50% (10.84, 38. 100.00% (73.54, 10 0% % (24.67, 31.39)						
Source of funding	None							
Limitations	Risk of bias: index test, reference standard Indirectness: none							
Comments								
Reference	Haroun 201	10						
Study type	Retrospectiv	ve cohort study						
Study methodology		e: Not reported						
	Recruitmen	t: Not reported						
Number of patients	n = 156							
Patient characteristics	Age, mean	(SD): male patients	51 (16) years; female pa	atients 46 (18) years				
	Gender (ma	ale to female ratio):	102:54					

Ethnicity: Not reported

Setting: Not reported

Country: Jordan

Inclusion criteria: patients who underwent UHCT scan and US for suspicion of urolithiasis

Reference	Haroun 2010					
		ria: Not reported				
Target condition(s)	Urolithiasis					
Index test(s) and reference standard	Index test					
	Reference standard CT. UHCT scan was performed with a Somatom Plus 4 machine (Siemens, Germany). The images were obtained with the patient in supine position during breath-hold plus quiet breathing. The explored area extended from the upper poles of both kidneys down to pubic symphysis using five mm collimation with a table speed of 7.5 mm/second giving a pitch of 1.5:1 The images were obtained with a 0.75-second gantry rotation using 120 KVp and 206 mA giving 155 mAs. Multiplanar reformation (MPR) in coronal oblique direction was used when the location of stone was uncertain. CT scan images were reported by consultant radiologists on hard copy films. Time between measurement of index test and reference standard: Not reported					
2×2 table		Reference standard +	Reference standard –	Total		
	Index test +	34	9	43		
	Index test – Total	25 59	88 97	113 156		
Statistical measures		63% (44.07, 70.39) 72% (83.12, 95.67) 66.14, 87.96) 72.19, 82.68) 1, 12.01)				
Source of funding	Not reported					
Limitations	Risk of bias: in Indirectness: n	dex test, reference test, fl one	ow and timing			
Comments						

Reference	Kanno 2017
Study type	Retrospective cohort study
Study methodology	Data source: database review
	Recruitment: Not reported
Number of patients	n = 822
Patient characteristics	Age, median: 60 years
	Gender (male to female ratio): 553:269
	Ethnicity: Not reported
	Setting: Not reported
	Country: Japan
	Inclusion criteria: Indications for imaging included acute flank pain, hematuria, or a history of urinary stones as previously described. Whereas new patients routinely underwent US for screening of the urinary tract at our institution, we also performed both KUB and NCCT in patients with acute flank pain and suspected urolithiasis to get information such as stone size, mean stone density, and skin- to-stone distance, except for patients who underwent NCCT in other hospitals and were referred to our institution.
	Exclusion criteria: Patients with solitary kidney, staghorn calculi, and urinary diversion were excluded from our analysis.
Target condition(s)	Renal stones
Index test(s)	Index test
and reference standard	US. All US examinations were performed by 7 experienced sonographers who are specialized in handling urologic US. Echogenic foci (with or without acoustic shadowing) that were seen in the renal pelvis or calices on US were diagnosed as renal stones, because small
Standard	stones may not cast an acoustic shadow as described previously
	KUB. No further details reported.
	Reference standard

Reference	Kanno 2017 CT. NCCT (Toshiba Aquilion ONE 640, Toshiba Medical Systems, Tokyo, Japan) was performed from the upper abdomen to the pelvis with images reconstructed at 1- to 2-mm intervals. US of the kidneys and bladder was performed using grayscale sonography (Toshiba SSA550A) with a 3.5-MHz convex transducer.					
	Time between	measurement of index tes	st and reference standar	d: same day		
2×2 table - US		Reference standard +	Reference standard -	Total		
	Index test +	882	206	1088		
	Index test -	112	444	556		
	Total	994	650	1644		
2×2 table -		Reference standard +	Reference standard -	Total		
KUB	Index test +	488	6	494		
	Index test -	506	644	1150		
	Total	994	650	1644		
Statistical measures	Specificity: 68 PPV: 81.07% NPV: 79.86% PLR: 2.80 (2.5 NLR: 0.16 (0.7 Index text: KU Sensitivity: 49 Specificity: 99 PPV: 98.79%	.73% (86.60, 90.63) .31% (64.58, 71.87) (79.24, 82.77) (76.77, 82.63) 50, 3.14) 14, 0.20) <u>B</u> .09% (45.94, 52.25) .08% (98.00, 99.66) (97.34, 99.45) (54.48, 57.51) 3.92, 118.25)				
Source of funding	Not reported	, , ,				
Limitations		atient selection history of stones				
Comments						

Reference	Kanno 2014a #1274
Study type	Retrospective cohort study
Study methodology	Data source: database review
	Recruitment: Not reported
Number of patients	n = 428
Patient characteristics	Age, median: Not reported
	Gender (male to female ratio): Not reported
	Ethnicity: Not reported
	Setting: Not reported
	Country: Japan
	Inclusion criteria: Indications for imaging included acute flank pain, haematuria, or a history of urinary stones
Townst	Exclusion criteria: Patients with solitary kidney, staghorn calculi, and urinary diversion were excluded
Target condition(s)	Renal stones
Index test(s) and reference standard	Index test US. US was performed using gray scale sonography (SSA550A; Toshiba) with a 3.5-MHz convex transducer. All US examinations were performed by 4 experienced sonographers who are specialized in handling urologic US. Echogenic foci (with or without acoustic shadowing) that were seen in the renal pelvis or calices on US were diagnosed as renal stones because small stones may not cast an acoustic shadow.
	Reference standard CT. NCCT (Aquilion ONE 640; Toshiba) was performed from the upper abdomen to the pelvis with images reconstructed at 1 or 2 mm intervals.

Time between measurement of index test and reference standard: same day

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50

Reference	Kanno 2014a #	1274				
2×2 table –		Reference standard +	Reference standard -	Total		
individual	Index test +	285	80	365		
stone	Index test -	76	412	488		
	Total	361	492	853		
Statistical measures	Index text: US – individual stones Sensitivity: 78.95% (74.37, 83.04) Specificity: 83.74 % (80.18, 86.89) PPV: 78.08% (74.33, 81.43) NPV: 84.43% (81.56, 86.92) PLR: 4.86 (3.95, 5.97) NLR: 0.25 (0.21, 0.31)					
Source of funding	No financial inte	erests				
Limitations	Risk of bias: reference standard Indirectness: history of stones					
Comments						
Reference	Kanno 2014					
Study type		ohort study				
Study methodology	Retrospective cohort study Data source: database review					
	Recruitment: No	ot reported				
Number of patients	n = 428					
Patient characteristics	Age, median: N					
	Gender (male to	o female ratio): Not repor	ted			
	Ethnicity: Not re	ported				
	Setting: Not rep	orted				

Reference	Kanno 2014								
	Country: Japa	n							
	Inclusion oritor	Inclusion criteria: Indications for imaging included acute flank pain, baomaturia, or a history of urinary stones							
	Inclusion criteria: Indications for imaging included acute flank pain, haematuria, or a history of urinary stones								
	Exclusion crite	eria: Patients with solitary k	kidney, staghorn calculi,	and urinary diversion	were excluded				
Target condition(s)	Ureteric stone	s		·					
Index test(s) and reference standard	Index test US. US was performed using gray scale sonography (SSA550A; Toshiba) with a 3.5-MHz convex transducer. All US examinations were performed by 4 experienced sonographers who are specialized in handling urologic US. Echogenic foci (with or without acoustic shadowing) that were seen in the renal pelvis or calices on US were diagnosed as renal stones because small stones may not cast an acoustic shadow. <u>Reference standard</u> CT. NCCT (Aquilion ONE 640; Toshiba) was performed from the upper abdomen to the pelvis with images reconstructed at 1 or 2 mm intervals. Time between measurement of index test and reference standard: same day								
2×2 table –		Reference standard +	Reference standard -	Total					
individual	Index test +	98	73	171					
stone	Index test -	17	668	685					
	Total	115	741	856					
Statistical measures	Index text: US Sensitivity: 85.22% (77.39, 91.15) Specificity: 90.15% (87.77, 92.20) PPV: 57.31% (51.59, 62.84) NPV: 97.52% (96.20, 98.39) PLR: 8.65 (6.87, 10.89) NLR: 0.16 (0.11, 0.25)								
Source of funding	No financial in	terests							

Defenses	
Reference	Kanno 2014
Limitations	Risk of bias: reference standard
•	Indirectness: none
Comments	
Reference	Kielar 2012
Study type	Prospective cohort study
Study	Data source: patients from an Emergency Department
methodology	
	Recruitment: Not reported
Number of	n = 55
patients	
Patient	Age, mean (range): 49 years (28–81 years)
characteristics	
	Gender (male to female ratio): 38:17
	Ethnicity: Not reported
	Setting: Not reported
	Country: Canada
	Inclusion criteria: Deeple with flenk pain in whom an unenhanced CT of the abdemon and polyie was requested
	Inclusion criteria: People with flank pain in whom an unenhanced CT of the abdomen and pelvis was requested
	Evolution criteria: Not reported
Torret	Exclusion criteria: Not reported
Target	Urolithiasis
condition(s)	
Index test(s) and reference	Index test
standard	US. The patient directly underwent a limited sonographic scan of the kidneys, ureters, and bladder (iU22; Philips Healthcare, Bothell, WA). This examination was performed for research purposes and was not considered the usual standard of care. It was performed with
Stanuaru	a standard ultrasound unit, which is always situated in the emergency radiology department, in a dedicated room next to the emergency
	radiology CT scanner. The examination was performed by a trained sonographer using a curved low-frequency probe (2–5 MHz) and a
	high pulse repetition frequency, with the machine's scale in the range of 60 to 70 cm/s. The pulse repetition frequency is defined as the
	number of pulses sent per second
	Reference standard

Reference	Kielar 2012	Kielar 2012					
	CT. All CT scans were performed on a 64-slice CT scanner (Lightspeed VCT; GE Healthcare, Milwaukee, WI). All scans were performed with a low-dose, unenhanced "renal colic protocol." The images were sent to a picture archiving and communication sys at the original axial 1.25mmslice thickness in addition to the reconstructed5-mm axial images and2-mmcoronal images. The1.25-m raw data were reviewed to eliminate the possibility of missing small calculi because of volume averaging. The post processing techniques do not expose the patient to any additional radiation.						
2×2 table –		Reference standard +	Reference standard -	Total			
individual	Index test +	74	8	82			
stone	Index test -	40	5	45			
	Total	114	13	127			
Statistical measures	Index text: US Sensitivity: 64.91% (55.41, 73.62) Specificity: 38.46% (13.86, 68.42) PPV: 90.24% (85.50, 93.55) NPV: 11.11% (5.67, 20.62) PLR: 1.05 (0.67, 1.66) NLR: 0.91 (0.44, 1.90)						
Source of funding	No financial in	nterests					
Limitations	Risk of bias: patient selection Indirectness: none						
Comments							
Reference	Levine 1997						
Study type	Retrospective	cohort study					
Study		retrospective review of rec	ords				

Levine 1997
Retrospective cohort study
Data source: retrospective review of records
Recruitment: Not reported
n = 178

Reference	Levine 1997								
Patient characteristics	Age, mean (range): 49 years (28–81 years)								
	Gender (male to female ratio): 38:17								
	Ethnicity: Not reported								
	Setting: Not rep	ported							
	Country: USA								
	Inclusion criteria	a: People with acute flan	k pain who had undergor	ne plain abdominal rad	diography before CT				
	Exclusion criter	ia: Not reported							
Target condition(s)	Urolithiasis								
Index test(s) and reference standard	Index test Plain abdominal radiograph. No details reported Reference standard CT. All CT. examinations were performed with a HiSpeed Advantage scenner (CE Medical Systems, Milwaukee, Mic), Images were								
	CT. All CT examinations were performed with a HiSpeed Advantage scanner (GE Medical Systems, Milwaukee, Wis). Images were obtained from the superior poles of the kidneys to the bladder base. Helical data acquisition was used with 5mm thick sections and a pitch of 1. Image clusters of 15-20 sections were obtained during suspended respiration.								
	Time between r	measurement of index tes	st and reference standar	d: 4 hours					
2×2 table –		Reference standard +	Reference standard -	Total					
individual	Index test +	39	25	70					
stone	Index test -	27	60	87					
	Total	72	85	157					

Reference	Levine 1997
Statistical measures	Index text: US Sensitivity: 59.09% (46.29, 71.05) Specificity: 70.59% (59.71, 79.98) PPV: 60.94% (51.47, 69.64) NPV: 68.97% (61.72, 75.39) PLR: 2.01 (1.37, 2.95) NLR: 0.58 (0.42, 0.80)
Source of funding	Not reported
Limitations	Risk of bias: reference standard Indirectness: none
Comments	

Reference	Passerotti 2009
Study type	Prospective cohort study
Study methodology	Data source: emergency department
	Recruitment: Consecutive patients
Number of patients	n = 50
Patient characteristics	Age, mean (range): 13.1 (2–18 years)
	Gender (male to female ratio): 25:25
	Ethnicity: Not reported
	Setting: Not reported
	Country: USA
	Inclusion criteria: People younger than 18 years who presented to the emergency department or the urology clinic with signs, symptoms or a history suggestive of urolithiasis.

Reference	Passerotti 2009						
	Exclusion criteria: Not reported						
Target condition(s)	Urinary calculi						
Index test(s) and reference standard	Index test US. Renal US using a standard departmental protocol with a Sequoia [™] 512 or an IU22 machine (Philips, Bothell, Washington) with variable combination of curved and vector array transducers, and frequencies best suited to the patient. Stones were defined as echogenic foci with posterior acoustic shadowing, confirmed in 2 planes and measured in 3 planes. Reference standard CT. Noncontrast CT of the abdomen and pelvis using 5 mm cuts with reconstruction at 1.5 mm was done according to standard departmental renal stone protocol using 1 of 3 scanners, including a LightSpeed [™] Ultra 16, a LightSpeed CT 32 or a Somatom® Sensation 64. No patient required sedation. Peak kV and mA settings were device specific, and based on patient weight and age. Images were obtained in the axial plane and reconstructed in the coronal and sagittal planes. Stones were defined as hyper dense foci see the kidneys, ureters or bladder. Time between measurement of index test and reference standard: 0.5-8 hours						
	Time between	measurement of index tes	st and reference standa	d: 0.5-8 hours			
2×2 table –	Time between						
2×2 table – individual		Reference standard +	Reference standard -	Total			
2×2 table – individual stone	Time between Index test + Index test -		Reference standard - 0	Total 26			
individual	Index test +	Reference standard + 26	Reference standard -	Total			
individual	Index test + Index test - Total Index text: US Sensitivity: 76. Specificity: 100 PPV: 100.00%	Reference standard + 26 8 34 47% (58.83, 89.25) 0.00% (79.41, 100.00) (52.188, 78.57)	Reference standard – 0 16	Total 26 24			
individual stone Statistical	Index test + Index test - Total Index text: US Sensitivity: 76. Specificity: 100 PPV: 100.00% NPV: 66.67% (PLR: -	Reference standard + 26 8 34 47% (58.83, 89.25) 0.00% (79.41, 100.00) (52.188, 78.57)	Reference standard – 0 16	Total 26 24			

Reference	Passerotti 2009
	Indirectness: history of stones
Comments	
Reference	Resorlu 2015
Study type	Retrospective cohort study
Study type	Data source: retrospective analysis of records
methodology	Data source. retrospective analysis of records
memodelegy	Recruitment: not reported
Number of patients	n = 500
Patient characteristics	Age, mean (SD): 49.8 (16.9) years
	Gender (male to female ratio): 297:203
	Ethnicity: Not reported
	Setting: Not reported
	Country: Turkey
	Inclusion criteria: People who underwent an initial urinary US, followed by urinary NCCT as part of their investigation for acute flank pain
	during working hours (between 08:00 a.m. and 05:00p.m.—Mondays to Fridays)
	Exclusion criteria: Patients who passed their stone in the interval between US and NCCT and those requiring stone treatment in terms of shock wave lithotripsy (SWL) or endoscopic intervention (ureterorenoscopic stone extraction) were excluded from the study.
Target condition(s)	Urinary stones
Index test(s) and reference standard	Index test US. Urinary US was performed by radiologists with grayscale ultrasound machines (Toshiba® Aplio XG and General Electric Logiq 9) using two convex transducers with 3.5, 4.0 MHz frequency. The presence of stone was defined as an echogenic image with or without posterior acoustic shadowing, clearly located within the urinary tract.
	Reference standard

Reference	Resorlu 2015							
	CT. All NCCT were performed with a 4-multidetector CT scanner (Toshiba® Asteion TSX-021B) without intravenous or oral contrast medium. Slices of 3-mm thickness with 1-mm reconstructed intervals were obtained, beginning from the superior aspect of the kidneys to the inferior aspect of the pubic symphysis. Stones were defined as hyperdense foci seen in the renal pelvis, calices, or ureters							
	Time between	measurement of index tes	st and reference standar	d: 10 days				
2×2 table	Reference standard + Reference standard - Total							
	Index test +	91	42	133				
	Index test -	111	256	367				
	Total	202	298	500				
Statistical measures		.05% (38.06, 52.19) .91% (81.43, 89.65) (61.16, 74.88) (66.88, 72.49) 32, 4.40)						
Source of funding	Not reported							
Limitations	Risk of bias: index test, reference standard Indirectness: none							
Comments								
Reference	Semins 2013							
Study type	Prospective cohort study							
Study methodology	Data source: e	emergency department						
	Recruitment: r	not reported						
Number of	n = 22							

patients

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D (0 1 0010								
Reference	Semins 2013								
Patient characteristics	Age, median (range): Not reported Gender (male to female ratio): Not reported								
	Ethnicity: Not reported								
	Setting: Not reported								
	Country: USA								
	Inclusion criteria: male or female patients with acute renal colic, age greater than 18 years, and ability to complete all necessar components of study								
		ia: inability to give inform omen who were pregnant			or unwillingness to comply with completion of etermined				
Target condition(s)	Obstructing stone								
Index test(s) and reference standard	Index test MRU. Non-contrast HASTE (Half-Fourier single shot turbo spin-echo) MRU with a 3-T scanner Reference standard CT. Non-contrast CT. No further details.								
	Time between measurement of index test and reference standard: MRU was performed immediately after CT								
2×2 table		Reference standard +	Reference standard -	Total					
	Index test +	16	0	16					
	Index test -	3	3	6					
	Total	19	3	22					

Reference	Semins 2013
Statistical	Index text: US
measures	Sensitivity: 84.21% (60.42, 96.62) Specificity: 100% (29.24, 100.00) PPV: 100.00% NPV: 50.00% (26.15, 73.85) PLR: NLR: 0.16 (0.06, 0.45)
Source of funding	Not reported
Limitations	Risk of bias: patient selection, index test, reference standard Indirectness: population
Comments	

Reference	Sternberg 2016
Study type	Retrospective cohort study
Study methodology	Data source: retrospective review of records from three institutions (University of Vermont Medical Center, Massachusetts General Hospital, Dartmouth-Hitchcock Medical Center)
	Recruitment: not reported
Number of patients	n = 155
Patient characteristics	Age, mean (range): Not reported
	Gender (male to female ratio): Not reported
	Ethnicity: Not reported
	Setting: Not reported
	Country: Lebanon
	Inclusion criteria: adult patients >18 years of age. Only formal radiologic US, not bedside-US, were included.

Reference	Sternberg 2016							
	Exclusion criteria: images were obtained >1 day apart, if imaging was of poor quality for interpretation, and/or if staghorn calculi were							
	present							
Target	, Urinary calculi							
condition(s)								
Index test(s)	Index test							
and reference	US. No further o	letails.						
standard								
	Reference stand	<u>dard</u>						
	CT. A standard	protocol was followed us	ing abdominal windows	and zooming in to bes	t visualize the stone of interest. Three			
	measurements	were made (length, width	n, height) using axial, saថ	gittal, and coronal sect	ions.			
	Time between n	neasurement of index tes	st and reference standard	d: 1 day				
2×2 table		Reference standard +	Reference standard -	Total				
	Index test +	79	2	81				
	Index test -	58	16	74				
	Total	137	18	155				
Statistical	Index text: US							
measures		6% (48.94, 66.05)						
		9% (65.29, 98.62)						
		PPV: 97.53% (91.39, 99.32)						
	NPV: 21.62% (1							
	PLR: 5.19 (1.39							
	NLR: 0.48 (0.37	, 0.61)						
Source of	Not reported							
funding	Notreponeu							
Limitations	Risk of bias: pat	tient selection, index test	reference standard					
	Indirectness: no							
Comments			e Group, and a Consulta	ant for Boston Scientifi	c, Bard, Retrophin, and Olympus.			
		One of the authors is owner of the Ravine Group, and a Consultant for Boston Scientific, Bard, Retrophin, and Olympus.						

Appendix E: Coupled sensitivity and specificity forest plots and sROC curves

E.1 Coupled sensitivity and specificity forest plots

Figure 2: Sensitivity and specificity of index test ultrasound for condition renal or ureteric stones in adults

Study	TP	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
de Souza 2007	9	0	31	12	0.23 [0.11, 0.38]	1.00 [0.74, 1.00]		
Haroun 2010	34	9	25	88	0.58 [0.44, 0.70]	0.91 [0.83, 0.96]		-
Kanno 2014a	285	80	76	412	0.79 [0.74, 0.83]	0.84 [0.80, 0.87]	-	-
Kanno 2014b	98	17	73	668	0.57 [0.50, 0.65]	0.98 [0.96, 0.99]		•
Kanno 2017	882	206	112	444	0.89 [0.87, 0.91]	0.68 [0.65, 0.72]	•	-
Resorlu 2015	91	42	111	256	0.45 [0.38, 0.52]	0.86 [0.81, 0.90]		-
Sternberg 2016	79	2	58	16	0.58 [0.49, 0.66]	0.89 [0.65, 0.99]	0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 3: Sensitivity and specificity of index test plain abdominal radiograph for condition renal or ureteric stones in adults

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Chan 2008	39	2	20	39	0.66 [0.53, 0.78]	0.95 [0.83, 0.99]		
Kanno 2017	488	6	506	644	0.49 [0.46, 0.52]	0.99 [0.98, 1.00]	-	•
Levine 1997	39	25	27	60	0.59 [0.46, 0.71]			
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 4: Sensitivity and specificity of index test MRI for condition renal or ureteric stones in adults

Study	ΤР	FP	FN	ΤN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Cifci 2016 (observer 1)	56	3	29	71	0.66 [0.55, 0.76]	0.96 [0.89, 0.99]		
Cifi 2016 (observer 2)	61	0	24	74	0.72 [0.61, 0.81]	1.00 [0.95, 1.00]		-
Semins 2013	16	0	3	3	0.84 [0.60, 0.97]			

Figure 5: Sensitivity and specificity of index test ultrasound for condition renal or ureteric stones in children

		opecificity (55% Of)	Sensitivity (95% CI)	IN	FN	FP	IP	Study
			0.76 [0.59, 0.89]	16	8	0	26	Passerotti 2009
	0.2 0.4 0.6 0.8 1		0.10 [0.00, 0.00]		Ŭ	Ŭ	20	1 40001011 2000

E.2 ROC curves

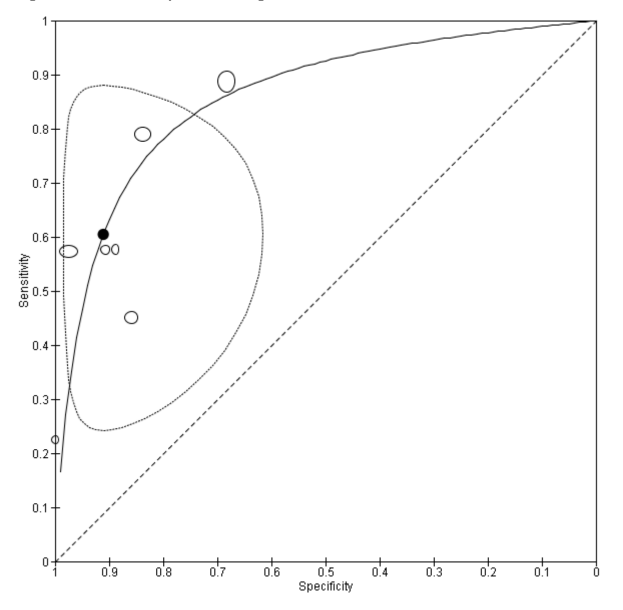
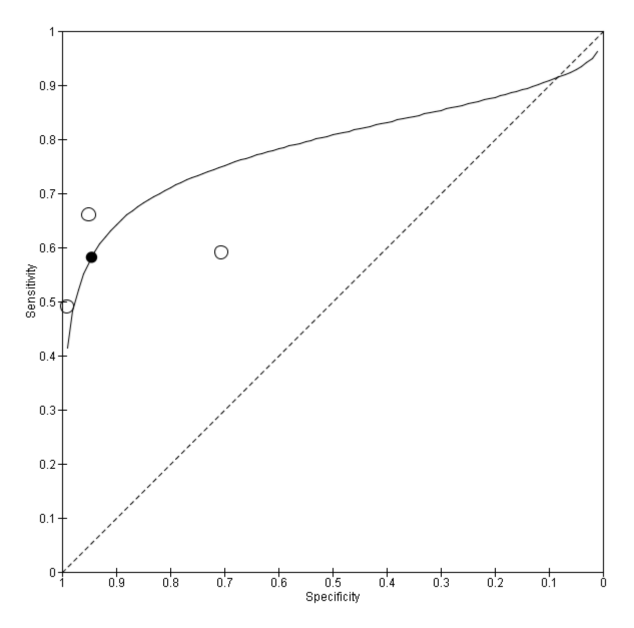


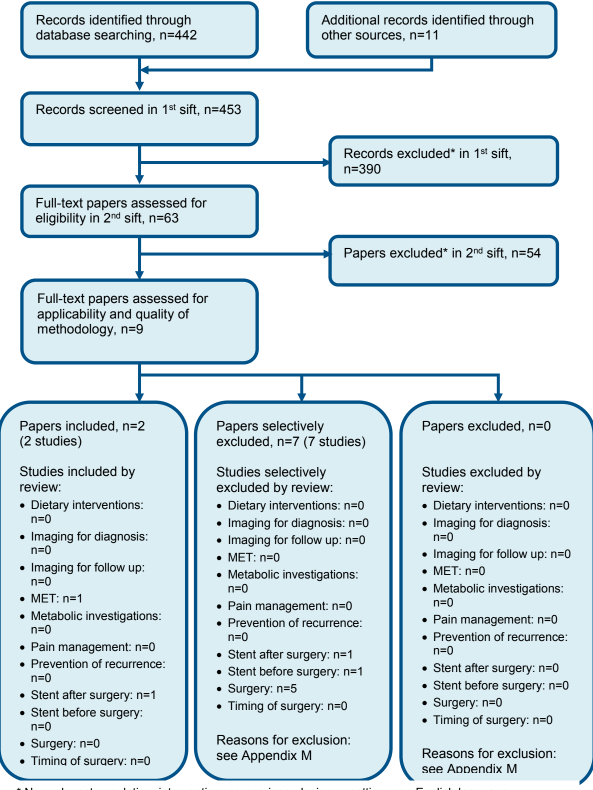
Figure 6: Pooled with prediction region: ultrasound





Appendix F: Health economic evidence selection

Figure 8: Flow chart of economic study selection for the guideline



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

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Appendix G: Health economic evidence tables

None

Appendix H: Excluded studies

H.1 Excluded clinical studies

Table 10: Studies excluded from the clinical review

Table 10: Studies excluded I	
Reference	Reason for exclusion
Abdel-Gawad 2014 ¹	Incorrect reference standard
Abdel-Gawad 2016 ²	Incorrect reference standard
Ahmad 2014 ³	No outcomes
Ahn 2002 ⁴	Incorrect population
Albani 2007 ⁵	Incorrect study design
Andresen 1997 ⁶	Incorrect reference standard
Arif 2013 ⁷	Incorrect reference standard
Assi 2000 ⁸	Incorrect reference standard
Ather 2004 ⁹	Incorrect reference standard
Ben Nakhi 2010 ¹⁰	Incorrect index test
Blandino 2004 11	Incorrect reference standard
Bozdar 2016 ¹²	No outcomes
Cabrera 2016 ¹³	Incorrect index test
Catalano 2002 ¹⁴	Incorrect reference standard
Cauberg 2011 ¹⁵	Incorrect population
Chen 1999 ¹⁷	Incorrect reference standard
Cochon 2017 ¹⁹	Incorrect study design
Dillman 2011 ²¹	Incorrect index test
Dorio 1999 22	Not available
Drake 2014 ²³	Systematic review checked for references
Dundee 2006 ²⁴	No outcomes
Edmonds 2010 ²⁵	Incorrect population
Ege 2004 ²⁶	Incorrect population
Eikefjord 2008 ²⁷	Incorrect reference standard
Ekici 2012 ²⁸	Incorrect reference standard
Eray 2003 ²⁹	Incorrect reference standard
Eshed 2002 ³⁰	Incorrect study design
Feroze 2007 ³¹	Incorrect reference standard
Foell 2013 ³²	Incorrect population
Fowler 2011 ³³	Incorrect reference standard
Fowler 2002 ³⁴	Incorrect reference standard
Gaspari 2005 ³⁵	Incorrect target condition
German 2002 36	No outcomes
Gliga 2017 ³⁷	Incorrect reference standard
Graumann 2012 38	Incorrect reference standard
Gurel 2006 39	Incorrect index test
Hamm 2001 40	Incorrect reference standard
Hamm 2001 41	Not in English
Herbst 2014 43	Incorrect target condition

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Reference	Reason for exclusion
Homer 2001 44	Incorrect index test
Hu 2010 ⁴⁵	Incorrect reference standard
Ibrahim 2016 46	Unclear population?
Jackman 2000 47	Incorrect population
Jeng 2001 ⁴⁸	Not in English
Johnston 2009 49	Incorrect index test
Joshi 2014 50	Incorrect reference standard
Jung 2010 ⁵¹	Incorrect population
Kennish 2008 55	Incorrect study design
Khan 2012 56	No outcomes
Kluner 2006 58	Incorrect reference standard
Korkmaz 2014 59	Incorrect population
Kravchick 2006 60	Incorrect reference standard
Lee 2015 61	Incorrect study design
Leo 2017 62	Incorrect target condition
Lew 2017 64	Incorrect population
Lin 2016 65	Incorrect reference standard
Lisanti 2014 66	Incorrect target condition
Liu 2000 67	Not available
Longo 2001 68	Incorrect reference standard
Lorberboym 2000 69	No outcomes
MacEjko 2009 70	Incorrect study design
Malaki 2014 71	Incorrect population
Marumo 2002 72	Incorrect study design
Masch 2016 73	Incorrect index test
Matani 2007 74	Incorrect index test
May 2016 ⁷⁵	Incorrect population
Meagher 2001 76	Incorrect population
Melnikow 2016 77	Incorrect study design
Mendelson 2003 78	Incorrect index test
Mermuys 2010 ⁷⁹	No outcomes
Middleton 1988 ⁸⁰	Incorrect reference standard
Miller 1998 ⁸¹	Incorrect reference standard
Mitterberger 2009 ⁸²	Incorrect population
Mitterberger 2007 ⁸³	Incorrect reference standard
Moak 2012	Incorrect index test
Moesbergen 2011 ⁸⁴	Incorrect population
Mos 2010 ⁸⁵	Incorrect population
Niall 1999 88	Incorrect reference standard
Nishiura 2009 89	Incorrect population
O'Kane 2016 90	Incorrect population
Olcott 1997 91	Incorrect population
Oner 2004 92	Incorrect reference standard
Palmer 2005 93	Incorrect population
Park 2008 94	Incorrect reference standard

Reference	Reason for exclusion
Patlas 2001 ⁹⁶	Incorrect reference standard
Pepe 2005 ⁹⁷	Incorrect reference standard
Pfister 2003 98	Incorrect reference standard
Pichler 2012 99	Incorrect study design
Poletti 2006 100	Incorrect study design
Quirke 2011 ¹⁰¹	Incorrect study design
Rajaie 2006 102	Incorrect index test
Ray 2010 ¹⁰³	No outcomes
Rengifo 2010 ¹⁰⁴	Not in English
Richards 1999 ¹⁰⁶	Incorrect index test
Riddell 2014 ¹⁰⁷	Incorrect population
Ripolles 2004 ¹⁰⁸	Incorrect reference standard
Ripolles 2013 ¹⁰⁹	Incorrect reference standard
Rosen 1998 110	Incorrect reference standard
Rosser 2000 111	Incorrect reference standard
Rowland 2001 ¹¹²	Incorrect target condition
Ryu 2001 ¹¹³	Incorrect reference standard
Sade 2017 ¹¹⁴	No outcomes
Sarofim 2016 ¹¹⁵	Incorrect reference standard
Sattar 2011 116	Not available
Schwartz 1984 ¹¹⁷	Incorrect index text and reference standard
Selberherr 2017 ¹¹⁸	Incorrect population
Sen 2017 120	Incorrect population
Sheafor 2000 ¹²¹	Incorrect reference standard
Shokeir 2001 122	Incorrect reference standard
Smith-Bindman 2014 ¹²³	Incorrect reference standard
Sudah 2002 125	Incorrect reference standard
Thomson 2001 ¹²⁶	Incorrect index test
Ulusan 2007 ¹²⁷	No usable outcomes
Unal 2003 ¹²⁸	Incorrect reference standard
Uraiqat 2007 ¹²⁹	Incorrect reference standard
Valencia 2014 ¹³⁰	Incorrect study design
Vallone 2013 ¹³¹	Incorrect population
Van Appledorn 2003 ¹³²	Incorrect population
Van Beers 2001 ¹³³	Incorrect reference standard
Vieweg 1998 ¹³⁴	Incorrect reference standard
Viprakasit 2012 135	Time between tests, no usable outcomes
Vrtiska 1992 ¹³⁶	Incorrect population
Wang 2003 137	Incorrect reference standard
Wang 2008 138	Incorrect reference standard
Wang 2004 139	Incorrect reference standard
Wang 2017 140	Incorrect study design
Watkins 2007 141	Incorrect target condition
Westergreen 2017 ¹⁴²	No outcomes
Winkel 2012 ¹⁴³	Incorrect index test

Reference	Reason for exclusion
Wong 2001 144	Incorrect index test
Yap 2012 145	Incorrect population
Yavuz 2015 146	Incorrect index test
Yilmaz 1998 147	Incorrect reference standard
Zilberman 2011 148	Incorrect study design

H.2 Excluded health economic studies

None