## National Institute for Health and Care Excellence

Consultation

# Renal and ureteric stones: assessment and management

**Medical expulsive therapy** 

NICE guideline
Intervention evidence review
July 2018

Consultation

This evidence review was developed by the National Guideline Centre



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#### 1 Medical expulsive therapy

### 2 1.1 Review question: Is medical expulsive therapy clinically and cost-effective in managing people with ureteric

#### 4 stones?

#### 1.2 Introduction

Most acute stone episodes are initially treated with a period of observation as spontaneous passage of a stone often occurs. The passage of the stone is influenced by the size and site of the stone , the smaller stones <5mm having the greatest chance of stone passage along with stones in the distal ureter as this site is closest to the bladder. The majority of stones are expelled in 4-6 weeks but during this period the patient will often experience deterioration in quality of life, as they have concerns about episodes of severe pain and admission to hospital as well as the economic implications of not being able to work. There would therefore be considerable benefit to patients and the health system if this potential time to stone passage in suitable patients could be reduced by medical expulsive therapy which is the medication used to enhance the passage of stones or stone fragments . A similar benefit to promoting stone passage may also be present if medical expulsive therapy is used following active stone treatment, SWL and ureteroscopy to remove residual fragments. It has been shown that both alpha blockers and calcium channel blockers may have a role in medical expulsive therapy though there are no clear guidelines on their use in initial conservative management or following definitive stone treatment .

#### 21 1.3 PICO table

For full details see the review protocol in appendix A.

#### Table 1: PICO characteristics of review question

Population	People with ureteric stones
Intervention(s)	<ul> <li>Alpha blockers (Tamsulosin, Alfuzosin, Doxazosin, Silodosin, Naftopidil, Terazosin)</li> <li>Calcium channel blocker (Nifedipine)</li> </ul>
Comparison(s)	Compared to:  • Each other  • Placebo  • No treatment  • Steroids
Outcomes	<ul> <li>Critical outcomes:</li> <li>Time to stone passage</li> <li>Stone passage</li> <li>Use of healthcare services/Hospitalisation</li> <li>Quality of life</li> <li>Adverse events (hypotension, dizzy spells, falls, floppy iris, retrograde ejaculation, headaches, flushing)</li> <li>Important outcomes:</li> <li>Pain intensity (visual analogue scale, verbal ratings, descriptive scales, time to pain relief, need to rescue medication)</li> <li>Analgesic use</li> </ul>

Study design

- Randomised controlled trials (RCTs), systematic reviews of RCTs
- If no RCTs are available, non-randomised comparative studies (prospective and retrospective observational studies) will be included

#### 1 1.4 Clinical evidence

#### 1.4.1 Included studies

A search was conducted for randomised trials comparing the effectiveness of alpha blockers or calcium channel blockers versus each other, placebo, no treatment or steroids alone or as an adjunctive therapy to surgery for people with ureteric stones. Seventy studies (71 papers) were included in the review; 1, 3, 5-9, 14-17, 20-22, 24, 28-31, 41, 42, 48, 56, 57, 59-61, 63, 66-68, 74, 83, 88, 89, 91, 99, 109, 111, 120, 125, 131, 132, 134, 136, 139, 142, 144-146, 157, 162, 163, 166, 168, 171, 178, 179, 186, 188, 189, 191, 198, 200, 205, 207, 208, 210, 211, 214, 216 these are summarised in Table 2 below. Evidence from these studies is summarised in the clinical evidence summary below (Table 7-23).

In adults with distal ureteric stones <10mm, 7 studies compared alpha blockers versus calcium channel blockers, 31 studies compared alpha blockers versus no treatment, 13 studies compared alpha blockers versus placebo, 3 studies compared calcium channel blockers versus no treatment and 1 study compared calcium channel blockers versus placebo.

In adults with mid ureteric stones <10mm, 1 study compared alpha blockers versus calcium channel blockers, 1 study compared alpha blockers versus no treatment and 2 studies compared alpha blockers versus placebo. No evidence was identified comparing calcium blockers versus no treatment or placebo for mid ureteric stones.

In adults with proximal ureteric stones <10mm, 1 study compared alpha blockers versus calcium channel blockers, 3 studies compared alpha blockers versus no treatment and 2 studies compared alpha blockers versus placebo. No evidence was identified comparing calcium blockers versus no treatment or placebo for proximal ureteric stones.

Three studies compared alpha blockers versus no treatment, and 2 studies compared alpha blockers versus placebo in children with distal ureteric stones <10mm. No evidence was identified comparing alpha blockers versus calcium channel blockers, calcium blockers versus no treatment or calcium channel blockers versus placebo in children. No evidence was identified for mid or proximal ureteric stones in children.

No evidence was identified for medical expulsive therapy alone (not as an adjunct to surgery) for ureteric stones >10mm in adults or children.

In adults with distal ureteric stones, 6 studies compared alpha blockers as adjunctive therapy to surgery versus surgery only for stones <10mm, 1 study compared alpha blockers as adjunctive therapy to surgery versus surgery only for stones 10-20mm and 1 study compared alpha blockers as adjunctive therapy to surgery versus placebo and surgery for stones <10mm. No evidence was identified comparing alpha blockers versus calcium channel blockers as adjunctive therapy to surgery, or calcium channel blockers as adjunctive therapy to surgery versus placebo or surgery only.

In adults with mid ureteric stones 10-20mm, 1 study compared alpha blockers as adjunctive therapy to surgery versus surgery only. No evidence was identified for alpha blockers versus calcium channel blockers as adjunctive therapy to surgery, alpha blockers versus placebo as an adjunctive therapy to surgery, or calcium channel blockers as adjunctive therapy to surgery versus placebo or surgery only. No evidence was identified for mid ureteric stones <10mm.

- In adults with proximal ureteric stones, 6 studies compared alpha blockers as adjunctive therapy to surgery versus surgery only for stones <10mm, 4 studies compared alpha blockers as adjunctive therapy to surgery versus surgery only for stones 10-20mm, and 1 study compared alpha blockers as adjunctive therapy to surgery versus placebo and surgery for stones <10mm. No evidence was identified comparing alpha blockers versus calcium channel blockers as adjunctive therapy to surgery or calcium channel blockers as adjunctive therapy to surgery versus placebo or surgery only.
- No evidence was identified for medical expulsive therapy as an adjunctive therapy to surgery for ureteric stones in children.
- See also the study selection flow chart in appendix C, study evidence tables in appendix D, forest plots in appendix E and GRADE tables in appendix H.

#### 12 1.4.2 Excluded studies

See the excluded studies list in appendix I.

#### 14 1.4.3 Heterogeneity

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For the comparison of alpha blockers versus Calcium channel blockers for distal ureteric stones <10mm in adults, there was substantial heterogeneity between the studies when they were meta-analysed for the outcome of stone passage. For the comparison of alpha blockers versus no treatment (pain management only) for distal ureteric stones <10mm in adults, there was substantial heterogeneity between the studies when they were meta-analysed for the outcomes of time to stone passage, stone passage, pain intensity (number of pain episodes) and analgesic use (number of times and diclofenac dose). For the comparison alpha blockers versus placebo for distal ureteric stones <10mm in adults, there was substantial heterogeneity between the studies when they were meta-analysed for the outcomes of stone passage and analgesic use (number of people using analgesics and diclofenac dose). For the comparison alpha blockers versus no treatment (pain management only) for distal ureteric stones <10mm in children, there was substantial heterogeneity between the studies when they were meta-analysed for the outcome of time to stone passage. For the comparison alpha blockers versus placebo for distal ureteric stones <10mm in children, there was substantial heterogeneity between the studies when they were metaanalysed for the outcome of time to stone passage and pain intensity (daily pain episodes). For the comparison alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only for proximal ureteric stones <10mm in adults, there was substantial heterogeneity between the studies when they were meta-analysed for the outcomes pain intensity (VAS), time to stone passage and analgesic use (number of people using analgesia). Where pre-specified subgroup analyses (see Appendix A:) were either unable to be performed, or did not explain the heterogeneity, a random effects meta-analysis was applied to these outcomes, and the evidence was downgraded for inconsistency in GRADE.

#### 1.4.4 Summary of clinical studies included in the evidence review

Table 2: Summary of studies included in the evidence review

Study	Intervention and comparison	Population	Outcomes	Comments
Abdelaziz 2017 <sup>3</sup>	Intervention (n=51): Tamsulosin 0.4mg daily before URS for 1 week.  Concurrent medication/care: URS and NSAIDs  Comparison (n=47): ureterorenoscopy.  Concurrent medication/care: NSAIDs	n=98  People with a single, radio opaque, lower ureteral stone, 5-10mm in maximum diameter  Mean (SD) age: 36.27 (6.7)  Male to female ratio 64:34  Saudi Arabia	Stone passage (2 weeks)  Use of healthcare services/hospitalisation (2 weeks): defined as length of stay, days	
Abdel-Meguid 2010 <sup>1</sup>	Intervention (n=75): Tamsulosin 0.4mg oral tablets once daily. Duration up to 4 weeks. Concurrent medication/care: hydration and analgesia (diclofenac 100mg) as needed, patients with non-symptomatic urinary tract infections given antibiotics  Comparison (n=75): Placebo. Duration up to 4 weeks.  Concurrent medication/care: hydration and analgesia (diclofenac 100mg) as needed, patients with non-symptomatic urinary tract infections given antibiotics	n=150  People with newly diagnosed single, unilateral, distal ureteral 4-10mm stones  >18 years  Male to female ratio 103:47  Saudi Arabia	Stone passage (4 weeks)  Pain intensity (4 weeks): defined as number of pain episodes	
Agarwal 2009 <sup>5</sup>	Intervention (n=20): Tamsulosin 0.4mg daily starting just before the session of SWL. SWL performed a maximum of 4 sessions for any significant ureteric fragment, ureteroscopy offered if stone did not show adequate	n=40  People with a single upper ureteric stone <15mm electing SWL	Time to stone passage (5 weeks)  Stone passage (5 weeks)	Included 14 patients with stones <10mm, 20 with 10mm stones and 10 with stones >10mm. Included in

Study	Intervention and comparison	Population	Outcomes	Comments
	fragmentation after 2 sessions. Duration up to 3 months.  Concurrent medication/care: over-the-counter NSAIDs, antispasmodics or Tramadol on demand  Comparison (n=20): SWL performed a maximum of 4 sessions for any significant ureteric fragment, ureteroscopy offered if stone did not show adequate fragmentation after 2 sessions. Duration up to 5 weeks.  Concurrent medication/care: over-the-counter NSAIDs, antispasmodics or Tramadol on demand	Mean (SD) age: alpha blocker group 32.4 (8.7); SWL only group 35.5 (15.4)  Male to female ratio 31:9  India	Pain intensity (5 weeks): defined as visual analogue scale (0-10)	the <10mm stones analysis and downgraded for indirectness.
Agrawal 2009 <sup>6</sup>	Intervention (n=34): Tamsulosin 0.4mg once daily. Duration up to 4 weeks. Concurrent medication/care: instructions to drink at least 3L fluids daily, diclofenac injection (75mg) intramuscularly on demand  Intervention (n=34): Alfuzosin 10mg once daily. Duration up to 4 weeks. Concurrent medication/care: instructions to drink at least 3L fluids daily, diclofenac injection (75mg) intramuscularly on demand  Comparison (n=34): Placebo. Duration up to 4 weeks. Concurrent medication/care: instructions to drink at least 3L fluids daily, diclofenac injection (75mg) intramuscularly on demand	n=102  People with a stone <10mm located in the distal part of the ureter  15-60 years  Male to female ratio 78:24 India	Stone passage (4 weeks)  Adverse events (4 weeks): hypotension, retrograde ejaculation	
Ahmad 2015 <sup>7</sup>	Intervention (n=50): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac Sodium 50mg 8 hourly on required basis	n=100	Stone passage (4 weeks)  Use of healthcare services/hospitalisation (4	

Study	Intervention and comparison	Population	Outcomes	Comments
	Comparison (n=50): Placebo 1 capsule daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac Sodium 50mg 8 hourly on required basis.	People with a stone size 8mm or smaller in distal third of the ureter  >18 years  Gender not reported  Pakistan	weeks): defined as hospitalisation  Adverse events (4 weeks)  Analgesic use (4 weeks)	
Ahmed 2010 <sup>8</sup>	Intervention (n=29): Tamsulosin 0.4mg once	n=87	Stone passage (30 days)	
	daily. Duration up to 30 days. Concurrent medication/care: diclofenac injection (75mg) intramuscularly as needed (up to twice a day)  Intervention (n=30): Alfuzosin 10mg once daily. Duration up to 30 days. Concurrent medication/care: diclofenac injection (75mg) intramuscularly as needed (up to twice a day)  Comparison (n=28): no intervention. Duration up to 30 days. Concurrent medication/care: diclofenac injection (75mg) intramuscularly as needed (up to twice a day)	People with acute renal colic and a distal ureteral stone ≤10 mm  ≥18 years  Male to female ratio 56:31  Saudi Arabia	Time to stone passage  Use of healthcare services/hospitalisation (30 days): hospital readmission  Adverse events (30 days): retrograde ejaculation  Pain intensity (30 days): number of pain attacks	
Ahmed 2017 <sup>9</sup>	Intervention (n=91): Tamsulosin 0.4mg daily before ureteroscopy. Duration 1 week.	n=183	Stone passage (4 weeks)	
	Concurrent medication/care: not reported  Comparison (n=92): Ureteroscopy. Duration procedure time. Concurrent medication/care: not reported.	People with proximal ureteral stones ≥10mm scheduled for URS lithotripsy ≥18 years	Use of healthcare services/hospitalisation; (8 weeks): defined as initial procedure hospitalisation time	

Study	Intervention and comparison	Population	Outcomes	Comments
Study	mervention and comparison	Male to female ratio 98:67  Saudi Arabia	Outcomes	Comments
Al-Ansari 2010 <sup>14</sup>	Intervention (n=50): Tamsulosin 0.4mg once daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 75mg injection on demand and advice to drink a minimum of 2 L of water daily  Comparison (n=50): Placebo. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 75mg injection on demand and advice to drink a minimum of 2 L of water daily	n=100  People with ureteral stones 10mm or smaller located below the common iliac vessels as assessed on non-contrast computed tomography  >18 years  Male to female ratio 67:33  Qatar	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Adverse events (4 weeks)  Pain intensity (4 weeks)  Analgesic use (4 weeks)	
Aldaqadossi 2015 <sup>15</sup>	Intervention (n=33): Tamsulosin 0.4mg for patients >5 years and 0.2mg for younger patients. Duration up to 4 weeks. Concurrent medication/care: Ibuprofen 4-10mg/kg orally every 6-8 hours as needed; in the case of intractable pain, Ketorolac 0.5-1mg/kg intramuscularly  Comparison (n=34): Ibuprofen 4-10mg/kg every 6-8 hours as needed; in the case of intractable pain Ketorolac 0.5-1mg/kg intramuscularly. Duration 4 weeks.  Concurrent medication/care: NA	Children presenting with a distal ureteric stone of <1cm below the common iliac vessels as assessed by enhanced CT  Mean (SD) age: tamsulosin group: 7.7 years (3.02); pain management only (NSAIDs) group 7.25 years (2.7)  Male to female ratio 36:27  Egypt	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Adverse events (4 weeks)  Pain intensity (4 weeks)  Analgesic use (4 weeks)	

Study	Intervention and comparison	Population	Outcomes	Comments
Aldemir 2011 <sup>16</sup>	Intervention (n=31): Tamsulosin 0.4mg once daily. Duration up to 10 days. Concurrent	n=60	Stone passage (10 days)	
	medication/care: Diclofenac as needed and advice to drink at least 2 L of water daily	People with stones located in the distal ureter with a size of	Adverse events (10 days)	
	Comparison (n=29): Diclofenac 100mg once	<10mm in largest diameter	Pain intensity (10 days)	
	daily. Duration up to 10 days. Concurrent medication/care: advice to drink at least 2 L of water daily	>17 years	Analgesic use (10 days)	
	water daily	Male to female ratio 58:32		
		Turkey		
Alizadeh 2014 <sup>17</sup>	Intervention (n=50): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent	n=96	Time to stone passage (4 weeks)	
	medication/care: Indomethacin 100mg as needed and advice to drink 2 L of water daily	People with renal colic (3-6mm ureteral stone of distal ureteral or UVj)	Stone passage (4 weeks)	
	Comparison (n=46): Indomethacin 100mg as needed. Duration up to 4 weeks. Concurrent medication/care: advice to drink 2 L of water	18-60 years of age	Adverse events (4 weeks)	
	daily	Male to female ratio 61:35	Analgesic use (4 weeks)	
		Iran		
Arrabal-Martin 2010 <sup>20</sup>	Intervention (n=35): Tamsulosin 0.4mg daily. Duration 3 weeks. Concurrent	n=70	Stone passage (30 days)	
	medication/care: Ibuprofen 600mg every 12 hours, 2 L of water daily and Tramadol in	Age not reported	Adverse events (30 days)	
	case of pain	Gender not reported	Analgesic use (30 days)	
	Comparison (n=35): Ibuprofen 600mg every 12 hours. Duration 3 weeks. Concurrent medication/care: 2 L of water daily and Tramadol in case of pain	People with ureteral lithiasis below the S3 and S4 levels and a calculus size of 4-10mm		

Study	Intervention and comparison	Population	Outcomes	Comments
		Spain		
Ates 2012 <sup>21</sup>	Intervention (n=35): Doxazosin controlled release 4mg daily within 24 hours before SWL, if stone was not fragmented into pieces ≥6mm a second session was performed 3 days after the first procedure. Duration up to 14 days. Concurrent medication/care: oral Diclofenac on demand and advice to drink at least 2L of fluid daily  Comparison (n=44): SWL, if stone was not fragmented into pieces ≥6mm a second session was performed 3 days after the first procedure. Duration procedure time.  Concurrent medication/care: oral Diclofenac on demand and advice to drink at least 2 L of fluid daily	n=79  People with radio-opaque upper ureteral stones  Mean (SD) age: doxazosin + SWL group: 38.35 (11.41); SWL group: 30.95 (9.68)  Male to female ratio 58:21  Turkey	Time to stone passage (14 days)  Stone passage (14 days)  Use of healthcare services/hospitalisation (14 days)  Pain intensity (time-point unclear)  Analgesic use (14 days)	Included stones < and >10mm but mean diameter <10mm in both groups. Included in <10mm analysis and downgraded for indirectness.
Autorino 2005 <sup>22</sup> De Sio 2006 <sup>48</sup>	Intervention (n=32): Tamsulosin 0.4mg daily. Duration up to 2 weeks. Concurrent medication/care: Diclofenac 100mg daily, Aescin 80mg daily, advice to drink 2 L of water daily, Omeprazole 20mg daily for the treatment period and Levofloxacin 250mg daily for the first week  Comparison (n=32): Diclofenac 100mg daily and Aescin 80mg daily. Duration up to 2 weeks. Concurrent medication/care: advice to drink 2 L of water daily, Omeprazole 20mg daily for the treatment period and Levofloxacin 250mg daily for the first week	n=64  People with unilateral distal ureteral calculi  Mean (SD not reported) age: tamsulosin group: 45; NSAID group: 43  Male to female ratio 62:34  Italy	Time to stone passage (2 weeks)  Stone passage (4 weeks)  Use of healthcare services/hospitalisation (2 weeks)  Adverse events (2 weeks)  Analgesic use (2 weeks)	
Aydogdu 2009 <sup>24</sup>	Intervention (n=19): Doxazosin 0.03mg/kg once daily administered at bedtime. Duration up to 3 weeks. Concurrent medication/care:	n=39	Time to stone passage (3 weeks)	

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Study	Intervention and comparison	Population	Outcomes	Comments
	Comparison (n=25): Diclofenac 50mg when required. Duration up to 4 weeks. Concurrent medication/care: advice to drink 2-2.5 L of water daily			
Basri 2013 <sup>30</sup>	Intervention (n=59): Tamsulosin 0.4mg daily immediately after shock wave lithotripsy. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 75mg injected intramuscularly on demand, gastro protective therapy 40mg Pantoprazole once daily and instruction to drink a minimum of 2L of water daily  Comparison (n=64): Shock wave lithotripsy. Duration unclear. Concurrent medication/care: Diclofenac 75mg injected intramuscularly on demand, gastro protective therapy 40mg Pantoprazole daily and instruction to a minimum of 2L of water daily	n=123  People with solitary ureteral stone 6-15mm located in the upper, mid or lower ureter  Mean (SD) age: tamsulosin + SWL group: 44.66 (13.25); SWL group: 42.19 (13.17)  Male to female ratio 98:25  Turkey	Time to stone passage (4weeks)  Pain intensity (4 weeks)	Results for distal, mid and proximal ureteric stones analysed separately.  Included stones < and >10mm but mean stone size was >10mm. Included in 10-20mm stone analysis and downgraded for indirectness.
Bayraktar 2017 <sup>31</sup>	Intervention (n=60): Tamsulosin 0.4 mg daily. Duration up to 4 weeks.  Concurrent medication/care: recommended daily intake of liquids to urinate at least 1.5-2L, and 75mg diclofenac was injected when needed  Comparison (n=64): Diclofenac 75mg injected when needed. Duration up to 4 weeks.  Concurrent medication/care: recommended daily intake of liquids to urinate at least 1.5-2L	n=124  People with radiopaque distal ureter stones 5-10mm  Age >18 years  Males only  Turkey	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Analgesic use (4 weeks): defined as number of daily analgesic injections	
Chau 2011 <sup>41</sup>	Intervention (n=33): Alfuzosin slow release 10mg daily. Duration 4 weeks. Concurrent medication/care: Dologesic (Paracetamol + Dextropropoxyphene) four tablets daily on demand for 2 weeks and Diclofenac slow	n=67  People with acute ureteric stone 5-10mm	Stone passage (5 weeks)  Adverse events (5 weeks)	

Study	Intervention and comparison	Population	Outcomes	Comments
	release 100mg daily on demand for 2 weeks in case of suboptimal pain control by Dologesic	Mean (SD) age: 47.7 (12.3)  Male to female ratio 41:26		
	Comparison (n=34): Dologesic (Paracetamol + Dextropropoxyphene) four tablets daily on demand for 2 weeks and Diclofenac slow release 100mg daily on demand for 2 weeks in case of suboptimal pain control by Dologesic. Duration 2 weeks. Concurrent medication/care: not reported	China		
Cho 2013 <sup>42</sup>	Intervention (n=41): ESWL then Alfuzosin 10mg daily, if the ureter stone remained and was larger than 5mm at the next follow up visit then additional ESWL was performed. Duration up to 42 days. Concurrent medication/care: Loxoprofen 68.1mg as needed and recommendation to drink at least 2L hydration daily  Comparison (n=43): ESWL, if the ureter stone remained and was larger than 5mm at the next follow up visit then additional ESWL was performed. Duration up to 42 days.  Concurrent medication/care: Loxoprofen 68.1mg as needed and recommendation to drink at least 2L hydration daily	n=84  People with radio-opaque ureter stones; 5-10mm in diameter  Mean (SD) age: alfuzosin + SWL group: 47.4 (12.6); SWL 47.7 (12.1)  Male to female ratio 60:24  South Korea	Time to stone passage (42 days)  Stone passage (42 days)  Adverse events (42 days)  Pain intensity (time-point unclear)  Analgesic use (42 days)	Included distal and proximal stones, >80% were proximal stones. Included in proximal analysis and downgraded for indirectness.
El Said 2015 <sup>56</sup>	Intervention (n=28): Alfuzosin sustained release 5mg twice daily after meals. Duration up to 4 weeks. Concurrent medication/care: oral hydration with ≥2 L of water daily, Diclofenac 75mg intramuscularly on demand and education from the clinical pharmacist about potential adverse events, methods of reporting adverse events, self-reporting of pain on the visual analogue scale, importance	n=54  People presenting with radio- opaque stones ≤10mm and located in the distal third of the ureter  >18 years	Stone passage (4 weeks)  Use of healthcare services/hospitalisation (4 weeks)  Adverse events (4 weeks)	

Study	Intervention and comparison	Population	Outcomes	Comments
	of adherence to medications and daily water intake	Male to female ratio 34:20		
	Comparison (n=26): Oral hydration with ≥2 L of water daily and Diclofenac 75mg intramuscularly on demand. Duration up to 4 weeks. Concurrent medication/care: education by the clinical pharmacist on potential adverse events, methods of reporting adverse events, self-reporting of pain on the visual analogue scale, importance of adherence to medications and daily water intake	Egypt		
Elgalaly 2017 <sup>57</sup>	Intervention (n=20): Silodosin 4mg at bedtime. Duration up to 4 weeks. Concurrent medication/care: ibuprofen 20mg/kg/day was divided into two doses for pain episodes, fluids were encouraged  Comparison (n=20): Placebo taken at bedtime. Duration up to 4 weeks. Concurrent medication/care: ibuprofen 20mg/kg/day was divided into two doses for pain episodes, fluids were encouraged	n=40  Children with unilateral radiopaque distal ureteric stones <10mm  < 18 years  Male to female ratio 27:13  Egypt	Time to stone passage (4 weeks) (days)  Stone passage (4 weeks): defined as visual confirmation of stone passage  Pain intensity (4 weeks): defined as number of pain episodes	
Elkoushy 2012 <sup>59</sup>	Intervention (n=63): SWL repeated every 3 weeks until the patient became stone free, Tamsulosin 0.4mg daily starting immediately after SWL. Duration up to 3 months. Concurrent medication/care: Diclofenac 50mg tablets or 75mg intramuscular injection on demand	n=126  People with single radio- opaque renal or upper ureteral stones <2cm in largest diameter	Time to stone passage (3 months)  Stone passage (3 months)	Reports results for renal and proximal ureteric stones separately. Data extracted for ureteric stones only.
	Comparison (n=63): SWL repeated every 3 weeks until the patient became stone free,	Mean (SD) age: tamsulosin + SWL group: 52.8 (8.2); SWL + placebo group: 49.4 (11.3)		Included stones < and >10mm but mean stone diameter was

Study	Intervention and comparison	Population	Outcomes	Comments
	placebo daily starting immediately after SWL. Duration up to 3 months. Concurrent medication/care: Diclofenac 50mg tablets or 75mg intramuscular injection on demand	Male to female ratio 72:54 Egypt		<10mm. Included in <10mm analysis and downgraded for indirectness.
Erturhan 2007 <sup>61</sup>	Intervention (n=30): Tamsulosin 0.4mg daily. Duration up to 3 weeks. Concurrent medication/care: prophylactic antibiotic therapy (Ceforoxime axetil 250mg daily) and 2.5 L hydration daily, injectable Diclofenac (max 200mg/day) recommended for routine use during pain episodes  Comparison (n=30): Injectable Diclofenac (max 200mg/day) recommended for routine use during pain episodes. Duration up to 3 weeks. Concurrent medication/care: prophylactic antibiotic therapy (Cefuroxime axetil 250mg daily) and 2.5 L hydration daily	n=60  People with distal ureteral stones <10mm and allowing urinary flow  Mean (range) age: 31.5 (19-51)  Male to female ratio 64:56  Turkey	Stone passage (4 weeks)  Use of healthcare services/hospitalisation (4 weeks)  Adverse events (4 weeks)	
Erturhan 2013 <sup>60</sup>	Intervention (n=24): Doxazosin 0.03mg/kg daily. Duration up to 3 weeks. Concurrent medication/care: Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum 40mg/kg daily divided in to 4 equal doses in the case of intractable pain  Comparison (n=21): Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum of 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Duration up to 3 weeks. Concurrent medication/care: NA	n=45  People with a single radiopaque lower ureteral stone  Mean (SD) age: 6.65 (3.78)  Male to female ratio 24:26  Turkey	Stone passage (3 weeks)	
Eryildirim 2016 <sup>63</sup>	Intervention (n=40): SWL and Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 75mg if needed	n=80	Stone passage (4 weeks)	

Study	Intervention and comparison	Population	Outcomes	Comments
	Comparison (n=40): SWL. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 75mg if needed	People with 5-10mm single radio-opaque upper ureteral stones  Mean (SD) age: 39.41 (12.99)  Male to female ratio 36:18	Use of healthcare services/hospitalisation (4 weeks)  Quality of life (4 weeks)  Pain intensity (4 weeks)	
		Turkey	Analgesic use (4 weeks)	
Ferre 2009 <sup>66</sup>	Intervention (n=39): Tamsulosin 0.4mg daily. Duration 10 days. Concurrent medication/care: Ibuprofen 800mg 3 times a day and Oxycodone 5010mg every 4-6 hours as needed for pain  Comparison (n=41): Ibuprofen 800mg 3 times a day and Oxycodone 5-10mg every 4-6 hours as needed for pain. Duration up to 14 days. Concurrent medication/care: NA	n=80  People with CT confirmed diagnosis of a single calculus in the distal third of the ureter (distal to the internal iliac vessels) inconsistent with phleboliths as determined by a board-certified radiologist  ≥18 years of age  Male to female ratio 56:21  USA	Stone passage (14 days)  Use of healthcare service/hospitalisation s (14 days)  Adverse events (14 days)  Pain intensity (14 days)  Analgesic use (14 days)	
Furyk 2016 <sup>67</sup>	Intervention (n=198): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: analgesia at the discretion of the treating physician - recommended regimens were Indomethacin 25-50mg 3 times daily and Oxycodone 5-10mg 3 times daily as required for breakthrough	n=393  People with symptoms suggestive of ureteric colic; calculus demonstrated in the distal ureter (distal to the sacroiliac joint)	Stone passage (4 weeks)  Use of healthcare services/hospitalisation (4 weeks)	

Study	Intervention and comparison	Population	Outcomes	Comments
	Comparison (n=195): Placebo. Duration up to 4 weeks. Concurrent medication/care: analgesia at the discretion of the treating physician - recommended regimens were Indomethacin 25-50mg 3 times daily and Oxycodone 5-10mg 3 times daily as required for breakthrough	>18 years  Male to female ratio 320:73  Australia	Pain intensity (1, 2, 3 and 4 weeks)	
Gandhi 2013 <sup>68</sup>	Intervention (n=64): Nifedipine 30mg slow-release daily. Duration up to 4 weeks. Concurrent medication/care: oral prednisolone 30mg daily for a maximum of 10 days, Diclofenac 75mg intramuscularly on demand and ≥2 L of water daily  Comparison (n=64): Tamsulosin 0.4mg. Duration up to 4 weeks. Concurrent medication/care: oral prednisolone 30mg daily for a maximum of 10 days, Diclofenac 75mg intramuscularly on demand and ≥2 L of water daily	n=128  People with a solitary stone in the distal ureter at the juxtavesical tract or vesicoureteric junction of 5-15mm  Mean (SD) age nifedipine group: 30.4 (11.36); tamsulosin group; 34 (12.83)  Male to female ration nifedipine group 1.48:1; tamsulosin group 1.28:1  Nepal	Stone passage (4 weeks)  Adverse events (4 weeks)  Analgesic use (4 weeks)	Included stones < and >10mm but mean stone diameter was <10mm in both groups. Included in <10mm stones analysis and downgraded for indirectness.
Gravas 2007 <sup>74</sup>	Intervention (n=30): ESWL then Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: hydration of at least 2 L daily and Diclofenac 50mg on demand  Comparison (n=31): ESWL. Duration up to 4 weeks. Concurrent medication/care: hydration of at least 2 L daily and Diclofenac 50mg on demand	n=61  People with a single radiopaque distal ureteral stone (below the sacral-iliac joint), ≥6mm in diameter undergoing ESWL for the first time	Stone passage (4 weeks)  Adverse events (4 weeks)	Included stones < and >10mm but mean stone diameter was <10mm in both groups. Included in <10mm stones analysis and downgraded for indirectness.

Study	Intervention and comparison	Population	Outcomes	Comments
		Mean (range) age: tamsulosin + SWL group 48.8 (27-73); SWL group: 49.2 (30-72)		
		Male to female ratio 38:23		
		Greece		
Hermanns 2009 <sup>83</sup>	Intervention (n=50): Tamsulosin 0.4mg daily. Duration up to 3 weeks. Concurrent medication/care: after initial analgesia for acute pain management, no regular analgesic medication was maintained. Oral Diclofenac (up to 3 X 50mg) as first line and oral Metamizole (up to 4 X 1g) as second line ondemand analgesics were prescribed	n=100  People with acute renal colic with a single ureteral stone ≤7mm below the common iliac vessels as assessed by CT ≥18 years	Time to stone passage (3 weeks)  Stone passage (3 weeks)  Use of healthcare services/hospitalisation (3 weeks)	
	Comparison (n=50): Placebo. Duration up to 3 weeks. Concurrent medication/care: after initial analgesia for acute pain management, no regular analgesic medication was maintained. Oral Diclofenac (up to 3 X 50mg) as first-line and oral Metamizole (up to 4 X 1g) as second-line on demand analgesics were prescribed	Male to female ratio 75:15  Switzerland	Adverse events (3 weeks)	
Islam 2012 <sup>89</sup>	Intervention (n=33): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: prophylactic antibiotic therapy (Ciprofloxacin 500mg twice daily), 2.5 L hydration daily and Diclofenac recommended for routine use during pain episodes  Comparison (n=33): Nifedipine 20mg (slow	n=98  People with distal ureteral stones (juxtavesical tract and ureterovesical junction) ≤1cm in size  Mean (SD not reported) age:	Stone passage (4 weeks)  Use of healthcare services/hospitalisation (4 weeks)  Adverse events (4 weeks)	
	release) daily. Duration up to 4 weeks. Concurrent medication/care: prophylactic	tamsulosin group: 46.6;		

Study	Intervention and comparison	Population	Outcomes	Comments
	antibiotic therapy (Ciprofloxacin 500mg twice daily), 2.5 L hydration daily and Diclofenac recommended for routine use during pain episodes  Comparison (n=32): No treatment. Duration up to 4 weeks. Concurrent medication/care: prophylactic antibiotic therapy (Ciprofloxacin 500mg twice daily), 2.5 L hydration daily and Diclofenac recommended for routine use during pain episodes	nifedipine group 47.4; no treatment group: 42.8 Male to female ratio 58:33 Bangladesh		
Ibrahim 2013 <sup>88</sup>	Intervention (n=50): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: diclofenac potassium 50mg given orally and/or diclofenac sodium 75mg given intramuscularly. Duration up to 4 weeks.  Intervention (n=50): Alfuzosin 10mg daily. Duration up to 4 weeks. Concurrent medication/care: diclofenac potassium 50mg given orally and/or diclofenac sodium 75mg given intramuscularly. Duration up to 4 weeks.  Comparison (n=50): Diclofenac potassium 50mg given orally and/or diclofenac sodium 75mg given intramuscularly. Duration up to 4 weeks.	n=150  People with symptomatic ureteric stone or <10mm >18 years  Male to female ratio 91: 21  Iraq	Stone passage (4 weeks): not defined	Included proximal, mid and distal ureteral stones and results were reported separately
Itoh 2011 <sup>91</sup>	Intervention (n=89): Silodosin 8mg daily. Duration up to 8 weeks. Concurrent medication/care: instruction to drink 2 L of water daily	n=187  People with symptomatic unilateral ureteral calculi <10mm in diameter	Time to stone passage (8 weeks)  Stone passage (8 weeks)	Included proximal, mid and distal ureteral stones and results were reported separately

Study	Intervention and comparison	Population	Outcomes	Comments
	Comparison (n=92): No treatment. Duration up to 8 weeks. Concurrent medication/care: instruction to drink 2 L of water daily	Mean (SD) age: silodosin group: 57.2 (12.7); no treatment group: 56.5 (10.1)  Male participants only  Japan	Analgesic use (8 weeks): number of times of analgesic use	
Ketabchi 2014 <sup>99</sup>	Intervention (n=52): Tamsulosin 0.4mg daily starting one day before URS. Duration up to 2 weeks. Concurrent medication/care: recommendation to drink 2 L of water daily, those with moderate to severe pain (>5 VAS) consumed Pethidine 25mg intravenously after the procedure in the recovery room and Indomethacin 500mg suppository daily  Comparison (n=50): Placebo daily starting one day before URS. Duration up to 2 weeks. Concurrent medication/care: recommendation to drink 2 L of water daily, those with moderate to severe pain (>5 VAS) consumed Pethidine 25mg intravenously after the procedure in the recovery room and Indomethacin 500mg suppository daily	n=102  People with a single radio opaque lower ureteral stone with 5-10mm diameter  Mean (SD) age: tamsulosin + URS group: 24 (6.5); placebo + URS group: 27 (8.8)  Male to female ratio 77:25  Iran	Stone passage (2 weeks)  Pain intensity (2 weeks)  Analgesic use (2 weeks)	
Kupeli 2004 <sup>109</sup>	Intervention (n=15): Tamsulosin 0.4mg daily. Duration 15 days. Concurrent medication/care: conventional treatment - oral hydration and oral Diclofenac 100mg daily  Comparison (n=15): Oral Diclofenac 100mg daily. Duration 15 days. Concurrent medication/care: oral hydration  Comparison (n=24): Tamsulosin 0.4mg daily beginning after shock wave lithotripsy.	n=78  People with lower ureteral stones within the distal 5cm of the ureter that ranged between 3 and 15mm in size  Mean (range) age: 42.9 (21-67)	Stone passage (15 days)  Adverse events (15 days)	Stone size <5mm given Tamsulosin or conventional treatment, stone size 6-15mm given SWL + conventional treatment or SWL + Tamsulosin + conventional treatment

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Study	Intervention and comparison	Population	Outcomes	Comments
	Duration 15 days. Concurrent medication/care: conventional treatment – oral hydration and oral Diclofenac100mg daily  Comparison (n=24): Shock wave lithotripsy. Concurrent medication/care: conventional treatment – oral hydration and oral Diclofenac 100mg daily	Male to female ratio 56:22 Turkey		Adjunctive therapy groups included 3 patients with stones >10mm. Included in the <10mm stones analysis and downgraded for indirectness.
Lee 2014 <sup>111</sup>	Intervention (n=54): Tamsulosin 0.2mg daily. Duration up to 4 weeks. Concurrent medication/care: instruction to drink 2 L of water daily and oral painkiller (Ultracet® combination of Tramadol and Acetaminophen) on demand  Comparison (n=54): No treatment. Duration up to 4 weeks. Concurrent medication/care: instruction to drink 2 L of water daily and oral painkiller (Ultracet® combination of Tramadol and Acetaminophen) on demand	n=108  People presenting with renal colic, with single, unilateral radiopaque, proximal ureteral calculi ≤6mm in diameter  ≥18 years  Male to female ratio 68:40  South Korea	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Quality of life (4 weeks): EuroQoL  Analgesic use (4 weeks): requirement of oral analgesics	
Lojanapiwat 2008 <sup>120</sup>	Intervention (n=50): Tamsulosin 0.2mg or 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 50mg twice daily for 10 days and Diclofenac 75mg infection if renal colic developed during treatment  Comparison (n=25): Diclofenac 50mg twice daily. Duration 10 days. Concurrent medication/care: Diclofenac 75mg injection if renal colic developed	n=75  People with distal ureteric stones of 4-10mm; measured by plain KUB; gave informed consent; interviewed prior to taking part  Mean (SD) age: tamsulosin 0.2mg group: 48 (15.74); tamsulosin 0.4mg group: 46.71 (12.2); pain management only (NSAID): 46.52 (13.63)	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Adverse events (4 weeks): hypotension; retrograde ejaculation  Analgesic use (4 weeks): number of people using analgesia	

Study	Intervention and comparison	Population	Outcomes	Comments
		Male to female ratio 55:20 Thailand		
Lv 2014 <sup>125</sup>	Intervention (n=35): Naftopidil 50mg daily. Duration up to 2 weeks. Concurrent medication/care: instruction to drink at least 2 L of fluids daily  Comparison (n=35): Naftopidil 50mg daily and Celecoxib 400mg immediately then 200mg every 12 hours. Duration up to 2 weeks. Concurrent medication/care: instruction to drink at least 2 L of fluids daily  Comparison (n=33): Celecoxib 400mg immediately then 200mg every 12 hours. Duration up to 2 weeks. Concurrent medication/care: instruction to drink at least 2 L of fluids daily	n=105  People with a distal ureteral stone 4-9mm  Mean (SD) age: naftopidil group: 31.4 (2.94); naftopidil + celecoxib group: 33.2 (5.28); celecoxib group: 33.75 (5.24)  Male to female ratio 59:44  China	Time to stone passage (2 weeks)  Stone passage (2 weeks)  Adverse events (2 weeks): headache; retrograde ejaculation  Pain intensity (2 weeks): defined as number of pain episodes; visual analogue scale	
Mokhless 2012 <sup>132</sup>	Intervention (n=33): Tamsulosin 0.4mg for age ≥4 years and 0.2 mg for age <4 years. Duration up to 4 weeks. Concurrent medication/care: standard analgesia (ibuprofen)  Comparison (n=28): Placebo. Duration up to 4 weeks. Concurrent medication/care: standard analgesia (ibuprofen)	n=61  Children with radiopaque lower ureteral stones of 12mm or smaller  Mean (SD) age: 8.1 (6.8)  Male to female ratio 36:25  Egypt	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Adverse events (4 weeks): hypotension, headache  Pain intensity (4 weeks): defined as number of pain episodes	

Study	Intervention and comparison	Population	Outcomes	Comments
			Analgesia use (4 weeks): need for analgesia	
Mohseni 2006 <sup>131</sup>	Intervention (n=32): Indomethacin. Duration up to 4 weeks. Concurrent medication/care: intravenous Pethidine in cases of incomplete pain control  Comparison (n=32): Terazosin 10mg daily. Duration up to 4 weeks. Concurrent medication/care: Indomethacin and intravenous Pethidine in cases of incomplete pain control	n=64  People with a lower ureteral stone  Mean (SD) age: terazosin group: 44.2 (12.9); indomethacin group: 39.3 (14.2)  Male to female ratio 44:20  Iran	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Adverse events (4 weeks): hypotension  Analgesic use (4 weeks): pain analgesia dose	
Moursy 2010 <sup>134</sup>	Intervention (n=44): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: Indomethacin 100mg suppository on demand and encouragement to drink a minimum 2.5 L of water daily  Comparison (n=44): Pain management only. Duration up to 4 weeks. Concurrent medication/care: Indomethacin 100mg suppository on demand and encouragement to drink a minimum 2.5 L of water daily	n=88  People with unilateral steinstrasse after SWL  >18 years  Mean (SD) age: tamsulosin group: 35.6 (9.95); pain management only group: 33.9 (9.71)  Egypt	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Use of healthcare services/hospitalisation (4 weeks)  Adverse events (4 weeks)  Analgesic use (4 weeks)	
Mustafa 2016 <sup>136</sup>	Intervention (n=64): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: conventional treatment -	n=128	Stone passage (4 weeks)	

Study	Intervention and comparison	Population	Outcomes	Comments
	hydration with minimum 2 L of water daily, physical exertion and analgesics (Diclofenac 50mg suppository with H2 blocker) if required  Comparison (n=64): No treatment. Duration up to 4 weeks. Concurrent medication/care: hydration with minimum 2 L of water daily, physical exertion and analgesics (Diclofenac 50mg suppository with H2 blocker) if required	People with unilateral, juxtavesical ureteral stone; normal functioning kidney; absence of clinical and laboratory signs of urinary tract infection; stone size up to 8mm >18 years  Gender not reported  Bangladesh	Pain intensity (4 weeks): defined as number of pain episodes	
Ochoa-Gomez 2011 <sup>139</sup>	Intervention (n=32): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: instruction to drink at least 2 L of water daily  Comparison (n=33): Placebo. Duration up to 4 weeks. Concurrent medication/care: instruction to drink at least 2 L of water daily	n=65  People with reno-ureteral stones 5-10mm determined by plain abdominal film and kidney ultrasound  >18 years  Male to female ratio 36:29  Mexico	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Adverse events (4 weeks): dizziness, retrograde ejaculation	
Park 2013 <sup>142</sup>	Intervention (n=48): Tamsulosin 0.2mg once daily, starting just before ESWL. Duration up to 3 weeks. Concurrent medication/care: Aceclofenac 100mg on demand and asked to drink 1.5-2L of water daily  Comparison (n=48): ESWL. Duration up to 3 weeks. Concurrent medication/care:	n=96  People with symptomatic, unilateral, single, proximal ureteral stone 6-20mm in longest axis  18-70 years	Stone passage (3 weeks)  Adverse events (3 weeks)	

Study	Intervention and comparison	Population	Outcomes	Comments
	Aceclofenac 100mg on demand and asked to drink 1.5-2L of water daily	Male to female ratio 57:31  South Korea		
Pedro 2008 <sup>144</sup>	Intervention (n=34): Alfuzosin daily. Duration up to 4 weeks. Concurrent medication/care: not reported  Placebo (n=35). Duration up to 4 weeks. Concurrent medication/care: not reported	n=69  People with a distal ureteral calculus  Mean (SD) age: alfuzosin group: 36.69 (13.06); placebo group: 42.03 (12.85)  Male to female ratio 55:14  USA	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Adverse events (4 weeks)  Analgesic use (4 weeks)	
Pickard 2015 <sup>145,</sup> 146	Intervention (n=391): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: standard care - analgesics, antiemetics, advice on adequate fluid intake and resumption of normal activity  Comparison (n=387): Nifedipine 30mg daily. Duration up to 4 weeks. Concurrent medication/care: standard care - analgesics, antiemetics and advice on adequate fluid intake and resumption of normal activity  Comparison (n=389): Placebo. Duration up to 4 weeks. Concurrent medication/care: standard care - analgesics, antiemetics and advice on adequate fluid intake and resumption of normal activity	n=1167  People presenting acutely with ureteric colic, with a stone ≤ 10 mm confirmed by noncontrast CT KUB, within any segment of the ureter  ≥ 18 years to ≤ 65 years  Male to female ratio 931:219  UK	Stone passage (4 weeks)	Included proximal, mid and distal ureteric stones. Results reported separately.

Study	Intervention and comparison	Population	Outcomes	Comments
Rahim 2012 <sup>157</sup>	Intervention (n=45): Terazosin 2mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 50mg twice daily  Comparison (n=45): Diclofenac 50mg twice daily. Duration up to 4 weeks. Concurrent medication/care: NA	n=90  People with 4-7mm stones in the distal segment of the ureter confirmed on ultrasound  16-63 years  Male to female ratio 63:27  Pakistan	Time to stone passage (4 weeks)  Stone passage (4 weeks)	
Resim 2005 <sup>162</sup>	Intervention (n=30): Tamsulosin 0.4mg daily. Duration up to 6 weeks. Concurrent medication/care: conservative treatment - hydration and Tenoxicam 20mg daily  Comparison (n=30): Conservative treatment - hydration and Tenoxicam 20mg daily. Duration up to 6 weeks. Concurrent medication/care: NA	n=60  People with lower ureteral calculi  Mean (SD) age: tamsulosin group: 35.3 (10.9); pain management only (NSAID): 33.5 (9.7)  Male to female ratio 45:15  Turkey	Stone passage (6 weeks)  Adverse events (6 weeks): headache, dizziness, abnormal ejaculation, hypotension	Included stones < and >10mm but mean stone diameter in both groups was <10mm. Included in <10mm analysis and downgraded for indirectness.
Resim 2005 <sup>163</sup>	Intervention (n=32): Tamsulosin 0.4mg daily. Duration up to 6 weeks. Concurrent medication/care: hydration and Tenoxicam 20mg daily  Comparison (n=35): Pain management only. Duration up to 6 weeks. Concurrent medication/care: hydration and Tenoxicam 20mg daily	n=67  People with steinstrasse in the lower ureter (juxtavesical or intramural portion) after undergoing ESWL  ≥ 18 years	Stone passage (6 weeks)  Adverse events (6 weeks)	

Study	Intervention and comparison	Population	Outcomes	Comments
		Male to female ratio 43:24  Turkey		
Sameer 2014 <sup>166</sup>	Intervention (n=35): Nifedipine 30mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 50mg every 12 hours for 1 week, Diclofenac 75mg injection as needed and Tramadol 100mg injection for persistent pain  Intervention (n=35): Alfuzosin 10mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 50mg every 12 hours for 1 week, Diclofenac 75mg injection as needed and Tramadol 100mg injection for persistent pain  Comparison (n=35): Diclofenac 50mg every 12 hours for 1 week. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 75mg injection as needed and Tramadol 100mg injection for persistent pain	n=105  People with single, unilateral ureteral stone of ≤10mm; distal defined as the segment from the lower border of the sacroiliac joint to the vesicoureteric junction  ≥8 years  Male to female ratio: 68:37  India	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Use of healthcare services/hospitalisation (4 weeks): re-admission  Pain intensity (4 weeks): defined as number of pain episodes	
Sayed 2008 <sup>168</sup>	Intervention (n=45): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: hydration (at least 2 L of water daily) and Diclofenac 100mg injection on demand  Comparison (n=45): No treatment. Duration up to 4 weeks. Concurrent medication/care: hydration (at least 2 L of water daily) and Diclofenac 100mg injection on demand	n=90  People with radiopaque stones 5-10mm in diameter in the distal ureter  >18 years  Male to female ratio 69:21  Egypt	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Adverse events (4 weeks): unspecified  Pain intensity (4 weeks): defined as number of pain episodes	

Study	Intervention and comparison	Population	Outcomes	Comments
			Analgesic use (4 weeks): number of times analgesic was used	
Sen 2017 <sup>171</sup>	Intervention (n=25): Doxasozin 4mg. Duration up to 3 weeks. Concurrent medication/care: diclofenac 100mg and daily 1500-2000 cc hydration  Intervention (n=22): Doxasozin 8mg. Duration up to 3 weeks. Concurrent medication/care: diclofenac 100mg and daily 1500-2000 cc hydration  Comparison (n=19): Diclofenac 100mg up to 3 weeks. Concurrent medication/care: daily 1500-2000 cc hydration	n=66  People with radio-opaque distal ureteral stones ≤10mm  Mean (SD) age: doxazosin group: 33.7 (10.4); pain management only (NSAID): 33 (11.3)  Male to female ratio 46:20  Turkey	Time to stone passage (3 weeks)  Stone passage (3 weeks)  Pain intensity (3 weeks): defined as number of pain episodes  Adverse events (3 weeks): hypotension	
Singh 2011 <sup>179</sup>	Intervention (n=59): Tamsulosin 0.4mg daily beginning just before the session of SWL, SWL repeated every 3 weeks for incomplete fragmented calculus. Duration up to 3 months. Concurrent medication/care: advice to drink 2.5L of fluid daily and Diclofenac on demand  Comparison (n=58): SWL repeated every 3 weeks for incomplete fragmented calculus up to 3 sessions. Duration up to 3 months. Concurrent medication/care: advice to drink 2.5L of fluid daily and Diclofenac on demand	n=120  People with symptomatic, unilateral and solitary upper (between the peli-ureteral junction and sacroiliac joint) ureteral calculi 6-15mm in major axis  18-70 years  Gender not reported  India	Time to stone passage (3 months)  Stone passage (3 months)  Pain intensity (3 months)	Results for stones 6- 10mm and 11-15mm analysed separately for primary outcome (stone passage). Included in the <10mm stones analysis and downgraded for indirectness for other outcomes.
Singh 2011 <sup>178</sup>	Intervention (n=60): Tamsulosin 0.4mg daily from the day of ESWL just before the session.	n=120	Time to stone passage (4 weeks)	Included stones < and >10mm but the

Study	Intervention and comparison	Population	Outcomes	Comments
	Duration up to 4 weeks. Concurrent medication/care: advice to drink 2.5L of fluid daily, antibiotics and Diclofenac on demand  Comparison (n=59): ESWL and placebo.  Duration up to 4 weeks. Concurrent medication/care: advice to drink 2.5L of fluid daily, antibiotics and Diclofenac on demand	People with symptomatic unilateral solitary lower ureteric calculus 4-12mm in major axis >18 years  Male to female ratio 84:35  India	Stone passage (4 weeks)  Analgesic use (4 weeks)	majority were <10mm. Included in the <10mm stones analysis and downgraded for indirectness.
Su 2016 <sup>186</sup>	Intervention (n=76): Tamsulosin 0.4mg daily. Duration up to 2 weeks. Concurrent medication/care: Ketorolac 10mg three times daily, Buprenorphine 0.2mg on demand and encouragement to drink a minimum of 2 L of water daily  Intervention (n=79): Silodosin 8mg daily. Duration up to 2 weeks. Concurrent medication/care: Ketorolac 10mg three times daily, Buprenorphine 0.2mg on demand and encouragement to drink a minimum of 2 L of water daily  Comparison (n=82): Placebo. Duration up to 2 weeks. Concurrent medication/care: Ketorolac 10mg three times daily, Buprenorphine 0.2mg on demand and encouragement to drink a minimum of 2 L of water daily	n=272  People with radiopaque distal ureteral stones <10mm  Mean (SD) age: tamsulosin group: 50.74 (10.08); silodosin group: 51.58 (8.27); placebo group: 52.16 (9.2)  Male to female ratio 122:82  Taiwan	Time to stone passage (2 weeks)  Stone passage (2 weeks)  Adverse events (2 weeks)  Analgesic use (2 weeks)	
Sun 2009 <sup>188</sup>	Intervention (n=30): Naftopidil 50mg daily. Duration up to 2 weeks. Concurrent medication/care: instruction to drink a minimum of 2 L of water daily and	n=60  People with unilateral distal (below the lower border of the sacroiliac joint) ureteral stones	Stone passage (2 weeks)  Use of healthcare services/hospitalisation (2 weeks)	

Study	Intervention and comparison	Population	Outcomes	Comments
	Indomethacin suppository to control acute episodes of ureteral colic if present  Comparison (n=30): Watchful waiting.  Duration up to 2 weeks. Concurrent medication/care: instruction to drink a minimum of 2 L of water daily and Indomethacin suppository used to control acute episodes of ureteral colic if present	18-65 years Male to female ratio 50:10 China	Adverse events (2 weeks): dizziness  Pain intensity (2 weeks): defined as episodes of renal colic	
Sur 2015 <sup>189</sup>	Intervention (n=115): Silodosin 8mg. Duration up to 4 weeks. Concurrent medication/care: Oxycodone 5mg to provide analgesia for renal colic and us concomitant pre-enrolment medications that would not confound study results  Comparison (n=117): Placebo. Duration up to 4 weeks. Concurrent medication/care: Oxycodone 5mg to provide analgesia for renal colic and use of other concomitant pre-enrolment medications that would not confound study results	n=239  People with unilateral calculus ≥4mm and ≤10mm in any location of the ureter ≥18 years  Male to female ratio 152:87  USA	Stone passage (4 weeks): visualisation of the stone or imaging  Adverse events (4 weeks): retrograde ejaculation, dizziness, headache	
Thapa 2014 <sup>191</sup>	Intervention (n=35): Tamsulosin 0.4mg daily. Duration up to 3 weeks. Concurrent medication/care: advice to have high fluid intake more than 3 L daily and Diclofenac 50mg 3 times daily for 5 days, then on demand  Comparison (n=35): Diclofenac 50mg 3 times daily for 5 days, then on demand. Duration up to 3 weeks. Concurrent medication/care: advice to have high fluid intake more than 3 L daily	n=70  People with symptomatic, unilateral, solitary lower ureteral stones (located below sacroiliac joint) of 5-10mm  >15 years  Male to female ratio 41:29  Nepal	Stone passage (3 weeks)	

Study	Intervention and comparison	Population	Outcomes	Comments
Wang 2008 <sup>198</sup>	Intervention (n=32): Tamsulosin 0.4mg daily. Duration up to 2 weeks. Concurrent medication/care: Ketorolac 10mg 3 times daily, sublingual Buprenorphine 0.2mg as needed and a minimum of 2 L of water daily  Intervention (n=32): Terazosin 2mg daily. Duration up to 2 weeks. Concurrent medication/care: Ketorolac 10mg 3 times daily, sublingual Buprenorphine 0.2mg on demand and a minimum of 2 L of water daily  Comparison (n=31): Ketorolac 10mg 3 times daily. Duration up to 2 weeks. Concurrent medication/care: sublingual Buprenorphine 0.2mg as needed and a minimum of 2 L of water daily	n=95  People with radiopaque lower ureteral stones  Mean (SD) age: tamsulosin group: 50.4 (9.7); terazosin group: 51.4 (8.6); pain management only (NSAID) group: 50.9 (9.6)  Male to female ratio 66:29  China	Time to stone passage (2 weeks)  Stone passage (2 weeks)  Adverse events (2 weeks): unspecified  Pain intensity (2 weeks): defined as number of colic episodes  Analgesic use (2 weeks): average pain relief consumption (mg)	
Wang 2014 <sup>205</sup>	Intervention (n=48): Tamsulosin 0.4mg daily after URS. Duration up to 6 weeks. Concurrent medication/care: 2-3L hydration and Diclofenac 75mg on demand  Comparison (n=46): URS only. Duration up to 6 weeks. Concurrent medication/care: 2-3L hydration and Diclofenac 75mg on demand	n=94  People with symptomatic stone; 10-15mm in size; located in the proximal ureter (between the ureteropelvic junction and sacroiliac joint); associated with moderate hydroureteronephrosis  Age not reported  Gender not reported  China	Time to stone passage (6 weeks)  Stone passage (6 weeks)  Adverse events (6 weeks)  Pain intensity (6 weeks)	
Wang 2016 <sup>200</sup>	Intervention (n=71): Silodosin 8mg daily. Duration up to 2 weeks. Concurrent	n=141	Time to stone passage (2 weeks)	

Study	Intervention and comparison	Population	Outcomes	Comments
	medication/care: Ketorolac three times daily, sublingual Buprenorphine 0.2mg on demand and encouragement to drink a minimum of 2 L of water daily  Comparison (n=70): Placebo. Duration up to 2 weeks. Concurrent medication/care: Ketorolac 10mg three times daily, sublingual Buprenorphine 0.2mg on demand and encouragement to drink a minimum of 2 L of water daily	People with radiopaque distal stones <10mm  28-72 years  Male to female ratio 83:40  Taiwan	Stone passage (2 weeks): no residual fragments  Adverse events(2 weeks): unspecified  Pain intensity (2 weeks): defined as number of renal colic episodes  Analgesic use (2 weeks): average pain relief consumption (mg)	
Ye 2011 <sup>207</sup>	Intervention (n=1596): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: encouragement to maintain a water intake of 2-2.5 L daily, Levofloxacin 0.2g twice daily and Diclofenac 50mg suppository on demand  Comparison (n=1593): Nifedipine 10mg 3 times daily. Duration up to 4 weeks. Concurrent medication/care: encouragement to maintain a water intake of 2-2.5 L daily, Levofloxacin 0.2g twice daily and Diclofenac 50mg suppository on demand	n=3189  People with emergency admission for renal colic; radiopaque or radiolucent single distal ureteric stone (juxtavesical or intramural portion) of 4-7mm  18-50 years  Male to female ratio 1987:1202  China	Stone passage (4 weeks): stone free on non-contrast CT  Adverse events (4 weeks): not specified  Analgesic use (4 weeks): number of participants using pain relief therapy	
Ye 2018 <sup>208</sup>	Intervention (n=1695): Tamsulosin 0.4mg (two capsules of 0.2mg). Duration until spontaneous stone passage, up to 28 days. Concurrent medication/care: 2L water per	n=3390	Time to stone passage (28 days)  Stone passage (28 days)	

Study	Intervention and comparison	Population	Outcomes	Comments
	day. 50mg sodium diclofenac suppository on demand  Comparison (n=1695): Placebo. Duration until spontaneous stone passage, up to 28 days. Concurrent medication/care: 2L water per day. 50mg sodium diclofenac suppository on demand	People with a stone in the distal ureter with a dimension of 4-7mm  18-60 years  Male to female ratio 2135:1161  China	Adverse events (28 days): retrograde ejaculation, dizziness, headache  Pain intensity (28 days): defined as rate of pain relief therapy  Analgesic use (28 days): average dose of diclofenac	
Yilmaz 2005 <sup>210</sup>	Intervention (n=28): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: symptomatic therapy with Diclofenac 75mg injections on demand and consumption of a minimum of 2 L of water daily  Intervention (n=28): Terazosin 5mg daily. Duration up to 4 weeks. Concurrent medication/care: symptomatic therapy with Diclofenac 75mg injections on demand and consumption of a minimum of 2 L of water daily  Intervention (n=29): Doxazosin 4mg daily. Duration up to 4 weeks. Concurrent medication/care: symptomatic therapy with Diclofenac 75mg injections on demand and consumption of a minimum of 2 L of water daily	n=114  People with radiopaque stones ≤10mm located in the distal tract of the ureter (juxtavesical tract and ureterovesical junction)  18-65 years old  Male to female ratio 46:68  Turkey	Time to stone passage (4 weeks)  Stone passage (4 weeks)  Adverse events (4 weeks): unspecified  Pain intensity (4 weeks): defined as number of pain episodes  Analgesic use (4 weeks): analgesic dose required	

Study	Intervention and comparison	Population	Outcomes	Comments
	Comparison (n=28): Symptomatic therapy with Diclofenac 75mg injections on demand. Duration up to 4 weeks. Concurrent medication/care: consumption of a minimum of 2 L of water daily			
Yuksel 2015 <sup>211</sup>	Intervention (n=35): Silodosin 4mg daily. Duration up to 3 weeks. Concurrent medication/care: Diclofenac 75mg daily as necessary, advice to remain active and drink at least 2 L of water daily  Comparison (n=35): Diclofenac 75mg daily as necessary. Duration up to 3 weeks. Concurrent medication/care: advice to remain active and drink at least 2 L of water daily	n=70  People with a distal ureteral stone 4-10mm  18-65 years old  Male to female ratio 39:31  Turkey	Time to stone passage (3 weeks)  Stone passage (3 weeks)  Pain intensity (3 weeks): defined as renal colic episodes  Analgesic use (3 weeks): analgesic dosage	
Zhang 2009 <sup>214</sup>	Intervention (n=102): Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: 2.5 L hydration daily, Levofloxacin 0.1g twice daily for the first 7 days and Diclofenac 75mg injection daily if needed  Comparison (n=97): Nifedipine 30mg 3 times daily. Duration up to 4 weeks. Concurrent medication/care: 2.5 L hydration daily, Levofloxacin 0.1g twice daily for the first 7 days and Diclofenac 75mg injection daily if needed	n=199  People with distal ureteral stones  Mean (SD) age: tamsulosin group: 34.6 (11.4); nifedipine group: 36.3 (9.7)  Male to female ratio 131:58  China	Stone passage (4 weeks): absence of any stone on x-ray	
Zhou 2011 <sup>216</sup>	Intervention (n=43): Naftopidil 10mg daily. Duration up to 2 weeks. Concurrent medication/care: instruction to drink at least 2 L of fluids daily and an Indomethacin	n=131 People with distal ureteral stones ≤9mm to >4mm	Time to stone passage (2 weeks)  Stone passage (2 weeks)	

Comments

5

# Quality assessment of clinical studies included in the evidence review

Intervention and comparison

Duration up to 2 weeks. Concurrent

Comparison (n=43): Watchful waiting. Duration up to 2 weeks. Concurrent

episodes

episodes

episodes

See appendix D for full evidence tables.

suppository recommended for use during pain

Comparison (n=45): Tamsulosin 0.4mg daily.

medication/care: instruction to drink at least 2

L of fluids daily and Indomethacin suppository recommended for routine use during pain

medication/care: instruction to drink at least 2 L of fluids daily and Indomethacin suppository recommended for routine use during pain

#### 4 1.4.5.1 Distal ureteric stones, <10mm, adults

Study

Table 3: Clinical evidence summary: Alpha blockers versus placebo for distal ureteric stones <10mm in adults

	No of		Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo (<10mm)	Risk difference with Alpha blockers (95% CI)
Time to stone passage mean number of days for spontaneous stone passage	3669 (5 studies) 2-4 weeks	⊕⊕⊖⊝ LOW3 due to risk of bias		The mean time to stone passage in the control groups was 12.31	The mean time to stone passage in the intervention groups was 4.13 lower (4.32 to 3.94 lower)

**Population** 

China

Mean (SD) age: naftopidil

wait group: 34.79 (9.63)

Male to female ratio 79:52

group: 33.73 (8.84); tamsulosin

group: 34.42 (8.64); watch and

**Outcomes** 

pain episodes

Pain intensity (2 weeks):

defined as number of

	No of			Anticipated absolute effects	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo (<10mm)	Risk difference with Alpha blockers (95% CI)
Time to stone passage mean number of hours for spontaneous stone passage	80 (1 study) 3 weeks	⊕⊖⊖ VERY LOW2,3 due to risk of bias, imprecision	HR 0.99 (0.55 to 1.78)	Moderate 0 per 1000	Not estimable <sup>8</sup>
Stone passage	5154	$\oplus \oplus \ominus \ominus$	RR 1.19	Moderate	
number of people spontaneously passing stones during follow up	(13 studies) 1-4 weeks		(1.09 to 1.29)	609 per 1000	116 more per 1000 (from 55 more to 177 more)
Hospitalisation	580	$\oplus \oplus \ominus \ominus$	RR 0.99	Moderate	
number of people hospitalised during follow up	(3 studies) 3-4 weeks	LOW2 due to imprecision	(0.59 to 1.64)	44 per 1000	0 fewer per 1000 (from 18 fewer to 28 more)
Use of healthcare	393	$\oplus \oplus \ominus \ominus$	RR 0.87	Moderate	
services (re-presentation to ED) number of people who re- presented to ED during follow up	(1 study) 4 weeks	LOW2 due to imprecision	(0.56 to 1.36)	180 per 1000	23 fewer per 1000 (from 79 fewer to 65 more)
Adverse events	363 (3 studies)	$\oplus \ominus \ominus \ominus$	RR 5.65	Moderate	
(unspecified) number of people experiencing adverse events during follow up	2-4 weeks	VERY LOW3 due to risk of bias	(1.5 to 21.29)	0 per 1000	70 more per 1000 (from 29 more to 112 more)5
Adverse events	3728	$\oplus \oplus \ominus \ominus$	Peto OR	Moderate	
	(6 studies) 3-4 weeks	, , , , , , , , , , , , , , , , , , ,	1.73 (1.23 to 2.43)	0 per 1000	20 more per 1000 (from 8 more to 32 more)5
	3957		RR 1.28	Moderate	

	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo (<10mm)	Risk difference with Alpha blockers (95% CI)	
Adverse events (dizziness) number of people experiencing dizziness during follow up	(7 studies) 1-4 weeks	⊕⊖⊝ VERY LOW2,3 due to risk of bias, imprecision	(0.92 to 1.79)	22 per 1000	6 more per 1000 (from 2 fewer to 17 more)	
Adverse events	3733	$\oplus \oplus \ominus \ominus$	RR 1.06	Moderate		
(headache) number of people experiencing headache during follow up	(4 studies) 4 weeks	LOW2 due to imprecision	(0.72 to 1.56)	29 per 1000	2 more per 1000 (from 8 fewer to 16 more)	
Adverse events	198	$\oplus \ominus \ominus \ominus$	Peto OR	Moderate		
(hypotension) number of people experiencing hypotension during follow up	number of people 4 weeks due imp	VERY LOW2,3 due to risk of bias, imprecision	6.82 (0.13 to 344.93)	0 per 1000	9 more per 1000 (from 18 fewer to 35 more)5	
Pain intensity (pain	150	$\oplus \oplus \oplus \ominus$	RR 0.34	Moderate		
episodes) number of people experiencing episodes of renal colic	(1 study) 4 weeks	MODERATE3 due to risk of bias	(0.23 to 0.51)	773 per 1000	510 fewer per 1000 (from 379 fewer to 595 fewer)	
Pain intensity (pain episodes) mean number of pain episodes	219 (2 studies) 2-4 weeks	⊕⊕⊝⊝ LOW2,3 due to risk of bias, imprecision		The mean pain intensity (pain episodes) in the control groups was 2.53	The mean pain intensity (pain episodes) in the intervention groups was 0.51 lower (0.86 to 0.15 lower)	
Pain intensity (pain score	367	$\oplus \oplus \oplus \oplus$	RR 0.98	Moderate		
>0) at 1 week verbal numeric pain scale	(1 study) 1 weeks	HIGH	(0.88 to 1.09)	786 per 1000	16 fewer per 1000 (from 94 fewer to 71 more)	
Pain intensity (pain score	353	$\oplus \oplus \ominus \ominus$	RR 1.04	Moderate		
>0) at 2 weeks verbal numeric pain scale	(1 study) 2 weeks	LOW2 due to imprecision	(0.77 to 1.4)	328 per 1000	13 more per 1000 (from 75 fewer to 131 more)	

	No of			Anticipated absolute effects	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo (<10mm)	Risk difference with Alpha blockers (95% CI)
Pain intensity (pain score	343	000	RR 0.94	Moderate	
>0) at 3 weeks verbal numeric pain scale	(1 study) 3 weeks	LOW2 due to imprecision	(0.62 to 1.42)	214 per 1000	13 fewer per 1000 (from 81 fewer to 90 more)
Pain intensity (pain score	347	$\oplus \oplus \ominus \ominus$	RR 0.93	Moderate	
>0) at 4 weeks verbal numeric pain scale	(1 study) 4 weeks	LOW2 due to imprecision	(0.57 to 1.53)	161 per 1000	11 fewer per 1000 (from 69 fewer to 85 more)
Analgesic use	3393	$\oplus \oplus \ominus \ominus$	RR 0.29	Moderate	
number of people using analgesics during follow up period	(2 studies) 4 weeks	LOW6 due to inconsistency	(0.13 to 0.66)	245 per 1000	174 fewer per 1000 (from 83 fewer to 213 fewer)
Analgesic use (number of times) mean number of times analgesics were used during follow up	165 (2 studies) 4-12 weeks	⊕⊕⊕⊖ MODERATE3 due to risk of bias		The mean analgesic use (number of times) in the control groups was 5.61	The mean analgesic use (number of times) in the intervention groups was 0.9lower (1.35 to 0.45 lower)
Analgesic use (Buprenorphine dose) mean dose (mg) of Buprenorphine used during follow up	267 (2 studies)	⊕⊕⊖ LOW3 due to risk of bias		The mean analgesic use (buprenorphine dose) in the control groups was 0.47	The mean analgesic use (buprenorphine dose) in the intervention groups was 0.06 lower (0.12 lower to 0 higher)
Analgesic use (Ketorolac dose) mean dose (mg) of Ketorolac used during follow up	315 (2 studies) 2 weeks	⊕⊕⊖ LOW3 due to risk of bias		The mean analgesic use (ketorolac dose) in the control groups was 337.87	The mean analgesic use (ketorolac dose) in the intervention groups was 97.44 lower (124.25 to 70.62 lower)
Analgesic use (Diclofenac dose) mean dose (mg) of Diclofenac used during follow up	3392 (2 studies) 4 weeks	⊕⊕⊕⊖ LOW7 due to inconsistency		The mean analgesic use (mean dose of drug) - diclofenac dose in the control groups was 181.5	The mean analgesic use (mean dose of drug) - diclofenac dose in the intervention groups was 149.03 lower (152.37 to 145.68 lower)

	No of			Anticipated absolute effects		
	Participants (studies)	Quality of the evidence	Relative effect		Risk difference with Alpha	
Outcomes	Follow up	(GRADE)	(95% CI)	Risk with Placebo (<10mm)	blockers (95% CI)	

- 1 Downgraded by 1 or 2 increments because heterogeneity, I2= 71%, p= > 0.1, unexplained by subgroup analysis
- 2 Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs
- 3 Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias
- 4 Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)
- 5 Risk difference calculated in Review Manager
- 6 Downgraded by 1 or 2 increments because heterogeneity, I2= 78%, p= > 0.1, unexplained by subgroup analysis
- 7 Downgraded by 1 or 2 increments because heterogeneity, I2= 97%, p= > 0.1, unexplained by subgroup analysis
- 8 Could not be calculated

Table 4: Clinical evidence summary: Alpha blockers versus no treatment (pain management only) for distal ureteric stones <10mm in adults

	No of			Anticipated absolute effects	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with no treatment (pain management only) (<10mm)	Risk difference with Alpha blockers (95% CI)
Time to stone passage (days) (mean number of days for spontaneous stone passage)	1542 (17 studies)2-8 weeks	⊕⊖⊖ VERY LOW1,4 due to risk of bias, inconsistency		The mean time to stone passage (days) in the control groups was 12.66 days	The mean time to stone passage (days) in the intervention groups was 4.14 lower (5.23 to 3.04 lower) )
Stone passage	2430	⊕⊝⊝⊝	RR 1.64	Moderate	
number of people spontaneously passing stones during follow up	(31 studies) 10 days - 8 weeks	VERY LOW1,6 due to risk of bias, inconsistency	(1.48 to 1.82)	511 per 1000	327 more per 1000 (from 245 more to 419 more)
Hospitalisation	487	$\oplus \oplus \ominus \ominus$	RR 0.27	Moderate	
number of people admitted to hospital during follow up		(0.15 to 0.46)	115 per 1000	84 fewer per 1000 (from 62 fewer to 98 fewer)	
				Moderate	

	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with no treatment (pain management only) (<10mm)	Risk difference with Alpha blockers (95% CI)	
Use of healthcare services (return to ED/primary care visit) number of people returning to ED or having an unscheduled primary care visit	77 (1 study) 2 weeks	⊕⊖⊖ VERY LOW1,3 due to risk of bias, imprecision	RR 0.77 (0.29 to 2.01)	205 per 1000	47 fewer per 1000 (from 146 fewer to 207 more)	
Adverse events (unspecified)	716	$\oplus \oplus \ominus \ominus$	Peto OR 5.89	Moderate		
number of people experiencing adverse events during follow up	(9 studies) 10 days - 4 weeks	LOW1 (due to risk of bias	(1.57 to 22.13)	0 per 1000	25 more per 1000 (from 8 more to 41 more)5	
Adverse events (dizziness)	514	⊕⊝⊝⊝	RR 1.34	Moderate		
number of people experiencing dizziness during follow up	during follow up 2-6 weeks due to risk of bia	VERY LOW1,3 due to risk of bias, imprecision	of bias,	0 per 1000	37 more per 1000 (from 6 fewer to 79 more)5	
Adverse events (hypotension)	508	508		Moderate		
number of people experiencing hypotension during follow up	(7 studies)	LOW1 due to risk of bias	(1.52 to 23.69)	0 per 1000	30 more per 1000 (from 9 more to 51 more)5	
Adverse events (retrograde	246	$\oplus \ominus \ominus \ominus$	RR 1.09	Moderate		
ejaculation) number of people experiencing retrograde ejaculation during follow up	(4 studies) 2-8 weeks	VERY LOW1,3 due to risk of bias, imprecision	(0.21 to 5.67)	0 per 1000	8 more per 1000 (from 23 fewer to 39 more)5	
Adverse events (headache)	163	$\oplus \ominus \ominus \ominus$	RR 1.48	Moderate		
number of people experiencing headache during follow up	(2 studies) 2-6 weeks	VERY LOW1,2,3 due to risk of bias, indirectness, imprecision	o risk of bias, ctness,	67 per 1000	32 more per 1000 (from 36 fewer to 47 more)	
Pain intensity	240	$\oplus \ominus \ominus \ominus$	RR 0.77	Moderate		
number of people experiencing pain during follow up	(3 studies) 10 days-4 weeks	VERY LOW1,3 due to risk of bias, imprecision	V1,3 (0.64 to 0.94) of bias,	793 per 1000	182 fewer per 1000 (from 48 fewer to 285 fewer)	

	No of			Anticipated absolute eff	ects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with no treatment (pain management only) (<10mm)	Risk difference with Alpha blockers (95% CI)		
Pain intensity (colicky pain episodes) mean number of colicky pain episodes	72 (1 study) 2 weeks	⊕⊖⊖ VERY LOW1,3 due to risk of bias, imprecision		The mean pain intensity (colicky pain episodes) in the control groups was 7.9	The mean pain intensity (colicky pain episodes) in the intervention groups was 0.05 lower (4.81 lower to 4.71 higher)		
Pain intensity (pain episodes) mean number of pain episodes during follow up	977 (10 studies)2-4 weeks	⊕⊖⊖ VERY LOW1,3,7 due to risk of bias, inconsistency, imprecision		The mean pain intensity (pain episodes) in the control groups was 2.21	The mean pain intensity (pain episodes) in the intervention groups was 0.65 lower (0.93 to 0.37 lower)		
Pain intensity (VAS score) at 3 days visual analogue scale	103 (1 study) 3 days	⊕⊕⊖⊖ LOW1 due to risk of bias		The mean pain intensity (VAS score) in the control groups was 3.06	The mean pain intensity (VAS score) in the intervention groups was 1.37 higher (0.84 to 1.90 higher)		
Pain intensity (VAS score) at 7 days visual analogue scale	103 (1 study) 7 days	⊕⊕⊖⊖ LOW1 due to risk of bias		The mean pain intensity (VAS score) in the control groups was 1.57	The mean pain intensity (VAS score) in the intervention groups was 1.63 higher (1.2 to 2.06 higher)		
Analgesic use	301	$\oplus \oplus \ominus \ominus$			RR 0.42	Moderate	
number of people using analgesics	(4 studies) 10 days-4 weeks	LOW1 due to risk of bias	(0.29 to 0.62)	485 per 1000	281 fewer per 1000 (from 184 fewer to 344 fewer)		
Analgesic use (number of times) mean number of times analgesics were used during follow up	421 (4 studies)	⊕⊖⊖⊖ VERY LOW1,3,9 due to risk of bias, inconsistency, imprecision		The mean analgesic use (number of times) in the control groups was 1.995	The mean analgesic use (number of times) in the intervention groups was 1.18 lower (2.49 lower to 0.13 higher)		
Analgesic use (Diclofenac dose) mean Diclofenac dose (mg) during follow up	234 (3 studies) 3-4 weeks	⊕⊖⊖ VERY LOW1,8 due to risk of bias, inconsistency		The mean analgesic use (diclofenac dose) in the control groups was 582.19	The mean analgesic use (diclofenac dose) in the intervention groups was 169.99 lower (314.6 to 25.37 lower)		

	No of			Anticipated absolute eff	ects
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with no treatment (pain management only) (<10mm)	Risk difference with Alpha blockers (95% CI)
Analgesic use (days) mean number of days analgesics were used	77 (1 study) 2 weeks	⊕⊖⊖ VERY LOW1,3 due to risk of bias, imprecision		The mean analgesic use (days) in the control groups was 4.3	The mean analgesic use (days) in the intervention groups was 4.94 lower (12.04 lower to 2.16 higher)
Analgesic use (Pethidine dose) mean dose (mg) of Pethidine used during follow up	64 (1 study) 4 weeks	⊕⊕⊖⊖ LOW1 due to risk of bias		The mean analgesic use (pethidine dose) in the control groups was 62.1	The mean analgesic use (pethidine dose) in the intervention groups was 27.7 lower (33.41 to 21.99 lower)
Analgesic use (Ketorolac dose) mean dose (mg) of Ketorolac used during follow up	95 (1 study) 2 weeks	⊕⊕⊖⊖ LOW1 due to risk of bias		The mean analgesic use (ketorolac dose) in the control groups was 347	The mean analgesic use (ketorolac dose) in the intervention groups was 103.5 lower (149.92 to 57.08 lower)
Analgesic use (Buprenorphine dose) mean dose (mg) of Buprenorphine during follow up	65 (1 study) 2 weeks	⊕⊕⊝⊝ LOW1 due to risk of bias		The mean analgesic use (buprenorphine dose) in the control groups was 0.39	The mean analgesic use (buprenorphine dose) in the intervention groups was 0.01 lower (0.16 lower to 0.14 higher)

Renal and ureteric stones: Medical expulsive therapy

- 1 Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias
- 2 Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population or the majority of the evidence had indirect outcomes
- 3 Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs
- 4 Downgraded by 1 or 2 increments because heterogeneity, I2= 91%, p= > 0.1, unexplained by subgroup analysis
- 5 Risk difference calculated in Review Manager
- 6 Downgraded by 1 or 2 increments because heterogeneity, I2= 57%, p= > 0.1, unexplained by subgroup analysis
- 7 Downgraded by 1 or 2 increments because heterogeneity, I2= 75%, p= > 0.1, unexplained by subgroup analysis
- 8 Downgraded by 1 or 2 increments because heterogeneity, I2= 92%, p= > 0.1, unexplained by subgroup analysis
- 9 Downgraded by 1 or 2 increments because heterogeneity, I2= 93%, p= > 0.1, unexplained by subgroup analysis

Table 5: Clinical evidence summary: Calcium channel blockers versus placebo for distal ureteric stones <10mm in adults

	No of Participants Quality of	Quality of the	evidence effect	Anticipated absolute effects	
Outcomes	(studies) Follow up	evidence eff		Risk with placebo (<10mm)	Risk difference with Calcium channel blockers (95% CI)
Stone passage	493	⊕⊕⊕⊕ RR 1.06 M	Moderate		
number of people spontaneously passing stones during follow up	(1 study) 28-45 days	HIGH	(0.98 to 1.14)	821 per 1000	49 more per 1000 (from 16 fewer to 115 more)

Table 6: Clinical evidence summary: Calcium channel blockers versus no treatment (pain management only) for distal ureteric stones <10mm in adults

	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with no treatment (pain management only) (<10mm)	Risk difference with Calcium channel blockers (95% CI)	
Time to stone passage mean number of days for spontaneous stone passage	70 (1 study) 4 weeks	⊕⊕⊕⊝ MODERATE1 due to risk of bias		The mean time to stone passage in the control groups was 12.29	The mean time to stone passage in the intervention groups was 0.29 lower (4.13 lower to 3.55 higher)	
Stone passage	179 ⊕⊕⊕⊝	udies) MODERATE1 (	RR 1.95	Moderate		
number of people spontaneously passing stones during follow up	(3 studies) 4 weeks		(1.4 to 2.71)	360 per 1000	342 more per 1000 (from 144 more to 616 more)	
Hospitalisation	129	$\oplus \oplus \oplus \ominus$		RR 0.41	Moderate	
number of people admitted to hospital during follow up	(2 studies) 4 weeks	MODERATE1 due to risk of bias	(0.24 to 0.69)	386 per 1000	228 fewer per 1000 (from 120 fewer to 293 fewer)	
Adverse events (hypotension)	59	$\oplus \ominus \ominus \ominus$	Peto OR	Moderate		
number of people experiencing hypotension during follow up	(1 study) 4 weeks	VERY LOW1,2 6. due to risk of bias, (0		0 per 1000	32 more per 1000 (from 55 fewer to 120 more)3	
Adverse events (dizziness)	50	$\oplus \oplus \ominus \ominus$	Not	Moderate		
number of people experiencing dizziness during follow up	(1 study) 4 weeks	LOW1 due to risk of bias	estimable bias 4	0 per 1000	0 more per 1000 (from 7 fewer to 7 more)3	

	No of		Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with no treatment (pain management only) (<10mm)	Risk difference with Calcium channel blockers (95% CI)
Pain intensity (pain episodes) mean number of pain episodes during follow up	70 (1 study) 4 weeks	⊕⊕⊖⊖ LOW1,2 due to risk of bias, imprecision		The mean pain intensity (pain episodes) in the control groups was 2.82	The mean pain intensity (pain episodes) in the intervention groups was 0.09 higher (0.41 lower to 0.59 higher)
Analgesic use (Diclofenac dose) mean Diclofenac dose (mg) during follow up	50 (1 study) 4 weeks	⊕⊕⊖⊝ LOW1 due to risk of bias		The mean analgesic use (diclofenac dose) in the control groups was 1408	The mean analgesic use (diclofenac dose) in the intervention groups was 806 lower (1103.31 to 508.69 lower)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 7: Clinical evidence summary: Alpha blockers versus Calcium channel blockers for distal ureteric stones <10mm in adults

	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Calcium channel blockers (<10mm)	Risk difference with Alpha blockers (95% CI)	
Time to stone passage (mean number of days for spontaneous stone passage)	70 (1 study) 4 weeks	⊕⊕⊕⊝ MODERATE1 due to risk of bias		The mean time to stone passage in the control groups was 12 days	The mean time to stone passage in the intervention groups was 0 higher (3.13 lower to 3.13 higher)	
Stone passage	4189	$\oplus \ominus \ominus \ominus$	RR 1.2	Moderate		
number of people spontaneously passing stones during follow up	ple (7 studies) VERY LOW1,2,3 passing stones 4 weeks due to risk of bias,	(1.05 to 1.39)	680 per 1000	136 more per 1000 (from 34 more to 265 more)		

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>3</sup> Risk difference calculated in Review Manager

<sup>4</sup> Could not be calculated as there were no events in the intervention or comparison group

	No of	No of		Anticipated absolut	te effects
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Calcium channel blockers (<10mm)	Risk difference with Alpha blockers (95% CI)
		inconsistency, imprecision			
Hospitalisation	133	$\oplus \oplus \ominus \ominus$	RR 0.45	Moderate	
number of people requiring Hospitalisation during follow up	(2 studies) 4 weeks	LOW1,3 due to risk of bias, imprecision	(0.18 to 1.17)	157 per 1000	86 fewer per 1000 (from 129 fewer to 27 more)
Adverse events (headache)	122	<b>0000</b>	RR 1.16	Moderate	
number of people experiencing headache during follow up	(1 study) 4 weeks	VERY LOW1,3 due to risk of bias, imprecision	(0.79 to 1.7)	431 per 1000	69 more per 1000 (from 91 fewer to 302 more)
Adverse events (dizziness)	172	4000		Moderate	
number of people experiencing dizziness during follow up		due to risk of bias,	(1.62 to 14.56)	26 per 1000	100 more per 1000 (from 16 more to 353 more)
Adverse events (hypotension)	63	$\oplus \ominus \ominus \ominus$	Peto OR 0.13	Moderate	
number of people experiencing hypotension during follow up	(1 study) 4 weeks	VERY LOW1,3 due to risk of bias, imprecision	(0 to 6.61)	32 per 1000	28 fewer per 1000 (from 32 fewer to 147 more)
Adverse events (not specified)	3189	$\oplus \ominus \ominus \ominus$	RR 0.92	Moderate	
number of people experiencing adverse events during follow up	(1 study) 4 weeks	VERY LOW1,3 due to risk of bias, imprecision	(0.69 to 1.21)	62 per 1000	5 fewer per 1000 (from 19 fewer to 13 more)
Adverse events (flushing)	122	$\oplus \ominus \ominus \ominus$	Peto OR 0.12	Moderate	
number of people experiencing flushing during follow up	(1 study) 4 weeks	VERY LOW1,3,4 due to risk of bias, indirectness, imprecision	(0.01 to 1.16)	52 per 1000	45 fewer per 1000 (from 51 fewer to 8 more)
Pain intensity (pain episodes) mean number of pain episodes	70 (1 study) 4 weeks	⊕⊕⊕⊝ MODERATE1 due to risk of bias		The mean pain intensity (pain episodes) in the	The mean pain intensity (pain episodes) in the intervention groups was

2

	No of			Anticipated absolute	e effects
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Calcium channel blockers (<10mm)	Risk difference with Alpha blockers (95% CI)
				control groups was 2.91 episodes	1.11 lower (1.54 to 0.68 lower)
Analgesic use (mg) mean Diclofenac mg used during follow up	50 (1 study) 4 weeks	⊕⊖⊖ VERY LOW1,3 due to risk of bias, imprecision		The mean analgesic use (mg) in the control groups was 602 mg	The mean analgesic use (mg) in the intervention groups was 58 lower (315.47 lower to 199.47 higher)
Analgesic use	3189	$\oplus \oplus \oplus \ominus$	RR 0.31	Moderate	
number of people using analgesics during follow up	(1 study) 4 weeks	MODERATE1 due to risk of bias	(0.2 to 0.49)	48 per 1000	33 fewer per 1000 (from 24 fewer to 38 fewer)
Analgesic use mean number of diclofenac injections	122 (1 study) 4-12 weeks	⊕⊕⊖⊖ LOW1 due to risk of bias,		The mean analgesic use in the control groups was 1.19	The mean analgesic use in the intervention groups was 0.77 lower (0.93 to 0.61 lower)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

#### 1 1.4.5.2 Mid ureteric stones <10mm in adults

Table 8: Clinical evidence summary: Alpha blockers versus placebo for mid ureteric stones <10mm in adults

	No of			Anticipated absolute effects	
Outcomes	Participant s (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo (<10mm)	Risk difference with Alpha blockers (95% CI)
				Moderate	

<sup>2</sup> Downgraded by 1 or 2 increments because heterogeneity, I2=88%, p=>0.1, unexplained by subgroup analysis

<sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>4</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments) or the majority of the evidence had indirect outcomes

	No of			Anticipated abso	lute effects	
Outcomes	s Quality of the (studies) evidence	Relative effect (95% CI)	Risk with Placebo (<10mm)	Risk difference with Alpha blockers (95% CI)		
Stone passage number of people spontaneously passing stones during follow up	126 (2 studies) 4 weeks	⊕⊕⊖⊖ LOW1,2 due to risk of bias, imprecision	RR 0.86 (0.67 to 1.09)	647 per 1000	91 fewer per 1000 (from 214 fewer to 58 more)	

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 9: Clinical evidence summary: Alpha blockers versus no treatment (pain management only) for mid ureteric stones <10mm in adults

	No of Participants	Quality of the	Relative	Anticipated absolute	Anticipated absolute effects		
Outcomes	(studies) Follow up	evidence (GRADE)	effect (95% CI)	Risk with No treatment (<10mm)	Risk difference with Alpha blockers (95% CI)		
Time to stone passage mean number of days for spontaneous stone passage	16 (1 study) 8 weeks	⊕⊕⊝ LOW1 due to risk of bias		The mean time to stone passage in the control groups was 21	The mean time to stone passage in the intervention groups was 12.33 lower (17.26 to 7.4 lower)		
Stone passage	27	$\oplus \ominus \ominus \ominus$	P⊝⊝ RR 4.09 Moderate				
number of people spontaneously passing stones during follow up	(2 studies) 4-8 weeks	VERY LOW1,2 due to risk of bias, imprecision	(1.09 to 15.33)	163 per 1000	504 more per 1000 (from 15 more to 1000 more)		
Analgesic use mean number of times analgesics were used during follow up	16 (1 study) 8 weeks	⊕⊖⊖ VERY LOW1,2 due to risk of bias, imprecision		The mean analgesic use in the control groups was 1.3	The mean analgesic use in the intervention groups was 1.2 lower (2.67 lower to 0.27 higher)		

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 10: Clinical evidence summary: Calcium channel blockers versus placebo for mid ureteric stones <10mm in adults

No of		Anticipated absolute effects			
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo	Risk difference with Calcium channel blockers (95% CI)
Stone passage	84	$\oplus \oplus \ominus \ominus$	RR 0.98	Moderate	
	(1 study) 4 weeks	LOW1,2 due to risk of bias, imprecision	(0.79 to 1.2)	818 per 1000	16 fewer per 1000 (from 172 fewer to 164 more)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 11: Clinical evidence summary: Alpha blockers versus Calcium channel blockers for mid ureteric stones <10mm in adults

				Anticipated absolute effects	
Outcomes	No of Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Calcium channel blockers (<10mm)	Risk difference with Alpha blockers (95% CI)
Stone passage	81	$\oplus \oplus \ominus \ominus$	RR 0.88	Moderate	
number of people spontaneously passing stones during follow up	(1 study) 4 weeks	LOW1,2 due to risk of bias, imprecision	(0.69 to 1.14)	800 per 1000	96 fewer per 1000 (from 248 fewer to 112 more)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

#### 1 1.4.5.3 Proximal ureteric stones <10mm in adults

Table 12: Clinical evidence summary: Alpha blockers versus placebo for proximal ureteric stones <10mm in adults

			Relative	Anticipated abso	solute effects	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	effect (95% CI)	Risk with Placebo	Risk difference with Alpha blockers (95% CI)	
Stone passage	257	$\oplus \oplus \ominus \ominus$	RR 0.96	Moderate		
number of people spontaneously passing stones during follow up	(2 studies) 4 weeks	LOW1,2 due to risk of bias, imprecision	(0.79 to 1.15)	568 per 1000	23 fewer per 1000 (from 119 fewer to 85 more)	

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 13: Clinical evidence summary: Alpha blockers versus no treatment (pain management only) for proximal ureteric stones <10mm in adults

	No of	No of		Anticipated absolute	effects	
Outcomes	Participants Quality of the (studies) evidence (GRADE)		Relative effect (95% CI)	Risk with No treatment (pain management only)	Risk difference with Alpha blockers (95% CI)	
Time to stone passage mean number of days for spontaneous stone passage	133 (2 studies) 4-8 weeks	⊕⊖⊖ VERY LOW1,2 due to risk of bias, imprecision		The mean time to stone passage in the control groups was 19.17	The mean time to stone passage in the intervention groups was 5.29 lower (8.43 to 2.16 lower)	
Stone passage	213		RR 1.57	RR 1.57 Moderate		
number of people spontaneously passing stones during follow up	(4 studies) 4-8 weeks		(1.2 to 2.03)	357 per 1000	203 more per 1000 (from 71 more to 368 more)	
Quality of life (EuroQoL) mean score on EuroQol	79 (1 study) 4 weeks	⊕⊖⊖ VERY LOW1,2 due to risk of bias, imprecision		The mean quality of life (EuroQoL) in the control groups was 5.5	The mean quality of life (EuroQoL) in the intervention groups was 0.1 lower (0.42 lower to 0.22 higher)	

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

	No of		Α	Anticipated absolute	Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with No treatment (pain management only)	Risk difference with Alpha blockers (95% CI)		
Analgesic use mean number of times analgesics were used	133 (2 studies) 4-8 weeks	⊕⊕⊖ LOW1 due to risk of bias		The mean analgesic use in the control groups was 3.25	The mean analgesic use in the intervention groups was 0.55 lower (2.06 lower to 0.97 higher)		

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 14: Clinical evidence summary: Calcium channel blockers versus placebo for proximal ureteric stones <10mm in adults

No of				Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo	Risk difference with Calcium channel blockers (95% CI)	
Stone passage	181	$\oplus \oplus \ominus \ominus$	RR 0.86	Moderate		
	(1 study) 4 weeks	LOW1,2 due to risk of bias, imprecision	(0.71 to 1.06)	730 per 1000	102 fewer per 1000 (from 212 fewer to 44 more)	

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 15: Clinical evidence summary: Alpha blockers versus Calcium channel blockers for proximal ureteric stones <10mm in adults

No of Participants Quality of the	Quality of the	Relative	Anticipated absolute effects			
Outcomes	(studies) Follow up	evidence		Risk with Calcium channel blockers	Risk difference with Alpha blockers (95% CI)	
Stone passage	ssage 180 $\oplus \oplus \ominus \ominus$ F		RR 1.12	Moderate		
number of people spontaneously passing stones during follow up	(1 study) 4 weeks	LOW1,2 due to risk of bias, imprecision	(0.91 to 1.37)	630 per 1000	76 more per 1000 (from 57 fewer to 233 more)	

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

	No of Participants	evidence	effect	Anticipated absolute effects	
Outcomes	(studies) Follow up			Risk with Calcium channel blockers	Risk difference with Alpha blockers (95% CI)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

#### 4.5.4 Distal ureteric stones <10mm in children

Table 16: Clinical evidence summary: Alpha blockers versus placebo for distal ureteric stones <10mm in children

	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo	Risk difference with Alpha blockers (95% CI)	
Time to stone passage (days)	98 (2 studies) (4 weeks)	⊕⊖⊖ VERY LOW1,2,3 due to risk of bias, inconsistency, imprecision		The mean time to stone passage (days) in the control groups was 12.45 days	The mean time to stone passage (days) in the intervention groups was 4.89 lower (7.73 to 2.05 lower)	
Stone passage	98	$\oplus \oplus \ominus \ominus$	RR 1.3	Moderate		
	4 weeks due	LOW1,2 due to risk of bias, imprecision	(1.04 to 1.62)	690 per 1000	207 more per 1000 (from 28 more to 428 more)	
Adverse events	37	$\oplus \oplus \ominus \ominus$		Peto OR	Moderate	
(headaches/dizziness)	(1 study) 4 weeks	LOW1,2 due to risk of bias, imprecision	'	0 per 1000	167 more per 1000 (from 21 fewer to 354 more)4	
Adverse events (headaches)	61	$\oplus \ominus \ominus \ominus$	RR 0.85	Moderate		
		(0.06 to 12.95)	36 per 1000	5 fewer per 1000 (from 34 fewer to 430 more)		
Adverse events	61	$\oplus \ominus \ominus \ominus$	Not	Moderate		
(hypotension)		estimable6	0 per 1000	0 fewer per 1000 (from 62 fewer to 62 more)4		

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

	No of				Anticipated absolute effects	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with Placebo	Risk difference with Alpha blockers (95% CI)	
		due to risk of bias, imprecision				
Pain intensity (number of pain episodes)	98 (2 studies) 4 weeks	⊕⊖⊖ VERY LOW1,2,5 due to risk of bias, inconsistency, imprecision		The mean pain intensity (number of pain episodes) in the control groups was 3.45	The mean pain intensity (number of pain episodes) in the intervention groups was 1.49 lower (3.04 lower to 0.06 higher)	

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

- 2 Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.
- 3 Downgraded by 1 or 2 increments because heterogeneity, I2= 73%, p= > 0.1, unexplained by subgroup analysis
- 4 Risk difference calculated in Review Manager
- 5 Downgraded by 1 or 2 increments because heterogeneity, I2= 77%, p= > 0.1, unexplained by subgroup analysis
- 6 Could not be calculated as there were no events in the intervention or comparison group

Table 17: Clinical evidence summary: Alpha blockers versus no treatment (pain management only) for distal ureteric stones <10mm in children

	No of			Anticipated absolute effects		
ts (studie		Participan ts Quality of the (studies) evidence Follow up (GRADE)		Risk with No treatment	Risk difference with Alpha blockers (95% CI)	
Time to stone passage mean number of days for spontaneous stone passage	102 (2 studies) 3-4 weeks	⊕⊖⊖ VERY LOW1,2,3 due to risk of bias, inconsistency, imprecision		The mean time to stone passage in the control groups was 12.05	The mean time to stone passage in the intervention groups was 5.26 lower (15.16 lower to 4.63 higher)	
Stone passage	one passage 147 ⊕⊖⊝		RR 1.45	Moderate		
number of people spontaneously passing stones (3 studies) VE 3-4 weeks	VERY LOW1,2 due to risk of bias, imprecision	(1.14 to 1.84)	625 per 1000	281 more per 1000 (from 87 more to 525 more)		

Outcomes	Participan ts (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with No treatment	Risk difference with Alpha blockers (95% CI)	
Adverse events	102	$\oplus \oplus \ominus \ominus$	Not	Moderate		
number of people experiencing adverse events (unspecified)	4400		0 per 1000	0 more per 1000 (from 50 fewer to 50 more)4		
Pain intensity (daily pain episodes) mean number of daily pain episodes during follow up	63 (1 study) 4 weeks	⊕⊖⊖ VERY LOW1,2 due to risk of bias, imprecision		The mean pain intensity (daily pain episodes) in the control groups was 2.5	The mean pain intensity (daily pain episodes) in the intervention groups was 0.9 lower (1.77 to 0.03 lower)	
Analgesic use mean number of times analgesics were used during follow up	63 (1 study) 4 weeks	⊕⊖⊖ VERY LOW1,2 due to risk of bias, imprecision		The mean analgesic use in the control groups was 1.8	The mean analgesic use in the intervention groups was 1.25 lower (1.87 to 0.63 lower)	

**Anticipated absolute effects** 

Renal and ureteric stones: Medical expulsive therapy

No of

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>3</sup> Downgraded by 1 or 2 increments because the point estimate varies widely across studies, the confidence intervals across studies show minimal or no overlap, or heterogeneity, I2=99%, p<0.04, unexplained by subgroup analysis.

<sup>4</sup> Risk difference calculated in Review Manager

<sup>5</sup> Could not be calculated as there were no events in the intervention or comparison group

# 1 1,4.5.5 Adjunctive therapy: distal ureteric stones <10mm in adults

Table 18: Clinical evidence summary: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only for distal ureteric stones <10mm in adults

	No of Participants	Quality of the		Anticipated absol	ute effects
Outcomes	(studies) Follow up	evidence (GRADE)	Relative effect (95% CI)	Risk with SWL	Risk difference with Alpha blockers + SWL (95% CI)
Time to stone passage number of days for stone passage	207 (2 studies) 4-6 weeks	⊕⊖⊖ VERY LOW1,3 due to risk of bias, imprecision		The mean time to stone passage in the control groups was 14.65	The mean time to stone passage in the intervention groups was 2.21 lower (3.35 to 1.08 lower)
Stone passage	383	$\oplus \ominus \ominus \ominus$	RR 1.28	Moderate	
number of people stone free at the end of follow up	(5 studies) 15 days - 6 weeks	VERY LOW1,2,3 due to risk of bias, indirectness, imprecision	(1.11 to 1.48)	568 per 1000	159 more per 1000 (from 62 more to 273 more)
Hospitalisation	oitalisation 88 ⊕⊕⊝⊝ RR (	RR 0.63	Moderate		
number of people hospitalized during follow up	(1 study) 4 weeks	LOW1,3 due to risk of bias, imprecision	(0.35 to 1.14)	432 per 1000	160 fewer per 1000 (from 281 fewer to 60 more)
Adverse events (dizziness)	206	$\oplus \oplus \ominus \ominus$	Peto OR 8.4	Moderate	
number of people experiencing dizziness during follow up	(3 studies) 15 days - 6 weeks	LOW1 due to risk of bias	(1.86 to 37.87)	0 per 1000	69 more per 1000 (from 17 more to 122 more)4
Adverse events (abnormal	98	$\oplus \oplus \ominus \ominus$	Peto OR 8.56	Moderate	
ejaculation) number of people experiencing abnormal ejaculation during follow up	(2 studies) 4-6 weeks	LOW1 due to risk of bias	(1.83 to 40.08)	0 per 1000	142 more per 1000 (from 40 more to 246 more)4
Adverse events (headache)	155	$\oplus \ominus \ominus \ominus$	RR 4.03	Moderate	
number of people experiencing headache during follow up	(2 studies) 4-6 weeks	VERY LOW1,3 due to risk of bias, imprecision	(1.04 to 15.72)	29 per 1000	88 more per 1000 (from 1 more to 427 more)
				Moderate	

	No of Participants	Quality of the		Anticipated absol	ute effects
Outcomes	(studies) Follow up	evidence (GRADE)	Relative effect (95% CI)	Risk with SWL	Risk difference with Alpha blockers + SWL (95% CI)
Adverse events (hypotension) number of people experiencing hypotension during follow up	67 (1 study) 6 weeks	⊕⊕⊖⊖ LOW1 due to risk of bias		0 per 1000	0 more per 1000 (from 60 fewer to 60 more)4
Analgesic use mean number of times analgesics were used during follow up	88 (1 study) 4 weeks	⊕⊖⊖⊖ VERY LOW1,3 due to risk of bias, imprecision		The mean analgesic use in the control groups was 6.11	The mean analgesic use in the intervention groups was 1.72 lower (2.88 to 0.56 lower)
Analgesic use (dosage) mean dosage (mg) of Diclofenac during follow up	119 (1 study) 4 weeks	⊕⊖⊖⊖ VERY LOW1,2 due to risk of bias, indirectness		The mean analgesic use (dosage) in the control groups was 116.1	The mean analgesic use (dosage) in the intervention groups was 50.27 lower (68.87 to 31.67 lower)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 19: Clinical evidence summary: Alpha blockers as adjunctive therapy to ureteroscopy versus ureteroscopy only for distal ureteric stones <10mm in adults

	No of			Anticipated absolute	e effects
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with URS	Risk difference with Alpha blockers + URS (95% CI)
				Moderate	

<sup>2</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)

<sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>4</sup> Risk difference calculated in Review Manager

	No of			Anticipated absolute	e effects
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with URS	Risk difference with Alpha blockers + URS (95% CI)
Stone passage number of people stone-free at the end of follow up	98 (1 study) 2 weeks	⊕⊕⊕⊝ MODERATE1 due to risk of bias	RR 1.08 (0.95 to 1.23)	872 per 1000	70 more per 1000 (from 44 fewer to 201 more)
Use of healthcare services length of hospital stay	98 (1 study)	⊕⊕⊖⊖ LOW1,2 due to risk of bias, imprecision		The mean use of healthcare services in the control groups was 1.7	The mean use of healthcare services in the intervention groups was 0.5 lower (0.81 to 0.19 lower)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 20: Clinical evidence summary: Alpha blockers as adjunctive therapy to ureteroscopy versus placebo and ureteroscopy for distal ureteric stones <10mm in adults

	No of		Quality of the Relative effect	Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)		Risk with placebo + URS	Risk difference with Alpha blockers + URS (95% CI)	
Stone passage	102	$\oplus \ominus \ominus \ominus$	RR 1.35	Moderate		
number of people stone free at the end of follow up	(1 study) 2 weeks	VERY LOW1,2 due to risk of bias, imprecision	(1.11 to 1.63)	700 per 1000	245 more per 1000 (from 77 more to 441 more)	
Pain intensity (colic episodes) mean number of colic episodes during follow up	102 (1 study) 2 weeks	⊕⊕⊖ LOW1 due to risk of bias		The mean pain intensity (colic episodes) in the control groups was	The mean pain intensity (colic episodes) in the intervention groups was 5 lower (5.99 to 4.01 lower)	
Analgesic use	102	$\oplus \ominus \ominus \ominus$	9⊝⊝⊝ RR 0.32			
number of people using (1 study) VERY LOW1,2 analgesics during follow up 2 weeks due to risk of bias, imprecision	(0.11 to 0.93)	240 per 1000	163 fewer per 1000 (from 17 fewer to 214 fewer)			

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

CONSULTATION

	No of			Anticipated absolute effects	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with placebo + URS	Risk difference with Alpha blockers + URS (95% CI)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

#### Adjunctive therapy: distal ureteric stones 10-20mm in adults

Table 21: Clinical evidence summary: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only for distal ureteric stones 10-20mm in adults

	No of			Anticipated absolu	te effects
Outcomes	Participants (studies) Follow up	evidence	Relative effect (95% CI)	Risk with SWL	Risk difference with Alpha blockers + SWL (95% CI)
Time to stone passage number of days for stone passage	38 (1 study) unclear	⊕⊖⊖ VERY LOW1,2,3 due to risk of bias, indirectness, imprecision		The mean time to stone passage in the control groups was 12.42	The mean time to stone passage in the intervention groups was 2.56 lower (7.78 lower to 2.66 higher)
Pain intensity (VAS) visual analogue scale. Scale from: 0 to 10.	38 (1 study) unclear	⊕⊖⊖⊖ VERY LOW1,2,3 due to risk of bias, indirectness, imprecision		The mean pain intensity (vas) in the control groups was	The mean pain intensity (vas) in the intervention groups was 1.21 lower (2.88 lower to 0.46 higher)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>2</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)

<sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

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#### 1 1,4.5.7 Adjunctive therapy: mid ureteric stones 10-20mm in adults

Table 22: Clinical evidence summary: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only for mid ureteric stones 10-20mm in adults

	No of	<b>2</b> 114 641	5.1.0	Anticipated absolute effects	
Outcomes	Participants (studies) Follow up	evidence	Relative effect (95% CI)	Risk with SWL	Risk difference with Alpha blockers + SWL (95% CI)
Time to stone passage number of days for stone passage	28 (1 study) unclear	⊕⊖⊖ VERY LOW1,2,3 due to risk of bias, indirectness, imprecision		The mean time to stone passage in the control groups was 10.75	The mean time to stone passage in the intervention groups was 1.5 lower (8.23 lower to 5.23 higher)
Pain intensity (VAS) visual analogue scale. Scale from: 0 to 10.	28 (1 study) unclear	⊕⊖⊖ VERY LOW1,2,3 due to risk of bias, indirectness, imprecision		The mean pain intensity (VAS) in the control groups was	The mean pain intensity (VAS) in the intervention groups was 0.62 lower (3.13 lower to 1.89 higher)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

#### 4 1.4.5.8 Adjunctive therapy: proximal ureteric stones <10mm in adults

Table 23: Clinical evidence summary: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only for proximal ureteric stones <10mm in adults

	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with SWL	Risk difference with Alpha blockers + SWL (95% CI)	
Time to stone passage number of days for stone passage	320 (4 studies) 2-12 weeks	⊕⊖⊖ VERY LOW1,2		The mean time to stone passage in the	The mean time to stone passage in the intervention groups was	

<sup>2</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)

<sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

	No of			Anticipated absolute e	ffects
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with SWL	Risk difference with Alpha blockers + SWL (95% CI)
		due to risk of bias, inconsistency		control groups was 23.12	4.32 lower (9.85 lower to 1.21 higher)
Stone passage	405	$\oplus \oplus \oplus \ominus$	RR 1.11	Moderate	
number of people stone free at the end of follow up	(6 studies) 2-12 weeks	MODERATE1 due to risk of bias	(1.03 to 1.21)	848 per 1000	93 more per 1000 (from 25 more to 178 more)
Hospitalisation mean number of Hospitalisations	79 (1 study) 2 weeks	⊕⊖⊖ VERY LOW1,4 due to risk of bias, indirectness		The mean Hospitalisation in the control groups was 0.52	The mean Hospitalisation in the intervention groups was 0.01 lower (0.31 lower to 0.29 higher)
Use of healthcare services (ED visits) mean number of ED visits during follow up	54 (1 study) 4 weeks	⊕⊖⊖ VERY LOW1,3 due to risk of bias, imprecision		The mean use of healthcare services (ED visits) in the control groups was 1.42	The mean use of healthcare services (ED visits) in the intervention groups was 0.6 lower (1.13 to 0.07 lower)
Quality of life (EQ5D) mean score on EQ5D. Scale from: 0 to 1.	54 (1 study) 4 weeks	⊕⊖⊖ VERY LOW1,3 due to risk of bias, imprecision		The mean quality of life (EQ5D) in the control groups was 0.78	The mean quality of life (EQ5D) in the intervention groups was 0.04 higher (0.01 lower to 0.09 higher)
Quality of life (EQ5D VAS) mean score on EQ5D visual analogue scale . Scale from: 0 to 100.	54 (1 study) 4 weeks	⊕⊖⊖ VERY LOW1,3 due to risk of bias, imprecision		The mean quality of life (EQ5D VAS) in the control groups was 73.65	The mean quality of life (EQ5D VAS) in the intervention groups was 6.71 higher (1.49 to 11.93 higher)
Adverse events (dizziness)	172	$\oplus \ominus \ominus \ominus$	Peto OR	Moderate	
number of people experiencing dizziness during follow up	(2 studies) 3-6 weeks	VERY LOW1,3 due to risk of bias, imprecision	7.76 (0.8 to 75.32)	0 per 1000	35 more per 1000 (from 9 fewer to 80 more)5
Adverse events (retrograde	84	$\oplus \oplus \ominus \ominus$	Not	Moderate	
ejaculation) number of people experiencing retrograde ejaculation during follow up	on) (1 study) LOW1 of people experiencing 6 weeks due to risk of bias de ejaculation during	estimable8	0 per 1000	0 more per 1000 (from 45 fewer to 45 more)5	

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	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with SWL	Risk difference with Alpha blockers + SWL (95% CI)	
Pain intensity (VAS) visual analogue scale . Scale from: 0 to 10.	374 (5 studies) 2-12 weeks	⊕⊖⊖ VERY LOW1,3,6 due to risk of bias, imprecision, inconsistency		The mean pain intensity (vas) in the control groups was 5.54	The mean pain intensity (vas) in the intervention groups was 0.89 lower (1.68 to 0.1 lower)	
Pain intensity (renal colic episodes) mean number of renal colic episodes during follow up	54 (1 study) 4 weeks	⊕⊖⊝ VERY LOW1,3 due to risk of bias, imprecision		The mean pain intensity (renal colic episodes) in the control groups was 4.92	The mean pain intensity (renal colic episodes) in the intervention groups was 2.38 lower (3.89 to 0.87 lower)	
Analgesic use (dosage) mean dosage (mg) of Diclofenac used during follow up	54 (1 study) 4 weeks	⊕⊖⊖ VERY LOW1,3 due to risk of bias, imprecision		The mean analgesic use (dosage) in the control groups was 431.7	The mean analgesic use (dosage) in the intervention groups was 189.7 lower (309.2 to 70.2 lower)	
Analgesic use	163	$\oplus \ominus \ominus \ominus$	RR 0.96	Moderate		
analgesia during follow up 2-6 weeks due to indirect impre	VERY LOW1,3,4,7 due to risk of bias, indirectness, imprecision, inconsistency	(0.49 to 1.91)	492 per 1000	20 fewer per 1000 (from 251 fewer to 448 more)		

- 1 Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias
- 2 Downgraded by 1 or 2 increments because the point estimate varies widely across studies, the confidence intervals across studies show minimal or no overlap or heterogeneity, I2=77%, p= > 0.1, unexplained by subgroup analysis.
- 3 Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs
- 4 Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)
- 5 Risk difference calculated in Review Manager
- 6 Downgraded by 1 or 2 increments because the point estimate varies widely across studies, the confidence intervals across studies show minimal or no overlap or heterogeneity, I2=86%, p= > 0.1, unexplained by subgroup analysis
- 7 Downgraded by 1 or 2 increments because the point estimate varies widely across studies, the confidence intervals across studies show minimal or no overlap or heterogeneity, I2=67%, p= > 0.1, unexplained by subgroup analysis

			Anticipated absolute e	TTECTS
Partic (studi Outcomes Follow	<b>/</b>	Relative effect (95% CI)	Risk with SWL	Risk difference with Alpha blockers + SWL (95% CI)

<sup>8</sup> Could not be calculated as there were no events in the intervention or comparison group

Table 24: Clinical evidence summary: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus placebo and shock wave lithotripsy for proximal ureteric stones <10mm in adults

	No of	rticipants Quality of the udies) evidence		Anticipated absolute effects	
Outcomes	Participants (studies) Follow up			Risk with Placebo + SWL	Risk difference with Alpha blockers + SWL (95% CI)
Time to stone passage number of days for stone passage	49 (1 study) 3 months	⊕⊕⊖⊖ LOW1,3 due to risk of bias, indirectness		The mean time to stone passage in the control groups was 7.5	The mean time to stone passage in the intervention groups was 3.3 lower (4.47 to 2.13 lower)
Stone passage	49	9		Moderate	
number of people stone free at the end of follow up	or of people stone free at the (1 study) LOW1,2		(1.06 to 1.97)	667 per 1000	300 more per 1000 (from 40 more to 647 more)

<sup>1</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>3</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

#### 1 1,4.5.9 Adjunctive therapy: proximal ureteric stones 10-20mm in adults

Table 25: Clinical evidence summary: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only for proximal ureteric stones 10-20mm in adults

	No of			Anticipated absolute effects	
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with SWL	Risk difference with Alpha blockers + SWL (95% CI)
Time to stone passage number of days to stone passage	57 (1 study) unclear	⊕⊖⊖ VERY LOW1,2,3 due to risk of bias, indirectness, imprecision		The mean time to stone passage in the control groups was 13.54	The mean time to stone passage in the intervention groups was 6.44 lower (10.3 to 2.58 lower)
Stone passage	57	$\oplus \ominus \ominus \ominus$	RR 1.09	Moderate	
number of people stone free at the end of follow up	(1 study) 3 months	VERY LOW1,2,3 due to risk of bias, indirectness, imprecision	of bias,	821 per 1000	74 more per 1000 (from 99 fewer to 287 more)
Pain intensity (VAS) visual analogue scale. Scale from: 0 to 10.	57 (1 study) unclear	⊕⊖⊖ VERY LOW1,2,3 due to risk of bias, indirectness, imprecision		The mean pain intensity (vas) in the control groups was 4	The mean pain intensity (vas) in the intervention groups was 1.1 lower (2.34 lower to 0.14 higher)

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 26: Clinical evidence summary: Alpha blockers as adjunctive therapy to ureteroscopy versus ureteroscopy only for proximal ureteric stones 10-20mm in adults

	No of			Anticipated absolute effects		
Outcomes	Participants (studies) Follow up	Quality of the evidence (GRADE)	Relative effect (95% CI)	Risk with URS	Risk difference with Alpha blockers + URS (95% CI)	
Time to stone passage number of days for stone passage	89 (1 study) 6 weeks	⊕⊖⊖ VERY LOW1,2		The mean time to stone passage in	The mean time to stone passage in the intervention groups was	

<sup>2</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)

<sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

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<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>3</sup> Risk difference calculated in Review Manager

See appendix F for full GRADE tables.

# 1.5 Economic evidence

2	1.5.1	included studies	

- One health economic study was identified in adults with the relevant comparison and has been included in this review. This is summarised in the health economic evidence profile
- 5 below (Table 27) and the health economic evidence table in appendix H.
- 6 No relevant health economic studies were identified in children.

#### 7 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 G.

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### 1 **21.5.3** Summary of studies included in the economic evidence review

Table 27: Health economic evidence profile: MET (tamsulosin or nifedipine) versus placebo and tamsulosin versus nifedipine

Study	Applicability	Limitations	Other comments	Incremental cost	Incremental effects	Cost effectiveness	Uncertainty
Pickard 2015 [UK] <sup>145</sup>	Directly applicable (a)	Potentially serious limitations (b)	Within trial analysis based on an RCT of 12 weeks. No extrapolation. Population is adult patients with ureteric stones. Interventions were MET (tamsulosin 400 µg and nifedipine 30 mg groups combined) for up to 4 weeks, and placebo. Tamsulosin versus Nifedipine also compared in the analysis. The study perspective is the NHS but also patient costs were collected from patients. These costs are difficult to be separated from the rest of NHS costs. Health related quality of life measures were collected by participant completed EQ-5D questionnaires. SF-36 also collected.	Placebo vs MET: £42  Nifedipine vs Tamsulosin: £87	Placebo vs MET: 0.001 Nifedipine vs Tamsulosin: 0.002	Placebo vs MET: £42,000 (d)  Nifedipine vs Tamsulosin: £43,500 (e)	Used non-parametric boostrapping to get 1000 estimates of the ICERs.  One-way sensitivity analyses using extreme values were performed around the QALY estimates.  An alternative measure was used for QoL; SF-36 responses were mapped on the SF-6D measure using the algorithm from another study to validate the estimate of utility value for each time point derived from the EQ-5D.  = Placebo now cost effective instead of MET. Tamsulosin still cost effective.  Also a Sensitivity analysis using imputed EQ-5D assuming the imputed values are the highest estimates was conducted.  = Placebo now cost effective instead of MET. Nifedipine now cost effective.

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- (a) Within trial analysis based on UK RCT. Uses an NHS perspective and EQ-5D. Included some participants costs that are not NHS costs related, and these were reported as part of NHS costs that they account for significant % of total costs of intervention; so it is difficult to separate participants' costs from the NHS costs in order to determine whether their magnitude is significant compared to the total costs of interventions. The categories where the patient reported outcomes fall include costs that are of similar amount in both interventions (MET, placebo), so unlikely changing the cost effectiveness results.
- (b) Study didn't meet the quality criteria around the choice of time horizon being 12 weeks and not longer. That was justified by the authors as there weren't many people who still needed interventions at the end of the trial. However there were no extrapolation and therefore assumptions made about what this treatment would be which could impact incremental costs and effects because different numbers of people are stone free in each arm.
- (c) Utilities for clinical response were derived using trial data and the EQ-5D questionnaire
- (d) This has been calculated by the health economist as there is an error in the study. This was reported as cost saving per QALY lost for MET versus placebo because MET was both cheaper and less effective. However for ease of interpretation in cases like this the intervention should be switched around i.e. to compare placebo versus MET so that the less effective intervention is used as the comparator and so the ICER can be interpreted as it normally would (if less than £20,000 then intervention is cost effective versus the comparison).
- (e) Similar to note d. Nifedipine is less expensive and less effective than tamsulosin, so the ICER of nifedipine versus tamsulosin is presented for ease of interpretation.

#### 1.5.4 Unit costs

Where several studies evaluated the same intervention in different doses we used the highest dose reported. Calculation for tablets and capsules for tamsulosin and nifedipine were made as capsules formulation of these drugs was identified in the cost utility analysis

Table 28: UK costs of alpha blockers and calcium channel blockers

Drug	Daily dose (mg)	Cost (per unit)	Daily cost	Cost – monthly	Cost- annual	Source of dose				
Alpha blockers										
Tamsulosin hydrocholoride TABLETS	0.4 per day	0.4mg tablet (Pack of 30) = £10.47	£0.35	£10.65	£127.39	Clinical review				
Tamsulosin hydrocholoride CAPSULES	0.4 per day	0.4mg capsule (Pack of 30) = £3.89	£0.13	£3.94	£47.33	Pickard 2015 <sup>145</sup>				
Alfuzosin hydrocholoride	10mg per day	10 mg tablet (pack of 30) =£12.51	£0.42	£12.68	£152.21	Clinical review				
Doxazosin	4mg per day	4mg tablet (pack of 28) = £5	£0.18	£5.43	£65.18	Clinical review				
Terazosin	10mg per day	10mg tablet (pack of 28) = £7.87	£0.28	£8.55	£102.59	Clinical review				
Calcium channel blockers										
Nifedipine TABLETS	30mg per day	30 mg tablet (pack of 28) = £6.85	£0.24	£7.44	£89.29	Clinical review				
Nifedipine CAPSULES	30mg per day	30 mg tablet (pack of 28 =£4.89	£0.17	£5.31	£63.74	Pickard 2015 <sup>145</sup>				

Source: BNF "Drug Tariff" price, DATE; September 2017 95

## 1.6 Resource costs

The recommendations made in this review are likely to have a substantial impact on resources.

Additional savings are likely to be made for the following reasons: MET are very inexpensive drugs, the cost of providing these would be outweighed by the savings from downstream resource use avoided because of the effectiveness of MET at helping stones to pass. Further work is being carried out to quantify the potential resource impact in this area.

<sup>(</sup>a) The cost of other alpha blockers, naftopodil, silodosin is not provided by BNF site

# 1.7 Evidence statements

#### 1.7.1 Clinical evidence statements

#### Distal ureteric stones

Seven studies compared alpha blockers to calcium channel blockers in adults with distal ureteric stones <10mm and the evidence suggested a clinically important benefit in favour of alpha blockers for the outcome stone passage (7 studies; 4189). One study reported the outcome time to stone passage and this evidence suggested no clinical difference between alpha blockers and calcium channel blockers (1 study; n=70). Reduction in the number of hospitalisations was reported by two studies and suggested a clinically important benefit in favour of alpha blockers (2 studies; n=133). For the outcome of adverse events, the evidence suggested a clinically important benefit in favour of calcium channel blockers for dizziness and headache, but no clinical difference for hypotension, flushing or unspecified adverse events (1-2 studies: n=63-3189). The evidence suggested a clinically important benefit in favour of alpha blockers for reducing the number of analgesic injections used, but no clinical difference in analgesic dosage or the number of people using analgesia (1-2 studies; n=50-3189). One study reported reduction in the number of pain episodes and this evidence suggested a clinically important benefit of alpha blockers (1 study; n=70). The quality of the evidence was Moderate to Very Low. The main reasons for downgrading the evidence were risk of bias and imprecision. In addition, two adverse event outcomes were downgraded for indirectness and one outcome for stone passage was downgraded for inconsistency.

Thirty-one studies compared alpha blockers to no treatment in adults with distal ureteric stones <10mm. For the outcomes of stone passage and time to stone passage, the evidence suggested a clinically important benefit in favour of alpha blockers (17-31 studies; n=1542-2430). For the outcome of adverse events (dizziness, headache, hypotension, retrograde ejaculation, and unspecified), the evidence suggested no clinical difference (2-9 studies; n=163-716). The evidence suggested a clinically important benefit in favour of alpha blockers for reducing hospitalisations, but no clinical difference between interventions in terms of reducing use of healthcare services (return to emergency department/primary care visit) (1-7 studies; n=77-487). Ten studies reported reduction in the number of pain episodes and the evidence suggested a clinically important benefit in favour of alpha blockers (10 studies; n=977). Three studies reported the number of people experiencing pain and this evidence also suggested a clinically important benefit in favour of alpha blockers (3 studies; n=240). In terms of colicky pain episodes, the evidence from one study suggested no clinical difference between interventions(1study: n=72). One study reported pain intensity measured by visual analogue scale and the evidence suggested a clinically important benefit in favour of no treatment (pain management only) (1 study; n=103). For reducing the number of people using analgesics, the average number of days of analgesic use, and the dose of analgesics (Diclofenac, Ketorolac and Pethidine), the evidence suggested a clinically important benefit in favour of alpha blockers (1-4 studies; n=64-301) but no clinical difference for average number of times analgesics were used or Buprenorphine dose (1-4 studies; 65-421). The quality of the evidence was Low to Very Low. The main reasons for downgrading the evidence were risk of bias, imprecision and inconsistency. One outcome for adverse events (headache) was downgraded for indirectness.

Thirteen studies compared alpha blockers to placebo in adults with distal ureteric stones <10mm. For the outcomes of stone passage and time to stone passage, the evidence suggested a clinically important benefit in favour of alpha blockers (5-13 studies; n=3369-5154). For the outcomes of hospitalisation and use of healthcare services (emergency department), the evidence suggested no clinical difference (1-3 studies; n=393-580). The evidence suggested a clinically important benefit in favour of placebo in terms of unspecified adverse events, and no clinical difference for all other adverse event outcomes (2-7 studies; n=198-3728). For reducing the number of people experiencing pain episodes, the evidence

suggested a clinically important benefit in favour of alpha blockers (1 study; n=150). In terms of the average number of pain episodes and pain intensity measured by verbal numeric pain scale, the evidence suggested no clinical difference (1-2 studies; n=219-367). The evidence suggested a clinically important benefit in favour of alpha blockers for reducing the number of people using analgesics and analgesic dose (Ketorolac and Diclofenac),but no clinical difference in the average number of episodes of analgesic use or Buprenorphine dose (2 studies; n=165-3392). The quality of the evidence was High to Very Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision. In addition, three outcomes for stone passage and analgesic use (number of people using analgesics and Diclofenac dose) were downgraded for inconsistency.

Three studies compared calcium channel blockers versus no treatment in adults with distal ureteric stones <10mm. This evidence suggested a clinically important benefit in favour of calcium channel blockers for stone passage but no clinical difference for time to stone passage (1-3 studies; n=70-179). For reducing hospitalisations, the evidence suggested a clinically important benefit of calcium channel blockers(2 studies; n=129), but no clinical difference in the average number of pain episodes (1 study; n=70). For the outcome of adverse events, the evidence suggested no clinical difference in hypotension or dizziness (1 study; n=50-59). The evidence suggested a clinical benefit in favour of calcium channel blockers for reducing the dose of analgesic (Diclofenac) (1 study; n=50). The quality of the evidence was Moderate to Very Low. The main reason for downgrading the quality of the evidence was risk of bias. In addition, two outcomes for adverse events (hypotension) and pain intensity (pain episodes) were downgraded for imprecision.

One study compared calcium channel blockers to placebo in adults with distal ureteric stones <10mm. This evidence suggested no clinical difference between interventions in terms of stone passage. The quality of the evidence was High.

#### Mid ureteric stones

One study compared alpha blockers to calcium channel blockers in adults with mid ureteric stones <10mm. This evidence suggested a clinically important benefit of calcium channel blockers for stone passage (n=81). The quality of the evidence was Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision.

Two studies compared alpha blockers to no treatment (pain management only) in adults with mid ureteric stones <10mm. For the outcome of stone passage, the evidence showed a benefit of alpha blockers (2 studies; n=27). In terms of reducing the time to stone passage, the evidence also suggested a clinically important benefit in favour of alpha blockers (1 study; n=16). For reducing the average number of episodes of analgesic use, the evidence suggested a clinically important benefit in favour of alpha blocker (1 study; n=16). The quality of the evidence was Low to Very Low. The main reason for downgrading the quality of the evidence was risk of bias. In addition, the outcomes for stone passage and analgesic use were downgraded for imprecision.

Two studies compared alpha blockers to placebo in adults with mid ureteric stones <10mm. This evidence suggested a clinically important benefit in favour of placebo for stone passage (n=126). The quality of the evidence was Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision.

One study compared calcium channel blockers to placebo in adults with mid ureteric stones <10mm. This evidence suggested no clinical difference between interventions for stone passage (n=181). The quality of the evidence was Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision.

#### Proximal ureteric stones

One study compared alpha blockers to calcium channel blockers in adults with proximal ureteric stones <10mm. This evidence suggested a clinically important benefit in favour of

alpha blockers for stone passage (n=180). The quality of the evidence was Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision.

Four studies compared alpha blockers to no treatment in adults with proximal ureteric stones <10mm. This evidence suggested a clinically important benefit in favour of alpha blockers for stone passage (4 studies; n=213). For reducing time to stone passage, the evidence also suggested a clinically important benefit in favour of alpha blockers (2 studies; n=133). The evidence suggested no clinical difference for outcomes of quality of life and analgesic use (1-2 studies; n=79-133). The quality of the evidence was Low to Very Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision.

Two studies compared alpha blockers to placebo in adults with proximal ureteric stones <10mm. This evidence suggested no clinical difference between interventions for the outcome of stone passage (n=257). The quality of the evidence was Low. The reasons for downgrading the quality of the evidence were risk of bias and imprecision.

One study compared calcium channel blockers placebo in adults with proximal ureteric stones <10mm. This evidence suggested a clinically important benefit in favour of placebo for stone passage (n=181). The quality of the evidence was Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision.

#### Children

Three studies compared alpha blockers to no treatment in children with distal ureteric stones <10mm. This evidence suggested a clinically important benefit in favour of alpha blockers for stone passage, time to stone passage and analgesic use (average number of episodes of analgesic use) (1-3 studies; n=63-147). The evidence suggested no clinical difference between interventions in terms of unspecified adverse events (2 studies; n=102). The evidence also suggested no clinical difference in average number of daily pain episodes (1 study; n=63). The quality of the evidence was Low to Very Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision. In addition, the outcome for time to stone passage was downgraded for inconsistency.

Two studies compared alpha blockers to placebo in children with distal ureteric stones <10mm. This evidence suggested a clinically important benefit in favour of alpha blockers for stone passage, time to stone passage and the number of pain episodes (2 studies; n=98). The evidence suggested a clinically important benefit in favour of placebo for headaches/dizziness, but no clinical difference between the interventions for headaches or hypotension (1 study; 1=37-61). The quality of the evidence was Low to Very Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision. In addition, outcomes for time to stone passage and pain intensity were downgraded for inconsistency.

### MET as an adjunctive therapy to surgery

#### Distal ureteric stones

Five studies compared alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only in adults with distal ureteric stones <10mm. This evidence suggested a clinically important benefit in favour of alpha blockers for stone passage, but no clinical difference in time to stone passage (2-5 studies; n=207-383). For the outcome of adverse events, the evidence suggested a clinically important benefit of SWL only for dizziness, abnormal ejaculation and headache, but no clinical difference for hypotension (1-2 studies; n=67-206). For reducing hospitalisations, the evidence suggested a clinically important benefit in favour of alpha blockers (1 study; n=88). The evidence suggested a clinically important benefit in favour of alpha blockers for reducing analgesic use (average number of episodes of analgesic use and dose of Diclofenac) (1 study; n=88-119). The quality of the evidence was Low to Very Low. The main reasons for downgrading the quality

of the evidence were risk of bias and imprecision. In addition, the outcomes for stone passage and analgesic use (dose of Diclofenac) were downgraded for indirectness.

One study compared alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only in adults with distal ureteric stones 10-12mm. This evidence suggested no clinical difference between interventions in terms of the time to stone passage or pain intensity measured by visual analogue scale(n=38). The quality of the evidence was Very Low. The main reasons for downgrading the quality of the evidence were risk of bias, indirectness and imprecision.

One study compared alpha blockers as adjunctive therapy to ureteroscopy versus ureteroscopy only in adults with distal ureteric stones <10mm. This evidence suggested a clinically important benefit in favour of alpha blockers for stone passage (n=98). The evidence also showed a benefit of alpha blockers for reducing length of hospital stay (n=98). The quality of the evidence was Moderate to Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision.

One study compared alpha blockers as adjunctive therapy to ureteroscopy versus placebo and ureteroscopy in adults with distal ureteric stones <10mm. This evidence suggested a clinically important benefit in favour of alpha blockers for stone passage and reducing the number of people using analgesics (n=102). The evidence also suggested a clinically important benefit in favour of alpha blockers for reducing the average number of pain episodes (n=102). The quality of the evidence was Low to Very Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision.

#### Mid ureteric stones

One study compared alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only in adults with mid ureteric stones 10-20mm. This evidence suggested no clinical difference between interventions in terms of the time to stone passage or pain intensity measured by visual analogue scale (n=28). The quality of the evidence was Very Low. The main reasons for downgrading the quality of the evidence were risk of bias, indirectness and imprecision.

#### **Proximal ureteric stones**

Six studies compared alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only in adults with proximal ureteric stones <10mm. This evidence a clinically important benefit in favour of alpha blockers for stone passage (6 studies; n=405). The evidence also suggested a clinically important benefit in favour of alpha blockers in terms of outcomes for quality of life, emergency department visits, pain intensity (visual analogue scale and average number of pain episodes) and analgesic dose (Diclofenac) (1-6 studies; n=54-405), but no clinical difference in terms of the time to stone passage, hospitalisation or the number of people using analgesia (1-4 studies; n=54-320). For the outcome of adverse events, the evidence suggested no clinically important difference between interventions in terms of retrograde ejaculation or dizziness (1-2 studies; n=84-172). The quality of the evidence was Moderate to Very Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision. In addition, outcomes for time to stone passage and pain intensity were downgraded for inconsistency, and analgesic use (in terms of the number of people using analgesia) was downgraded for inconsistency and indirectness.

Two studies compared alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only in adults with proximal ureteric stones 10-20mm. This evidence suggested a clinically important benefit in favour of alpha blockers for time to stone passage and stone passage, but no clinical difference in pain intensity (visual analogue scale) (1-2 studies; n=57). The quality of the evidence was Very Low. The main reasons for downgrading the quality of the evidence were risk of bias, indirectness and imprecision.

One study compared alpha blockers as adjunctive therapy to shock wave lithotripsy versus placebo and shock wave lithotripsy in adults with proximal ureteric stones <10mm. This evidence suggested a clinically important benefit in favour of alpha blockers for stone passage and time to stone passage (n=49). The quality of the evidence was Low. The main reasons for downgrading the quality of the evidence were risk of bias, indirectness and imprecision.

Two studies compared alpha blockers as adjunctive therapy to ureteroscopy versus ureteroscopy only in adults with proximal ureteric stones 10-20mm. This evidence suggested a clinically important benefit in favour of alpha blockers for stone passage (2 studies; n=254). For the outcomes of time to stone passage and hospitalisation, the evidence suggested no clinical difference between interventions (1-2 studies; n=89-165). The evidence suggested no clinical difference between interventions for dizziness, but a clinically important benefit in favour of alpha blockers for reducing the number of people experiencing pain episodes (1 study; n=89). The quality of the evidence was Moderate to Very Low. The main reasons for downgrading the quality of the evidence were risk of bias and imprecision.

#### 1.7.2 Health economic evidence statements

- Interventions studied only in separate pairwise analyses:
  - One cost-utility analysis found that placebo was not cost effective compared to MET in people with symptomatic ureteric stones of ≤ 10 mm (ICER: £42,000 per QALY gained).
  - One cost-utility analysis found that Nifedipine was not cost effective compared to Tamsulosin in people with symptomatic ureteric stones of ≤ 10 mm (ICER: £43,500 per QALY gained)

# 1.8 Recommendations

- D1. Offer alpha blockers<sup>a</sup> to adults, children and young people with distal ureteric stones less than 10 mm.
- D2. Consider oral nifedipine<sup>b</sup> for adults with distal ureteric stones less than 10 mm if alpha blockers are contraindicated.

# MET as adjunct to surgery

D3. Consider alpha blockers<sup>c</sup> as adjunctive therapy for adults having SWL for ureteric stones less than 10 mm.

<sup>a</sup> At the time of consultation (July 2018), alpha blockers did not have a UK marketing authorisation for this indication. 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 Prescribing guidance: prescribing unlicensed medicines for further information.

<sup>b</sup> At the time of consultation (July 2018), nifedipine did not have a UK marketing authorisation for this indication. 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 Prescribing guidance: prescribing unlicensed medicines for further information.

c At the time of consultation (July 2018), alpha blockers did not have a UK marketing authorisation for this indication. 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 Prescribing guidance: prescribing unlicensed medicines for further information.

#### 1.8.1 Research recommendations

What is the clinical and cost effectiveness of tamsulosin as an adjunct to ureteroscopy?

# 1.9 Rationale and impact

# 1.9.1 Why the committee made the recommendations

# Medical expulsive therapy versus each other/placebo/no treatment

Evidence showed that in adults both alpha blockers and calcium channel blockers improved passage of distal ureteric stones of less than 10 mm compared with no treatment. Alpha blockers also improved stone passage when compared with placebo. Alpha blockers offered more benefit than calcium channel blockers in terms of stone passage, hospital stay and pain, but there was no difference in time to stone passage and adverse events. The committee agreed that alpha blockers should be offered to adults with small distal ureteric stones, but calcium channel blockers should be considered when alpha blockers are contraindicated. The committee noted that all the evidence for calcium channel blockers was for oral nifedipine and so specified this in the recommendation.

Limited evidence in children showed that alpha blockers improved stone passage and time to stone passage, and decreased pain compared with no treatment or placebo. They were not associated with any more adverse events so the committee agreed that alpha blockers could be offered for children and young people with distal ureteric stones less than 10 mm.

There was not enough evidence for the committee to make recommendations on alpha blockers or calcium channel blockers for proximal or mid-ureteric stones in adults, children and young people, or for calcium channel blockers for children and young people with distal ureteric stones less than 10 mm.

Medical expulsive therapy is low cost, and the savings from interventions avoided because of this therapy are likely to offset the cost of the therapy.

#### Medical expulsive therapy as an adjunctive therapy to surgery

Evidence showed a benefit in terms of stone passage when alpha blockers were used as adjunctive therapy for adults having SWL for small distal or proximal ureteric stones (less than 10 mm). There was no difference in adverse events. The committee agreed that alpha blockers could be considered as adjunctive therapy to SWL for adults with small ureteric stones in any location. There was no evidence for mid ureteric stones less than 10 mm, however the committee agreed that this is a small group of people and usual clinical practice often involves waiting to see if the stone progresses to the distal ureter. There was not enough evidence for the committee to make a recommendation for adjunctive therapy for other interventions or for larger ureteric stones of 10 to 20 mm.

Evidence showed a benefit of alpha blockers as adjunctive therapy to URS in terms of stone passage and some outcomes relating to pain for adults with small distal ureteric stones (less than 10 mm) and proximal ureteric stones (10 to 20 mm). The committee agreed that this is not usual practice and also noted that the evidence was based on single studies. They agreed that further research on the use of alpha blockers, particularly tamsulosin, as adjunctive to URS for any stone less than 20 mm would be beneficial to inform future practice, so decided to make a research recommendation.

## 1.9.2 Impact of the recommendations on practice

Medication versus each other/placebo/no treatment

Current practice is varied, but many healthcare professionals do not offer alpha blockers for managing symptomatic ureteric stones. However, recently published evidence has called into question the established approach in the UK, and this has been confirmed by the committee's review of the evidence. Up to 2015, medical expulsive therapy was recommended practice in the UK to aid the passage of small ureteric stones. This changed after the SUSPEND trial (Pickard et al 2015), the largest RCT on this subject, concluded that there was no benefit in using alpha blockers. The guideline committee reviewed all the available evidence, some of which was more recent than the SUSPEND trial, and agreed that alpha blockers can help the passage of small ureteric stones and the management of pain. The committee agreed that prescribing alpha blockers to people with distal ureteric stones less than 10 mm or as an adjunct to shockwave lithotripsy for small ureteric stones less than 10 mm may mean a change in practice but also a potential reduction in the dose of analgesics prescribed and the length of time they are used for.

#### Medication as an adjunctive therapy to surgery

Alpha blockers are not widely used as an adjunct to SWL for ureteric stones so this will represent a change in practice. The small cost of the alpha blockers is likely to be outweighed by the saving related to improved stone clearance.

# 1.10 The committee's discussion of the evidence

# 1.10.1 Interpreting the evidence

#### 1.10.1.1 The outcomes that matter most

The committee agreed that time to stone passage, stone passage, use of healthcare services/hospitalization, quality of life, and adverse events (hypotension, dizzy spells, falls, floppy iris, retrograde ejaculation, headaches, flushing) were the outcomes that were critical for decision-making. Pain intensity and analgesic use were also considered as an important outcome.

Evidence was reported for all of the critical and important outcomes.

### 1.10.1.2 The quality of the evidence

For the majority of evidence included in this review, the quality ranged from a GRADE rating of moderate to very low. This was due to lack of blinding, presence of selection bias, and risk of measurement bias, resulting in a high or very high risk of bias rating. Additionally, the imprecise nature of the results extracted and analysed in this review, and the presence of heterogeneity for some outcomes, further downgraded the quality of the evidence. It was also difficult to classify several studies according to the strata specified in the protocols, because the results were not stratified by stone size; for this reason, some of the evidence was downgraded for indirectness.

#### 1.10.1.3 Benefits and harms

Evidence for people with both symptomatic and asymptomatic stones was searched for, however only 1 study included a population of people with asymptomatic stones, and this evidence suggested that there was no difference in the outcomes between people with symptomatic or asymptomatic stones. As there was no other evidence for people with asymptomatic stones, the committee were not confident that those with asymptomatic stones can be treated the same as those with symptomatic stones. They concluded that the recommendations should only apply to those with symptomatic stones.

It is important to note that the population that MET would be appropriate for would generally be people who were symptomatic but whose pain is not ongoing after treatment with analgesia.

#### Medication versus each other/placebo/no treatment

#### Distal ureteric stones <10mm

#### Alpha blockers

When alpha blockers were compared to placebo, the committee noted that there was a benefit of alpha blockers in terms of stone passage, time to stone passage, pain when measured as the number of people experiencing pain episodes, and analgesic use when measured as the number of people using analgesics during follow-up and average ketorolac or diclofenac dose. There was no difference between the groups in terms of all other pain and analgesic use outcomes or hospitalisation. In terms of adverse events, there was a benefit of placebo in terms of unspecified adverse events, but no difference between interventions for all other outcomes. This suggests that alpha blockers are not associated with an increased risk of experiencing adverse effects.

When compared to no treatment (pain relief only), there was a benefit of alpha blockers in terms of stone passage and time to stone passage based on two meta-analyses of 31 and 17 studies respectively. There was also a benefit of alpha blockers in terms of hospitalisation, and analgesic use (number of people using analgesics and average dose of analgesia). The committee noted that evidence from one study demonstrated a potential benefit of no treatment (pain relief only) in terms of pain intensity when measured on a visual analogue scale, but that this conflicted with other meta-analyses demonstrating a benefit of alpha blockers in terms of the number of pain episodes and the number of people experiencing pain. The committee considered that overall, there was some evidence that alpha blockers may have an analgesic effect as well as improving stone passage, which may have implications for the patient as well as from a resource use perspective, as it may reduce the amount of analgesia required. There was no difference between the interventions in terms of adverse events, which again demonstrated that alpha blockers are not associated with an increased risk of adverse effects. The committee agreed that this evidence demonstrates that there is a benefit of treating people with distal ureteric stones <10mm with alpha blockers compared to no treatment.

#### Calcium channel blockers

When compared to placebo, there was no difference between interventions in terms of stone passage. No other outcomes were reported. The committee noted that this evidence was from a single study; however they did consider that this study is from the UK and has been influential in terms of shaping current practice.

When calcium channel blockers were compared to no treatment, there was a benefit of calcium channel blockers for stone passage, hospitalisation, and analgesic use (Diclofenac dose). There was no clinical difference between interventions in terms of time to stone passage or adverse events. The committee considered that compared to no treatment, there did seem to be a benefit of calcium channel blockers. When compared to placebo, the evidence was less convincing, but they committee noted that the evidence was from a single study.

## Alpha blockers versus calcium channel blockers

The committee noted that there were more stones passed, fewer patients requiring hospitalisation, fewer diclofenac injections and fewer pain episodes for those receiving alpha blockers, compared to calcium channel blockers. The committee noted that the group receiving calcium channel blockers experienced fewer episodes of dizziness and headaches, but there was no difference between the interventions in terms of other reported outcomes,

such as time to stone passage, all other adverse events and analgesic use in terms of length of use and dose. The committee considered that the only outcomes demonstrating a clinical benefit of calcium channel blockers over alpha blockers were dizziness and headaches, and agreed that these were not very serious adverse events. The committee discussed that the benefits of alpha blockers in terms of increased stone passage, less hospitalisation and less pain, outweighed the experience of dizziness and headache. The committee considered that these benefits would reduce requirements for pain medication and may reduce the need for surgical intervention.

The committee noted that current practice is partly based on the findings of the SUSPEND trial, a large UK study which showed no benefit of alpha blocker or calcium channel blockers when compared to placebo. They considered that this is the only UK study included in the review, and so may best represent UK practice and the UK population. However, they also noted that including this study in the analysis still led to an overall benefit of alpha blockers. Therefore they agreed that this single study does not outweigh the body of evidence suggesting that alpha blockers and calcium channel blockers may be beneficial, especially considering that there are no significant harms associated with either drug. Therefore the committee agreed that alpha blockers should be offered to people with small distal ureteric stones. The committee considered the evidence for calcium channel blockers and that although there was no benefit when compared to placebo, this was based on a single study, and there did seem to be evidence of a benefit when compared to no treatment. They also considered that in the head-to-head comparison with alpha blockers, alpha blockers were favoured, so the committee agreed that calcium channel blockers should only be considered when alpha blockers are not an option. The committee noted that all of the evidence for calcium channel blockers was based on oral nifedipine. There was no evidence for other routes or administration or preparations of calcium channel blockers. Therefore they agreed to recommend oral nifedipine as a specific calcium channel blocker.

#### Mid ureteric stones <10mm

#### Alpha blockers

When compared to placebo, there was a harm of alpha blockers for stone passage; however the committee again noted concerns regarding the quality and limited evidence with only two studies with 126 participants found for this comparator.

Compared to no treatment (pain relief only), there was a benefit of alpha blockers for all of the reported outcomes, including stone passage, time to stone passage, and analgesic use. The committee noted that the evidence came from a very small single study of just 16 people, and was low and very low quality, and agreed that based on this, they did not have confidence in extrapolating this data to clinical practice.

#### Calcium channel blockers

When compared to placebo, there was no difference between interventions for stone passage. The committee noted that this evidence came from a single study.

#### Alpha blockers versus calcium channel blockers

There was a benefit of calcium channel blockers for stone passage. The committee discussed that this evidence was from a single study of 81 participants and was of low quality, and therefore they agreed that they did not have confidence in the findings.

The committee noted that all of the evidence for the mid ureteric stones population was of low or very low quality, and that all of the evidence came from a very small number of studies with very small participant numbers. The committee therefore agreed that there was a lack of sufficient convincing evidence to make a recommendation. The committee also discussed that this population would be a small number of cases in clinical practice, and that there is no

consensus on how these stones should be treated. Based on this, the committee decided not to make a recommendation.

#### Proximal ureteric stones <10mm

#### Alpha blockers

When compared to placebo, there was no clinical difference between the two groups in terms stone passage, and no other outcomes were reported.

When compared to no treatment, there was a benefit of alpha blockers for stone passage and time to stone passage. There was no clinical difference for quality of life (EuroQoL) and analgesic use (mean number of times analgesics were used). The committee noted that the evidence was of very low and low quality, and involved a small number of participants.

#### Calcium channel blockers

When compared to placebo, there was a benefit of placebo for stone passage, although noted that the evidence came from a single study.

Alpha blockers versus calcium channel blockers

The committee noted that alpha blockers appeared to be more clinically effective than calcium channel blockers in terms of stone passage. No other outcomes were reported. The committee considered that this evidence came from a single study and was low quality; therefore, the committee agreed that the evidence was not strong enough to draw conclusions from.

The committee noted that the majority of comparisons for this population were based on evidence from small studies or single studies, and all was low or very low quality. The committee discussed that the evidence for alpha blockers versus calcium channel blockers suggested a benefit of alpha blockers, and that there was a lot of uncertainty surrounding the outcomes due to low quality evidence and small participant numbers. They agreed that overall, there was a lack of convincing evidence, and so no recommendation could be made for this group.

#### Children and young people, distal ureteric stones <10mm

## Alpha blockers

The committee considered the evidence for children, and noted that all of the evidence was for distal ureteric stones <10mm. When alpha blockers (tamsulosin and doxazosin) were compared to no treatment (pain management only), the evidence suggested a benefit of alpha blockers in terms of stone passage, time to stone passage and analgesic use but no difference between the groups in terms of the number of pain episodes and unspecified adverse events. The committee noted from clinical experience that children may spontaneously pass stones more easily than adults.

When compared to placebo, there was a benefit of alpha blockers in terms of stone passage, time to stone passage and the number of pain episodes. There was a benefit of placebo in terms of headaches/dizziness, but no difference between interventions for all other adverse events.

The committee also considered that current practice for alpha blockers is varied, but that they are considered much safer for children than calcium channel blockers. Overall, the committee considered that this evidence suggests that conservative management is more likely to succeed with the use of alpha blockers, which may make the need for surgery less likely. The committee agreed that given the benefits of alpha blockers in terms of increasing stone passage and reducing the time to stone passage, as well as the potential analgesic effects and implications in terms of reducing the need for further intervention and no

evidence of increased risk of harms, alpha blockers should be offered to children and young people with distal ureteric stones <10 mm.

#### MET as an adjunctive therapy to surgery

The committee considered the evidence for MET as an adjunctive therapy to surgery. It was noted that for all comparisons, the MET was alpha blockers, and there was no evidence for calcium channel blockers or other MET drugs. It was also noted that all of the evidence for MET as adjunctive was in an adult population, and there was no evidence for the paediatric population.

#### Distal ureteric stones <10mm

Alpha blockers as adjunctive to SWL versus SWL

When MET was adjunctive to SWL in people with stones less than 10mm, the committee noted that there was a benefit of alpha blockers as adjunctive to SWL for outcomes relating to stone passage, hospitalisation and pain, but a benefit of SWL alone for most adverse events outcomes. The committee considered that because the adverse events were not serious, the benefit of adjuvant alpha blockers in terms of stone passage outweighed the experience of such adverse events.

The committee agreed to make a recommendation to consider alpha blockers as adjuvant therapy when people are having SWL. This was because the added potential benefit of MET was potentially significant, and there was a lack of serious associated harms. Current practice for people with these stones is SWL without the use of MET, and therefore use of MET would be a change in practice.

Alpha blockers as adjunctive to URS versus URS

The committee also considered the evidence for alpha blockers as adjunctive to URS. The evidence demonstrated a benefit of alpha blockers as adjuvant to URS for stone passage and length of stay, compared to URS alone. The committee discussed that the evidence for stone passage was unusual, as it was agreed that when performing a URS most UK surgeons would either fragment the stones to fragments <2-3mm, which would be expected to pass, or remove all the fragments during the procedure. Therefore the committee agreed that the use of adjuvant alpha blockers is likely to add very little benefit to UK practice. They also considered that the evidence came from a single study of 98 people.

Alpha blockers as adjunctive to URS versus placebo + URS

Evidence also demonstrated a benefit of alpha blockers as adjuvant to URS when compared to placebo as adjuvant to URS, in terms of stone passage and pain related outcomes. The committee were concerned that the evidence was based on a single study and was low and very low quality. It was further noted that the study used a ballistic method during URS, rather than laser, which does not reflect UK practice and may make stone fragments more difficult to pass, therefore potentially overestimate the effect of alpha blockers.

Overall, the committee agreed that evidence for alpha blockers as adjunctive to URS (with or without placebo) was not convincing and not sufficient on which to base a recommendation. They considered that a research recommendation investigating the use of alpha blockers as adjunctive to URS may be beneficial in terms of providing high quality evidence to help address this gap in the evidence and inform future practice.

#### Distal ureteric stones 10-20mm

Alpha blockers as adjunctive to SWL versus SWL

The committee noted that evidence from a single study of 38 participants demonstrated no difference between the interventions. Further, this evidence was very low quality. Therefore

the committee agreed that there was not convincing evidence of a benefit of adjuvant MET for people with 10-20mm stones, and decided not make a recommendation.

#### Mid ureteric stones 10-20mm

Alpha blockers as adjunctive to SWL versus SWL

There was evidence from one study in a population of mid ureteric stones. This study demonstrated no clinical difference between alpha blockers as adjunctive therapy to SWL and SWL only. The committee considered that this evidence came from a single study of 28 people, and was very low quality. The committee also considered that this was a small patient group and are not normally treated with SWL in UK clinical practice. They agreed that there was insufficient evidence to make a recommendation for this population.

#### Proximal ureteric stones <10mm

Alpha blockers as adjunctive to SWL versus SWL

When MET as adjunctive to SWL was compared to SWL alone in people with <10mm proximal ureteral stones, the committee noted a clinical benefit for alpha blockers for stone passage outcomes, quality of life outcomes, most pain outcomes, and use of healthcare services in terms of the number of ED visits at follow up. There was no clinical difference between interventions in terms retrograde ejaculation, dizziness, analgesic use and hospitalisation. The committee considered that the two adverse events are not serious and would not outweigh the benefits of increased stone passage and improved pain and quality of life.

Overall, the committee considered that the evidence for stone passage came from a number of studies and was of moderate quality; this was a key outcome of success and would lead to reduced downstream resource use. The benefits of the treatment were also thought to far outweigh any minor risks, therefore the committee made a consider recommendation for ureteric stones <10mm.

Alpha blockers as adjunctive to SWL versus placebo + SWL

When alpha blockers adjuvant to SWL was compared to placebo + SWL, the evidence demonstrated a clinical benefit for alpha blockers in terms of stone passage and time to stone passage. Although the evidence came from a single study, the committee noted that this supported the evidence for the comparison of alpha blockers adjuvant to SWL versus SWL alone.

#### Proximal ureteric stones 10-20mm

Alpha blockers as adjunctive to SWL versus SWL

Alpha blockers as adjuvant to SWL were also compared to SWL alone in people with 10-20mm stones. For this comparison the committee noted that the evidence was not conclusive. Although there was a clinical benefit of alpha blockers in terms of time to stone passage and stone passage, there was no difference between interventions in terms of pain. The committee noted that this was based on single studies and very low quality evidence. They agreed not to make a recommendation for alpha blockers as adjunctive to SWL for this stone size.

Alpha blockers as adjunctive to URS versus URS

When alpha blockers as adjuvant to URS were compared to URS alone in people with stones between 10-20mm, the committee noted that there was conflicting evidence. There was a suggested benefit of alpha blockers for stone passage and colic episodes, but no difference in terms of time to stone passage and outcomes relating to hospitalisation. The committee considered the evidence and discussed that usually during URS, the surgeon

either removes all residual stones, or breaks them down to <2-3mm fragments to pass spontaneously. Therefore alpha blockers may increase the passage rate of residual stones when the latter method is used. The committee considered that although alpha blockers may be beneficial in terms in reducing the need for pain relief and increasing passage of residual stones, most of the evidence was from single studies, which limited the degree of confidence the committee could place in the results.

The committee agreed that as with the <10mm group, the evidence for MET as an adjunct to URS was not considered sufficient to make recommendations, and is not commonly used in current practice, so a research recommendation would be beneficial to inform future practice. They agreed that the research recommendation should include all stones less than 20 mm, and include any location within the ureter.

### 1.10.2 Cost effectiveness and resource use

One cost utility analysis (Pickard 2015) was identified from the literature and presented to the committee. This was a within trial, cost utility analysis based on SUSPEND, an RCT conducted in the UK, that compared two medical expulsive therapies (tamsulosin or nifedipine) to each other and then combined the groups to compare medical expulsive therapy, in general, to placebo. There was no economic evidence identified for the use of medical expulsive therapy as adjunctive to surgery.

The study was assessed as directly applicable, as it was a UK cost utility analysis taking the NHS perspective. The study also reported values of health effects expressed in terms of QALYs and used EQ-5D data collected directly from patients. The study was rated as having potentially serious limitations because the time horizon was only the 12 week period of the RCT and no extrapolation of study results took place beyond that period; so effects and costs from any stones that might have needed treatment after this period wouldn't be captured by the analysis. Also, the estimates of relative treatment effects and resource use were not derived from a systematic literature review but from the study effectiveness data and records.

The study found that the use of medical expulsive therapy was associated with cost savings but also less QALYs (only slightly, so a negligible difference in QALYs). The cost savings are because the resource use involved in the MET group was overall lower (e.g. admission days, interventions undertaken), and is consistent with what we would think about the intervention, because if more people are passing their stone with MET, then there is less downstream resource use being consumed, such as time in hospital or other interventions.

We can change round the comparators for ease of interpretation of the ICER, so the more expensive and effective alternative (placebo – with its slightly higher QALYs) is compared to the less effective alternative (MET). This shows that the use of placebo compared to medical expulsive therapy was not cost effective (ICER of £42,000), therefore the alternative of medical expulsive therapy is a cost effective option because we are only comparing two alternatives, so if placebo is not cost effective according to the NICE threshold then that means the comparator is the cost effective choice. In effect, the placebo strategy involved more resource use overall (making it more costly), and there was a negligible difference in quality of life between the two strategies. When comparing tamsulosin to nifedipine, the study also found that tamsulosin was associated with cost savings but also less QALYs (again a negligible difference in QALYs). Tamsulosin was a cost effective option compared to nifedipine. The study results were sensitive to any changes in QoL values.

The Pickard study was included in the clinical review, and provided a conservative estimate of medical expulsive therapy's effectiveness when compared to the other studies pooled in the review for the stone passage of alpha blockers versus placebo. The point estimate was very close to the no difference line, while the pooled estimate was further on the left, favouring alpha blockers. Higher effectiveness of alpha blockers could impact cost effectiveness of medical expulsive therapy compared to placebo, making the choice of alpha

blockers (MET) even more cost effective than what the Pickard study estimated. The committee agreed that the magnitude of cost effectiveness of medical expulsive therapy compared to placebo is likely to be higher than the Pickard study demonstrated if the effectiveness of alpha blockers is in fact higher.

Unit costs of the interventions identified from the clinical review divided into alpha blockers and calcium blockers were presented using BNF prices and doses from clinical review data. Costs were presented monthly because from the trials people tended to take MET for around 4 weeks (although the committee noted that 2 weeks is also used in practice). The drug formulation was that of modified release tablets or capsules with alpha blockers represented by more drug options and calcium channel blockers represented only by nifedipine. There were differences between drug prices between the two categories and an attempt to identify the most and least expensive drug from the unit costs data was made; doxazosin (alpha blocker) was found to be the cheapest option and alfuzosin (alpha blocker) the most expensive one among alpha blockers. The GC members highlighted that the most commonly prescribed alpha blocker, tamsulosin was shown to be less expensive than nifedipine in the capsules formulation, but more expensive as a tablet.

Resource impact data were also presented, using an average monthly cost of medical expulsive therapy of £10.65 (similar to the tamsulosin tablet monthly cost), and the population with ureteric stones from HES hospital admitted activity 2015-16 data (calculus of ureter finished consultant episodes; 24,589). Even at the extreme scenario of medical expulsive therapy that would be recommended for use for all the people diagnosed with ureteric stones at hospitals, the resource impact wasn't expected to meet the NICE threshold of 'significant', as the results showed that the annual NHS spending would be around £262,000. The data from HES may well underestimate the population with ureteric stones because there may be people coping with their stone who haven't been admitted to hospital, but on the other hand the HES data is probably a mix of stone sizes whereas the recommendations are mainly for smaller stone size groups.

Passing the stone earlier will also have a QoL improvement, as an individual does not have a stone anymore (e.g. if pass a stone at 2 weeks instead of 4 weeks then that is an extra 2 weeks where the individual has returned to their normal QoL level). The time to stone passage for alpha blockers versus placebo was also shown to be clinically significant. The issue around short term pain and any associated improvement in quality of life from passing a stone earlier (or conversely the loss in quality of life from having a stone for a few more days if they didn't have MET to pass the stone earlier) was discussed. An ICER example was provided using data from the clinical review showing MET (alpha blockers specifically) would help you pass your stone on average 4 days quicker (given 4 weeks of treatment costing around £10); using quality of life data derived from the Health Survey for England 2014 as the utility level for those who don't have a stone (0.874), and the utility of patients with stones was from baseline data in the Pickard study (0.684 –(EQ-5D)). This showed that helping you pass your stone 4 days earlier would have an ICER of around £5,000. This is cost effective taking into account only a few days of pain avoided, and this is because the drug is so cheap. It was highlighted that avoiding pain of short duration wasn't expected to contribute a significant improvement in the quality of life for people achieving stone passage, therefore the committee agreed with the incremental QALY estimates presented that were very small. Discussion indicated that in practice the cheapest drug is likely to be given, which is Tamsulosin in a capsule form, and is much cheaper than the tablet form (£4 a month versus £10 a month respectively), therefore the estimates used above are likely to be overestimates.

The above example has only taken into account the people who would pass their stone *quicker* with the drug, but not the large proportion of people who would pass their stone if they used MET (compared to if they didn't), and what downstream treatment they could therefore avoid. The committee recognised that the use of MET could contribute in avoiding further downstream costs, such as surgery, from more people that passed their stone using MET.

More specifically; using as a reference point the clinical review data for the stone passage achieved with alpha blockers compared to no treatment for distal ureteric stones <10mm in adults (Table 4 in the evidence report);

- 327 more patients per 1000 that used alpha blockers passed their stones compared to the no treatment group.
- It was assumed otherwise these 327 patients would undergo a lithotripsy (a conservative estimate, as some of the patients would undergo URS that is more costly, but some patients given more time may just pass the stone and not need treatment).
- Therefore the cost from the interventions avoided considering a unit cost of £452 for an SWL session, were estimated to be £452 x 327 = £147,084. This is a conservative estimate considering only the cost of the intervention, without any retreatment or ancillary procedure cost needed for an unsuccessful first lithotripsy.
- The cost of providing alpha blockers for 1 month for 1000 people, to avoid the 327 lithotripsies, would be around £10,000 (a conservative estimate assuming a cost of the drug of £10 a month, but this could be less as mentioned in the previous paragraph).
- This makes an overall incremental saving from those additional people passing their stone with MET equal to £147,084 £10,000 = £137,084 for every 1,000 people that medical expulsive therapy is provided for.

This saving would actually allow MET to be provided to over 13,000 people. The committee were confident that this recommendation has potential to be a cost saving recommendation because of the costs offset. Not everyone in the under 10mm stone size group would go on to need an intervention to clear their stone as some may pass spontaneously with more time. The Pickard study reported that for the placebo arm the proportion requiring no further intervention at 4 weeks was 86% in the <=5mm group and 61% in the >5mm group. Breaking this down even further by size was not possible but committee opinion was that stones of between 4-7mm are the ones where clinicians would be uncertain if they would pass, and <4mm would be given more time to pass and >7mm would usually require intervention. If treating 1000 people with MET costs £10,000, then this only has to avoid 22 sessions of SWL (which would be for 2.5% of the 1000 people (assuming one session per person)) or around 5 URS's to make the intervention cost neutral. This is likely to be achievable given the low numbers, and so even if only a proportion of people go on to avoid treatment it is still likely that MET is cost saving.

The above is an illustrative calculation which is rather simplistic. As well as the interventions unit costs, the cost of other resources should be considered; such as appointments with staff including GPs and consultants, for review of medication therapy and any monitoring of adverse events. Additionally, the clinical review showed that MET was associated with fewer hospitalisations when compared to no treatment. MET had more adverse events, but mainly dizziness and headache, which the committee considered to be minor adverse events.

Used as an adjunct to surgery, alpha blockers were also shown to be effective at improving stone passage, which means further treatments could be avoided.

The committee agreed that MET is likely to be a cost effective if not cost saving treatment, and recommended alpha blockers to adults and children with distal ureteric stones <10mm, and the consideration of calcium channel blockers when alpha blockers are contraindicated, as well as recommending alpha blockers an adjunct to SWL for ureteric stones <10mm.

#### 1.10.3 Other factors the committee took into account

The committee noted that both alpha blockers and calcium channel blockers are not licensed specifically for renal stones, but they are licensed for other conditions. Alpha blockers are mainly used for men with symptomatic lower urinary tract symptoms and the management of

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acute retention of urine, with some also indicated for hypertension. Calcium channel blocker nifedipine is primarily licensed for Raynaud's syndrome. Alpha blockers and calcium channel blockers are not licensed for children.

The committee noted that the evidence included studies that used Silodosin, and that this is not available in the UK.

The committee were aware that the SUSPEND trial reported other quality of life data, however noted that due to the way in which it was reported, it did not meet the protocol and so could not be considered.

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# **Appendices**

# Appendix A: Review protocols

Table 29: Review protocol: Medical expulsive therapy

Field	Content
Review question	Is medical expulsive therapy clinically and cost-effective in managing people with ureteric stones?
Type of review question	Intervention 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.
Objective of the review	To determine whether medical expulsive therapy enhances stone passage in people with ureteric stones.
	Key issues and questions from the scope:
	2 Pharmacological management of symptomatic renal and ureteric stones (for example, non-steroidal anti-inflammatory drugs, opioids and alpha-blockers).
	2.1 What are the most clinical and cost-effective drugs for managing symptomatic renal or ureteric stones?
	4 Managing asymptomatic renal and ureteric stones.
	4.1 What is the most clinically and cost-effective management (surgical and non-surgical) of asymptomatic renal and ureteric stones?
Eligibility criteria – People (adults, children and young people) with symptomatic ureteric stones condition / issue / domain	
Eligibility criteria – intervention(s) / exposure(s) / prognostic factor(s)	<ul> <li>Medical expulsive therapy:</li> <li>Alpha blockers (Tamsulosin, Alfuzosin, Doxazosin, Naftopidil, Silodosin, Terazosin)</li> <li>Calcium channel blocker (Nifedipine)</li> </ul>
Eligibility criteria –	Compared to:
comparator(s) / control or reference (gold) standard	Each other
reference (gold) standard	Placebo
	<ul><li>No treatment</li><li>Steroids</li></ul>
Outcomes and	Critical outcomes:
prioritisation	Time to stone passage
	Stone passage
	<ul><li>Use of healthcare services/Hospitalisation</li><li>Quality of life</li></ul>
	<ul> <li>Quality of life</li> <li>Adverse events (hypotension, dizzy spells, falls, floppy iris,</li> </ul>
	retrograde ejaculation, headaches, flushing) Important outcomes:
	<ul> <li>Pain intensity (visual analogue scale, verbal ratings, descriptive scales, time to pain relief, need to rescue medication)</li> </ul>
	Analgesic use
Eligibility criteria – study design	Randomised controlled trials (RCTs), systematic reviews of RCTs.  If no RCT evidence is available, non-randomised comparative studies, prospective and retrospective search for observational studies.

Field	Content
Other inclusion exclusion	Bladder stones
criteria	Open surgery for renal (kidney and ureteric) stones Non-English language studies
Proposed sensitivity / subgroup analysis, or meta-regression	Strata:  Population
Selection process – duplicate screening / selection / analysis	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.
Data management (software)	Pairwise meta-analyses were performed using Cochrane Review Manager (RevMan5).  GRADEpro was used to assess the quality of evidence for each outcome.  Endnote for bibliography, citations, sifting and reference management Data extractions performed using EviBase, a platform designed and maintained by the National Guideline Centre (NGC)
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

Field	Content
Data collection process – forms / duplicate	A standardised evidence table format will be used, and published as appendix D of the evidence report.
Data items – define all variables to be collected	For details please see evidence tables in Appendix D (clinical evidence tables) or H (health economic evidence tables).
Methods for assessing bias at outcome / study level	Standard study checklists were used to critically appraise individual studies. For details please see section 6.2 of Developing NICE guidelines: the manual  The risk of bias across all available evidence was evaluated for each outcome using an adaptation of the 'Grading of Recommendations Assessment, Development and Evaluation (GRADE) toolbox' developed by the international GRADE working group http://www.gradeworkinggroup.org/
Criteria for quantitative synthesis	For details please see section 6.4 of Developing NICE guidelines: the manual.
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.
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 30: 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	<ul> <li>Populations, interventions and comparators must be as specified in the individual review protocol above.</li> </ul>

- 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. 138

#### 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 over whether it should be included.

## Where there is discretion

The health economist will make a decision based on the relative applicability and quality of the available evidence for that question, in discussion with the 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 applicable studies and to selectively exclude the remaining studies. All studies excluded on the basis of applicability or 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.

Table 31: Database date parameters and filters used

Database	Dates searched	Search filter used
Medline (OVID)	1946 – 21 March 2018	Exclusions Randomised controlled trials Systematic review studies Observational studies
Embase (OVID)	1974 – 21 March 2018	Exclusions Randomised controlled trials Systematic review studies Observational studies
The Cochrane Library (Wiley)	Cochrane Reviews to 2018 Issue 3 of 12 CENTRAL to 2018 Issue 2 of 12 DARE, and NHSEED to 2015 Issue 2 of 4 HTA to 2016 Issue 2 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.	(expuls* adj3 (therap* or treatment* or intervention*)).ti,ab.	
28.	((calculus or calculi or stone*) adj3 (expuls* or pass*)).ti,ab.	
29.	exp Adrenergic alpha-Antagonists/ or exp Adrenergic alpha-1 Receptor Antagonists/ or exp Adrenergic alpha-2 Receptor Antagonists/	
30.	(alpha* adj3 blocker*).ti,ab.	
31.	(tamsulosin or alfuzosin or doxazosin).ti,ab.	
32.	(Cositam or Contiflo or Diffundox or Faramsil or Flectone or Flomax or Flomaxtra or Galebon or Losinate or Pamsvax or Petyme or Pinexel or Prosurin or Tabphyn or Tamfrex or Tamurex or Combodart or Urimax or Vesomni or Besavar or Uroxatral or Xatral or Fuzatal or Varsan or Larbex or Cardozin or Cardura or Doxadura or Raporsin or Slocinx).ti,ab.	
33.	exp Calcium Channel Blockers/	
34.	(calcium channel blocker* or c-channel blocker* or Ca channel blocker* or CCB).ti,ab.	
35.	exp Nifedipine/	
36.	nifedipine.ti,ab.	
37.	(Adalat or Adipine or Calchan or Coracten or Cordipin or Cordipine or Corinfar or Fenigidin or Fortipine or Korinfar or Nifangin or Nifedipress or Nimodrel or Procardia or Tenif or Tensipine or Valni or Vascard).ti,ab.	
38.	or/27-37	
39.	26 and 38	
40.	randomized controlled trial.pt.	
41.	controlled clinical trial.pt.	
	•	

40	and and the different
42.	randomi#ed.ti,ab.
43.	placebo.ab.
44.	randomly.ti,ab.
45.	Clinical Trials as topic.sh.
46.	trial.ti.
47.	or/40-46
48.	39 and 47
49.	Meta-Analysis/
50.	exp Meta-Analysis as Topic/
51.	(meta analy* or metanaly* or meta regression).ti,ab.
52.	((systematic* or evidence*) adj3 (review* or overview*)).ti,ab.
53.	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.
54.	(search strategy or search criteria or systematic search or study selection or data extraction).ab.
55.	(search* adj4 literature).ab.
56.	(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.
57.	cochrane.jw.
58.	((multiple treatment* or indirect or mixed) adj2 comparison*).ti,ab.
59.	or/49-58
60.	39 and 59
61.	Epidemiologic studies/
62.	Observational study/
63.	exp Cohort studies/
64.	(cohort adj (study or studies or analys* or data)).ti,ab.
65.	((follow up or observational or uncontrolled or non randomi#ed or epidemiologic*) adj (study or studies or data)).ti,ab.
66.	((longitudinal or retrospective or prospective or cross sectional) and (study or studies or review or analys* or cohort* or data)).ti,ab.
67.	Controlled Before-After Studies/
68.	Historically Controlled Study/
69.	Interrupted Time Series Analysis/
70.	(before adj2 after adj2 (study or studies or data)).ti,ab.
71.	or/61-70
72.	exp case control study/
73.	case control*.ti,ab.
74.	or/72-73
75.	71 or 74
76.	Cross-sectional studies/
77.	(cross sectional and (study or studies or review or analys* or cohort* or data)).ti,ab.
78.	or/76-77
79.	71 or 78
80.	71 or 74 or 78
81.	39 and 80
82.	48 or 60
83.	81 or 82

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.	(expuls* adj3 (therap* or treatment* or intervention*)).ti,ab.	
26.	((calculus or calculi or stone*) adj3 (expuls* or pass*)).ti,ab.	
27.	exp alpha adrenergic receptor blocking agent/ or exp alpha 1 adrenergic receptor blocking agent/ or exp alpha 2 adrenergic receptor blocking agent/	
28.	(alpha* adj3 blocker*).ti,ab.	
29.	(tamsulosin or alfuzosin or doxazosin).ti,ab.	
30.	(Flomax or Flomaxtra or Urimax or Besavar or Uroxatral or Xatral or Cardozin or Cardura or Doxadura or Raporsin or Slocinx).ti,ab.	
31.	exp calcium channel blocking agent/	
32.	(calcium channel blocker* or c-channel blocker* or Ca channel blocker* or CCB).ti,ab.	
33.	exp nifedipine/	
34.	nifedipine.ti,ab.	
35.	(Adalat or Adipine or Calchan or Coracten or Cordipin or Cordipine or Corinfar or Fenigidin or Fortipine or Korinfar or Nifangin or Nifedipress or Nimodrel or Procardia or Tensipine or Valni or Vascard).ti,ab.	
36.	or/25-35	
37.	24 and 36	
38.	random*.ti,ab.	
39.	factorial*.ti,ab.	
40.	(crossover* or cross over*).ti,ab.	

41. 42. 43. 44.	((doubl* or singl*) adj blind*).ti,ab.  (assign* or allocat* or volunteer* or placebo*).ti,ab.	
43.		
44.	crossover procedure/	
	single blind procedure/	
45.	randomized controlled trial/	
46.	double blind procedure/	
47.	or/38-46	
48.	37 and 47	
49.	systematic review/	
50.	meta-analysis/	
51.	(meta analy* or metanaly* or metaanaly* or meta regression).ti,ab.	
52.	((systematic* or evidence*) adj3 (review* or overview*)).ti,ab.	
53.	(reference list* or bibliograph* or hand search* or manual search* or relevant journals).ab.	
54.	(search strategy or search criteria or systematic search or study selection or data extraction).ab.	
55.	(search* adj4 literature).ab.	
56.	(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.	
57.	cochrane.jw.	
58.	((multiple treatment* or indirect or mixed) adj2 comparison*).ti,ab.	
59.	or/49-58	
60.	37 and 59	
61.	Clinical study/	
62.	Observational study/	
63.	family study/	
64.	longitudinal study/	
65.	retrospective study/	
66.	prospective study/	
67.	cohort analysis/	
68.	follow-up/	
69.	cohort*.ti,ab.	
70.	68 and 69	
71.	(cohort adj (study or studies or analys* or data)).ti,ab.	
72.	((follow up or observational or uncontrolled or non randomi#ed or epidemiologic*) adj (study or studies or data)).ti,ab.	
73.	((longitudinal or retrospective or prospective or cross sectional) and (study or studies or review or analys* or cohort* or data)).ti,ab.	
74.	(before adj2 after adj2 (study or studies or data)).ti,ab.	
75.	or/61-67,70-74	
76.	exp case control study/	
77.	case control*.ti,ab.	
78.	or/76-77	
79.	75 or 78	
80.	cross-sectional study/	
81.	(cross sectional and (study or studies or review or analys* or cohort* or data)).ti,ab.	
82.	or/80-81	

83.	75 or 82
84.	75 or 78 or 82
85.	37 and 84
86.	48 or 60
87.	85 or 86

Cochrane Library (Wiley) search terms

#1.	MeSH descriptor: [Urolithiasis] explode all trees	
#2.	(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.	(expuls* near/3 (therap* or treatment* or intervention*)):ti,ab	
#8.	((calculus or calculi or stone*) near/3 (expuls* or pass*)):ti,ab	
#9.	MeSH descriptor: [Adrenergic alpha-Antagonists] explode all trees	
#10.	MeSH descriptor: [Adrenergic alpha-1 Receptor Antagonists] explode all trees	
#11.	MeSH descriptor: [Adrenergic alpha-2 Receptor Antagonists] explode all trees	
#12.	(alpha* near/3 blocker*):ti,ab	
#13.	(tamsulosin or alfuzosin or doxazosin):ti,ab	
#14.	(Cositam or Contiflo or Diffundox or Faramsil or Flectone or Flomax or Flomaxtra or Galebon or Losinate or Pamsvax or Petyme or Pinexel or Prosurin or Tabphyn or Tamfrex or Tamurex or Combodart or Urimax or Vesomni or Besavar or Uroxatral or Xatral or Fuzatal or Varsan or Larbex or Cardozin or Cardura or Doxadura or Raporsin or Slocinx):ti,ab	
#15.	MeSH descriptor: [Calcium Channel Blockers] explode all trees	
#16.	(calcium channel blocker* or c-channel blocker* or Ca channel blocker* or CCB):ti,ab	
#17.	MeSH descriptor: [Nifedipine] explode all trees	
#18.	nifedipine:ti,ab	
#19.	(Adalat or Adipine or Calchan or Coracten or Cordipin or Cordipine or Corinfar or Fenigidin or Fortipine or Korinfar or Nifangin or Nifedipress or Nimodrel or Procardia or Tenif or Tensipine or Valni or Vascard):ti,ab	
#20.	(or #7-#19)	
#21.	#6 and #20	

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

Table 32: Database date parameters and filters used

Database	Dates searched	Search filter used
Medline	2014 – 9 March 2018	Exclusions Health economics studies

Database	Dates searched	Search filter used
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

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/
26.	exp economic evaluation/
27.	exp health care cost/
28.	exp fee/
29.	budget/

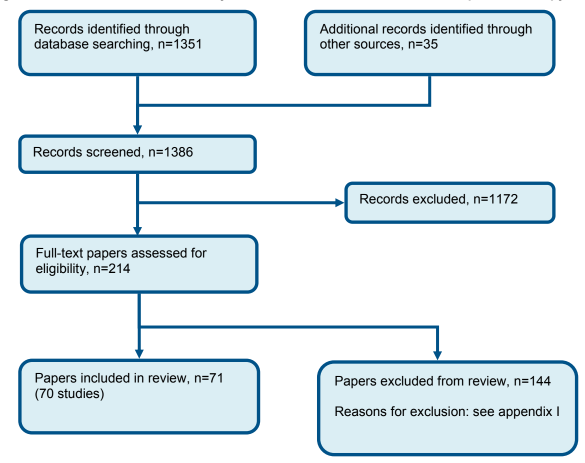
30.	funding/
31.	budget*.ti,ab.
32.	cost*.ti.
33.	(economic* or pharmaco?economic*).ti.
34.	(price* or pricing*).ti,ab.
35.	(cost* adj2 (effective* or utilit* or benefit* or minimi* or unit* or estimat* or variable*)).ab.
36.	(financ* or fee or fees).ti,ab.
37.	(value adj2 (money or monetary)).ti,ab.
38.	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 medical expulsive therapy



# **Appendix D: Clinical evidence tables**

Study	Abdelaziz 2017 <sup>3</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=98)
Countries and setting	Conducted in Saudi Arabia; Setting: not reported
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 1 week + 2 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: medical history, physical examination and laboratory investigations, abdominal x-rays for KUB, urinary ultrasonography, intravenous urography and/or abdominal computed tomography
Stratum	Adults (≥16 years), ureteric stone <1 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	≥18 years; single, radio opaque, lower ureteral stone, 5-10mm in maximum diameter
Exclusion criteria	pregnant women; history of endoscopic or open ureteral surgery, persistent renal pain; urinary tract infection; renal impairment; solitary kidney; bilateral ureteral stones; high grade hydronephrosis; hypersensitivity to alpha-blockers
Recruitment/selection of patients	consecutive patients meeting the inclusion criteria during recruitment period
Age, gender and ethnicity	Age - Mean (SD): 36.27 (6.7). Gender (M:F): 64/34. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness: NA
Interventions	(n=51) Intervention 1: Alpha blockers and URS. Tamsulosin 0.4mg daily before URS. Duration 1 week. Concurrent medication/care: ureterorenoscopy and NSAIDs. Indirectness: No indirectness; Indirectness comment: NA
	(n=47) Intervention 2: Surgery - URS. URS. Duration procedure time. Concurrent medication/care: NSAIDs. Indirectness: No indirectness; Indirectness comment: NA

Funding No funding

#### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND URS versus URS

Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: length of stay at during Hospitalisation; Group 1: mean 1.2 days (SD 0.6); n=51, Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, stone location (left/right) or size; Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone free rate at 2 weeks; Group 1: 48/51, Group 2: 41/47 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, stone location (left/right) or size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Adverse events; Pain intensity; Analgesic use; Time to stone passage

Renal and ureteric stones: Medical expulsive therapy

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Study	Abdel-Meguid 2010 <sup>1</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=150)
Countries and setting	Conducted in Saudi Arabia; Setting: Department of Urology, University Hospital
Line of therapy	1st line
Duration of study	Intervention + follow up: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	>18 years; single, unilateral, newly diagnosed, 4-10mm in transverse diameter, distal ureteral stones; in paired kidneys patients with minimal or no ipsilateral hydronephrosis, normal contralateral kidney and normal overall renal functions; stones evident in either KUB x-ray or ultrasonography or both
Exclusion criteria	history of ipsilateral ureteral endoscopic or surgical manipulations or ESWL; patients with symptomatic urinary tract infections; pregnant or lactating women; patients already receiving alpha blockers, beta blockers, calcium channel antagonists or corticosteroids; patients with serious medical conditions
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Median (range): Group A 36 years (19-72), Group B 34 years (20-67). Gender (M:F): 103/47. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=75) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg oral tablets once daily. Duration up to 4 weeks. Concurrent medication/care: hydration and analgesia (diclofenac 100mg) as needed, patients with non-symptomatic urinary tract infections were given antibiotics. Indirectness: No indirectness
	(n=75) Intervention 2: Placebo. Placebo. Duration up to 4 weeks. Concurrent medication/care: hydration and analgesia (diclofenac 100mg) as needed, patients with non-symptomatic urinary tract infections were given antibiotics. Indirectness: No indirectness
Funding	Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus PLACEBO

#### Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage at 4 weeks; Group 1: 61/75, Group 2: 42/75
Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: comparable for sex, age and stone size; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: episodes of renal colic at 4 weeks; Group 1: 20/75, Group 2: 58/75
Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: comparable for sex, age and stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Time to stone passage; Adverse events; Analgesic use ; Hospitalisation/ Use of healthcare
study	services

Study	Agrawal 2009 <sup>6</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=102)
Countries and setting	Conducted in India; Setting: Not reported
Line of therapy	1st line
Duration of study	Follow up (post intervention): 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Plain x-rays of the abdomen, ultrasonography of the urinary system, intravenous urography and non-contrast CT in selected patients
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	Patients presenting with stone <1cm in size located in the distal part of the ureter (juxtavesical part and ureterovesical junction)
Exclusion criteria	Urinary tract infection, sever hydroureteronephrosis, diabetes mellitus, multiple stones, hypotension, pregnancy, previous spontaneous stone expulsion, distal ureteral surgery and history of intake of any of the following: warfarin, $\alpha$ -adrenergic blockers, calcium antagonist, steroids, cimetidine

CONSULTATION

Recruitment/selection of patients	Not reported
Age, gender and ethnicity	Age - Range: 15-60. Gender (M:F): 78/24. Ethnicity: Not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not stated / Unclear 3. Obesity /skin-to-stone distance: Not stated / Unclear 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not stated / Unclear 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=68) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin once daily 0.4mg or 10mg alfuzosin once daily. Duration 4 weeks. Concurrent medication/care: Instructions to drink at least 3 L of fluids daily, diclofenac injection (75mg) intramuscularly on demand for pain relief. Indirectness: No indirectness (n=34) Intervention 2: Placebo. Placebo. Duration 4 weeks. Concurrent medication/care: Instructions to drink at least 3 L of fluids daily, diclofenac injection (75mg) intramuscularly on demand for pain relief. Indirectness: No indirectness
Funding	Funding not stated

### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN/ALFUZOSIN versus PLACEBO

### Protocol outcome 1: Stone passage at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Stone expulsion at 4 weeks; Group 1: 52/68, Group 2: 12/34 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Adverse events at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Postural hypotension at 4 weeks; Group 1: 0/68, Group 2: 0/34 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Retrograde ejaculation at 4 weeks; Group 1: 3/68, Group 2: 0/34 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Dizziness at 4 weeks; Group 1: 9/68, Group 2: 2/34 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Headache at 4 weeks; Group 1: 8/68, Group 2: 1/34 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,

Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:	

Protocol outcomes not reported by the study	Quality of life at Define; Time to stone passage at Define; Pain intensity at Define; Analgesic use at Define; Hospitalisation/ Use of healthcare services at Define

Study	Agarwal 2009 <sup>5</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=40)
Countries and setting	Conducted in India; Setting: department of urology, single centre
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 3 months
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: radiological and metabolic evaluation
Stratum	Adults (≥16 years), ureteric stone <1 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	single upper ureteric stone <15mm electing SWL
Exclusion criteria	extremes of ages; serum creatinine >2mg/dL; concomitant stones in ipsilateral kidney; radiolucent stones; history of previous unsuccessful SWL; active urinary tract infection; diabetes; concomitant treatment with calcium channel blockers, alpha-blockers and/or corticosteroids; previous pyeloureteral surgery; severe vertebral malformation; morbid obesity; pregnancy; aortic and/or renal artery aneurysm; uncorrected coagulopathy; ureteral stent
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): intervention group: 32.4 (8.7), control group: 35.5 (15.4). Gender (M:F): 31/9. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Upper ureteric stones
Indirectness of population	Serious indirectness: included 14 patients with stones <10mm, 20 with 10mm stones and 10 with stones >10mm
Interventions	(n=20) Intervention 1: Alpha blockers and SWL. Tamsulosin 0.4mg daily starting from the day of SWL, just before the session. SWL performed up to a maximum of 4 sessions for any significant ureteric fragment, ureteroscopy offered if stone did not show adequate fragmentation after 2 sessions. Duration up to 3

	months. Concurrent medication/care: over-the-counter NSAIDs, antispasmodics or Tramadol on demand. Indirectness: No indirectness; Indirectness comment: NA  (n=20) Intervention 2: Surgery - SWL. SWL performed up to a maximum of 4 sessions for any significant ureteric fragment, ureteroscopy offered if stone did not show adequate fragmentation after 2 sessions.
	Duration up to 5 weeks. Concurrent medication/care: over-the-counter NSAIDs, antispasmodics or Tramadol on demand. Indirectness: No indirectness; Indirectness comment: NA
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL

#### Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone clearance at the end of the study at study duration; Group 1: 19/20, Group 2: 18/20 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, weight, height, BMI or stone size; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: days to stone clearance at study duration; Group 1: mean 30.7 days (SD 19.6); n=20, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, weight, height, BMI or stone size; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 3: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: VAS at unclear; Group 1: mean 25.3 (SD 17.9); n=20,
Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, weight, height, BMI or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Adverse events; Analgesic use; Hospitalisation/ Use of healthcare services
study	

Study	Ahmad 2015 <sup>7</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=100)
Countries and setting	Conducted in Pakistan; Setting: Armed Forces Institute of Urology
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	age >18 years; stone size 8mm or smaller in distal third of the ureter
Exclusion criteria	ureteric obstruction; distal ureteric stricture; previous ureteral surgery; solitary kidney; aberrant ureteral anatomy; urinary tract infection; radiolucent stone
Recruitment/selection of patients	consecutive meeting the inclusion/exclusion criteria during the study period (10 months)
Age, gender and ethnicity	Age - Mean (range): . Gender (M:F): not reported . Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=50) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac Sodium 50mg 8 hourly on required basis. Indirectness: No indirectness  (n=50) Intervention 2: Placebo. Placebo 1 capsule daily. Duration up to 4 weeks. Concurrent
Funding	medication/care: Diclofenac Sodium 50mg 8 hourly on required basis. Indirectness: No indirectness Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus PLACEBO

Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number requiring hospitalisation at 4 weeks; Group 1: 0/49, Group 2: 1/48 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,

Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size and stone lateralisation; Group 1 Number missing: 1; Group 2 Number missing: 2

#### Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 4 weeks; Group 1: 42/49, Group 2: 26/48
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size and stone
lateralisation; Group 1 Number missing: 1; Group 2 Number missing: 2

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: drug side effects at 4 weeks; Group 1: 0/49, Group 2: 0/48
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size and stone
lateralisation; Group 1 Number missing: 1; Group 2 Number missing: 2

### Protocol outcome 4: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number requiring analgesic (diclofenac) at 4 weeks; Group 1: 9/49, Group 2: 19/48
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size and stone
lateralisation; Group 1 Number missing: 1; Group 2 Number missing: 2

Protocol outcomes not reported by the study

Quality of life; Pain intensity; Time to stone passage

Study	Ahmed 2017 <sup>9</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=183)
Countries and setting	Conducted in Saudi Arabia; Setting: department of urology, 3 centres
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 1 week + 8 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: unenhanced abdominal CT
Stratum	Adults (≥16 years), ureteric stone 1-2 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	adult patients (≥18 years); proximal ureteral stones ≥10mm; scheduled for URS lithotripsy
Exclusion criteria	pregnancy; persistent moderate/severe pain; bilateral ureteral stones; solitary kidney; renal impairment; ureteral stricture and/or history of previous ureteral surgery or endoscopy
Recruitment/selection of patients	consecutive patients meeting the inclusion criteria during the recruitment period
Age, gender and ethnicity	Age - Mean (SD): 36.7 (11.1). Gender (M:F): 98/67. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Upper ureteric stones
Indirectness of population	No indirectness: NA
Interventions	(n=91) Intervention 1: Alpha blockers and URS. Tamsulosin 0.4mg daily before URS. Duration 1 week. Concurrent medication/care: not reported. Indirectness: No indirectness; Indirectness comment: NA (n=92) Intervention 2: Surgery - URS. URS. Duration procedure time. Concurrent medication/care: not reported. Indirectness: No indirectness; Indirectness comment: NA
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND URS versus URS

Protocol outcome 1: Hospitalisation/ Use of healthcare services

<sup>-</sup> Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: Hospitalisation time at initial procedure; Group 1: mean 1.2 days (SD 0.3); n=81, Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, BMI, stone density,

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stone size or location (left/right); Group 1 Number missing: 10, Reason: non-compliance with medication (4), lost to follow up/did not complete investigation (6); Group 2 Number missing: 8, Reason: lost to follow up/did not complete investigations (8)

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: readmission at 8 weeks; Group 1: 3/81, Group 2: 5/84
Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, BMI, stone density,
stone size or location (left/right); Group 1 Number missing: 10, Reason: non-compliance with medication (4), lost to follow up/did not complete
investigation (6); Group 2 Number missing: 8, Reason: lost to follow up/did not complete investigations (8)

#### Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: stone free rate at 4 weeks; Group 1: 74/81, Group 2: 67/84
Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, BMI, stone density,
stone size or location (left/right); Group 1 Number missing: 10, Reason: non-compliance with medication (4), lost to follow up/did not complete
investigation (6); Group 2 Number missing: 8, Reason: lost to follow up/did not complete investigations (8)

Protocol outcomes not reported by the	Quality of life; Adverse events; Pain intensity; Analgesic use; Time to stone passage
study	

Study	Al-ansari 2010 <sup>14</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=100)
Countries and setting	Conducted in Qatar; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	ureteral stones 10mm or smaller located below the common iliac vessels as assessed on non-contrast computed tomography

Exclusion criteria	age <18 years; non-radiopaque stones; multiple stones; urinary tract infections; severe hydronephrosis; pregnancy; hypotension; peptic ulcer; history of endoscopic or open ureteral surgery; taking calcium antagonist medications; refusal to participate
Recruitment/selection of patients	consecutive patients meeting the inclusion/exclusion criteria during the study period
Age, gender and ethnicity	Age - Mean (SD): 36.7 (9.35), range 21-55 years. Gender (M:F): 67/33. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=50) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg once daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 75mg injection on demand and advice to drink a minimum of 2 L of water daily. Indirectness: No indirectness
	(n=50) Intervention 2: Placebo. Placebo. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 75mg injection on demand and advice to drink a minimum of 2 L of water daily. Indirectness: No indirectness
Funding	Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus PLACEBO

# Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 4 weeks; Group 1: 41/50, Group 2: 28/46
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover
- Low, Subgroups Low; Indirectness of outcome: No indirectness; Baseline details: comparable regarding age, sex, stone size and stone location; Group 1 Number missing: 0; Group 2 Number missing: 4

### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: day the patient reported the passage of the stone, confirmed by absence of radiopaque calculi shadow on x-ray at 4 weeks; Group 1: mean 6.4 days (SD 2.77); n=50,
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low; Indirectness of outcome: No indirectness; Baseline details: comparable regarding age, sex, stone size and stone location; Group 1 Number missing: 0; Group 2 Number missing: 4

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: retrograde ejaculation at 4 weeks; Group 1: 1/32, Group 2: 0/35 Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,

Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: comparable regarding age, sex, stone size and stone location; Group 1 Number missing: 0; Group 2 Number missing: 4

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 4 weeks; Group 1: 2/50, Group 2: 2/46
  Risk of bias: All domain High, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement High,
  Crossover Low, Subgroups Low; Indirectness of outcome: No indirectness; Baseline details: comparable regarding age, sex, stone size and stone location; Group 1 Number missing: 0; Group 2 Number missing: 4
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: headache at 4 weeks; Group 1: 2/50, Group 2: 2/46
  Risk of bias: All domain High, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement High,
  Crossover Low, Subgroups Low; Indirectness of outcome: No indirectness; Baseline details: comparable regarding age, sex, stone size and stone location; Group 1 Number missing: 0; Group 2 Number missing: 4
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: postural hypotension at 4 weeks; Group 1: 1/50, Group 2: 0/46
  Risk of bias: All domain High, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement High,
  Crossover Low, Subgroups Low; Indirectness of outcome: No indirectness; Baseline details: comparable regarding age, sex, stone size and stone location; Group 1 Number missing: 0; Group 2 Number missing: 4

### Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of pain episodes at 4 weeks; Group 1: mean 1.6 pain episodes (SD 1.3); n=50, Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low, Subgroups - Low; Indirectness of outcome: No indirectness; Baseline details: comparable regarding age, sex, stone size and stone location; Group 1 Number missing: 0; Group 2 Number missing: 4

# Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: need for Diclofenac injection at 4 weeks; Group 1: mean 0.9 (SD 0.93); n=50, Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low; Indirectness of outcome: No indirectness; Baseline details: comparable regarding age, sex, stone size and stone location; Group 1 Number missing: 0; Group 2 Number missing: 4
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dose of Diclofenac injection at 4 weeks; Group 1: mean 67.5 mg (SD 69.8); n=50, Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover Low, Subgroups Low; Indirectness of outcome: No indirectness; Baseline details: comparable regarding age, sex, stone size and stone location; Group 1 Number missing: 0; Group 2 Number missing: 4

Protocol outcomes not reported by the study

Quality of life; Hospitalisation/ Use of healthcare services

Study	Aldaqadossi 2015 <sup>15</sup>
Study type	RCT ( randomised; Parallel)
Number of studies (number of participants)	1 (n=67)
Countries and setting	Conducted in Egypt; Setting: not reported
Line of therapy	1st line
Duration of study	Intervention + follow up: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis
Stratum	Children (<16 years)
Subgroup analysis within study	Not applicable
Inclusion criteria	children presenting with a distal ureteric stone of <1cm below the common iliac vessels as assessed by enhanced CT
Exclusion criteria	bilateral ureteric stones, multiple stones, marked hydronephrosis, urinary tract infection, urinary tract abnormalities, voiding dysfunction, any previous open or endoscopic ureteric surgery
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group 7.7 years (3.02), control group 7.25 years (2.7). Gender (M:F): 36/27. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=31) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg for patients >5 years and 0.2mg for younger patients. Duration up to 4 weeks. Concurrent medication/care: Ibuprofen 4-10mg/kg orally every 6-8 hours as needed; in the case of intractable pain, Ketorolac 0.5-1mg/kg intramuscularly. Indirectness: No indirectness
	(n=32) Intervention 2: Pain management only - NSAIDs. Ibuprofen 4-10mg/kg every 6-8 hours as needed; in the case of intractable pain Ketorolac 0.5-1mg/kg intramuscularly. Duration 4 weeks. Concurrent medication/care: NA. Indirectness: No indirectness
Funding	No funding
RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS	

#### Protocol outcome 1: Stone passage

- Actual outcome for Children (<16 years): stone-free rate at 4 weeks; Group 1: 25/31, Group 2: 20/32; Comments: numbers calculated from percentages Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 2: Time to stone passage

- Actual outcome for Children (<16 years): time to stone expulsion (days) at 4 weeks; Group 1: mean 7.7 days (SD 1.9); n=31, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Children (<16 years): major side effects at 4 weeks; Group 1: 0/31, Group 2: 0/32 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing; Group 2 Number missing;

#### Protocol outcome 4: Pain intensity

- Actual outcome for Children (<16 years): daily pain episodes at 4 weeks; Group 1: mean 1.6 mean number of daily pain episodes (SD 1.6); n=31, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 5: Analgesic use

- Actual outcome for Children (<16 years): mean number of Ketorolac injections during the study at 4 weeks; Group 1: mean 0.55 (SD 0.8); n=31, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Hospitalisation/ Use of healthcare services

Study	Aldemir 2011 <sup>16</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=90)
Countries and setting	Conducted in Turkey; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 10 days
Method of assessment of guideline condition	Adequate method of assessment/diagnosis
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	older than 17 years; stones located in the distal ureter with a size of <10mm in largest diameter
Exclusion criteria	urinary tract infection; solitary kidney; severe hydronureteroephrosis; renal insufficiency; diabetes; multiple stones; bilateral stones; hypotension; pregnancy; previous spontaneous stone expulsion; previous distal ureteral surgery; history of intake of nifedipine, alpha-adrenergic blockers, calcium antagonists or steroids
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 42.4 (16.1), control group: 43.5 (16.6). Gender (M:F): 58/32. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=31) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg once daily. Duration up to 10 days. Concurrent medication/care: Diclofenac as needed and advice to drink at least 2 L of water daily. Indirectness: No indirectness
	(n=29) Intervention 2: Pain management only - NSAIDs. Diclofenac 100mg once daily. Duration up to 10 days. Concurrent medication/care: advice to drink at least 2 L of water daily. Indirectness: No indirectness
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS

Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate at 10 days ; Group 1: 25/31, Group 2: 11/29; Comments: numbers calculated from percentages

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size, stone location or stone site; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: significant adverse events at 10 days; Group 1: 0/31, Group 2: 0/29
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: Serious indirectness; Baseline details: no significant difference in age, gender, stone size, stone location or stone site; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: incidence of ureteral colic at 10 days; Group 1: 20/31, Group 2: 23/29
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size, stone location or stone
site; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 4: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: additional analgesic requirement at 10 days; Group 1: 10/31, Group 2: 18/29
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size, stone location or stone
site; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Time to stone passage; Hospitalisation/ Use of healthcare services

National Inetitute for Health and Pare Eventence 2017

Radiology Centre or emergency department at a single centre
al stone of distal ureteral or UVj)
(UB; acute hydronephrosis (grades 2 and 3) in sonography; olic blood pressure <100; taking calcium antagonist reter; single renal patients; creatinine >1.4 for males and >1.2 atment; NSAID intolerance or adverse effects of Tamsulosin; he study; pregnancy
nnicity: not reported
cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to- omen: Non-pregnant 5. Stone composition/hounsfield units: Not ones
llosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. 0mg as needed and advice to drink 2 L of water daily.
- NSAIDs. Indomethacin 100mg as needed. Duration up to 4 drink 2 L of water daily. Indirectness: No indirectness

#### Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous expulsion at 4 weeks; Group 1: 41/50, Group 2: 30/46 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone size; Group 1 Number missing; Group 2 Number missing:

#### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 4 weeks; Group 1: mean 3.7 days (SD 5.07); n=50, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone size; Group 1 Number missing; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: symptoms such as UTI, fever, severe obstructive uropathy, worsening of symptoms and side effects of Tamsulosin or Indomethacin that require discontinuation at 4 weeks; Group 1: 0/50, Group 2: 0/46 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: Serious indirectness; Baseline details: no significant difference in age or stone size; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 4: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; average analgesic consumption at 4 weeks; Group 1; mean 1.48 number of times (SD 2.15); n=50,

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone size; Group 1 Number missing; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Pain intensity; Hospitalisation/ Use of healthcare services

Study	Arrabal-martin 2010 <sup>20</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=70)
Countries and setting	Conducted in Spain; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention + follow up: 30 days
Method of assessment of guideline condition	Adequate method of assessment/diagnosis
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	ureteral lithiasis below the S3 and S4 levels and a calculus size of 4-10mm
Exclusion criteria	urinary infection; abdominal alterations; multiple lithiases; urinary derivation (double-J catheter in the ureter or percutaneous nephrostomy); other factors hindering the removal of calculi
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age: . Gender (M:F): not reported . Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=35) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration 3 weeks. Concurrent medication/care: Ibuprofen 600mg every 12 hours, 2 L of water daily and Tramadol in case of pain. Indirectness: No indirectness  (n=35) Intervention 2: Pain management only - NSAIDs. Ibuprofen 600mg every 12 hours. Duration 3 weeks. Concurrent medication/care: 2 L of water daily and Tramadol in case of pain. Indirectness: No indirectness
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS

Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion at 30 days; Group 1: 30/35, Group 2: 19/35
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,

Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in sex, age or lithiasis size; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: fever >37.5•c or side effects concerning Tamsulosin at 30 days ; Group 1: 0/35, Group 2: 0/35

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - High, Measurement - High, Crossover - Low; Indirectness of outcome: Serious indirectness; Baseline details: no significant difference in sex, age or lithiasis size; Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 3: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: use of Tramadol at 30 days ; Group 1: 9/35, Group 2: 21/35; Comments: numbers calculated from percentages

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in sex, age or lithiasis size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Time to stone passage; Pain intensity; Hospitalisation/ Use of healthcare services
study	

Study	Ates 2012 <sup>21</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=79)
Countries and setting	Conducted in Turkey; Setting: 4 urology departments at 3 centres
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 2 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: history, physical evaluation, urinary analysis, laboratory findings, ultrasonography
Stratum	Adults (≥16 years), ureteric stone <1 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	radio-opaque upper ureteral stones

Exclusion criteria	abnormal renal anatomy and function; use of medications that may lead to stone formation; pregnancy or suspicion of pregnancy; distal obstruction; history of previous urinary stone surgery; hydronephrosis >grade 1; presence of coagulopathy; active urinary tract infection; history of hypersensitivity to Doxazosin; serum creatinine level >2mg/dL; existence of >1 ureteral stone; hypotension; pain that could not be controlled with an analgesic
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Doxazosin group: 38.35 (11.41), control group: 30.95 (9.68). Gender (M:F): 58/21. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Upper ureteric stones
Indirectness of population	Serious indirectness: includes stones < and > 10mm
Interventions	(n=35) Intervention 1: Alpha blockers and SWL. Doxazosin controlled release 4mg daily within 24 hours before SWL, if stone was not influenced or fragmented into pieces ≥6mm a second session was performed 3 days after the first procedure. Duration up to 14 days. Concurrent medication/care: oral Diclofenac on demand and advice to drink at least 2L of fluid daily. Indirectness: No indirectness; Indirectness comment: NA
	(n=44) Intervention 2: Surgery - SWL. SWL, if stone was not influenced or fragmented into pieces ≥6mm a second session was performed 3 days after the first procedure. Duration procedure time. Concurrent medication/care: oral Diclofenac on demand and advice to drink at least 2 L of fluid daily. Indirectness: No indirectness; Indirectness comment: NA
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL

Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of hospital admissions at 14 days; Group 1: mean 0.51 admissions (SD 0.7); n=35, Group 2: mean 0.52 admissions (SD 0.62); n=44; VAS 0-10 Top=High is poor outcome

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: Doxazosin group were older; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone free rate at 14 days; Group 1: 33/35, Group 2: 35/44

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: Doxazosin group were older; Group 1 Number missing; Group 2 Number missing:

#### Protocol outcome 3: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to stone passage at 14 days; Group 1: mean 4.14 days (SD 1.78); n=35, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: Doxazosin group were older; Group 1 Number missing; Group 2 Number missing:

#### Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: VAS at unclear; Group 1: mean 6.89 (SD 1.02); n=35, Group 2: mean 6.59 (SD 1.58); n=44: VAS 0-10 Top=High is poor outcome

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: Doxazosin group were older; Group 1 Number missing; Group 2 Number missing:

# Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: need for analgesics at 14 days; Group 1: 29/35, Group 2: 30/44 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: Doxazosin group were older; Group 1 Number missing; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Adverse events

Study (subsidiary papers)	Autorino 2005 <sup>22</sup> (De sio 2006 <sup>48</sup> )
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=64)
Countries and setting	Conducted in Italy; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention + follow up: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: unenhanced CT scan
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	unilateral distal ureteral calculi
Exclusion criteria	urinary tract infection; severe hydronephrosis; diabetes; ulcer; hypotension or hypertension when in treatment with alpha-blockers or calcium-antagonists; pregnancy; multiple stones; history of spontaneous stone expulsion or ureteral stricture
Recruitment/selection of patients	consecutive patients meeting the inclusion/exclusion criteria during the study period
Age, gender and ethnicity	Age - Other: Tamsulosin group mean: 45, control group mean: 43. Gender (M:F): 62/34. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=50) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 2 weeks. Concurrent medication/care: Diclofenac 100mg daily, Aescin 80mg daily, advice to drink 2 L of water daily, Omeprazole 20mg daily for the treatment period and Levofloxacin 250mg daily for the first week. Indirectness: No indirectness
	(n=46) Intervention 2: Pain management only - NSAIDs. Diclofenac 100mg daily and Aescin 80mg daily. Duration up to 2 weeks. Concurrent medication/care: advice to drink 2 L of water daily, Omeprazole 20mg daily for the treatment period and Levofloxacin 250mg daily for the first week . Indirectness: No indirectness
Funding	Funding not stated
RESULTS (NUMBERS ANALYSED) AND R	ISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hospitalisation for recurrent colic at 2 weeks; Group 1: 5/50, Group 2: 11/46
Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender or stone size; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate at 2 weeks; Group 1: 45/50, Group 2: 27/46
Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender or stone size; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 3: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 2 weeks; Group 1: mean 4.4 days (SD 2.1); n=50, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender or stone size; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 4: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 2 weeks; Group 1: 1/50, Group 2: 0/46
  Risk of bias: All domain Very high, Selection Low, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High,
  Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender or stone size; Group 1 Number
  missing: Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension at 2 weeks; Group 1: 2/50, Group 2: 0/46
  Risk of bias: All domain Very high, Selection Low, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High,
  Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender or stone size; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of patients requiring different analgesics from those used in the standard treatment regimen at 2 weeks; Group 1: 5/50, Group 2: 17/46

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Pain intensity

Study	Aydogdu 2009 <sup>24</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=39)
Countries and setting	Conducted in Turkey; Setting: paediatric urology unit, single centre
Line of therapy	1st line
Duration of study	Intervention time: up to 3 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis
Stratum	Children (<16 years)
Subgroup analysis within study	Not applicable
Inclusion criteria	radiopaque lower ureteral stones 2-10mm in diameter
Exclusion criteria	anatomical abnormalities; previously diagnosed reflux; voiding dysfunction; history of ureteral surgery or steinstrasse formed after ESWL; receiving calcium channel blockers
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): 5.6 (2.6). Gender (M:F): 21/18. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=19) Intervention 1: Alpha blockers - Doxazosin. Doxazosin 0.03mg/kg once daily administered at bedtime. Duration up to 3 weeks. Concurrent medication/care: Ibuprofen 20mg/kg daily divided in 2 equal doses for pain episodes. Indirectness: No indirectness  (n=20) Intervention 2: Pain management only - NSAIDs. Ibuprofen 20mg/kg daily divided in 2 equal doses
Funding	for pain episodes. Duration up to 3 weeks. Concurrent medication/care: none. Indirectness: No indirectness  Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: DOXAZOSIN versus NSAIDS

Protocol outcome 1: Stone passage

- Actual outcome for Children (<16 years): stone expulsion at 3 weeks; Group 1: 16/19, Group 2: 14/20
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,

CONSULTATION

Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender or stone size; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Time to stone passage

- Actual outcome for Children (<16 years): time to expulsion at 3 weeks; Group 1: mean 5.9 days (SD 2.1); n=19,

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender or stone size; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Children (<16 years): adverse events including hypotension, asthenia, syncope and palpitations at 3 weeks; Group 1: 0/19, Group 2: 0/20

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: Serious indirectness; Baseline details: no significant difference in age, gender or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare services
study	

Study	Bajwa 2013 <sup>28</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=60)
Countries and setting	Conducted in Pakistan; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	lower ureteric stone <1cm; sterile urine; symptom free
Exclusion criteria	obstruction; stone size >1cm; urinary tract infection
Recruitment/selection of patients	not reported

Age, gender and ethnicity	Age - Mean (SD): 33.15 (8.97). Gender (M:F): 37/23. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	<ul> <li>(n=30) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg once daily. Duration up to 4 weeks. Concurrent medication/care: not reported. Indirectness: No indirectness</li> <li>(n=30) Intervention 2: Pain management only - NSAIDs. Diclofenac 50mg 12 hourly. Duration up to 4 weeks. Concurrent medication/care: not reported. Indirectness: No indirectness</li> </ul>
	Concurrent medication/care. not reported. indirectness. No indirectness
Funding	Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS

# Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone discharged at 4 weeks; Group 1: 23/30, Group 2: 11/30 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: comparable for age, gender and stone size; Group 1 Number missing:; Group 2 Number missing:

# Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 4 weeks; Group 1: mean 15.7 days (SD 3.72); n=30, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: comparable for age, gender and stone size; Group 1 Number missing:; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Adverse events; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare services
study	

Study	Balci 2014 <sup>29</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=75)
Countries and setting	Conducted in Turkey; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: up to 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	stones of 5-10mm diameter in the lower third of the ureter (below the common iliac vessels)
Exclusion criteria	proximal or intramural part of the ureteral stone; active urinary tract infection; ureterohydronephrosis; acute renal failure; fever; multiple ureteral stones; history of surgery or endoscopic procedures for urolithiasis; chronic renal failure; diabetes; peptic ulcer; concomitant treatment with alpha-blocker and beta-blocker, calcium antagonists or nitrates; pregnancy; lactation; patient desire for immediate stone removal
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): 36.8 (11.3). Gender (M:F): 53/22. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=25) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 50mg when required and advice to drink 2-2.5 L of water daily. Indirectness: No indirectness
	(n=25) Intervention 2: Calcium channel blockers - Nifedipine. Nifedipine 30mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 50mg when required and advice to drink 2-2.5 L of water daily. Indirectness: No indirectness
	(n=25) Intervention 3: Pain management only - NSAIDs. Diclofenac 50mg when required. Duration up to 4 weeks. Concurrent medication/care: advice to drink 2-2.5 L of water daily. Indirectness: No indirectness
Funding	Funding not stated

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 4 weeks; Group 1: 19/25, Group 2: 16/25
Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size and Hounsfield Units;
Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 4 weeks; Group 1: 2/25, Group 2: 0/25
Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size and Hounsfield Units;
Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 3: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: analgesic use at 4 weeks; Group 1: mean 544 mg (SD 493); n=25, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size and Hounsfield Units; Group 1 Number missing: ; Group 2 Number missing:

## RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS

# Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 4 weeks; Group 1: 19/25, Group 2: 9/25
Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size and Hounsfield Units;
Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 4 weeks; Group 1: 2/25, Group 2: 0/25
Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size and Hounsfield Units;
Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 3: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: analgesic use at 4 weeks; Group 1: mean 544 mg (SD 493); n=25, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,

Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size and Hounsfield Units; Group 1 Number missing: ; Group 2 Number missing:

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NIFEDIPINE versus NSAIDS

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 4 weeks; Group 1: 16/25, Group 2: 9/25 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size and Hounsfield Units; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 4 weeks; Group 1: 0/25, Group 2: 0/25 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size and Hounsfield Units; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 3: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: analgesic use at 4 weeks; Group 1: mean 602 mg (SD 434); n=25, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, gender, stone size and Hounsfield Units; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Time to stone passage; Pain intensity; Hospitalisation/ Use of healthcare services

Study	Basri 2013 <sup>30</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=123)
Countries and setting	Conducted in Turkey; Setting: single centre
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: plain x-ray of the KUB and/or ultrasound imaging
Stratum	Adults (≥16 years), ureteric stone 1-2 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	solitary ureteral stone 6-15mm located in the upper, middle or lower ureter
Exclusion criteria	<18 years of age; weight <50kg or >100kg; severe skeletal malformation; pregnancy; aortic and/or renal artery aneurysm; history of drug or alcohol abuse; long-term use of drugs such as antidepressants, histamine blockers or anxiolytics; allergy to the study medications; concomitant treatment with calcium antagonists and/or an alpha adrenergic antagonist; concomitant renal stones; previous unsuccessful attempts at SWL; elevated serum creatinine >2 mg/dL; urinary tract infection; diabetes; peptic ulcer; history of spontaneous stone expulsion; hypotension; coagulopathy; urinary congenital abnormalities; previous nephroureteral surgery
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 44.66 (13.25), control group: 42.19 (13.17). Gender (M:F): 98/25. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Not stated / Unclear
Indirectness of population	Serious indirectness: included stones < and >10mm
Interventions	(n=59) Intervention 1: Alpha blockers and SWL. Tamsulosin 0.4mg daily immediately after SWL. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 75mg injected intramuscularly on demand, gastro protective therapy 40mg Pantoprazole once daily and instruction to drink a minimum of 2L of water daily. Indirectness: No indirectness; Indirectness comment: NA
	(n=64) Intervention 2: Surgery - SWL. SWL. Duration unclear. Concurrent medication/care: Diclofenac 75mg injected intramuscularly on demand, gastro protective therapy 40mg Pantoprazole daily and instruction to a

	minimum of 2L of water daily. Indirectness: No indirectness; Indirectness comment: NA
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL

Protocol outcome 1: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: time to stone clearance (upper stones) at unclear; Group 1: mean 7.1 days (SD 6.4); n=29, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: time to stone clearance (middle stones) at unclear; Group 1: mean 9.25 days (SD 9.95); n=16.

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: time to stone clearance (lower stones) at unclear; Group 1: mean 9.86 days (SD 6.94); n=14.

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcome 2: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: VAS (upper stones) at unclear; Group 1: mean 2.9 (SD 2.19); n=29, Group 2: mean 4 (SD 2.58); n=28; VAS 0-10 Top=High is poor outcome

Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: VAS (middle stones) at unclear; Group 1: mean 2.38 (SD 2.42); n=16, Group 2: mean 3 (SD 3.91); n=12; VAS 0-10 Top=High is poor outcome
- Risk of bias: All domain High, Selection Low, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover - Low: Indirectness of outcome: No indirectness, Comments; NA; Baseline details; no significant differences in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: VAS (lower stones) at unclear; Group 1: mean 2.79 (SD 2.42); n=14, Group 2: mean 4 (SD 2.71); n=24; VAS 0-10 Top=High is poor outcome

Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex or stone size; Group 1

Protocol outcomes not reported by the study

Quality of life; Stone passage; Adverse events; Analgesic use; Hospitalisation/ Use of healthcare services

Study	Bayraktar 2017 <sup>31</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=124)
Countries and setting	Conducted in Turkey; Setting: Not reported
Line of therapy	1st line
Duration of study	Follow up (post intervention): 4 weeks
Method of assessment of guideline condition	Method of assessment /diagnosis not stated
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	Males with radiopaque distal ureter stones 5-10mm
Exclusion criteria	Not reported
Recruitment/selection of patients	Not reported
Age, gender and ethnicity	Age - Mean (SD): Alpha blocker group 34.4 (13.5); control group 36.92 (12.4). Gender (M:F): All male. Ethnicity: Not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not stated / Unclear 3. Obesity /skin-to-stone distance: Not stated / Unclear 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not stated / Unclear 6. Uteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=70) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily as a single dose. Duration Unclear. Concurrent medication/care: As a standard medical therapy, all patients were recommended a daily intake of liquids to urinate at least 1.5-2 litres, and 75mg of diclofenac was injected when needed. Indirectness: No indirectness
	(n=71) Intervention 2: Pain management only - NSAIDs. No treatment. Duration Unclear. Concurrent medication/care: As a standard medical therapy, all patients were recommended a daily intake of liquids to urinate at least 1.5-2 litres, and 75mg of diclofenac was injected when needed. Indirectness: No indirectness

# Funding Funding not stated RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS Protocol outcome 1: Stone passage at Define - Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Stone expulsion at 2 weeks; Group 1: 42/60, Group 2: 18/64 Risk of bias: All domain - High, Selection - Very high, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 11; Group 2 Number missing: 6 - Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Stone expulsion at 4 weeks; Group 1: 49/60, Group 2: 33/64 Risk of bias: All domain - High, Selection - Very high, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 11; Group 2 Number missing: 6 Protocol outcome 2: Time to stone passage at Define - Actual outcome for Adults (≥16 years), ureteric stone <1 cm; Expulsion time at 4 weeks; Group 1; mean 9.3 days (SD 5.8); n=60, Group 2; mean 8.7 days (SD 6.4); n=64 Risk of bias: All domain - High, Selection - Very high, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 11; Group 2 Number missing: 6

## Protocol outcome 3: Pain intensity at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Number of NSAID injections at 4 weeks; Group 1: mean 1.3 (SD 0.4); n=60, Group 2: mean 1.4 (SD 0.4); n=64

Risk of bias: All domain - Very high, Selection - Very high, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 11; Group 2 Number missing: 6

Protocol outcomes not reported by the study

Quality of life at Define; Adverse events at Define; Analgesic use at Define; Hospitalisation/ Use of healthcare services at Define

Study type       RCT (Patient randomised; Parallel)         Number of studies (number of participants)       1 (n=67)         Countries and setting       Conducted in China; Setting: Urology division, Surgery, single centre         Line of therapy       1st line         Duration of study       Intervention + follow up: 5 weeks         Method of assessment of guideline condition       Adequate method of assessment/diagnosis: non-contrast computerised tomography used to confirm presence of radio-opaque stone         Stratum       Adults (≥16 years), ureteric stone <1 cm         Subgroup analysis within study       Not applicable         Inclusion criteria       acute ureteric stone 5-10mm         Exclusion criteria       radiolucent stone; paper thin cortex; non-functioning kidney; intolerance to Alfuzosin; renal insufficiency (serum creatinine >160umol/L); concurrent alpha-blocker/calcium channel blocker/steroid/Furosemide usage; pregnancy; hypotension; history of ureteral stricture; history of ureteric stone treatment; allergic reaction to the study medication; patient on double-J ureteric stenting or percutaneous nephrostomy drainage; uncontrolled urosepsis         Recruitment/selection of patients       consecutive         Age, gender and ethnicity       Age - Mean (SD): 47.7 (12.3). Gender (M:F): 41/26. Ethnicity: not reported	Study	Chau 2011 <sup>41</sup>
Countries and setting  Line of therapy  Duration of study  Method of assessment of guideline condition  Stratum  Subgroup analysis within study  Inclusion criteria  Exclusion criteria  Exclusion criteria  Exclusion to the study medication; hypotension; hypotension; hypotension; hypotension; hypotension; hypotension; patient on double-J ureteric stenting or percutaneous nephrostomy drainage; uncontrolled urosepsis  Recruitment/selection of patients  Conducted in China; Setting: Urology division, Surgery, single centre  1st line  Duration of study  Intervention + follow up: 5 weeks  Adequate method of assessment/diagnosis: non-contrast computerised tomography used to confirm presence of radio-opaque stone  Adults (≥16 years), ureteric stone <1 cm  Not applicable  acute ureteric stone 5-10mm  radiolucent stone; paper thin cortex; non-functioning kidney; intolerance to Alfuzosin; renal insufficiency (serum creatinine >160umol/L); concurrent alpha-blocker/calcium channel blocker/steroid/Furosemide usage; pregnancy; hypotension; history of ureteral stricture; history of ureteric stone treatment; allergic reaction to the study medication; patient on double-J ureteric stenting or percutaneous nephrostomy drainage; uncontrolled urosepsis  Recruitment/selection of patients	Study type	RCT (Patient randomised; Parallel)
Line of therapy  Duration of study  Method of assessment of guideline condition  Stratum  Adequate method of assessment/diagnosis: non-contrast computerised tomography used to confirm presence of radio-opaque stone  Stratum  Adults (≥16 years), ureteric stone <1 cm  Subgroup analysis within study  Inclusion criteria  Exclusion criteria  Exclusion criteria  Exclusion criteria  Exclusion criteria  Recruitment/selection of patients  1st line  Intervention + follow up: 5 weeks  Adequate method of assessment/diagnosis: non-contrast computerised tomography used to confirm presence of radio-opaque stone  Adults (≥16 years), ureteric stone <1 cm  Not applicable  Inclusion criteria  acute ureteric stone 5-10mm  Facilitation: non-functioning kidney; intolerance to Alfuzosin; renal insufficiency (serum creatinine >160umol/L); concurrent alpha-blocker/calcium channel blocker/steroid/Furosemide usage; pregnancy; hypotension; history of ureteral stricture; history of ureteric stone treatment; allergic reaction to the study medication; patient on double-J ureteric stenting or percutaneous nephrostomy drainage; uncontrolled urosepsis  Recruitment/selection of patients  consecutive	Number of studies (number of participants)	1 (n=67)
Duration of study         Intervention + follow up: 5 weeks           Method of assessment of guideline condition         Adequate method of assessment/diagnosis: non-contrast computerised tomography used to confirm presence of radio-opaque stone           Stratum         Adults (≥16 years), ureteric stone <1 cm	Countries and setting	Conducted in China; Setting: Urology division, Surgery, single centre
Method of assessment of guideline condition       Adequate method of assessment/diagnosis: non-contrast computerised tomography used to confirm presence of radio-opaque stone         Stratum       Adults (≥16 years), ureteric stone <1 cm	Line of therapy	1st line
condition       presence of radio-opaque stone         Stratum       Adults (≥16 years), ureteric stone <1 cm	Duration of study	Intervention + follow up: 5 weeks
Subgroup analysis within study Inclusion criteria acute ureteric stone 5-10mm  Exclusion criteria radiolucent stone; paper thin cortex; non-functioning kidney; intolerance to Alfuzosin; renal insufficiency (serum creatinine >160umol/L); concurrent alpha-blocker/calcium channel blocker/steroid/Furosemide usage; pregnancy; hypotension; history of ureteral stricture; history of ureteric stone treatment; allergic reaction to the study medication; patient on double-J ureteric stenting or percutaneous nephrostomy drainage; uncontrolled urosepsis  Recruitment/selection of patients consecutive	_	
Inclusion criteria acute ureteric stone 5-10mm  Exclusion criteria radiolucent stone; paper thin cortex; non-functioning kidney; intolerance to Alfuzosin; renal insufficiency (serum creatinine >160umol/L); concurrent alpha-blocker/calcium channel blocker/steroid/Furosemide usage; pregnancy; hypotension; history of ureteral stricture; history of ureteric stone treatment; allergic reaction to the study medication; patient on double-J ureteric stenting or percutaneous nephrostomy drainage; uncontrolled urosepsis  Recruitment/selection of patients consecutive	Stratum	Adults (≥16 years), ureteric stone <1 cm
Exclusion criteria radiolucent stone; paper thin cortex; non-functioning kidney; intolerance to Alfuzosin; renal insufficiency (serum creatinine >160umol/L); concurrent alpha-blocker/calcium channel blocker/steroid/Furosemide usage; pregnancy; hypotension; history of ureteral stricture; history of ureteric stone treatment; allergic reaction to the study medication; patient on double-J ureteric stenting or percutaneous nephrostomy drainage; uncontrolled urosepsis  Recruitment/selection of patients consecutive	Subgroup analysis within study	Not applicable
(serum creatinine >160umol/L); concurrent alpha-blocker/calcium channel blocker/steroid/Furosemide usage; pregnancy; hypotension; history of ureteral stricture; history of ureteric stone treatment; allergic reaction to the study medication; patient on double-J ureteric stenting or percutaneous nephrostomy drainage; uncontrolled urosepsis  Recruitment/selection of patients  consecutive	Inclusion criteria	acute ureteric stone 5-10mm
· · · · · · · · · · · · · · · · · · ·	Exclusion criteria	(serum creatinine >160umol/L); concurrent alpha-blocker/calcium channel blocker/steroid/Furosemide usage; pregnancy; hypotension; history of ureteral stricture; history of ureteric stone treatment; allergic reaction to the study medication; patient on double-J ureteric stenting or percutaneous nephrostomy
Age, gender and ethnicity Age - Mean (SD): 47.7 (12.3). Gender (M:F): 41/26. Ethnicity: not reported	Recruitment/selection of patients	consecutive
	Age, gender and ethnicity	Age - Mean (SD): 47.7 (12.3). Gender (M:F): 41/26. Ethnicity: not reported
	Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Not stated / Unclear
Indirectness of population No indirectness	Indirectness of population	No indirectness
Interventions  (n=33) Intervention 1: Alpha blockers - Alfuzosin. Alfuzosin slow release 10mg daily. Duration 4 weeks.  Concurrent medication/care: Dologesic (Paracetamol + Dextropropoxyphene) four tablets daily on demand for 2 weeks and Diclofenac slow release 100mg daily on demand for 2 weeks in case of suboptimal pain control by Dologesic. Indirectness: No indirectness  (n=34) Intervention 2: Pain management only - Opioids. Dologesic (Paracetamol + Dextropropoxyphene) four tablets daily on demand for 2 weeks and Diclofenac slow release 100mg daily on demand for 2 weeks in case of suboptimal pain control by Dologesic. Duration 2 weeks. Concurrent medication/care: not reported. Indirectness: No indirectness	Interventions	Concurrent medication/care: Dologesic (Paracetamol + Dextropropoxyphene) four tablets daily on demand for 2 weeks and Diclofenac slow release 100mg daily on demand for 2 weeks in case of suboptimal pain control by Dologesic . Indirectness: No indirectness  (n=34) Intervention 2: Pain management only - Opioids. Dologesic (Paracetamol + Dextropropoxyphene) four tablets daily on demand for 2 weeks and Diclofenac slow release 100mg daily on demand for 2 weeks in case of suboptimal pain control by Dologesic . Duration 2 weeks. Concurrent medication/care: not
Funding Funding not stated	Funding	Funding not stated

Renal and ureteric stones: CONSULTATION Medical expulsive therapy

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALFUZOSIN versus OPIOIDS

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone passage (upper ureteral stones) at 5 weeks; Group 1: 8/11, Group 2: 3/14 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: difference in serum creatinine level; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone passage (lower ureteral stones) at 5 weeks; Group 1: 19/22, Group 2: 14/20 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: difference in serum creatinine level; Group 1 Number missing; Group 2 Number missing:

#### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 5 weeks; Group 1: 2/33, Group 2: 0/34 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: difference in serum creatinine level; Group 1 Number missing; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Time to stone passage; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare
study	services

Study	Cho 2013 <sup>42</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=90)
Countries and setting	Conducted in South Korea; Setting: urology department, single centre
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: up to 42 days
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: plain KUB x-ray, urinalysis, physical examination, non-contrast CT
Stratum	Adults (≥16 years), ureteric stone <1 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	radio-opaque ureter stones; 5-10mm in diameter

Exclusion criteria	radiolucent stones; paper-thin cortex; non-functional kidney; previous genitourinary tract surgery; elevated serum creatinine >1.5mg/dL; severe obesity; pregnancy; concurrent alpha-blocker/calcium channel blocker/steroid/Frusemide usage; aortic or renal artery aneurysm; contraindications to alpha AR antagonist treatment
Recruitment/selection of patients	consecutive patients meeting the inclusion criteria during the recruitment period
Age, gender and ethnicity	Age - Mean (SD): Alfuzosin group: 47.4 (12.6), control group: 47.7 (12.1). Gender (M:F): 60/24. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Upper ureteric stones
Indirectness of population	Serious indirectness: included mainly upper but some lower stones
Interventions	(n=41) Intervention 1: Alpha blockers and SWL. ESWL then Alfuzosin 10mg daily, if the ureter stone remained and was larger than 5mm at the next follow up visit then additional ESWL was performed. Duration up to 42 days. Concurrent medication/care: Loxoprofen 68.1mg as needed and recommendation to drink at least 2L hydration daily. Indirectness: No indirectness; Indirectness comment: NA  (n=43) Intervention 2: Surgery - SWL. ESWL, if the ureter stone remained and was larger than 5mm at the next follow up visit then additional ESWL was performed. Duration up to 42 days. Concurrent
	medication/care: Loxoprofen 68.1mg as needed and recommendation to drink at least 2L hydration daily. Indirectness: No indirectness; Indirectness comment: NA
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL

# Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone free rate at 42 days; Group 1: 39/41, Group 2: 40/43 Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, stone size or stone location (left/right, upper/lower); Group 1 Number missing: 4, Reason: migration/discontinuation of medication/lost to follow up; Group 2 Number missing: 2, Reason: migration/lost to follow up

# Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to stone free at 42 days; Group 1: mean 9.5 days (SD 4.8); n=41, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, stone size or stone

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location (left/right, upper/lower); Group 1 Number missing: 4, Reason: migration/discontinuation of medication/lost to follow up; Group 2 Number missing: 2, Reason: migration/lost to follow up

## Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 42 days; Group 1: 2/41, Group 2: 0/43
- Risk of bias: All domain Very high, Selection High, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, stone size or stone location (left/right, upper/lower); Group 1 Number missing: 4, Reason: migration/discontinuation of medication/lost to follow up; Group 2 Number missing:
- 2, Reason: migration/lost to follow up
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: retrograde ejaculation at 42 days; Group 1: 0/41, Group 2: 0/43 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, stone size or stone location (left/right, upper/lower); Group 1 Number missing: 4, Reason: migration/discontinuation of medication/lost to follow up; Group 2 Number missing: 2, Reason: migration/lost to follow up

## Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: VAS at unclear; Group 1: mean 5.33 (SD 1.22); n=41, Group 2: mean 6.43 (SD 1.36); n=43; VAS 0-10 Top=High is poor outcome
- Risk of bias: All domain High, Selection High, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, stone size or stone location (left/right, upper/lower); Group 1 Number missing: 4, Reason: migration/discontinuation of medication/lost to follow up; Group 2 Number missing: 2, Reason: migration/lost to follow up

# Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of patients requiring analgesics at 42 days; Group 1: 8/41, Group 2: 13/43 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant differences in age, sex, stone size or stone location (left/right, upper/lower); Group 1 Number missing: 4, Reason: migration/discontinuation of medication/lost to follow up; Group 2 Number missing: 2, Reason: migration/lost to follow up

Protocol outcomes not reported by the study

Quality of life; Hospitalisation/ Use of healthcare services

Study	El said 2015 <sup>56</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=54)
Countries and setting	Conducted in Egypt; Setting: Urology outpatient department, single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: full medical history, physical and laboratory evaluation
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	>18 years; presenting with radio-opaque stones ≤10mm and located in the distal third of the ureter
Exclusion criteria	urinary tract infections; ureteral strictures; renal impairment; solitary functioning kidney; hepatic insufficiency; severe hydronephrosis; multiple stones; peptic ulcers; diabetes; hypotension; pregnancy; lactation; sensitivity to alpha-blockers; receiving alpha-blockers, nitrates, calcium channel blockers, steroids, beta blockers, sildenafil, ketoconazole, itraconazole or ritonavir
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Alfuzosin group: 32.8 (9.5), control group 32.1 (9.2). Gender (M:F): 34/20. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=28) Intervention 1: Alpha blockers - Alfuzosin. Alfuzosin sustained release 5mg twice daily after meals. Duration up to 4 weeks. Concurrent medication/care: oral hydration with ≥2 L of water daily, Diclofenac 75mg intramuscularly on demand and education from the clinical pharmacist about potential adverse events, methods of reporting adverse events, self-reporting of pain on the visual analogue scale, importance of adherence to medications and daily water intake. Indirectness: No indirectness  (n=26) Intervention 2: Pain management only - NSAIDs. Oral hydration with ≥2 L of water daily and Diclofenac 75mg intramuscularly on demand. Duration up to 4 weeks. Concurrent medication/care: education by the clinical pharmacist on potential adverse events, methods of reporting adverse events, self-reporting of pain on the visual analogue scale, importance of adherence to medications and daily water

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	intake. Indirectness: No indirectness
Funding	No funding

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALFUZOSIN versus NSAIDS

## Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Hospitalisation at 4 weeks; Group 1: 0/28, Group 2: 3/26 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate at 4 weeks; Group 1: 15/28, Group 2: 7/26 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: adverse events at 4 weeks; Group 1: 4/28, Group 2: 0/26; Comments: adverse events: headache (2), dizziness (1), hypotension (3) - all tolerable and did not result in discontinuation Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Pain intensity; Analgesic use; Time to stone passage
study	

Study	Elgalaly 2017 <sup>57</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=40)
Countries and setting	Conducted in Egypt; Setting: Not reported
Line of therapy	1st line
Duration of study	Intervention + follow up: 4 weeks

Method of assessment of guideline condition	Adequate method of assessment/diagnosis: All patients were evaluated by complete history taking and a thorough physical examination. Laboratory investigations included urine analysis and serum creatinine. Radiological assessment with plain abdominal radiograph of the kidneys, ureters and bladder (KUB) and abdomino-pelvic ultrasonography was done
Stratum	Children (<16 years)
Subgroup analysis within study	Not applicable
Inclusion criteria	Paediatric patients who presented with single, radiopaque DUS, age <18 years, single unilateral radiopaque DUS, and largest stone diameter of ≤10 mm
Exclusion criteria	Multiple, bilateral or recurrent stones, radiolucent stone, largest stone diameter >10 mm, UTI or urosepsis, anomalies of the ureter or the kidney, previous urinary tract endoscopy or surgery, marked hydronephrosis, and abnormal renal function
Recruitment/selection of patients	Not reported
Age, gender and ethnicity	Age - Mean (SD): Alpha blocker group 8.4 (3.1); placebo group 7.7 (2.3). Gender (M:F): 27/13. Ethnicity: Not reported
Further population details	1. Kidney pole: 2. Neuropathic/ cerebral-palsy /immobility: 3. Obesity /skin-to-stone distance: 4. Pregnant women: 5. Stone composition/hounsfield units: 6. Uteric stone:
Indirectness of population	No indirectness
Interventions	(n=20) Intervention 1: Alpha blockers - Silodosin. Silodosin 4 mg given at bed-time. For those who could not swallow the capsule, the capsule contents were emptied into a small amount of water or juice. Duration Unclear. Concurrent medication/care: Ibuprofen (20 mg/kg/day) was divided into two doses for pain episodes. Children were encouraged to take plenty of fluids. Indirectness: No indirectness (n=20) Intervention 2: Placebo. Placebo. Duration Unclear. Concurrent medication/care: Ibuprofen (20 mg/kg/day) was divided into two doses for pain episodes. Children were encouraged to take plenty of fluids. Indirectness: No indirectness
Funding	No funding

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: SILODOSIN versus PLACEBO

Protocol outcome 1: Stone passage at Define

- Actual outcome for Children (<16 years): Stone free at 2 weeks; Group 1: 13/18, Group 2: 11/19
Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2; Group 2 Number missing: 1
- Actual outcome for Children (<16 years): Stone free at 4 weeks; Group 1: 16/18, Group 2: 14/19

## Protocol outcome 2: Time to stone passage at Define

- Actual outcome for Children (<16 years): Time to stone expulsion at 4 weeks; Group 1: mean 7 days (SD 4.3); n=18, Group 2: mean 10.4 days (SD 4.7); n=19

Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2; Group 2 Number missing: 1

#### Protocol outcome 3: Adverse events at Define

- Actual outcome for Children (<16 years): Headache and dizziness at 4 weeks; Group 1: 3/18, Group 2: 0/19 Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2; Group 2 Number missing: 1

## Protocol outcome 4: Pain intensity at Define

- Actual outcome for Children (<16 years): Number of pain episodes at 4 weeks; Group 1: mean 2.3 (SD 1.4); n=18, Group 2: mean 4.7 (SD 2.6); n=19 Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 2; Group 2 Number missing: 1

Protocol outcomes not reported by the study

Quality of life at Define; Analgesic use at Define; Hospitalisation/ Use of healthcare services at Define

Study	Elkoushy 2012 <sup>59</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=126)
Countries and setting	Conducted in Egypt; Setting: Department of Urology, single centre
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 3 months
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: full history, clinical examination, laboratory investigations, plain abdominal film KUB, intravenous urography
Stratum	Adults (≥16 years), ureteric stone <1 cm: mean stone size 9.7 (2.6), 8.6 (1.7)
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	single radio-opaque renal or upper ureteral stones <2cm in largest diameter
Exclusion criteria	age <18 years; multiple stones; radiolucent stones; stones >2cm in largest diameter; previous SWL failure; history of spontaneous stone expulsion; urinary tract infection; distal obstruction; congenital renal or ureteral anomalies; serum creatinine ≥2mg/dl; uncorrectable bleeding disorders; hypotension; morbid obesity; pregnancy; concomitant use of calcium channel-blockers, alpha-adrenergic antagonists or corticosteroids
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 52.8 (8.2), control group: 49.4 (11.3). Gender (M:F): 72/54. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Upper ureteric stones
Indirectness of population	Serious indirectness: includes stones < and > 10mm
Interventions	(n=63) Intervention 1: Alpha blockers and SWL. SWL repeated every 3 weeks until the patient became stone free, Tamsulosin 0.4mg daily starting immediately after SWL. Duration up to 3 months. Concurrent medication/care: Diclofenac 50mg tablets or 75mg intramuscular injection on demand. Indirectness: No indirectness; Indirectness comment: NA  (n=63) Intervention 2: Surgery and placebo - SWL and placebo. SWL repeated every 3 weeks until the patient became stone free, placebo daily starting immediately after SWL. Duration up to 3 months. Concurrent medication/care: Diclofenac 50mg tablets or 75mg intramuscular injection on demand. Indirectness: No indirectness; Indirectness comment: NA
Funding	Funding not stated
	0

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL AND PLACEBO

Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone-free rate at 3 months; Group 1: 27/28, Group 2: 14/21; Comments: numbers calculated from percentages

Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, BMI, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to clearance at 3 months; Group 1: mean 4.2 weeks (SD 1.7); n=28, Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, BMI, stone size or stone location; Group 1 Number missing:; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Adverse events; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare services

Renal and ureteric stones: Medical expulsive therapy

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## Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hospitalisation at 4 weeks; Group 1: 1/30, Group 2: 2/30
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right);
Group 1 Number missing:; Group 2 Number missing:

## Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 4 weeks; Group 1: 22/30, Group 2: 12/30
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right);
Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: withdrawal from the study due to side effects caused by the medications at 4 weeks; Group 1: 0/30, Group 2: 0/30

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Pain intensity; Analgesic use; Time to stone passage
study	

Study	Erturhan 2013 <sup>60</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=50)
Countries and setting	Conducted in Turkey; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 3 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis
Stratum	Children (<16 years)
Subgroup analysis within study	Not applicable

Inclusion criteria single radiopaque lower ureteral stone  Exclusion criteria history of ureteral and/or bladder surgery; anatomic urinary system abnormality; vesicoureteral reflux; neurogenic/non-neurogenic voiding dysfunction; bilateral or nonopaque ureteral stones; severe hydronephrosis; colic pain attacks; use of diuretic and/or calcium channel blockers  Recruitment/selection of patients not reported  Age . Mean (SD): 6.65 (3.78). Gender (M:F): 24/26. Ethnicity: not reported  Further population details 1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones  Indirectness of population No indirectness  Interventions (n=25) Intervention 1: Alpha blockers - Doxazosin. Doxazosin 0.03mg/kg daily. Duration up to 3 weeks. Concurrent medication/care: lbuprofen 20mg/kg daily divided in to 2 equal doses or a maximum 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Indirectness: No indirectness  (n=25) Intervention 2: Pain management only - NSAIDs. Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum of 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Duration up to 3 weeks. Concurrent medication/care: NA. Indirectness: No indirectness  Funding Funding not stated		
neurogenic/non-neurogenic voiding dysfunction; bilateral or nonopaque ureteral stones; severe hydronephrosis; colic pain attacks; use of diuretic and/or calcium channel blockers  Recruitment/selection of patients  Age, gender and ethnicity  Age - Mean (SD): 6.65 (3.78). Gender (M:F): 24/26. Ethnicity: not reported  1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones  Indirectness of population  No indirectness  (n=25) Intervention 1: Alpha blockers - Doxazosin. Doxazosin 0.03mg/kg daily. Duration up to 3 weeks. Concurrent medication/care: Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Indirectness: No indirectness  (n=25) Intervention 2: Pain management only - NSAIDs. Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum of 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Duration up to 3 weeks. Concurrent medication/care: NA. Indirectness: No indirectness	Inclusion criteria	single radiopaque lower ureteral stone
Age - Mean (SD): 6.65 (3.78). Gender (M:F): 24/26. Ethnicity: not reported  1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones  Indirectness of population  No indirectness  (n=25) Intervention 1: Alpha blockers - Doxazosin. Doxazosin 0.03mg/kg daily. Duration up to 3 weeks. Concurrent medication/care: Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Indirectness: No indirectness or a maximum of 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Duration up to 3 weeks. Concurrent medication/care: NA. Indirectness: No indirectness	Exclusion criteria	neurogenic/non-neurogenic voiding dysfunction; bilateral or nonopaque ureteral stones; severe
Further population details  1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones  Indirectness of population  No indirectness  (n=25) Intervention 1: Alpha blockers - Doxazosin. Doxazosin 0.03mg/kg daily. Duration up to 3 weeks. Concurrent medication/care: Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Indirectness: No indirectness  (n=25) Intervention 2: Pain management only - NSAIDs. Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum of 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Duration up to 3 weeks. Concurrent medication/care: NA. Indirectness: No indirectness	Recruitment/selection of patients	not reported
stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones  Indirectness of population  Interventions  (n=25) Intervention 1: Alpha blockers - Doxazosin. Doxazosin 0.03mg/kg daily. Duration up to 3 weeks.  Concurrent medication/care: Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Indirectness: No indirectness  (n=25) Intervention 2: Pain management only - NSAIDs. Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum of 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Duration up to 3 weeks. Concurrent medication/care: NA. Indirectness: No indirectness	Age, gender and ethnicity	Age - Mean (SD): 6.65 (3.78). Gender (M:F): 24/26. Ethnicity: not reported
Interventions  (n=25) Intervention 1: Alpha blockers - Doxazosin. Doxazosin 0.03mg/kg daily. Duration up to 3 weeks. Concurrent medication/care: Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Indirectness: No indirectness  (n=25) Intervention 2: Pain management only - NSAIDs. Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum of 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Duration up to 3 weeks. Concurrent medication/care: NA. Indirectness: No indirectness	Further population details	stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not
Concurrent medication/care: Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Indirectness: No indirectness  (n=25) Intervention 2: Pain management only - NSAIDs. Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum of 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Duration up to 3 weeks. Concurrent medication/care: NA. Indirectness: No indirectness	Indirectness of population	No indirectness
	Interventions	Concurrent medication/care: Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Indirectness: No indirectness  (n=25) Intervention 2: Pain management only - NSAIDs. Ibuprofen 20mg/kg daily divided in to 2 equal doses or a maximum of 40mg/kg daily divided in to 4 equal doses in the case of intractable pain. Duration up to 3
Funding Funding not stated		
	Funding	Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: DOXAZOSIN versus NSAIDS

# Protocol outcome 1: Stone passage

- Actual outcome for Children (<16 years): expulsion rate at 3 weeks; Group 1: 17/24, Group 2: 6/21 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, body weight, or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Time to stone passage; Adverse events; Pain intensity; Analgesic use; Hospitalisation/ Use of
study	healthcare services

Study	Eryildirim 2016 <sup>63</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=80)
Countries and setting	Conducted in Turkey; Setting: urology clinic, single centre
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: history, uro-genital examination, biochemical evaluation, urinalysis tests, non-contrast CT
Stratum	Adults (≥16 years), ureteric stone <1 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	5-10mm single radio-opaque upper ureteral stones
Exclusion criteria	multiple stones; previous stone-related procedures; obstruction; stent placement; auxiliary procedures; congenital anomalies; active urinary tract infection; pregnancy; renal insufficiency
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): 39.41 (12.99). Gender (M:F): 36/18. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Upper ureteric stones
Indirectness of population	No indirectness: NA
Interventions	(n=40) Intervention 1: Alpha blockers and SWL. SWL and Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 75mg if needed. Indirectness: No indirectness; Indirectness comment: NA  (n=40) Intervention 2: Surgery - SWL. SWL. Duration up to 4 weeks. Concurrent medication/care: Diclofenac
Funding	75mg if needed. Indirectness: No indirectness; Indirectness comment: NA  Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL

Protocol outcome 1: Quality of life
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: EQ5D at 4 weeks ; Group 1: mean 0.82 (SD 0.11); n=28, Group 2: mean 0.78 (SD 0.09); n=26; EQ5D 0-1 Top=High is good outcome

Risk of bias: Crossover unit or degre required DJ - Actual outo (SD 8.43); no Risk of bias: Crossover -

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, BMI, stone size, hounsfeld unit or degree of hydronephrosis; Group 1 Number missing: 12, Reason: 7 required DJ stent placement; Group 2 Number missing: 14, Reason: 5 required DJ stent placement

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: EQ5D VAS at 4 weeks; Group 1: mean 80.36 (SD 11.05); n=28, Group 2: mean 73.65 (SD 8.43); n=26; EQ5D VAS 0-100 Top=High is good outcome

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, BMI, stone size, hounsfeld unit or degree of hydronephrosis; Group 1 Number missing: 12, Reason: 7 required DJ stent placement; Group 2 Number missing: 14, Reason: 5 required DJ stent placement

#### Protocol outcome 2: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of ED visits at 4 weeks; Group 1: mean 0.82 (SD 0.9); n=28, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, BMI, stone size, hounsfeld unit or degree of hydronephrosis; Group 1 Number missing: 12, Reason: 7 required DJ stent placement; Group 2 Number missing: 14, Reason: 5 required DJ stent placement

## Protocol outcome 3: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone free rate at 4 weeks; Group 1: 20/28, Group 2: 17/26
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, BMI, stone size, hounsfeld
unit or degree of hydronephrosis; Group 1 Number missing: 12, Reason: 7 required DJ stent placement; Group 2 Number missing: 14, Reason: 5
required DJ stent placement

# Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of renal colic at 4 weeks; Group 1: mean 2.54 (SD 2.55); n=28, Risk of bias: All domain Very high, Selection High, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, BMI, stone size, hounsfeld unit or degree of hydronephrosis; Group 1 Number missing: 12, Reason: 7 required DJ stent placement; Group 2 Number missing: 14, Reason: 5 required DJ stent placement
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: VAS during pain at 4 weeks ; Group 1: mean 5.86 (SD 1.41); n=28, Group 2: mean 6.65 (SD 1.57); n=26; visual analogue pain scale 0-10 Top=High is poor outcome

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, BMI, stone size, hounsfeld unit or degree of hydronephrosis; Group 1 Number missing: 12, Reason: 7 required DJ stent placement; Group 2 Number missing: 14, Reason: 5 required DJ stent placement

## Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: analgesic required at 4 weeks; Group 1: mean 242 mg (SD 196.6); n=28, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, BMI, stone size, hounsfeld unit or degree of hydronephrosis; Group 1 Number missing: 12, Reason: 7 required DJ stent placement; Group 2 Number missing: 14, Reason: 5 required DJ stent placement

Protocol outcomes not reported by the	
study	

Adverse events; Time to stone passage

Study	Ferre 2009 <sup>66</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=80)
Countries and setting	Conducted in USA; Setting: Department of Emergency Medicine, single centre
Line of therapy	1st line
Duration of study	Intervention + follow up: 14 days
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: computed tomography confirmed diagnosis
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	≥18 years of age; able to provide written informed consent; CT confirmed diagnosis of a single calculus in the distal third of the ureter (distal to the internal iliac vessels) inconsistent with phleboliths as determined by a board-certified radiologist
Exclusion criteria	allergy/sensitivity to the study drug; sulfa/sulfonamide allergy; lithiasis of the ureteral intramural tract; acute or chronic renal failure; fever; presence of multiple ureteral stones; peptic ulcer disease; liver failure; pregnancy; breastfeeding; history of urinary surgery; history of endoscopic treatment; concomitant treatment with alphalytic dugs, calcium channel antagonists, nitrates or vardenafil hydrochloride; inability to use the study pain scale; inability to read, write and speak the English language
Recruitment/selection of patients	convenience sampling
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin 47 (14), standard therapy 45 (12). Gender (M:F): 56/21. Ethnicity: white race Tamsulosin group 92.1%, standard therapy group 97.4%

Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=39) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration 10 days. Concurrent medication/care: Ibuprofen 800mg 3 times a day and Oxycodone 5010mg every 4-6 hours as needed for pain. Indirectness: No indirectness  (n=41) Intervention 2: Pain management only - Opioids. Ibuprofen 800mg 3 times a day and Oxycodone 5-10mg every 4-6 hours as needed for pain. Duration up to 14 days. Concurrent medication/care: NA. Indirectness: No indirectness
Funding	Academic or government funding (academic grant from the Maine Medical Center Mentored Research Committee )

## RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus OPIOIDS

Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: return to emergency department or unscheduled visit with primary care provider at 14 days ; Group 1: 6/38, Group 2: 8/39

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: more males in study group, no significant difference in race, age, BMI, stone size or emergency department length of stay; Group 1 Number missing:; Group 2 Number missing:

# Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous passage at 14 days; Group 1: 27/38, Group 2: 24/39 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: more males in study group, no significant difference in race, age, BMI, stone size or emergency department length of stay; Group 1 Number missing:; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: adverse medication effects (nausea, vomiting, dizziness, hypotension, ejaculatory abnormalities, diarrhea, headache, arthralgia, rash) at 14 days; Group 1: 0/38, Group 2: 0/39

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: more males in study group, no significant difference in race, age, BMI, stone size or emergency department length of stay; Group 1 Number missing; Group 2 Number missing;

#### Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: colicky pain episodes at 14 days; MD; -0.05 (95%CI -4.81 to 4.7); Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: more males in study group, no significant difference in race, age, BMI, stone size or emergency department length of stay; Group 1 Number missing:; Group 2 Number missing:

# Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: opioid used (days) at 14 days; MD; -4.94 (95%Cl -12.04 to 2.15); Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: more males in study group, no significant difference in race, age, BMI, stone size or emergency department length of stay; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study	Quality of life; Time to stone passage
Study	Furyk 2016 <sup>67</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=393)
Countries and setting	Conducted in Australia; Setting: 5 emergency departments
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: computed tomography of KUB
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	>18 years; symptoms suggestive of ureteric colic; calculus demonstrated in the distal ureter (distal to the sacroiliac joint)
Exclusion criteria	temperature >38 degrees; estimated glomerula filtration rate of <60mL/minute per 1.73m²; calculus >10mm; solitary kidney; transplanted kidney; history of ureteral stricture; known allergy to the study medication; current calcium channel blocker or alpha-blocker use; hypotension; pregnant or planning pregnancy
Recruitment/selection of patients	opportunity sampling by medical staff and screening of ED databases for any patient meeting inclusion/exclusion criteria during the recruitment period
Age, gender and ethnicity	Age - Median (IQR): Tamsulosin group: 45.5 (35-55), placebo group: 46 (37-55). Gender (M:F): 320/73. Ethnicity: not reported

Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	<ul> <li>(n=198) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: analgesia at the discretion of the treating physician - recommended regimens were Indomethacin 25-50mg 3 times daily and Oxycodone 5-10mg 3 times daily as required for breakthrough. Indirectness: No indirectness</li> <li>(n=195) Intervention 2: Placebo. Placebo. Duration up to 4 weeks. Concurrent medication/care: analgesia at the discretion of the treating physician - recommended regimens were Indomethacin 25-50mg 3 times daily and Oxycodone 5-10mg 3 times daily as required for breakthrough. Indirectness: No indirectness</li> </ul>
Funding	Academic or government funding (grant from the Queensland Emergency Medicine Research Foundation )

## RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus PLACEBO

Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: re-presentation to ED at 4 weeks; Group 1: 31/198, Group 2: 35/195 Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location or urine culture result;
- Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: admission to hospital at 4 weeks; Group 1: 20/198, Group 2: 23/195
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover
- Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location or urine culture result; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone passage at 4 weeks; Group 1: 140/161, Group 2: 127/155 Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location or urine culture result; Group 1 Number missing: 37; Group 2 Number missing: 40

# Protocol outcome 3: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: pain score >0 at 1 week; Group 1: 142/185, Group 2: 143/182; Comments: verbal numeric pain scale

Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: pain score >0 at 2 weeks; Group 1: 60/176, Group 2: 58/177

Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location or urine culture result; Group 1 Number missing: 22; Group 2 Number missing: 18

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: pain score >0 at 3 weeks; Group 1: 34/170, Group 2: 37/173

Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover

- Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location or urine culture result; Group 1 Number missing: 28; Group 2 Number missing: 22

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: pain score >0 at 4 weeks; Group 1: 26/173, Group 2: 28/174

Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover

- Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location or urine culture result; Group 1 Number missing: 25; Group 2 Number missing: 21

Protocol outcomes not reported by the study

Quality of life; Adverse events; Analgesic use; Time to stone passage

Protocol outcome 1: Stone passage

Study	Gandhi 2013 <sup>68</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=128)
Countries and setting	Conducted in Nepal; Setting: Department of General Surgery, single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: abdominal ultrasonography, IVU or CT when necessary
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	solitary stone in the distal ureter at the juxtavesical tract or vesico-ureteric junction of 5-15mm
Exclusion criteria	urinary tract infection; gross hydronephrosis; diabetes; peptic ulcer disease; hypersensitivity to Nifedipine or corticosteroid; history of spontaneous stone expulsion and hypotension; pregnant women; children
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Nifedipine group: 30.4 (11.36), Tamsulosin group: 34 (12.83). Gender (M:F): Nifedipine group: 1.48:1, Tamsulosin group 1.28:1. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	Serious indirectness: included stones < and > 10mm
Interventions	(n=64) Intervention 1: Calcium channel blockers - Nifedipine. Nifedipine 30mg slow-release daily. Duration up to 4 weeks. Concurrent medication/care: oral prednisolone 30mg daily for a maximum of 10 days, Diclofenac 75mg intramuscularly on demand and ≥2 L of water daily . Indirectness: No indirectness (n=64) Intervention 2: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg. Duration up to 4 weeks. Concurrent medication/care: oral prednisolone 30mg daily for a maximum of 10 days, Diclofenac 75mg intramuscularly
Funding	on demand and ≥2 L of water daily . Indirectness: No indirectness  No funding

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NIFEDIPINE

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: expulsion at 4 weeks; Group 1: 51/64, Group 2: 32/58 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness: Baseline details: no significant difference in age, sex, duration of pain, stone size or stone location (left/right); Group 1 Number missing: 0; Group 2 Number missing: 6

#### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm; headache at 4 weeks; Group 1: 32/64, Group 2: 25/58 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details; no significant difference in age, sex, duration of pain, stone size or stone location (left/right); Group 1 Number missing: 0; Group 2 Number missing: 6
- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: dizziness at 4 weeks; Group 1: 16/64, Group 2: 3/58 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low: Indirectness of outcome: No indirectness: Baseline details: no significant difference in age, sex, duration of pain, stone size or stone location (left/right); Group 1 Number missing: 0; Group 2 Number missing: 6
- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm; flushing at 4 weeks; Group 1: 0/64, Group 2: 3/58 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, duration of pain, stone size or stone location (left/right); Group 1 Number missing: 0; Group 2 Number missing: 6

# Protocol outcome 3: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: analgesic use at 4 weeks; Group 1: mean 0.42 (SD 0.14); n=64, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, duration of pain, stone size or stone location (left/right); Group 1 Number missing: 0; Group 2 Number missing: 6

Protocol outcomes not reported by the study

Quality of life; Time to stone passage; Pain intensity; Hospitalisation/ Use of healthcare services

Mational Institute for Health and Pare Evoellence 2017

Study	Gravas 2007 <sup>74</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=61)
Countries and setting	Conducted in Greece; Setting: Department of Urology, single centre
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: plain KUB x-ray
Stratum	Adults (≥16 years), ureteric stone <1 cm: stone size range 6-13mm
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	single radiopaque distal ureteral stone (below the sacral-iliac joint); ≥6mm in diameter undergoing ESWL for the first time
Exclusion criteria	hypotension; ulcer; therapy of benign prostatic obstruction with alpha-blockers; presence of a double J stent previously placed
Recruitment/selection of patients	consecutive patients meeting the inclusion criteria during the recruitment period
Age, gender and ethnicity	Age - Mean (range): Tamsulosin group: 48.8 (27-73), control group: 49.2 (30-72). Gender (M:F): 38/23. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	Serious indirectness: includes stones < and > 10mm
Interventions	(n=30) Intervention 1: Alpha blockers and SWL. ESWL then Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: hydration of at least 2 L daily and Diclofenac 50mg on demand. Indirectness: No indirectness; Indirectness comment: NA
	(n=31) Intervention 2: Surgery - SWL. ESWL. Duration up to 4 weeks. Concurrent medication/care: hydration of at least 2 L daily and Diclofenac 50mg on demand. Indirectness: No indirectness; Indirectness comment: NA
Funding	Funding not stated
RESULTS (NUMBERS ANALYSED) AND R	ISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone free rate at 4 weeks; Group 1: 19/30, Group 2: 16/31
Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone size or number of shock waves; Group 1 Number missing: 1; Group 2 Number missing: 2

#### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 4 weeks; Group 1: 2/30, Group 2: 0/31
Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone size or number of shock waves; Group 1 Number missing: 1; Group 2 Number missing: 2

Protocol outcomes not reported by the study

Quality of life; Time to stone passage; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare services

Study	Hermanns 2009 <sup>83</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=100)
Countries and setting	Conducted in Switzerland; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 3 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: non-contrast-enhanced abdominal computed tomography
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	≥18 years; acute renal colic; single ureteral stone ≤7mm below the common iliac vessels as assessed by CT
Exclusion criteria	multiple ureteral stones; renal insufficiency; urinary tract infection; solitary kidney; pregnancy; history of ureteral surgery or previous endoscopic procedures; hypersensitivity to Tamsulosin; current alpha-blocker, calcium antagonist or corticosteroid medication
Recruitment/selection of patients	consecutive
Age, gender and ethnicity	Age - Median (IQR): Tamsulosin group 36 (30-44), placebo group 41 (33-54). Gender (M:F): 75/15. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=50) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 3 weeks. Concurrent medication/care: after initial analgesia for acute pain management, no regular analgesic medication was maintained. Oral Diclofenac (up to 3 X 50mg) as first line and oral Metamizole (up to 4 X 1g) as second line on-demand analgesics were prescribed. Indirectness: No indirectness (n=50) Intervention 2: Placebo. Placebo. Duration up to 3 weeks. Concurrent medication/care: after initial analgesia for acute pain management, no regular analgesic medication was maintained. Oral Diclofenac (up
	to 3 X 50mg) as first-line and oral Metamizole (up to 4 X 1g) as second-line on demand analgesics were prescribed. Indirectness: No indirectness
Funding	No funding

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus PLACEBO

Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hospital readmission with consecutive intervention and discontinuation of medication due to uncontrollable pain or side effects at 3 weeks; Group 1: 6/45, Group 2: 2/45

Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone expulsion rate at 3 weeks; Group 1: 39/45, Group 2: 40/45 Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low: Indirectness of outcome: No indirectness: Baseline details: no significant difference in age, sex, stone size or stone location: Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 3: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to stone passage at 3 weeks; HR; 0.99 (95%CI 0.55 to 1.79) (p value: 0.97), Comments: multiple cox regression analysis for predictive factors - therapy;

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 4: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: retrograde ejaculation at 3 weeks; Group 1: 2/39, Group 2: 0/36 Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 3 weeks; Group 1: 0/45, Group 2: 1/45 Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Analgesic use; Pain intensity

Study	Ibrahim 2013 <sup>88</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=150)
Countries and setting	Conducted in Iraq; Setting: Not reported
Line of therapy	1st line
Duration of study	Intervention + follow up: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Urinary ultrasonography (US) and a plain abdominal X-ray. IVU or CT was used in a few patients depending on specific indications
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not stratified but pre-specified: Mid, upper and proximal stone location
Inclusion criteria	Symptomatic ureteric stone of <10 mm in diameter
Exclusion criteria	Acute infection, a solitary kidney, elevated levels in renal functional tests at presentation, severe hydronephrosis, bilateral ureteric stones, pregnancy or lactation, current use of a-blockers, calcium-channel blockers or steroids, age <18 years, and any allergic reaction to the study medication
Recruitment/selection of patients	Not reported
Age, gender and ethnicity	Age - Mean (SD): Alpha blocker group 37.34 (13.15); control 36.71 (11.64). Gender (M:F): 91/21. Ethnicity: Not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not stated / Unclear 3. Obesity /skin-to-stone distance: Not stated / Unclear 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not stated / Unclear 6. Uteric stone: Upper ureteric stones
Indirectness of population	No indirectness
Interventions	(n=22) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin capsule of 0.4 mg daily (n=30) or alfuzosin 10 mg daily (n=40). Duration 4 weeks. Concurrent medication/care: All the patients were given diclofenac potassium orally 50 mg and/or diclofenac sodium as an intramuscular injection of 75 mg on demand. Indirectness: No indirectness
	(n=4) Intervention 2: Pain management only - NSAIDs. No alpha blockers. Duration 4 weeks. Concurrent medication/care: All the patients were given diclofenac potassium orally 50 mg and/or diclofenac sodium as an intramuscular injection of 75 mg on demand. Indirectness: No indirectness
	(n=6) Intervention 3: Alpha blockers - Tamsulosin. As above. Duration 4 weeks. Concurrent medication/care: As above. Indirectness: No indirectness

Medical expulsive therapy

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(n= me (n= me	=5) Intervention 4: Pain management only - NSAIDs. As above. Duration 4 weeks. Concurrent edication/care: As above. Indirectness: No indirectness  =23) Intervention 5: Pain management only - NSAIDs. As above. Duration 4 weeks. Concurrent edication/care: As above. Indirectness: No indirectness  =52) Intervention 6: Alpha blockers - Tamsulosin. As above. Duration 4 weeks. Concurrent edication/care: As above. Indirectness: No indirectness
Funding	o funding

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN/ALFUZOSIN (UPPER) versus NSAIDS (UPPER)

Protocol outcome 1: Stone passage at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Stones passed at 4 weeks; Group 1: 13/22, Group 2: 1/4
Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN/ALFUZOSIN (MID) versus NSAIDS (MID)

Protocol outcome 1: Stone passage at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Stones passed at 4 weeks; Group 1: 5/6, Group 2: 1/5
Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing:

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN/ALDUZOSIN (LOWER) versus NSAIDS (LOWER)

Protocol outcome 1: Stone passage at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Stones passed at 4 weeks; Group 1: 46/52, Group 2: 12/23
Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life at Define; Time to stone passage at Define; Adverse events at Define; Pain intensity at Define; Analgesic use at Define; Hospitalisation/ Use of healthcare services at Define

Study	Islam 2012 <sup>89</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=98)
Countries and setting	Conducted in Bangladesh; Setting: not reported
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: abdominal ultrasonography, x-ray of the kidneys ureters and bladder and excretory urography
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	distal ureteral stones (juxtavesical tract and ureterovesical junction); ≤1cm in size
Exclusion criteria	urinary tract infection; severe hydronephrosis; solitary kidney; extra stone in the upper urinary system; previous surgery for a urinary system stone; nonopaque stone; disease such as diabetes or hypertension; pregnant; renal reserve reduced by >50%
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Other: Tamsulosin group mean: 46.6, Nifedipine group mean: 47.4, control group mean: 42.8. Gender (M:F): 58/33. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=33) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: prophylactic antibiotic therapy (Ciprofloxacin 500mg twice daily), 2.5 L hydration daily and Diclofenac recommended for routine use during pain episodes. Indirectness: No indirectness
	(n=33) Intervention 2: Calcium channel blockers - Nifedipine. Nifedipine 20mg (slow release) daily. Duration up to 4 weeks. Concurrent medication/care: prophylactic antibiotic therapy (Ciprofloxacin 500mg twice daily), 2.5 L hydration daily and Diclofenac recommended for routine use during pain episodes. Indirectness: No indirectness
	(n=32) Intervention 3: No treatment. No treatment. Duration up to 4 weeks. Concurrent medication/care: prophylactic antibiotic therapy (Ciprofloxacin 500mg twice daily), 2.5 L hydration daily and Diclofenac

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	recommended for routine use during pain episodes. Indirectness: No indirectness
Funding	Funding not stated

### RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NIFEDIPINE

## Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hospitalisation for recurrent colic at 4 weeks; Group 1: 0/32, Group 2: 0/31 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion at 4 weeks; Group 1: 27/32, Group 2: 22/31
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing:
; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension at 4 weeks; Group 1: 0/32, Group 2: 1/31
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing:
; Group 2 Number missing:

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NO TREATMENT

# Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hospitalisation for recurrent colic at 4 weeks; Group 1: 0/32, Group 2: 0/28
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing:
; Group 2 Number missing:

## Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion at 4 weeks; Group 1: 27/32, Group 2: 13/28
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing:
; Group 2 Number missing:

CONSULTATION

### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension at 4 weeks; Group 1: 0/32, Group 2: 0/28 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NIFEDIPINE versus NO TREATMENT

Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hospitalisation for recurrent colic at 4 weeks; Group 1: 0/31, Group 2: 0/28 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion at 4 weeks; Group 1: 22/31, Group 2: 13/28 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension at 4 weeks; Group 1: 1/31, Group 2: 0/28 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details; no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Pain intensity; Analgesic use; Time to stone passage

Study	Itoh 2011 <sup>91</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=187)
Countries and setting	Conducted in Japan; Setting: Department of Nephro-urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 8 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: unenhanced computed tomography
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	symptomatic unilateral ureteral calculi <10mm in diameter
Exclusion criteria	urinary tract infection; severe hydronephrosis; diabetes; ulcers; hypotension; multiple stones; ureteral stricture
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): silodosin: 57.2 (12.7), control: 56.5 (10.1). Gender (M:F): 187 males. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not applicable 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Not stated / Unclear
Indirectness of population	No indirectness
Interventions	(n=95) Intervention 1: Alpha blockers - Silodosin. Silodosin 8mg daily. Duration up to 8 weeks. Concurrent medication/care: instruction to drink 2 L of water daily. Indirectness: No indirectness (n=92) Intervention 2: No treatment. No treatment. Duration up to 8 weeks. Concurrent medication/care: instruction to drink 2 L of water daily. Indirectness: No indirectness
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: SILODOSIN versus NO TREATMENT

Protocol outcome 1: Stone passage
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate (proximal stones) at 8 weeks; Group 1: 15/26, Group 2: 15/28; Comments: numbers calculated form percentages

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, stone size, stone location, or stone composition; Group 1 Number missing; Group 2 Number missing;

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate (mid-ureteral stones) at 8 weeks; Group 1: 4/8, Group 2: 1/8; Comments: numbers calculated from percentages

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details; no significant difference in age, stone size, stone location, or stone composition; Group 1 Number missing:; Group 2 Number missing:

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate (distal stones) at 8 weeks; Group 1: 40/55, Group 2: 31/56; Comments: numbers calculated from percentages

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, stone size, stone location, or stone composition; Group 1 Number missing:; Group 2 Number missing:

## Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time (proximal stones) at 8 weeks; Group 1: mean 13.45 days (SD 13.48); n=26, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, stone size, stone location, or stone composition; Group 1 Number missing:; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time (mid-ureteral stones) at 8 weeks; Group 1: mean 8.67 days (SD 5.03); n=8, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, stone size, stone location, or stone composition; Group 1 Number missing:; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time (distal stones) at 8 weeks; Group 1: mean 9.29 days (SD 5.91); n=55, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, stone size, stone location, or stone composition; Group 1 Number missing; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: retrograde ejaculation at 8 weeks; Group 1: 3/95, Group 2: 0/92 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details; no significant difference in age, stone size, stone location, or stone composition; Group 1 Number missing:; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension at 8 weeks; Group 1: 1/95, Group 2: 0/92 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, stone size, stone location, or stone composition; Group 1 Number missing:; Group 2 Number missing:

Protocol outcome 4: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of times analgesics were required (proximal stones) at 8 weeks; Group 1: mean 2.3 (SD 6.6); n=26,

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, stone size, stone location, or stone composition; Group 1 Number missing: ; Group 2 Number missing:

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of times analgesics were required (mid-ureteral stones) at 8 weeks; Group 1: mean 0.1 (SD 0.3); n=8,

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, stone size, stone location, or stone composition; Group 1 Number missing: ; Group 2 Number missing:

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of times analgesics were required (distal stones) at 8 weeks; Group 1: mean 0.3 (SD 0.9); n=55.

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, stone size, stone location, or stone composition; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Pain intensity; Hospitalisation/ Use of healthcare services

Study	Ketabchi 2014 <sup>99</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=142)
Countries and setting	Conducted in Iran; Setting: Urology department, single centre
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 2 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: KUB x-ray, abdominal ultrasonography and intravenous urography
Stratum	Adults (≥16 years), ureteric stone <1 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	single radio opaque lower ureteral stone with 5-10mm diameter
Exclusion criteria	urinary tract infections; high grade hydronephrosis; diabetes; history of hypersensitivity to alpha-blockers; ureteral stricture; pregnant women; history of spontaneous stone expulsion; previous ureteral surgery; hypotension or systolic blood pressure <110mmHg
Recruitment/selection of patients	consecutive patients meeting the inclusion criteria during the recruitment period
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 24 (6.5), control group: 27 (8.8). Gender (M:F): 77/25. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness: NA
Interventions	(n=52) Intervention 1: Alpha blockers and URS. Tamsulosin 0.4mg daily starting one day before URS. Duration up to 2 weeks. Concurrent medication/care: recommendation to drink 2 L of water daily, those with moderate to severe pain (>5 VAS) consumed Pethidine 25mg intravenously after the procedure in the recovery room and Indomethacin 500mg suppository daily. Indirectness: No indirectness; Indirectness comment: NA
	(n=50) Intervention 2: Surgery and placebo - URS and placebo. Placebo daily starting one day before URS. Duration up to 2 weeks. Concurrent medication/care: recommendation to drink 2 L of water daily, those with moderate to severe pain (>5 VAS) consumed Pethidine 25mg intravenously after the procedure in the recovery room and Indomethacin 500mg suppository daily. Indirectness: No indirectness; Indirectness comment: NA

Renal and ureteric stones: CONSULTATION Medical expulsive therapy

CONSULTATION

Funding

Academic or government funding (physiology center of Kerman University of Medical Sciences)

## RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND URS versus URS AND PLACEBO

### Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 2 weeks; Group 1: 49/52, Group 2: 35/50 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone location (left/right) or stone size; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 2: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: colic episodes at 2 weeks; Group 1: mean 1 (SD 0.7); n=52, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low: Indirectness of outcome: No indirectness. Comments: NA: Baseline details: no significant difference in age, sex, stone location (left/right) or stone size; Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 3: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: need for analgesia at 2 weeks; Group 1: 4/52, Group 2: 12/50 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low: Indirectness of outcome: No indirectness. Comments: NA: Baseline details: no significant difference in age, sex, stone location (left/right) or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Time to stone passage; Adverse events; Hospitalisation/ Use of healthcare services

Study	Kupeli 2004 <sup>109</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=78)
Countries and setting	Conducted in Turkey; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 15 days
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: x-rays, intravenous pyelography, helical computed tomography etc.
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	lower ureteral stones within the distal 5cm of the ureter that ranged between 3 and 15mm in size
Exclusion criteria	signs and symptoms of urinary tract infection; pregnancy; severely impacted stones; multiple stones; nonopaque stones; severe hydronephrosis; hepatic dysfunction; non-functioning kidney; treatment with calcium antagonists; morbid obesity
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (range): 42.9 (21-67). Gender (M:F): 56/22. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness: adjunctive therapy groups included stones < and > 10mm
Interventions	(n=15) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration 15 days. Concurrent medication/care: conventional treatment - oral hydration and oral Diclofenac 100mg daily. Indirectness: No indirectness; Indirectness comment: NA
	(n=15) Intervention 2: Pain management only - NSAIDs. Oral Diclofenac 100mg daily. Duration 15 days. Concurrent medication/care: oral hydration. Indirectness: No indirectness; Indirectness comment: NA
	(n=24) Intervention 3: Alpha blockers and SWL. Tamsulosin 0.4mg daily after SWL. Duration 15 days. Concurrent medication/care: conventional treatment - oral hydration and oral Diclofenac 100mg daily. Indirectness: No indirectness; Indirectness comment: NA
	(n=24) Intervention 4: Surgery and pain management - SWL and pain management. SWL. Duration 15 days. Concurrent medication/care: conventional treatment - oral hydration and oral Diclofenac 100mg daily.

	Indirectness: No indirectness; Indirectness comment: NA
Funding	Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone-free rate at 15 days; Group 1: 8/15, Group 2: 3/15

Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover

- Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone diameter; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 15 days; Group 1: 1/39, Group 2: 0/39
Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone diameter; Group 1 Number missing:
; Group 2 Number missing:

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL AND PAIN MANAGEMENT

# Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone-free rate at 15 days ; Group 1: 17/24, Group 2: 8/24
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover
- Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone diameter; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study Quality of life; Time to stone passage; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare services

Number of studies (number of participants)  Countries and setting  Line of therapy  Duration of study  Method of assessment of guideline condition  Stratum  Subgroup analysis within study  Inclusion criteria  Exclusion criteria  Exclusion criteria  Exclusion criteria  Exclusion criteria  Recruitment/selection of patients  Age, gender and ethnicity  Age - Mean (SD): 45.8 (12.1). Gender (M:F): 68/40. Ethnicity: not reported  Age (Inclusion composition/hou applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hou applicable 6. Ureteric stone: Urology, 2 university hospitals  1 (n=108)  Conducted in South Korea; Setting: Department of Urology, 2 university hospitals  1 (n=108)  Conducted in South Korea; Setting: Department of Urology, 2 university hospitals  1 (n=108)  Conducted in South Korea; Setting: Department of Urology, 2 university hospitals  1 (n=108)  Conducted in South Korea; Setting: Department of Urology, 2 university hospitals  1 (n=108)  Conducted in South Korea; Setting: Department of Urology, 2 university hospitals  1 (n=108)  Conducted in South Korea; Setting: Department of Urology, 2 university hospitals  Adequate method of assessment/diagnosis: plain abdominal radiography kidney ureter blade contrast CT  Madequate method of assessment/diagnosis: plain abdominal radiography kidney ureter blade contrast CT  Adults (≥16 years), ureteric stone <1 cm  Adequate method of assessment/diagnosis: plain abdominal radiography kidney ureter blade contrast CT  Adults (≥16 years), ureteric stone <1 cm  Adequate method of assessment/diagnosis: plain abdominal radiography kidney ureter blade contrast CT  Adults (≥16 years), ureteric stone <1 cm  Adults (≥16 years), ureteri	
Countries and setting  Line of therapy  1st line  Duration of study  Method of assessment of guideline condition  Stratum  Subgroup analysis within study  Inclusion criteria  Exclusion criteria  Exclusion criteria  Exclusion criteria  Exclusion criteria  Recruitment/selection of patients  Age - Mean (SD): 45.8 (12.1). Gender (M:F): 68/40. Ethnicity: Not applicable 2. Neuropathic/ cerebral-palsy //immobility: Not applicable 3. Capplicable 4. Pregnant women: Non-pregnant 5. Stone composition/hou applicable 6. Ureteric stone: Upper ureteric stones	
Line of therapy Duration of study Intervention time: 4 weeks Method of assessment of guideline condition Stratum Adequate method of assessment/diagnosis: plain abdominal radiography kidney ureter bladd contrast CT Stratum Adults (≥16 years), ureteric stone <1 cm Subgroup analysis within study Inclusion criteria  ≥18 years; presenting with renal colic; diagnosed with single, unilateral radiopaque, proxima segment between the ureteropelvic junction and the upper border of the sacroiliac joint) uret ≤6mm in diameter; agreed to undergo conservative management  Exclusion criteria  ureteral calculi ≥7mm or multiple ureteral calculi; febrile urinary tract infection; single kidney; kidney; pregnancy; azotaemia (creatinine >1.8 mg/dl); ureteral stricture; severe hydronephrovate treatment with medications that could affect stone passage such as alpha-blockers, calcium blockers, steroids, or nitrates; patients wanting immediate stone removal because of colic  Recruitment/selection of patients  Age, gender and ethnicity  Age - Mean (SD): 45.8 (12.1). Gender (M:F): 68/40. Ethnicity: not reported  1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Calcium distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hou applicable 6. Ureteric stone: Upper ureteric stones	
Duration of study  Method of assessment of guideline condition  Stratum  Adequate method of assessment/diagnosis: plain abdominal radiography kidney ureter blade contrast CT  Stratum  Adults (≥16 years), ureteric stone <1 cm  Subgroup analysis within study  Inclusion criteria  ≥18 years; presenting with renal colic; diagnosed with single, unilateral radiopaque, proxima segment between the ureteropelvic junction and the upper border of the sacroiliac joint) uret ≤6mm in diameter; agreed to undergo conservative management  Exclusion criteria  ureteral calculi ≥7mm or multiple ureteral calculi; febrile urinary tract infection; single kidney; kidney; pregnancy; azotaemia (creatinine >1.8mg/dl); ureteral stricture; severe hydronephrot treatment with medications that could affect stone passage such as alpha-blockers, calcium blockers, steroids, or nitrates; patients wanting immediate stone removal because of colic  Recruitment/selection of patients  Age, gender and ethnicity  Age - Mean (SD): 45.8 (12.1). Gender (M:F): 68/40. Ethnicity: not reported  1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. C stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hou applicable 6. Ureteric stone: Upper ureteric stones	
Method of assessment of guideline condition       Adequate method of assessment/diagnosis: plain abdominal radiography kidney ureter blade contrast CT         Stratum       Adults (≥16 years), ureteric stone <1 cm	
condition  Stratum  Adults (≥16 years), ureteric stone <1 cm  Subgroup analysis within study  Inclusion criteria  ≥18 years; presenting with renal colic; diagnosed with single, unilateral radiopaque, proxima segment between the ureteropelvic junction and the upper border of the sacrolliac joint) uret ≤6mm in diameter; agreed to undergo conservative management  Exclusion criteria  ureteral calculi ≥7mm or multiple ureteral calculi; febrile urinary tract infection; single kidney; kidney; pregnancy; azotaemia (creatinine >1.8mg/dl); ureteral stricture; severe hydronephrot treatment with medications that could affect stone passage such as alpha-blockers, calcium blockers, steroids, or nitrates; patients wanting immediate stone removal because of colic  Recruitment/selection of patients  Age, gender and ethnicity  Age - Mean (SD): 45.8 (12.1). Gender (M:F): 68/40. Ethnicity: not reported  Further population details  1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Ostone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hou applicable 6. Ureteric stone: Upper ureteric stones	
Subgroup analysis within study       Not applicable         Inclusion criteria       ≥18 years; presenting with renal colic; diagnosed with single, unilateral radiopaque, proxima segment between the ureteropelvic junction and the upper border of the sacroiliac joint) uret ≤6mm in diameter; agreed to undergo conservative management         Exclusion criteria       ureteral calculi ≥7mm or multiple ureteral calculi; febrile urinary tract infection; single kidney; kidney; pregnancy; azotaemia (creatinine >1.8mg/dl); ureteral stricture; severe hydronephrosis treatment with medications that could affect stone passage such as alpha-blockers, calcium blockers, steroids, or nitrates; patients wanting immediate stone removal because of colic not reported         Recruitment/selection of patients       not reported         Age, gender and ethnicity       Age - Mean (SD): 45.8 (12.1). Gender (M:F): 68/40. Ethnicity: not reported         Further population details       1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Content of the sacroiliac composition/hou applicable 6. Ureteric stone: Upper ureteric stones	der and non-
Inclusion criteria  ≥18 years; presenting with renal colic; diagnosed with single, unilateral radiopaque, proxima segment between the ureteropelvic junction and the upper border of the sacroiliac joint) uret ≤6mm in diameter; agreed to undergo conservative management  Exclusion criteria  ureteral calculi ≥7mm or multiple ureteral calculi; febrile urinary tract infection; single kidney; kidney; pregnancy; azotaemia (creatinine >1.8mg/dl); ureteral stricture; severe hydronephrost treatment with medications that could affect stone passage such as alpha-blockers, calcium blockers, steroids, or nitrates; patients wanting immediate stone removal because of colic Recruitment/selection of patients  Age, gender and ethnicity  Age - Mean (SD): 45.8 (12.1). Gender (M:F): 68/40. Ethnicity: not reported  1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Constance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hou applicable 6. Ureteric stone: Upper ureteric stones	
segment between the ureteropelvic junction and the upper border of the sacroiliac joint) uret ≤6mm in diameter; agreed to undergo conservative management  Exclusion criteria ureteral calculi ≥7mm or multiple ureteral calculi; febrile urinary tract infection; single kidney; kidney; pregnancy; azotaemia (creatinine >1.8mg/dl); ureteral stricture; severe hydronephrost treatment with medications that could affect stone passage such as alpha-blockers, calcium blockers, steroids, or nitrates; patients wanting immediate stone removal because of colic not reported  Age, gender and ethnicity Age - Mean (SD): 45.8 (12.1). Gender (M:F): 68/40. Ethnicity: not reported  1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Constance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hou applicable 6. Ureteric stone: Upper ureteric stones	
kidney; pregnancy; azotaemia (creatinine >1.8mg/dl); ureteral stricture; severe hydronephrosis treatment with medications that could affect stone passage such as alpha-blockers, calcium blockers, steroids, or nitrates; patients wanting immediate stone removal because of colic not reported  Age, gender and ethnicity  Age - Mean (SD): 45.8 (12.1). Gender (M:F): 68/40. Ethnicity: not reported  1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Constance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hou applicable 6. Ureteric stone: Upper ureteric stones	
Age, gender and ethnicity  Age - Mean (SD): 45.8 (12.1). Gender (M:F): 68/40. Ethnicity: not reported  1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Constance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hou applicable 6. Ureteric stone: Upper ureteric stones	sis; current
Further population details  1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. C stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hou applicable 6. Ureteric stone: Upper ureteric stones	
stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hou applicable 6. Ureteric stone: Upper ureteric stones	
In Proceedings of a contraction of the Contraction	
Indirectness of population No indirectness	
Interventions  (n=54) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.2mg daily. Duration up to Concurrent medication/care: instruction to drink 2 L of water daily and oral painkiller (Ultrace of Tramadol and Acetaminophen) on demand. Indirectness: No indirectness  (n=54) Intervention 2: No treatment. No treatment. Duration up to 4 weeks. Concurrent medication instruction to drink 2 L of water daily and oral painkiller (Ultracet® combination of Tramadol at Acetaminophen) on demand. Indirectness: No indirectness	et® combination
Funding Study funded by industry (Korean Astellas Pharm, Co.)	

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NO TREATMENT

# Protocol outcome 1: Quality of life

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: post-trial EuroQoL at 4 weeks; Group 1: mean 5.4 (SD 0.6); n=44, Group 2: mean 5.5 (SD 0.8); n=35; EuroQoL 0-10 Top=High is good outcome

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, past stone history, baseline pain and QoL scores, stone site or stone size; Group 1 Number missing: 10, Reason: 6 lost to follow-up, 4 converted to active treatment; Group 2 Number missing: 19, Reason: 8 lost to follow-up, 11 converted to active treatment

### Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone passage at 4 weeks; Group 1: 40/54, Group 2: 25/54 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, past stone history, baseline pain and QoL scores, stone site or stone size; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 3: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to stone passage at 4 weeks; Group 1: mean 14.3 days (SD 7.9); n=44, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, past stone history, baseline pain and QoL scores, stone site or stone size; Group 1 Number missing: 10, Reason: 6 lost to follow-up, 4 converted to active treatment; Group 2 Number missing: 19, Reason: 8 lost to follow-up, 11 converted to active treatment

# Protocol outcome 4: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: requirement for oral analgesics at 4 weeks; Group 1: mean 3.5 unclear (SD 3.8); n=44, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, past stone history, baseline pain and QoL scores, stone site or stone size; Group 1 Number missing: 10, Reason: 6 lost to follow-up, 4 converted to active treatment; Group 2 Number missing: 19, Reason: 8 lost to follow-up, 11 converted to active treatment

Protocol outcomes not reported by the study

Adverse events; Pain intensity; Hospitalisation/ Use of healthcare services

Study	Lojanapiwat 2008 <sup>120</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=75)
Countries and setting	Conducted in Thailand; Setting: Division of Urology, Department of Surgery, 2 hospitals
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: plain kidney, ureter and bladder radiographs
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	distal ureteric stones of 4-10mm; measured by plain KUB; gave informed consent; interviewed prior to taking part
Exclusion criteria	urinary tract infection; severe hydronephrosis; history of ureteric surgery
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): control group: 46.52 (13.63), Tamsulosin 0.2mg: 48 (15.74), Tamsulosin 0.4mg: 46.71 (12.2). Gender (M:F): 55/20. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=50) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.2mg or 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 50mg twice daily for 10 days and Diclofenac 75mg infection if renal colic developed during treatment . Indirectness: No indirectness
	(n=25) Intervention 2: Pain management only - NSAIDs. Diclofenac 50mg twice daily. Duration 10 days. Concurrent medication/care: Diclofenac 75mg injection if renal colic developed. Indirectness: No indirectness
Funding	Equipment / drugs provided by industry (Astellas Pharma )

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS

Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate (0.2mg Tamsulosin) at 4 weeks; Group 1: 10/25, Group 2: 1/25 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, weight or stone size; Group 1 Number missing: : Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate (0.4mg Tamsulosin) at 4 weeks; Group 1: 17/25, Group 2: 1/25 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, weight or stone size; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time (0.2mg Tamsulosin) at 4 weeks; Group 1: mean 9.3 days (SD 6.06); n=25, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, weight or stone size; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time (0.4mg Tamsulosin) at 4 weeks : Group 1: mean 10.76 days (SD 7.52); n=25.

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, weight or stone size; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension (0.2mg Tamsulosin) at 4 weeks; Group 1: 0/25, Group 2: 0/25 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, weight or stone size; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension (0.4mg Tamsulosin) at 4 weeks; Group 1: 0/25, Group 2: 0/25 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, weight or stone size; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: retrograde ejaculation (0.2mg Tamsulosin) at 4 weeks : Group 1: 0/15, Group 2: 0/20 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, weight or stone size; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: retrograde ejaculation (0.4mg Tamsulosin) at 4 weeks; Group 1: 0/20, Group 2: 0/20 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, weight or stone size; Group 1 Number missing: ; Group 2 Number missing:

CONSULTATION

Protocol outcome 4: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Diclofenac injection (0.2mg Tamsulosin) at 4 weeks; Group 1: 1/25, Group 2: 0/25 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, weight or stone size; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Diclofenac injection (0.4mg Tamsulosin) at 4 weeks; Group 1: 0/25, Group 2: 0/25 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, weight or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported b	y the
study	

Quality of life; Pain intensity; Hospitalisation/ Use of healthcare services

Study	Lv 2014 <sup>125</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=105)
Countries and setting	Conducted in China; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 2 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: abdominal ultrasound, plain abdominal x-ray for KUB, intravenous urogram or unenhanced CT when necessary
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	distal ureteral stone 4-9mm
Exclusion criteria	multiple stones; history of distal ureteral surgery; renal colic for >24hours; urinary tract infection; severe hydronephrosis; voiding dysfunction; hypotension; cardiovascular and cerebrovascular diseases; hepatic and renal dysfunction; pregnancy; diabetes; ulcer disease; history of hypersensitivity to Naftopidil; subjects receiving treatment with cardiovascular drugs, other NSAIDs, alpha receptor antagonists or calcium antagonists
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Naftopidil group: 31.4 (2.94), Naftopidil + Celecoxib group: 33.2 (5.28), Celecoxib group: 33.75 (5.24). Gender (M:F): 59/44. Ethnicity: not reported

Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=35) Intervention 1: Alpha blockers - Naftopidil. Naftopidil 50mg daily. Duration up to 2 weeks. Concurrent medication/care: instruction to drink at least 2 L of fluids daily. Indirectness: No indirectness (n=35) Intervention 2: Alpha blockers - Naftopidil . Naftopidil 50mg daily and Celecoxib 400mg immediately then 200mg every 12 hours. Duration up to 2 weeks. Concurrent medication/care: instruction to drink at least 2 L of fluids daily. Indirectness: No indirectness (n=33) Intervention 3: Pain management only - NSAIDs. Celecoxib 400mg immediately then 200mg every 12 hours. Duration up to 2 weeks. Concurrent medication/care: instruction to drink at least 2 L of fluids daily. Indirectness: No indirectness
Funding	Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NAFTOPIDIL versus NSAIDS

# Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate (Naftopidil) at 2 weeks; Group 1: 29/35, Group 2: 20/33 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low: Indirectness of outcome: No indirectness: Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing:; Group 2 Number missing:

# Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to expulsion (Naftopidil) at 2 weeks; Group 1: mean 8 days (SD 2.07); n=35, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness (Naftopidil) at 2 weeks; Group 1: 4/35, Group 2: 8/33 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing:; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: headache (Naftopidil) at 2 weeks; Group 1: 2/35, Group 2: 0/33

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing:

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: retrograde ejaculation (Naftopidil) at 2 weeks; Group 1: 0/20, Group 2: 0/18
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or
baseline VAS score; Group 1 Number missing:; Group 2 Number missing:

### Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: pain episodes (Naftopidil) at 2 weeks; Group 1: mean 2.22 pain episodes (SD 0.94); n=35, Risk of bias: All domain Very high, Selection High, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: VAS score (Naftopidil) at 3 days; Group 1: mean 5.74 (SD 0.92); n=35, Group 2: mean 3.06 (SD 1.14); n=33; visual analogue scale 0-10 Top=High is poor outcome

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing:

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: VAS score (Naftopidil) at 7 days ; Group 1: mean 4.8 (SD 0.53); n=35, Group 2: mean 1.57 (SD 0.5); n=33; visual analogue scale 0-10 Top=High is poor outcome

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing:

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NAFTOPIDIL versus NSAIDS

# Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate (Naftopidil + Celecoxib) at 2 weeks; Group 1: 33/35, Group 2: 20/33 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to expulsion (Naftopidil + Celecoxib) at 2 weeks; Group 1: mean 7.7 days (SD 2.34); n=35.

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; dizziness (Naftopidil + Celecoxib) at 2 weeks; Group 1: 6/35, Group 2: 8/33 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; headache (Naftopidil + Celecoxib) at 2 weeks; Group 1: 2/35, Group 2: 0/33 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing:; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: retrograde ejaculation (Naftopidil + Celecoxib) at 2 weeks; Group 1: 1/21, Group 2: 0/18 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low: Indirectness of outcome: No indirectness: Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing:; Group 2 Number missing:

## Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: pain episodes (Naftopidil + Celecoxib) at 2 weeks; Group 1: mean 1.37 pain episodes (SD 1.33); n=35,

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing; Group 2 Number missing;

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: VAS score (Naftopidil + Celecoxib) at 3 days; Group 1: mean 3.11 (SD 0.63); n=35, Group 2: mean 3.06 (SD 1.14); n=33; visual analogue scale 0-10 Top=High is poor outcome
- Risk of bias: All domain Very high, Selection High, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing:; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: VAS score (Naftopidil + Celecoxib) at 7 days; Group 1: mean 1.6 (SD 0.6); n=35, Group 2: mean 1.57 (SD 0.5); n=33; visual analogue scale 0-10 Top=High is poor outcome

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness: Baseline details: no significant difference in age, sex, stone size, stone location (left/right) or baseline VAS score; Group 1 Number missing:; Group 2 Number missing:

Protocol outcomes not reported by the study	Quality of life; Analgesic use; Hospitalisation/ Use of healthcare services
Study	Mokhless 2012 <sup>132</sup>
Study type	RCT (Patient randomised; Parallel)

Number of studies (number of participants)	1 (n=61)
Countries and setting	Conducted in Egypt; Setting: Section of Pediatric Urology and Endourology, Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: ultrasound of urinary tract, plain x-ray of abdomen and pelvis and renal function tests, non-contrast CT when indicated
Stratum	Children (<16 years)
Subgroup analysis within study	Not applicable
Inclusion criteria	radiopaque lower ureteral stones of ≤12mm
Exclusion criteria	anatomical abnormalities; non-radiopaque stones; voiding dysfunction; urinary tract infection; severe hydronephrosis; history of endoscopic or open ureteral surgery
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): 8.1 (6.8). Gender (M:F): 36/25. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not applicable 5. Stone composition/hounsfield units: Not applicable 6. Uteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=33) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4 mg for children older than 4 years and 0.2 mg for younger children at bed time in addition to standard analgesia (ibuprofen). Those who could swallow the whole capsule were allowed to do so otherwise the content of the capsule was evacuated in water or juice
	. Duration 4 weeks. Concurrent medication/care: Standard analgesia (ibuprofen)
	(n=28) Intervention 2: Placebo. Placebo. Duration 4 weeks. Concurrent medication/care: Standard analgesia (ibuprofen)
Funding	No funding
RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus PLACEBO	

## Protocol outcome 1: Stone passage at Define

- Actual outcome for Children (<16 years): Expulsion rate at 4 weeks; Group 1: 29/33, Group 2: 18/28 Risk of bias: All domain - Very high, Selection - Very high, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 2: Time to stone passage at Define

- Actual outcome for Children (<16 years): Time to expulsion at 4 weeks; Group 1: mean 8.2 days (SD 3.4); n=33, Group 2: mean 14.5 days (SD 4.5); n=28

Risk of bias: All domain - Very high, Selection - Very high, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events at Define

- Actual outcome for Children (<16 years): Hypotension at 4 weeks; Group 1: 0/33, Group 2: 0/28

Risk of bias: All domain - Very high, Selection - Very high, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low. Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing; Group 2 Number missing;

## Protocol outcome 4: Pain intensity at Define

- Actual outcome for Children (<16 years): Pain episodes at 4 weeks; Group 1: mean 1.4 (SD 1.2); n=33, Group 2: mean 2.2 (SD 1.4); n=28 Risk of bias: All domain - Very high, Selection - Very high, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 5: Analgesic use at Define

- Actual outcome for Children (<16 years): Need for analgesia at 4 weeks; Group 1: mean 0.7 (SD 0.9); n=33, Group 2: mean 1.4 (SD 1.1); n=28 Risk of bias: All domain - Very high, Selection - Very high, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life at Define; Hospitalisation/ Use of healthcare services at Define

Study	Mohseni 2006 <sup>131</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=64)
Countries and setting	Conducted in Iran; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: abdominal sonography or kidney, ureter, bladder
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	lower ureteral stone
Exclusion criteria	urinary tract infection; severe hydronephrosis; elevated creatinine; hypertension; history of peptic ulcer disease; spontaneous stone passage; any previous intervention
Recruitment/selection of patients	consecutive
Age, gender and ethnicity	Age - Mean (SD): Terazosin group: 44.2 (12.9), control group: 39.3 (14.2). Gender (M:F): 44/20. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=32) Intervention 1: Pain management only - NSAIDs. Indomethacin. Duration up to 4 weeks. Concurrent medication/care: intravenous Pethidine in cases of incomplete pain control. Indirectness: No indirectness (n=32) Intervention 2: Alpha blockers - Terazosin. Terazosin 10mg daily. Duration up to 4 weeks. Concurrent medication/care: Indomethacin and intravenous Pethidine in cases of incomplete pain control. Indirectness: No indirectness
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TERAZOSIN versus NSAIDS

Protocol outcome 1: Stone passage - Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate at 4 weeks ; Group 1: 29/32, Group 2: 20/32

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone size; Group 1 Number missing; Group 2 Number missing:

### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to expulsion at 4 weeks; Group 1: mean 76.3 hours (SD 60); n=32. Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone size; Group 1 Number missing; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension at 4 weeks; Group 1: 3/32, Group 2: 0/32 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone size; Group 1 Number missing; Group 2 Number missing:

## Protocol outcome 4: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: amount of Pethidine administered at 4 weeks; Group 1: mean 34.4 mg (SD 12.7); n=32, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone size; Group 1 Number missing; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Pain intensity; Hospitalisation/ Use of healthcare services

Study	Moursy 2010 <sup>134</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=88)
Countries and setting	Conducted in Egypt; Setting: Urology department, single centre
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: KUB radiographs
Stratum	Adults (≥16 years), ureteric stone <1 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	>18 years with unilateral steinstrasse
Exclusion criteria	clinical and laboratory signs of urinary tract infection, severe hydronephrosis, alterations in creatininaemia, diabetes, ulcer disease or hypotension; concomitant usage of calcium antagonists; distal ureteral surgery
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 35.6 (9.95), control group: 33.9 (9.71). Gender (M:F): 55/33. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Not stated / Unclear
Indirectness of population	No indirectness: NA
Interventions	(n=44) Intervention 1: Alpha blockers and SWL. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: Indomethacin 100mg suppository on demand and encouragement to drink a minimum 2.5 L of water daily. Indirectness: No indirectness; Indirectness comment: NA
	(n=44) Intervention 2: Surgery - SWL. Pain management only. Duration up to 4 weeks. Concurrent medication/care: Indomethacin 100mg suppository on demand and encouragement to drink a minimum 2.5 L of water daily. Indirectness: No indirectness; Indirectness comment: NA
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL

Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Hospitalisation at 4 weeks; Group 1: 12/44, Group 2: 19/44 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone location, stone length or fragment size; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate at 4 weeks; Group 1: 32/44, Group 2: 25/44 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone location, stone length or fragment size; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 3: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; stone expulsion time at 4 weeks; Group 1; mean 12.67 days (SD 2.29); n=44. Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone location, stone length or fragment size; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 4: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: anejaculation at 4 weeks; Group 1: 6/28, Group 2: 0/27 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone location, stone length or fragment size; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: headache at 4 weeks; Group 1: 4/44, Group 2: 0/44 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone location, stone length or fragment size; Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of times analgesics used at 4 weeks; Group 1: mean 4.39 (SD 2.42); n=44, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone location, stone length or fragment size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Pain intensity

Study	Mustafa 2016 <sup>136</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=128)
Countries and setting	Conducted in Bangladesh; Setting: Outpatient Department of Urology
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: history, physical examination and investigations (e.g. ultrasonography)
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	>18 years; unilateral, juxtavesical ureteral stone; normal functioning kidney; absence of clinical and laboratory signs of urinary tract infection; stone size up to 8mm
Exclusion criteria	multiple stones; severe hydronephrosis; history of spontaneous stone expulsion; distal ureteral surgery; diabetes; peptic ulcer disease; hypotension;
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 37.7 (9.33), control group: 38.5 (10.05). Gender (M:F): not reported . Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=64) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: conventional treatment - hydration with minimum 2 L of water daily, physical exertion and analgesics (Diclofenac 50mg suppository with H2 blocker) if required. Indirectness: No indirectness
	(n=64) Intervention 2: No treatment. No treatment. Duration up to 4 weeks. Concurrent medication/care: hydration with minimum 2 L of water daily, physical exertion and analgesics (Diclofenac 50mg suppository with H2 blocker) if required. Indirectness: No indirectness
Funding	Funding not stated
RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NO TREATMENT	

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion at 4 weeks; Group 1: 51/60, Group 2: 32/60 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone size; Group 1 Number missing; Group 2 Number missing:

### Protocol outcome 2: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; pain episodes at 4 weeks; Group 1; mean 2.58 (SD 1.519); n=60, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone size; Group 1 Number missing; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of people suffering pain at 4 weeks; Group 1: 36/60, Group 2: 48/60 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age or stone size; Group 1 Number missing; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Time to stone passage; Adverse events; Analgesic use; Hospitalisation/ Use of healthcare services

Renal and ureteric stones: Medical expulsive therapy

CONSULTATION

Study	Ochoa-gomez 2011 <sup>139</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=65)
Countries and setting	Conducted in Mexico; Setting: Emergency room, single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: plain abdominal film and kidney ultrasound
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	>18 years; reno-ureteral stones 5-10mm determined by plain abdominal film and kidney ultrasound
Exclusion criteria	hydronephrosis; acute or chronic renal insufficiency, multiple ureteral lithiasis; history of surgery or endourologic procedures; large and impacted ureteral calculi; pregnancy; lactation; distal ureteral lithiasis in a single kidney; patients taking alpha- or beta-blockers, nitrates or calcium antagonists; patients who worked as airline pilots
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 38.5 (11.3), placebo group: 38.2 (12.4). Gender (M:F): 36/39. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=32) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: instruction to drink at least 2 L of water daily. Indirectness: No indirectness (n=33) Intervention 2: Placebo. Placebo. Duration up to 4 weeks. Concurrent medication/care: instruction to drink at least 2 L of water daily. Indirectness: No indirectness
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus PLACEBO

Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; stone expulsion at 4 weeks; Group 1: 22/32, Group 2: 23/33 Risk of bias: All domain - High, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness: Baseline details: no significant difference in age, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 4 weeks; Group 1: mean 22 days (SD 6.77); n=32, Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 4 weeks; Group 1: 2/32, Group 2: 0/33 Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details; no significant difference in age, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: retrograde ejaculation at 4 weeks; Group 1: 2/15, Group 2: 0/21 Risk of bias: All domain - Very high, Selection - High, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare services

Study	Park 2013 <sup>142</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=96)
Countries and setting	Conducted in South Korea; Setting: outpatient setting
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 3 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: plain abdominal KUB radiography and non-enhanced kidney CT
Stratum	Adults (≥16 years), ureteric stone <1 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	18-70 years with symptomatic, unilateral, single, proximal ureteral stone 6-20mm in longest axis
Exclusion criteria	active urinary tract infection; severe hydronephrosis; pregnancy; inadequate renal function (serum creatinine >2mg/dL); concomitant treatment with alpha blockers, calcium channel blockers or steroids; hypotension; multiple urinary stones; morbid obesity; stone on non-functioning kidney; history of previous failed ESWL; history of urinary tract surgery; uncorrected urinary tract obstruction
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Median (IQR): Tamsulosin group: 49.5 (34.25-57.75), control group: 50.5 (39.25-55.75). Gender (M:F): 57/31. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Upper ureteric stones
Indirectness of population	No indirectness: NA
Interventions	(n=48) Intervention 1: Alpha blockers and SWL. Tamsulosin 0.2mg once daily, starting just before ESWL. Duration up to 3 weeks. Concurrent medication/care: Aceclofenac 100mg on demand and asked to drink 1.5-2L of water daily. Indirectness: No indirectness; Indirectness comment: NA
	(n=48) Intervention 2: Surgery - SWL. ESWL . Duration up to 3 weeks. Concurrent medication/care: Aceclofenac 100mg on demand and asked to drink 1.5-2L of water daily. Indirectness: No indirectness; Indirectness comment: NA
Funding	Study funded by industry (Astellas Pharma Korea)
RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL	

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone free rate at 3 weeks; Group 1: 37/44, Group 2: 29/44 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (left/right) or stone size; Group 1 Number missing: 4; Group 2 Number missing: 4

Renal and ureteric stones: Medical expulsive therapy

CONSULTATION

## Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 3 weeks; Group 1: 1/44, Group 2: 0/44 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (left/right) or stone size; Group 1 Number missing: 4; Group 2 Number missing: 4

Protocol outcomes not reported by the	Quality of life; Time to stone passage; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare
study	services

Study	Pedro 2008 <sup>144</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=69)
Countries and setting	Conducted in USA; Setting: Department of Urology (patients recruited from emergency room), single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: CT
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	distal ureteral calculus
Exclusion criteria	stones >8mm; renal insufficiency (serum creatinine >1.8mg/dl); solitary kidney; urinary infection; current alpha-blocker use; pregnancy; history of ureteral stricture; allergic reaction to study medication
Recruitment/selection of patients	consecutive patients meeting the inclusion/exclusion criteria during the recruitment period
Age, gender and ethnicity	Age - Mean (SD): Alfluzosin group: 36.69 (13.06), placebo group: 42.03 (12.85). Gender (M:F): 55/14. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	<ul><li>(n=34) Intervention 1: Alpha blockers - Alfuzosin. Alfuzosin daily. Duration up to 4 weeks. Concurrent medication/care: not reported. Indirectness: No indirectness</li><li>(n=35) Intervention 2: Placebo. Placebo. Duration up to 4 weeks. Concurrent medication/care: not reported.</li></ul>
Funding	Indirectness: No indirectness  Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALFUZOSIN versus PLACEBO

Protocol outcome 1: Stone passage
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage at 4 weeks; Group 1: 25/34, Group 2: 27/35; Comments: numbers calculated from percentages

Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, blood pressure, degree of hydronephrosis or stone size, higher baseline pain score in Alfuzosin group; Group 1 Number missing; Group 2 Number missing;

### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to stone passage at 4 weeks; Group 1: mean 5.19 days (SD 4.82); n=34, Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, blood pressure, degree of hydronephrosis or stone size, higher baseline pain score in Alfuzosin group; Group 1 Number missing; Group 2 Number missing;

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: side effects (dizziness and hypotension) at 4 weeks; Group 1: 4/34, Group 2: 0/35 Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, blood pressure, degree of hydronephrosis or stone size, higher baseline pain score in Alfuzosin group; Group 1 Number missing; Group 2 Number missing;

### Protocol outcome 4: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of opioid derived medications consumed at 4 weeks; Group 1: mean 8.63 (SD 8.58): n=34.

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details; no significant difference in age, sex, blood pressure, degree of hydronephrosis or stone size, higher baseline pain score in Alfuzosin group; Group 1 Number missing; Group 2 Number missing;

Protocol outcomes not reported by the study

Quality of life; Pain intensity; Hospitalisation/ Use of healthcare services

Study	Rahim 2012 <sup>157</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=90)
Countries and setting	Conducted in Pakistan; Setting: Urology department, single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: ultrasound
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	4-7mm stones in the distal segment of the ureter confirmed on ultrasound
Exclusion criteria	urinary tract infection; severe hydronephrosis; pregnancy; ulcer disease; hypotension; patients on calcium channel blockers; serum creatinine >2mg/dl; multiple ureteral stones; bilateral distal ureteric stones; solitary kidney; ureteral stricture; patient desire for immediate stone retrieval
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): 33.12 (11.2). Gender (M:F): 63/27. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=45) Intervention 1: Alpha blockers - Terazosin. Terazosin 2mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 50mg twice daily. Indirectness: No indirectness (n=45) Intervention 2: Pain management only - NSAIDs. Diclofenac 50mg twice daily. Duration up to 4 weeks. Concurrent medication/care: NA. Indirectness: No indirectness
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TERAZOSIN versus NSAIDS

Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion at 4 weeks; Group 1: 37/45, Group 2: 22/45
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,

Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age; Group 1 Number missing:; Group 2 Number missing:

Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 4 weeks; Group 1: mean 13.3 days (SD 6.31); n=45, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Adverse events; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare services

Study	Resim 2005 <sup>162</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=60)
Countries and setting	Conducted in Turkey; Setting: outpatient Division of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 6 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: plain abdominal radiography and urinary ultrasonography
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	lower ureteral calculi
Exclusion criteria	solitary kidney; severe refractory pain; urinary tract infection; multiple stones; severe hydronephrosis
Recruitment/selection of patients	consecutive meeting the inclusion/exclusion criteria during the recruitment period
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 35.3 (10.9), control group 33.5 (9.7). Gender (M:F): 45/15. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	Serious indirectness: included stones < and > 10mm
Interventions	(n=30) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 6 weeks. Concurrent medication/care: conservative treatment - hydration and Tenoxicam 20mg daily. Indirectness: No indirectness  (n=30) Intervention 2: Pain management only - NSAIDs. Conservative treatment - hydration and Tenoxicam
	20mg daily. Duration up to 6 weeks. Concurrent medication/care: NA. Indirectness: No indirectness
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS

Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage at 6 weeks; Group 1: 26/30, Group 2: 22/30 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,

Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size, higher pain scores in control group; Group 1 Number missing; Group 2 Number missing:

Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: headache at 6 weeks ; Group 1: 4/30, Group 2: 4/30
- Risk of bias: All domain Very high, Selection High, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size, higher pain scores in control group; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 6 weeks; Group 1: 5/30, Group 2: 3/30
  Risk of bias: All domain Very high, Selection High, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size, higher pain scores in control group; Group 1 Number missing: Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: abnormal ejaculation at 6 weeks; Group 1: 0/22, Group 2: 1/23
  Risk of bias: All domain Very high, Selection High, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High,
  Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size, higher pain scores in
  control group; Group 1 Number missing: Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: orthostatic hypotension at 6 weeks; Group 1: 0/30, Group 2: 0/30
  Risk of bias: All domain Very high, Selection High, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High,
  Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size, higher pain scores in
  control group; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Time to stone passage; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare services

Protocol outcome 1: Stone passage

Study	Resim 2005 <sup>163</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=67)
Countries and setting	Conducted in Turkey; Setting: Department of Urology, single centre
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 6 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: plain abdominal radiography and urinary ultrasonography
Stratum	Adults (≥16 years), ureteric stone <1 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	steinstrasse in the lower ureter (juxtavesical or intramural portion) after undergoing ESWL
Exclusion criteria	<18 years; weight <50kg or >100kg; history of drug or alcohol abuse; ipsilateral ureteral surgery; chronic use of drugs such as antidepressants, histamine blockers and anxiolytics; allergy to one of the study medications
Recruitment/selection of patients	consecutive patients meeting the inclusion criteria during the recruitment period
Age, gender and ethnicity	Age - Median (range): Tamsulosin group: 39 (21-55), control group: 37 (23-57). Gender (M:F): 43/24. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Uteric stone: Lower ureteric stones
Indirectness of population	No indirectness: NA
Interventions	(n=32) Intervention 1: Alpha blockers and SWL. Tamsulosin 0.4mg daily. Duration up to 6 weeks. Concurrent medication/care: hydration and Tenoxicam 20mg daily. Indirectness: No indirectness; Indirectness comment: NA
	(n=35) Intervention 2: Surgery and pain management - SWL and pain management. Pain management only. Duration up to 6 weeks. Concurrent medication/care: hydration and Tenoxicam 20mg daily. Indirectness: No indirectness; Indirectness comment: NA
Funding	Funding not stated
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RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL AND PAIN MANAGEMENT

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; spontaneous passage at 6 weeks; Group 1: 24/32, Group 2: 23/35 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex or stone burden before ESWL; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; headache at 6 weeks; Group 1: 5/32, Group 2: 2/35 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex or stone burden before ESWL; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 6 weeks; Group 1: 4/32, Group 2: 0/35 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low: Indirectness of outcome: No indirectness. Comments: NA: Baseline details: no significant difference in age, sex or stone burden before ESWL; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: abnormal ejaculation at 6 weeks; Group 1: 1/21, Group 2: 0/22 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex or stone burden before ESWL; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: orthostatic hypotension at 6 weeks; Group 1: 0/32, Group 2: 0/35 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex or stone burden before ESWL; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Time to stone passage; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare services

Study	Sameer 2014 <sup>166</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=105)
Countries and setting	Conducted in India; Setting: single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: history, physical examination, X-rays KUB, ultrasonography, etc.
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	≥8 years; single, unilateral ureteral stone of ≤10mm; distal defined as the segment from the lower border of the sacroiliac joint to the vesico-ureteric junction
Exclusion criteria	previous surgery on the ipsilateral ureter; bilateral ureteric stones; multiple stones; solitary kidney; urinary tract infection; moderate or severe hydronephrosis; contraindications for non-steroidal anti-inflammatory drugs; known allergy to Tamsulosin or Alfuzosin; renal insufficiency; currently on alpha-blocker therapy; pregnant or lactating women
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Nifedipine group: 32.74 (9.58), Alfuzosin group: 30.82 (7.85), control group: 33.06 (8.76). Gender (M:F): 68/37. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=35) Intervention 1: Calcium channel blockers - Nifedipine. Nifedipine 30mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 50mg every 12 hours for 1 week, Diclofenac 75mg injection as needed and Tramadol 100mg injection for persistent pain. Indirectness: No indirectness  (n=35) Intervention 2: Alpha blockers - Alfuzosin. Alfuzosin 10mg daily. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 50mg every 12 hours for 1 week, Diclofenac 75mg injection as needed and Tramadol 100mg injection for persistent pain. Indirectness: No indirectness
	(n=35) Intervention 3: Pain management only - NSAIDs. Diclofenac 50mg every 12 hours for 1 week. Duration up to 4 weeks. Concurrent medication/care: Diclofenac 75mg injection as needed and Tramadol

jection for persistent pain. Indirectness: No indirectness
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# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NIFEDIPINE versus NSAIDS

Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hospital readmissions due to uncontrollable pain at 4 weeks ; Group 1: 11/35, Group 2: 27/35

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group 1 Number missing:

## Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate at 4 weeks; Group 1: 21/35, Group 2: 7/35

Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 3: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: duration of stone expulsion at 4 weeks; Group 1: mean 12 days (SD 6.69); n=35, Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group 1 Number missing:

# Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: episodes of pain at 4 weeks; Group 1: mean 2.91 days (SD 1.01); n=35, Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group 1 Number missing: ; Group 2 Number missing:

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALFUZOSIN versus NIFEDIPINE

# Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hospital readmissions due to uncontrollable pain at 4 weeks ; Group 1: 5/35, Group 2: 11/35

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate at 4 weeks; Group 1: 30/35, Group 2: 21/35
Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group 1 Number missing: ; Group 2 Number missing:

Protocol outcome 3: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: duration of stone expulsion at 4 weeks; Group 1: mean 12 days (SD 6.67); n=35, Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group 1 Number missing: ; Group 2 Number missing:

Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: episodes of pain at 4 weeks; Group 1: mean 1.8 days (SD 0.83); n=35, Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group 1 Number missing: ; Group 2 Number missing:

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALFUZOSIN versus NSAIDS

Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hospital readmissions due to uncontrollable pain at 4 weeks; Group 1: 5/35, Group 2: 27/35

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group 1 Number missing:

Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate at 4 weeks; Group 1: 30/35, Group 2: 7/35

Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group 1 Number missing: ; Group 2 Number missing:

Protocol outcome 3: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: duration of stone expulsion at 4 weeks; Group 1: mean 12 days (SD 6.67); n=35, Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,

Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: episodes of pain at 4 weeks; Group 1: mean 1.8 days (SD 0.83); n=35, Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or location (left/right); Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes	not reported	by the
study		

Quality of life; Analgesic use; Adverse events

Study	Sayed 2008 <sup>168</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=90)
Countries and setting	Conducted in Egypt; Setting: Urology department, single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: physical evaluation, urinalysis, abdominal ultrasound, KUB X-ray etc.
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	>18 years; radiopaque stones 5-10mm in diameter in the distal ureter
Exclusion criteria	urinary tract infection; severe hydronephrosis; multiple stones; pregnancy; lactation; hypotension; ureteral stricture or a history of spontaneous stone passage; concomitant treatment with anaphalytic drugs, beta-blockers or calcium antagonists; desire by patient for immediate stone removal
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): standard therapy group: 37.1 (9.8), Tamsulosin group: 39.3 (10.6). Gender (M:F): 69/21. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones

Indirectness of population	No indirectness
Interventions	(n=45) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: hydration (at least 2 L of water daily) and Diclofenac 100mg injection on demand. Indirectness: No indirectness  (n=45) Intervention 2: No treatment. no treatment. Duration up to 4 weeks. Concurrent medication/care: hydration (at least 2 L of water daily) and Diclofenac 100mg injection on demand. Indirectness: No indirectness
Funding	Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NO TREATMENT

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion rate at 4 weeks; Group 1: 40/45, Group 2: 23/45 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to expulsion at 4 weeks; Group 1: mean 7.32 days (SD 0.78); n=45, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension or other side effects requiring cessation of treatment at 4 weeks; Group 1: 0/45, Group 2: 0/45

Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of pain episodes at 4 weeks; Group 1: mean 1.53 (SD 0.25); n=45, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: : Group 2 Number missing:

Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: number of analgesic vials at 4 weeks; Group 1: mean 0.14 (SD 0.5); n=45, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Hospitalisation/ Use of healthcare services
study	

Study	Sen 2017 <sup>171</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=66)
Countries and setting	Conducted in Turkey; Setting: Not reported
Line of therapy	Unclear
Duration of study	Follow up (post intervention): 3 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Direct urinary system graphy, urinary system ultrasonography, and intravenous pyelography or unenhanced computed tomography
Stratum	Adults (≥16 years), ureteric stone <1 cm:
Subgroup analysis within study	Not applicable
Inclusion criteria	People with distal ureteral stones that were radio-opaque and ≤10mm
Exclusion criteria	Study discontinuation criteria included hypersensitivity to the agents used, advanced hydronephrosis, persistent pain despite proper and adequate analgesic use, urinary tract infection, low blood pressure
Recruitment/selection of patients	Not reported
Age, gender and ethnicity	Age - Mean (SD): Alpha blockers group 33.7 (10.4); control group 33 (11.3). Gender (M:F): Define. Ethnicity: Not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not stated / Unclear 3. Obesity /skin-to-stone distance: Not stated / Unclear 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not stated / Unclear 6. Uteric stone: Lower ureteric stones
Indirectness of population	No indirectness

Interventions	(n=47) Intervention 1: Alpha blockers - Doxazosin. Doxazosin: 25 participants received 4mg and 22 participants received 8mg. Duration 3 weeks. Concurrent medication/care: Diclofenac 100mg oral and daily 1500-2000 cc hydration . Indirectness: No indirectness  (n=19) Intervention 2: Pain management only - NSAIDs. No treatment. Duration 3 weeks. Concurrent medication/care: Diclofenac 100mg oral and daily 1500-2000 cc hydration . Indirectness: No indirectness
Funding	Funding not stated

## RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: DOXAZOSIN versus NSAIDS

## Protocol outcome 1: Stone passage at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Stone expulsion at 3 weeks; Group 1: 33/47, Group 2: 5/19
Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 2: Time to stone passage at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Stone expulsion rate at 3 weeks; Group 1: mean 13.51 days (SD 4.09); n=47, Group 2: mean 19.6 days (SD 4.2); n=19

Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Hypotension at 3 weeks; Group 1: 3/47, Group 2: 0/19
Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 4: Pain intensity at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Number of pain episodes at 3 weeks; Group 1: mean 0.64 (SD 0.33); n=47, Group 2: mean 1.3 (SD 0.5); n=19

Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	
study	

Quality of life at Define; Analgesic use at Define; Hospitalisation/ Use of healthcare services at Define

Study	Singh 2011 <sup>179</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=120)
Countries and setting	Conducted in India; Setting: outpatient department , single centre
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 3 months
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: KUB x-ray and ultrasonography of the KUB region
Stratum	Adults (≥16 years), ureteric stone <1 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	18-70 years; symptomatic, unilateral and solitary upper (between the peli-ureteral junction and sacroiliac joint) ureteral calculi 6-15mm in major axis
Exclusion criteria	active urinary tract infection; fever; acute renal failure; chronic renal failure; history of urinary tract surgery or endoscopic treatment; uncorrected distal obstruction; severe hydronephrosis; pregnancy; concomitant treatment with alpha-blockers, calcium channel blockers or steroids; morbid obesity; history of previous failed SWL
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 32.2 (12.22), control group: 36 (13.78). Gender (M:F): Define. Ethnicity: not stated
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Upper ureteric stones
Indirectness of population	Serious indirectness: included stones < and > 10mm, results reported separately for primary outcome (stone clearance)
Interventions	(n=59) Intervention 1: Alpha blockers and SWL. Tamsulosin 0.4mg daily beginning just before the session of SWL, SWL repeated every 3 weeks for incomplete fragmented calculus. Duration up to 3 months. Concurrent medication/care: advice to drink 2.5L of fluid daily and Diclofenac on demand. Indirectness: No indirectness; Indirectness comment: NA  (n=58) Intervention 2: Surgery - SWL. SWL repeated every 3 weeks for incomplete fragmented calculus up to 3 sessions. Duration up to 3 months. Concurrent medication/care: advice to drink 2.5L of fluid daily and Diclofenac on demand. Indirectness: No indirectness; Indirectness comment: NA

Renal and ureteric stones: CONSULTATION Medical expulsive therapy

Funding

Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND SWL versus SWL

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone clearance (6-10mm) at 3 months; Group 1: 28/30, Group 2: 27/30 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone clearance (11-15mm) at 3 months; Group 1: 26/29, Group 2: 23/28 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 3 months; Group 1: mean 26.78 days (SD 11.96); n=59, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 3: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: visual analogue pain scale at 3 months; Group 1: mean 24.92 days (SD 7.57); n=59, Group 2: mean 41.81 days (SD 17.24); n=58; visual analogue pain scale 0-100 Top=High is poor outcome Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the studv

Quality of life; Adverse events; Analgesic use; Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone clearance at 4 weeks; Group 1: 52/60, Group 2: 42/59
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 4 weeks; Group 1: mean 12.9 days (SD 7.5); n=60, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 3: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dose of analgesic at 4 weeks; Group 1: mean 65.83 mg (SD 48.26); n=60, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Adverse events; Pain intensity; Hospitalisation/ Use of healthcare services
study	

Study (subsidiary papers)	Spontaneous Urinary Stone Passage Enabled by Drugs (SUSPEND) trial: Pickard 2015 <sup>146</sup> (Pickard 2015 <sup>145</sup> )
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=1167)
Countries and setting	Conducted in United Kingdom; Setting: 24 hospitals
Line of therapy	1st line
Duration of study	Intervention + follow up: 12 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: non-contrast CT KUB
Stratum	Adults (≥16 years), ureteric stone <1 cm: upper middle and lower ureteral stones included, analysed as subgroups for primary outcome (stone passage)
Subgroup analysis within study	Not applicable

Inclusion criteria	presenting acutely with ureteric colic; ≥ 18 years to ≤ 65 years; stone confirmed by non-contrast CT KUB;
mousion chema	stone within any segment of the ureter; unilateral ureteric stone; largest dimension of the stone ≤ 10 mm; female participants willing to use two of the listed methods of contraception prior to taking any trial medication until at least 28 days after receiving the last dose of trial medication, who were post-menopausal or who had undergone permanent sterilisation; capable of giving written informed consent, which includes compliance with the requirements of the trial
Exclusion criteria	those requiring immediate intervention; sepsis; estimated glomerular filtration rate less than 30mL/min; already taking or unable to take alpha-blocker or calcium channel stabiliser; pregnancy; breastfeeding; women intending to become pregnant during study period; asymptomatic incidentally found ureteric stone; stone not previously confirmed by CT KUB; kidney stone without presence of ureteric stone; multiple stones within one ureter
Recruitment/selection of patients	consecutive patients meeting inclusion/exclusion criteria at participating sites during the recruitment period
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 43.1 (11.5), Nifedipine group: 42.3 (11), placebo group: 42.8 (12.3) . Gender (M:F): 931/219. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Not stated / Unclear (mixed).
Indirectness of population	Serious indirectness: population includes upper, middle and lower ureteric stones
Interventions	(n=391) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: standard care - analgesics, antiemetics, advice on adequate fluid intake and resumption of normal activity. Indirectness: No indirectness  (n=387) Intervention 2: Calcium channel blockers - Nifedipine. Nifedipine 30mg daily. Duration up to 4
	weeks. Concurrent medication/care: standard care - analgesics, antiemetics and advice on adequate fluid intake and resumption of normal activity. Indirectness: No indirectness
	(n=389) Intervention 3: Placebo. Placebo. Duration up to 4 weeks. Concurrent medication/care: standard care - analgesics, antiemetics and advice on adequate fluid intake and resumption of normal activity. Indirectness: No indirectness
Funding	Academic or government funding (UK National Institute for Health Research Health Technology Assessment Programme)

#### Protocol outcome 3: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (upper ureteric stones) at 4 weeks; Group 1: 62/88, Group 2: 58/92

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: similar age, sex, stone size, stone location, history of previous stone, duration of pain, pain score, analgesic use, antibiotic use, SF-36 physical and mental score between groups; Group 1 Number missing; Group 2 Number missing:

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (middle ureteric stones) at 4 weeks; Group 1: 29/41, Group 2: 32/40

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: similar age, sex, stone size, stone location, history of previous stone, duration of pain, pain score, analgesic use, antibiotic use, SF-36 physical and mental score between groups; Group 1 Number missing; Group 2 Number missina:

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (lower ureteric stones) at 4 weeks; Group 1: 216/249, Group 2: 214/247

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: similar age, sex, stone size, stone location, history of previous stone, duration of pain, pain score, analgesic use, antibiotic use, SF-36 physical and mental score between groups; Group 1 Number missing: ; Group 2 Number missing:

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus PLACEBO

Protocol outcome 3: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (upper ureteric stones) at 4 weeks; Group 1: 62/88, Group 2: 65/89

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: similar age, sex, stone size, stone location, history of previous stone, duration of pain, pain score, analgesic use, antibiotic use, SF-36 physical and mental score between groups; Group 1 Number missing: ; Group 2 Number missina:

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (middle ureteric stones) at 4 weeks; Group 1: 29/41, Group 2: 36/44

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: similar age, sex, stone size, stone location, history of previous stone, duration of pain, pain score, analgesic use, antibiotic use, SF-36 physical and mental score between groups; Group 1 Number missing: ; Group 2 Number missing:

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (lower ureteric stones) at 4 weeks; Group 1: 216/249, Group 2: 202/246

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: similar age, sex, stone size, stone location, history of previous stone, duration of pain, pain score, analgesic use, antibiotic use, SF-36 physical and mental score between groups; Group 1 Number missing: ; Group 2 Number missing:

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NIFEDIPINE versus PLACEBO

Protocol outcome 3: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (upper ureteric stones) at 4 weeks ; Group 1: 58/92, Group 2: 65/89

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: similar age, sex, stone size, stone location, history of previous stone, duration of pain, pain score, analgesic use, antibiotic use, SF-36 physical and mental score between groups; Group 1 Number missing: ; Group 2 Number missing:

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (middle ureteric stones) at 4 weeks ; Group 1: 32/40, Group 2: 36/44

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: similar age, sex, stone size, stone location, history of previous stone, duration of pain, pain score, analgesic use, antibiotic use, SF-36 physical and mental score between groups; Group 1 Number missing: Group 2 Number missing:

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (lower ureteric stones) at 4 weeks ; Group 1: 214/247, Group 2: 202/246

Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: similar age, sex, stone size, stone location, history of previous stone, duration of pain, pain score, analgesic use, antibiotic use, SF-36 physical and mental score between groups; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes	not reported	by t	he
study			

Study	Su 2016 <sup>186</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=272)
Countries and setting	Conducted in Taiwan; Setting: single centre
Line of therapy	1st line
Duration of study	Intervention time: 2 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: non-enhanced computed tomography
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	radiopaque distal ureteral stones <10mm
Exclusion criteria	urinary tract infections; high grade hydronephrosis; diabetes; peptic ulcers; history of hypersensitivity to alpha-blockers; pregnancy or nursing; history of spontaneous stone expulsion; hypotension; systolic blood pressure <110mmHg
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 50.74 (10.08), Silodosin group: 51.58 (8.27), placebo group: 52.16 (9.2). Gender (M:F): 122/82. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=76) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 2 weeks. Concurrent medication/care: Ketorolac 10mg three times daily, Buprenorphine 0.2mg on demand and encouragement to drink a minimum of 2 L of water daily. Indirectness: No indirectness
	(n=79) Intervention 2: Alpha blockers - Silodosin. Silodosin 8mg daily. Duration up to 2 weeks. Concurrent medication/care: Ketorolac 10mg three times daily, Buprenorphine 0.2mg on demand and encouragement to drink a minimum of 2 L of water daily. Indirectness: No indirectness
	(n=82) Intervention 3: Placebo. Placebo. Duration up to 2 weeks. Concurrent medication/care: Ketorolac 10mg three times daily, Buprenorphine 0.2mg on demand and encouragement to drink a minimum of 2 L of water daily . Indirectness: No indirectness

Renal and ureteric stones: CONSULTATION Medical expulsive therapy

CONSULTATION

Funding

Funding not stated

## RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus PLACEBO

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 2 weeks; Group 1: 40/47, Group 2: 29/49
Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 29; Group 2 Number missing: 33

## Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 2 weeks; Group 1: mean 6.28 days (SD 2.41); n=47, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 29; Group 2 Number missing: 33

# Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: adverse effects at 2 weeks; Group 1: 1/47, Group 2: 0/49; Comments: adverse effect not reported

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: Serious indirectness, Comments: adverse effect not reported; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 29; Group 2 Number missing: 33

# Protocol outcome 4: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Buprenorphine consumption at 2 weeks; Group 1: mean 0.36 mg (SD 0.19); n=47, Risk of bias: All domain Very high, Selection Low, Blinding High, Incomplete outcome data High, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 29; Group 2 Number missing: 33
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Ketorolac consumption at 2 weeks; Group 1: mean 230.87 mg (SD 114.69); n=47, Risk of bias: All domain Very high, Selection Low, Blinding High, Incomplete outcome data High, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 29; Group 2 Number missing: 33

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: SILODOSIN versus PLACEBO

# Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 2 weeks; Group 1: 38/48, Group 2: 29/49
Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - Low,

CONSULTATION

Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 29; Group 2 Number missing: 33

## Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 2 weeks; Group 1: mean 6.03 days (SD 2.72); n=47, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 29; Group 2 Number missing: 33

## Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: adverse effects at 2 weeks; Group 1: 6/48, Group 2: 0/49; Comments: adverse effects: transient hypotension, asthenia, syncope and retrograde ejaculation

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - High, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 29; Group 2 Number missing: 33

## Protocol outcome 4: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Buprenorphine consumption at 2 weeks; Group 1: mean 0.37 mg (SD 0.19); n=48, Risk of bias: All domain Very high, Selection Low, Blinding High, Incomplete outcome data High, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 29; Group 2 Number missing: 33
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Ketorolac consumption at 2 weeks; Group 1: mean 221.56 (SD 94.22); n=47, Risk of bias: All domain Very high, Selection Low, Blinding High, Incomplete outcome data High, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 29; Group 2 Number missing: 33

Protocol outcomes not reported by the study

Quality of life; Pain intensity; Hospitalisation/ Use of healthcare services

Study	Sun 2009 <sup>188</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=60)
Countries and setting	Conducted in China; Setting: Departments of Urology and Pharmacy, single centre
Line of therapy	1st line
Duration of study	Intervention time: 2 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: urinary system ultrasonography and KUB
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	18-65 years; unilateral distal (below the lower border of the sacroiliac joint) ureteral stones
Exclusion criteria	multiple stones; severe incarcerated stones; history of distal ureteral surgery or spontaneous stone expulsion; renal colic more than 24 hours in duration; urinary tract infection; severe hydronephrosis; voiding dysfunction; hypotension; cardiovascular and cerebrovascular diseases; hepatic and renal dysfunction; pregnancy; diabetes; ulcer disease; hypersensitivity to Naftopidil; receiving treatment with cardiovascular drugs, alpha-adrenergic receptor antagonists or calcium antagonists
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): watchful waiting group: 37.8 (10.2), Naftopidil group: 38.2 (12.6). Gender (M:F): 50/10. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=30) Intervention 1: Alpha blockers - Naftopidil. Naftopidil 50mg daily. Duration up to 2 weeks. Concurrent medication/care: instruction to drink a minimum of 2 L of water daily and Indomethacin suppository to control acute episodes of ureteral colic if present. Indirectness: No indirectness  (n=30) Intervention 2: No treatment - Watch and wait. Watchful waiting. Duration up to 2 weeks. Concurrent medication/care: instruction to drink a minimum of 2 L of water daily and Indomethacin suppository used to
Funding	control acute episodes of ureteral colic if present. Indirectness: No indirectness  Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NAFTOPIDIL versus WATCH AND WAIT

## Protocol outcome 1: Hospitalisation/ Use of healthcare services

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hospitalisation at 2 weeks; Group 1: 0/30, Group 2: 0/30 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details; no significant difference in age, sex or stone size; Group 1 Number missing; ; Group 2 Number missing:

## Protocol outcome 2: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion at 2 weeks; Group 1: 27/30, Group 2: 8/30 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness and fatigue at 2 weeks; Group 1: 2/30, Group 2: 0/30 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: significant ureteral colic at 2 weeks; Group 1: 0/30, Group 2: 0/30 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details; no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Analgesic use; Time to stone passage

Study	Sur 2015 <sup>189</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=239)
Countries and setting	Conducted in USA; Setting: 27 centres
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: KUB radiograph and/or non-contrast helical computed tomography
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not stratified but pre-specified: stone location
Inclusion criteria	≥18 years; unilateral calculus ≥4mm and ≤10mm in any location of the ureter
Exclusion criteria	multiple ureteral calculi; solitary kidney; refractory renal colic; nonopaque calculus; severe hydronephrosis
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Silodosin group: 47 (13), placebo 47 (15). Gender (M:F): 152/80. Ethnicity: white 210/232
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Not stated / Unclear
Indirectness of population	No indirectness
Interventions	(n=119) Intervention 1: Alpha blockers - Silodosin. Silodosin 8mg. Duration up to 4 weeks. Concurrent medication/care: Oxycodone 5mg to provide analgesia for renal colic and us concomitant pre-enrolment medications that would not confound study results. Indirectness: No indirectness  (n=120) Intervention 2: Placebo. Placebo. Duration up to 4 weeks. Concurrent medication/care: Oxycodone 5mg to provide analgesia for renal colic and use of other concomitant pre-enrolment medications that would not confound study results. Indirectness: No indirectness
Funding	Study funded by industry (Actavis Inc.)

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: SILODOSIN versus PLACEBO

Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (distal) at 4 weeks; Group 1: 36/52, Group 2: 27/59
Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover

- Low; Indirectness of outcome: No indirectness; Baseline details; no significant difference in age, sex, ethnicity, weight, height, BMI, stone size or location; Group 1 Number missing:; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (middle) at 4 weeks; Group 1: 8/20, Group 2: 10/21 Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, ethnicity, weight, height, BMI, stone size or location; Group 1 Number missing:; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage (proximal) at 4 weeks; Group 1: 16/43, Group 2: 15/37 Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, ethnicity, weight, height, BMI, stone size or location; Group 1 Number missing:; Group 2 Number missing:

#### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: retrograde ejaculation at 4 weeks; Group 1: 11/72, Group 2: 1/80 Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness: Baseline details: no significant difference in age, sex, ethnicity, weight, height, BMI, stone size or location; Group 1 Number missing:; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: dizziness at 4 weeks; Group 1: 8/119, Group 2: 2/120 Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, ethnicity, weight, height, BMI, stone size or location; Group 1 Number missing:; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; headache at 4 weeks; Group 1: 4/119, Group 2: 0/120 Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, ethnicity, weight, height, BMI, stone size or location; Group 1 Number missing; Group 2 Number missing;

Protocol outcomes not reported by the study	Quality of life; Time to stone passage; Pain intensity; Analgesic use; Hospitalisation/ Use of healthcare services
Study	Thapa 2014 <sup>191</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=70)
Countries and setting	Conducted in Nepal; Setting: Surgery outpatient department and emergency department, single centre
Line of therapy	1st line
Duration of study	Intervention time: 3 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: plain X-ray or ultrasound of the KUB

Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	>15 years; symptomatic, unilateral, solitary lower ureteral stones (located below sacroiliac joint) of 5-10mm
Exclusion criteria	urinary tract infection; renal failure; history of urinary surgery or endoscopic treatment; uncorrected distal obstruction; moderate to severe hydronephrosis; deranged renal function or intractable pain that couldn't be managed on outpatient basis; refusal to participate
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Range: 15-63. Gender (M:F): 41/29. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=35) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 3 weeks. Concurrent medication/care: advice to have high fluid intake more than 3 L daily and Diclofenac 50mg 3 times daily for 5 days, then on demand. Indirectness: No indirectness  (n=35) Intervention 2: Pain management only - NSAIDs. Diclofenac 50mg 3 times daily for 5 days, then on demand. Duration up to 3 weeks. Concurrent medication/care: advice to have high fluid intake more than 3 L
Funding	daily. Indirectness: No indirectness  Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS

# Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone clearance at 3 weeks; Group 1: 28/35, Group 2: 21/35 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Time to stone passage; Adverse events; Pain intensity; Analgesic use; Hospitalisation/ Use of
study	healthcare services

CONSULTATION

Funding

Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 2 weeks; Group 1: 26/32, Group 2: 17/31 Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 2 weeks; Group 1: mean 6.3 days (SD 2.4); n=32, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low: Indirectness of outcome: No indirectness: Baseline details: no significant difference in age, sex, stone size or stone location (left/right): Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: adverse events at 2 weeks; Group 1: 1/32, Group 2: 0/31; Comments: adverse event not reported

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: Serious indirectness, Comments: adverse effect not reported; Baseline details: no significant difference in age, sex, stone size or stone location (left/right); Group 1 Number missing:; Group 2 Number missing:

# Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; colic episodes at 2 weeks; Group 1; mean 1.97 (SD 1.45); n=32, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Ketorolac consumption at 2 weeks; Group 1: mean 231 mg (SD 112); n=32, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Buprenorphine consumption at 2 weeks; Group 1: mean 0.39 mg (SD 0.29); n=32, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right); Group 1 Number missing: : Group 2 Number missing:

# Protocol of

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TERAZOSIN versus NSAIDS

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 2 weeks; Group 1: 25/32, Group 2: 17/31

Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,

Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right);

Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 2 weeks; Group 1: mean 6.3 days (SD 2.1); n=32, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: adverse events at 2 weeks; Group 1: 5/32, Group 2: 0/31; Comments: adverse effects: transient hypotension, asthenia, syncope and palpitations

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: Serious indirectness, Comments: adverse effects: transient hypotension, asthenia, syncope and palpitations; Baseline details: no significant difference in age, sex, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: colic episodes at 2 weeks; Group 1: mean 1.84 (SD 1.51); n=32, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:

# Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Ketorolac consumption at 2 weeks; Group 1: mean 256 mg (SD 112); n=32, Risk of bias: All domain Very high, Selection Low, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right); Group 1 Number missing: ; Group 2 Number missing:
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Buprenorphine consumption at 2 weeks; Group 1: mean 0.36 mg (SD 0.3); n=32, Risk of bias: All domain Very high, Selection Low, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location (left/right); Group 1 Number missing:

Protocol outcomes not reported by the	Quality of life; Hospitalisation/ Use of healthcare services
1 Totocol outcomes not reported by the	Quality of file, Flospitalisation, Osc of Fleatificate Services
study	

Study	Wang 2014 <sup>205</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=94)
Countries and setting	Conducted in China; Setting: Department of urology, single centre
Line of therapy	Adjunctive to current care
Duration of study	Intervention + follow up: 6 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: ultrasound and/or KUB x-ray
Stratum	Adults (≥16 years), ureteric stone 1-2 cm: NA
Subgroup analysis within study	Not applicable: NA
Inclusion criteria	symptomatic stone; 10-15mm in size; located in the proximal ureter (between the ureteropelvic junction and sacroiliac joint); associated with moderate hydroureteronephrosis
Exclusion criteria	fever; leukocytosis; presence of ureteral stricture distal to the stone; co-existence of a kidney stone on ultrasound; proximal stone migration during ureteroscopic Ho:YAG laser lithotripsy
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Other: not reported. Gender (M:F): not reported . Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Upper ureteric stones
Indirectness of population	No indirectness: NA
Interventions	(n=48) Intervention 1: Alpha blockers and URS. Tamsulosin 0.4mg daily after URS. Duration up to 6 weeks. Concurrent medication/care: 2-3L hydration and Diclofenac 75mg on demand. Indirectness: No indirectness; Indirectness comment: NA
	(n=46) Intervention 2: Surgery - URS. URS only. Duration up to 6 weeks. Concurrent medication/care: 2-3L hydration and Diclofenac 75mg on demand. Indirectness: No indirectness; Indirectness comment: NA
Funding	Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: ALPHA BLOCKERS AND URS versus URS

# Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: stone free rate at 6 weeks; Group 1: 44/45, Group 2: 41/44; Comments: numbers calculated from percentages

Risk of bias: All domain - High, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone size or operative time; Group 1 Number missing: 3; Group 2 Number missing: 2

## Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: time of fragment expulsion at 6 weeks; Group 1: mean 7.86 days (SD 4.99); n=45, Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone size or operative time; Group 1 Number missing: 3; Group 2 Number missing: 2

## Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: dizziness at 6 weeks; Group 1: 2/45, Group 2: 0/44 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone size or operative time; Group 1 Number missing: 3; Group 2 Number missing: 2

# Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone 1-2 cm: ureteral colic rate at 6 weeks; Group 1: 2/45, Group 2: 10/44; Comments: numbers calculated from percentages

Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness, Comments: NA; Baseline details: no significant difference in age, sex, stone size or operative time; Group 1 Number missing: 3; Group 2 Number missing: 2

Protocol outcomes not reported by the study

Quality of life; Analgesic use; Hospitalisation/ Use of healthcare services

Study type	
	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=141)
Countries and setting	Conducted in Taiwan; Setting: Department of Surgery, Division of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 2 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: non-enhanced computed tomography
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
nclusion criteria	Radiopaque distal ureteral stones <10mm
Exclusion criteria	Urinary tract infections; high-grade hydronephrosis; diabetes; peptic ulcers; history of hypersensitivity to alpha-1 blockers; pregnancy or nursing; history of spontaneous stone expulsion; hypotension; systolic blood pressure <110mmHg
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Silodosin group: 51.42 (8.68), control group: 51.51 (10.03). Gender (M:F): Define. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
ndirectness of population	No indirectness
nterventions	(n=71) Intervention 1: Alpha blockers - Silodosin. Silodosin 8mg daily. Duration up to 2 weeks. Concurrent medication/care: Ketorolac three times daily, sublingual Buprenorphine 0.2mg on demand and encouragement to drink a minimum of 2 L of water daily. Indirectness: No indirectness
	(n=70) Intervention 2: Placebo. Placebo. Duration up to 2 weeks. Concurrent medication/care: Ketorolac 10mg three times daily, sublingual Buprenorphine 0.2mg on demand and encouragement to drink a minimum of 2 L of water daily. Indirectness: No indirectness
Funding	Funding not stated

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 2 weeks; Group 1: 48/62, Group 2: 33/61
Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 9, Reason: 5 missed primary outcome and 4 withdrew consent; Group 2 Number missing: 9, Reason: 5 missed primary outcome and 4 withdrew consent

## Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 2 weeks; Group 1: mean 6.31 days (SD 2.13); n=62, Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 9, Reason: 5 missed primary outcome and 4 withdrew consent; Group 2 Number missing: 9, Reason: 5 missed primary outcome and 4 withdrew consent

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: adverse effects at 2 weeks; Group 1: 10/62, Group 2: 2/61
Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High,
Crossover - Low; Indirectness of outcome: Serious indirectness, Comments: control group adverse effects not reported, Silodosin group adverse effects:
transient hypotension, asthenia, syncope and palpitations; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone
size; Group 1 Number missing: 9, Reason: 5 missed primary outcome and 4 withdrew consent; Group 2 Number missing: 9, Reason: 5 missed primary
outcome and 4 withdrew consent

# Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Colic episodes at 2 weeks; Group 1: mean 2.39 (SD 1.3); n=62, Risk of bias: All domain - High, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 9, Reason: 5 missed primary outcome and 4 withdrew consent; Group 2 Number missing: 9, Reason: 5 missed primary outcome and 4 withdrew consent

## Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Ketorolac consumption at 2 weeks; Group 1: mean 255.97 mg (SD 112.97); n=62, Risk of bias: All domain High, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone size; Group 1 Number missing: 9, Reason: 5 missed primary outcome and 4 withdrew consent; Group 2 Number missing: 9, Reason: 5 missed primary outcome and 4 withdrew consent
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Buprenorphine consumption at 2 weeks; Group 1: mean 0.47 mg (SD 0.27); n=62, Risk of bias: All domain High, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement High, Crossover Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, BMI, stone location (right/left) or stone

size; Group 1 Number missing: 9, Reason: outcome and 4 withdrew consent	5 missed primary outcome and 4 withdrew consent ; Group 2 Number missing: 9, Reason: 5 missed primary
Protocol outcomes not reported by the study	Quality of life; Hospitalisation/ Use of healthcare services

Chudy	Va 2044207
Study	Ye 2011 <sup>207</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=3189)
Countries and setting	Conducted in China; Setting: outpatient departments from 10 medical centres
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: plain abdominal radiography, urinary system ultrasonography, non-contrast CT and IVU
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	18-50 years; emergency admission for renal colic; radiopaque or radiolucent single distal ureteric stone (juxtavesical or intramural portion) of 4-7mm
Exclusion criteria	fever; urinary tract infection; severe hydronephrosis; renal insufficiency (estimated GFR <60mL/min per 1.73m²); multiple ureteric stones; urethrostenosis; ureteric stricture; gastric ulcer; diabetes; hypotension; pregnancy; current use of alpha-adrenoceptor antagonists, calcium-channel blockers or corticosteroids; history of ipsilateral ureteric surgery, spontaneous stone expulsion or known or suspected allergy to one of the study medications
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Median (range): Tamsulosin group: 30.7 (18-48), Nifedipine group: 34.5 (22-50). Gender (M:F): 1987/1202. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=1596) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: encouragement to maintain a water intake of 2-2.5 L daily, Levofloxacin 0.2g

	twice daily and Diclofenac 50mg suppository on demand . Indirectness: No indirectness
	(n=1593) Intervention 2: Calcium channel blockers - Nifedipine. Nifedipine 10mg 3 times daily. Duration up to 4 weeks. Concurrent medication/care: encouragement to maintain a water intake of 2-2.5 L daily, Levofloxacin 0.2g twice daily and Diclofenac 50mg suppository on demand . Indirectness: No indirectness
Funding	Study funded by industry (Astellas Pharmaceutical )

## RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NIFEDIPINE

## Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 4 weeks; Group 1: 1530/1596, Group 2: 1171/1593 Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: side effect incidence at 4 weeks; Group 1: 90/1596, Group 2: 98/1593 Risk of bias: All domain - Very high, Selection - Low, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: Serious indirectness, Comments: side effects not specified; Baseline details: no significant difference in age, sex, stone size or stone location; Group 1 Number missing:; Group 2 Number missing:

# Protocol outcome 3: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: rate of pain relief therapy at 4 weeks; Group 1: 24/1596, Group 2: 77/1593; Comments: numbers calculated from percentages
- Risk of bias: All domain Very high, Selection Low, Blinding High, Incomplete outcome data Low, Outcome reporting Low, Measurement High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details; no significant difference in age, sex, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the	Quality of life; Time to sto
study	

one passage; Pain intensity; Hospitalisation/ Use of healthcare services

Study	Ye 2018 <sup>208</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=3390)

Countries and setting	Conducted in China; Setting: Not reported
Line of therapy	1st line
Duration of study	Intervention + follow up: 28 days
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: Confirmed by plain abdominal radiography (kidney–ureters–bladder), urinary ultrasonography, and/or non-contrast computed tomography (CT)
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	Adults,18–60 yr; emergency admission for renal colic; presence of a single ureteral stone; a stone in the distal ureter, with a dimension of 4–7 mm; and a unilateral presentation
Exclusion criteria	Fever; urinary tract infections; severe hydronephrosis; renal insufficiency, defined by an estimated glomerular filtration rate of<60 ml/min per 1.73m2; abnormal anatomy, such as a solitary kidney, horseshoe kidney, or a duplex urinary system; urethrostenosis; a history of ureter strictures; diabetes mellitus; hypotension (systolic blood pressure<100 mmHg); known or suspected pregnancy; current use of a-adrenoceptor antagonists or corticosteroids; and a previous history of ipsilateral ureteral surgery, spontaneous stone expulsion, or known or suspected allergy to the study medications
Recruitment/selection of patients	Not reported
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin 40.1 (11.6); placebo 40.7 (12.3). Gender (M:F): 2135/1161. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not stated / Unclear 3. Obesity /skin-to-stone distance: Not stated / Unclear 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not stated / Unclear 6. Uteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=1695) Intervention 1: Alpha blockers - Tamsulosin. Two capsules of tamsulosin 0.2 mg taken daily until spontaneous stone passage, up to a maximum of 28 d or the need for intervention . Duration 28 days. Concurrent medication/care: Participants were instructed to drink 2 I water per day and to collect the urine stone after urine filtration using a sieve. Additionally, the patients were authorized to use pain relief therapy with a 50mg sodium diclofenac suppository on demand. Participants were asked to stop taking their medication use if stones were passed over the course of treatment. Indirectness: No indirectness
	(n=1695) Intervention 2: Placebo. Placebo, taken daily until spontaneous stone passage, up to a maximum of 28 d or the need for intervention. Duration 28 days. Concurrent medication/care: Participants were instructed to drink 2 I water per day and to collect the urine stone after urine filtration using a sieve. Additionally, the patients were authorized to use pain relief therapy with a 50mg sodium diclofenac suppository on demand. Participants were asked to stop taking their

	medication use if stones were passed over the course of treatment. Indirectness: No indirectness
Funding	Academic or government funding (Supported by health industry special scientific research projects, Ministry of Health of China (201002010). Astellas Pharma supported this study and was involved with preparation of the manuscript.)

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus PLACEBO

## Protocol outcome 1: Stone passage at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Stone expulsion at 28 days; Group 1: 1419/1642, Group 2: 1300/1654 Risk of bias: All domain - Low, Selection - Low, Blinding - Low, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover
- Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 53; Group 2 Number missing: 41

## Protocol outcome 2: Time to stone passage at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Time to stone passage at 28 days; Group 1: mean 148.3 hours (SD 63.2); n=1642, Group 2: mean 248.7 hours (SD 76.6); n=1654
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 53; Group 2 Number missing: 41

## Protocol outcome 3: Adverse events at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Retrograde ejaculation at 28 days; Group 1: 67/1642, Group 2: 48/1654
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover
- Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 53; Group 2 Number missing: 41
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Dizziness at 28 days; Group 1: 52/1642, Group 2: 50/1654
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover
- Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 53; Group 2 Number missing: 41
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Headache at 28 days; Group 1: 41/1642, Group 2: 46/1654
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover - Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 53; Group 2 Number missing: 41

# Protocol outcome 4: Analgesic use at Define

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Rate of pain relief therapy at 28 days; Group 1: 31/1642, Group 2: 155/1654
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover
- Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 53; Group 2 Number missing: 41
- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: Diclofenac dose at 28 days; Group 1: mean 86 mg (SD 32); n=1642, Group 2: mean 263 mg (SD 62); n=1654
- Risk of bias: All domain Low, Selection Low, Blinding Low, Incomplete outcome data Low, Outcome reporting Low, Measurement Low, Crossover

- Low; Indirectness of outcome: No indirectness; Group 1 Number missing: 53; Group 2 Number missing: 41	
Protocol outcomes not reported by the study	Quality of life at Define; Pain intensity at Define; Hospitalisation/ Use of healthcare services at Define

Study	Yilmaz 2005 <sup>210</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=114)
Countries and setting	Conducted in Turkey; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: X-rays of KUB and urinary system ultrasonography
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	18-65 years; radiopaque stones ≤10mm located in the distal tract of the ureter (juxtavesical tract and ureterovesical junction)
Exclusion criteria	urinary system infection; radiolucency stones; severe hydronephrosis; diabetes; ulcer disease; hypotension and having calcium antagonist medication; distal ureter surgery
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): control group: 41.6 (12.01), Tamsulosin group: 40.62 (10.27), Treazosin group: 41.67 (11.41), Doxazosin group: 42.13 (10.46). Gender (M:F): 46/68. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=29) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: symptomatic therapy with Diclofenac 75mg injections on demand and consumption of a minimum of 2 L of water daily. Indirectness: No indirectness
	(n=28) Intervention 2: Alpha blockers - Terazosin. Terazosin 5mg daily. Duration up to 4 weeks. Concurrent medication/care: symptomatic therapy with Diclofenac 75mg injections on demand and consumption of a

	minimum of 2 L of water daily. Indirectness: No indirectness
	(n=29) Intervention 3: Alpha blockers - Doxazosin. Doxazosin 4mg daily. Duration up to 4 weeks. Concurrent medication/care: symptomatic therapy with Diclofenac 75mg injections on demand and consumption of a minimum of 2 L of water daily. Indirectness: No indirectness
	(n=28) Intervention 4: Pain management only - NSAIDs. Symptomatic therapy with Diclofenac 75mg injections on demand. Duration up to 4 weeks. Concurrent medication/care: consumption of a minimum of 2 L of water daily. Indirectness: No indirectness
Funding	Funding not stated

# RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NSAIDS

#### Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage at 4 weeks; Group 1: 23/29, Group 2: 15/28
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to expulsion at 4 weeks; Group 1: mean 6.31 days (SD 0.88); n=29, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension or other side effects requiring cessation of medication at 4 weeks; Group 1: 0/29, Group 2: 0/28

Risk of bias: All domain - ; Indirectness of outcome: No indirectness

### Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: pain episodes at 4 weeks; Group 1: mean 1.72 (SD 0.88); n=29, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

## Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: analgesic requirement at 4 weeks; Group 1: mean 129.31 mg (SD 17.81); n=29, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TERAZOSIN versus NSAIDS

#### Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage at 4 weeks; Group 1: 22/28, Group 2: 15/28
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to expulsion at 4 weeks; Group 1: mean 5.75 days (SD 0.88); n=28, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension or other side effects requiring cessation of medication at 4 weeks ; Group 1: 0/28, Group 2: 0/28

Risk of bias: All domain - ; Indirectness of outcome: No indirectness

#### Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: pain episodes at 4 weeks; Group 1: mean 1.57 (SD 0.23); n=28, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: analgesic requirement at 4 weeks; Group 1: mean 117.85 mg (SD 17.85); n=28, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing: ; Group 2 Number missing:

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: DOXAZOSIN versus NSAIDS

#### Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: spontaneous stone passage at 4 weeks; Group 1: 22/29, Group 2: 15/28 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing:; Group 2 Number missing:

#### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: time to expulsion at 4 weeks; Group 1: mean 5.93 days (SD 0.59); n=29, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing:; Group 2 Number missing:

#### Protocol outcome 3: Adverse events

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: hypotension or other side effects requiring cessation of medication at 4 weeks; Group 1: 0/29, Group 2: 0/28

Risk of bias: All domain - ; Indirectness of outcome: No indirectness

### Protocol outcome 4: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: pain episodes at 4 weeks; Group 1: mean 1.67 (SD 0.17); n=29, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing:; Group 2 Number missing:

### Protocol outcome 5: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: analgesic requirement at 4 weeks; Group 1: mean 118.68 mg (SD 16.21); n=29, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, height, weight, stone size or stone location; Group 1 Number missing:; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life: Hospitalisation/ Use of healthcare services

Mational Institute for Health and Pare Evoellence 2017

Study	Yuksel 2015 <sup>211</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=70)
Countries and setting	Conducted in Turkey; Setting: Department of Urology outpatient clinic, single centre
Line of therapy	1st line
Duration of study	Intervention time: 3 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: urinary system x-ray, urinary system ultrasonography and low-dose abdominal tomography if necessary
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	detection of distal ureteral stone 4-10mm
Exclusion criteria	age <18 or >65 years; multiple stones; grade 3 or 4 hydronephrosis; solitary or transplanted kidney; urinary tract infection; recurrent and persistent renal colic in reaction to analgesic administration; renal failure; allergic reaction to NSAID or alpha-blocker; hypotension; current intake of alpha-blockers, calcium channel blockers or steroids
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Silodosin group: 35.31 (11.55), control group: 35.23 (11.2). Gender (M:F): 39/31. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Not stated / Unclear 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=35) Intervention 1: Alpha blockers - Silodosin. Silodosin 4mg daily. Duration up to 3 weeks. Concurrent medication/care: Diclofenac 75mg daily as necessary, advice to remain active and drink at least 2 L of water daily. Indirectness: No indirectness
	(n=35) Intervention 2: Pain management only - NSAIDs. Diclofenac 75mg daily as necessary. Duration up to 3 weeks. Concurrent medication/care: advice to remain active and drink at least 2 L of water daily. Indirectness: No indirectness
Funding	Funding not stated
RESULTS (NUMBERS ANALYSED) AND R	ISK OF BIAS FOR COMPARISON: SILODOSIN versus NSAIDS

#### Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; stone expulsion at 3 weeks; Group 1: 32/35, Group 2: 25/35 Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

Renal and ureteric stones: Medical expulsive therapy

CONSULTATION

#### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone expulsion duration at 3 weeks; Group 1: mean 8.03 days (SD 4.99); n=35, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: : Group 2 Number missing:

#### Protocol outcome 3: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; renal colic episodes at 3 weeks; Group 1; mean 1.17 (SD 1.44); n=35, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 4: Analgesic use

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm; analgesic dosage at 3 weeks; Group 1; mean 113.57 mg (SD 130.38); n=35, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Adverse events; Hospitalisation/ Use of healthcare services

Study	Zhang 2009 <sup>214</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=314)
Countries and setting	Conducted in China; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 4 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: plain abdominal X-rays, urinary ultrasonography and helical computed tomography
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	distal ureteral stones
Exclusion criteria	history of urinary system stone; previous surgery on urinary tract; multiple stones; nonopaque stones; urinary tract infection; severe hydronephrosis; solitary kidney; diabetes; peptic ulcers; hypotension or hypertension treated with alpha-adrenoceptor blocker or calcium-antagonists; severe obesity; kidney failure; pregnancy
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Tamsulosin group: 34.6 (11.4), Nifedipine group: 36.3 (9.7). Gender (M:F): 199/94. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones
Indirectness of population	No indirectness
Interventions	(n=102) Intervention 1: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 4 weeks. Concurrent medication/care: 2.5 L hydration daily, Levofloxacin 0.1g twice daily for the first 7 days and Diclofenac 75mg injection daily if needed. Indirectness: No indirectness
	(n=97) Intervention 2: Calcium channel blockers - Nifedipine. Nifedipine 30mg 3 times daily. Duration up to 4 weeks. Concurrent medication/care: 2.5 L hydration daily, Levofloxacin 0.1g twice daily for the first 7 days and Diclofenac 75mg injection daily if needed. Indirectness: No indirectness
Funding	Funding not stated
RESULTS (NUMBERS ANALYSED) AND R	ISK OF BIAS FOR COMPARISON: TAMSULOSIN versus NIFEDIPINE

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: stone free rate at 4 weeks; Group 1: 75/102, Group 2: 66/97
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex or stone size; Group 1 Number missing:
; Group 2 Number missing:

Renal and ureteric stones: Medical expulsive therapy

Protocol outcomes not reported by the	Quality of life; Time to stone passage; Adverse events; Pain intensity; Analgesic use; Hospitalisation/ Use of
study	healthcare services

Study	Zhou 2011 <sup>216</sup>
Study type	RCT (Patient randomised; Parallel)
Number of studies (number of participants)	1 (n=131)
Countries and setting	Conducted in China; Setting: Department of Urology, single centre
Line of therapy	1st line
Duration of study	Intervention time: 2 weeks
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: abdominal, ultrasonography and plain abdominal X-ray (kidney-ureter-bladder, IVU or unenhanced CT)
Stratum	Adults (≥16 years), ureteric stone <1 cm
Subgroup analysis within study	Not applicable
Inclusion criteria	distal ureteral stones ≤9mm to >4mm
Exclusion criteria	multiple stones; severe incarcerated stones; history of distal ureteral surgery; history of stone expulsion; renal colic for more than 24 hours; urinary tract infection; severe hydronephrosis; voiding dysfunction; hypotension; cardiovascular and cerebrovascular diseases; hepatic and renal dysfunction; pregnancy; diabetes; history of hypersensitivity to Naftopidil; subjects receiving treatment with cardiovascular drugs, alpha receptor antagonists or calcium antagonists
Recruitment/selection of patients	not reported
Age, gender and ethnicity	Age - Mean (SD): Naftopidil group: 33.73 (8.84), Tamsulosin group: 34.42 (8.64), control group: 34.79 (9.63). Gender (M:F): 79/52. Ethnicity: not reported
Further population details	1. Kidney pole: Not applicable 2. Neuropathic/ cerebral-palsy /immobility: Not applicable 3. Obesity /skin-to-stone distance: Not applicable 4. Pregnant women: Non-pregnant 5. Stone composition/hounsfield units: Not applicable 6. Ureteric stone: Lower ureteric stones

Indirectness of population	No indirectness
Interventions	(n=43) Intervention 1: Alpha blockers - Naftopidil . Naftopidil 10mg daily. Duration up to 2 weeks. Concurrent medication/care: instruction to drink at least 2 L of fluids daily and an Indomethacin suppository recommended for use during pain episodes. Indirectness: No indirectness  (n=45) Intervention 2: Alpha blockers - Tamsulosin. Tamsulosin 0.4mg daily. Duration up to 2 weeks. Concurrent medication/care: instruction to drink at least 2 L of fluids daily and Indomethacin suppository recommended for routine use during pain episodes. Indirectness: No indirectness  (n=43) Intervention 3: No treatment - Watch and wait. Watchful waiting. Duration up to 2 weeks. Concurrent medication/care: instruction to drink at least 2 L of fluids daily and Indomethacin suppository recommended
	for routine use during pain episodes. Indirectness: No indirectness
Funding	Funding not stated

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: NAFTOPIDIL versus WATCH AND WAIT

#### Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 2 weeks; Group 1: 31/43, Group 2: 13/43
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone location (left/right) or stone size;
Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 2 weeks; Group 1: mean 7.6 days (SD 2.26); n=43, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone location (left/right) or stone size; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 3: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: pain episodes at 2 weeks; Group 1: mean 1.3 (SD 1.18); n=43, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone location (left/right) or stone size; Group 1 Number missing: ; Group 2 Number missing:

RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: TAMSULOSIN versus WATCH AND WAIT

#### Protocol outcome 1: Stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion rate at 2 weeks; Group 1: 37/45, Group 2: 13/43
Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low,
Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone location (left/right) or stone size;
Group 1 Number missing: ; Group 2 Number missing:

#### Protocol outcome 2: Time to stone passage

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: expulsion time at 2 weeks; Group 1: mean 7.7 days (SD 1.94); n=45, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone location (left/right) or stone size; Group 1 Number missing: ; Group 2 Number missing:

### Protocol outcome 3: Pain intensity

- Actual outcome for Adults (≥16 years), ureteric stone <1 cm: pain episodes at 2 weeks; Group 1: mean 1.2 (SD 1.65); n=45, Risk of bias: All domain - Very high, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness; Baseline details: no significant difference in age, sex, stone location (left/right) or stone size; Group 1 Number missing: ; Group 2 Number missing:

Protocol outcomes not reported by the study

Quality of life; Adverse events; Analgesic use; Hospitalisation/ Use of healthcare services

# **Appendix E: Forest plots**

# 2 E.1 Distal ureteric stones <10mm in adults

# 3 E.1.1 Alpha blockers versus placebo

Figure 2: Time to stone passage (days)

_	Alpha blocker		Placebo			-	Mean Difference	Mean Difference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Al-Ansari 2010	6.4	2.77	50	9.87	5.4	46	1.2%	-3.47 [-5.21, -1.73]	<del></del> -
Ochoa-Gomez 2011	22	6.77	32	23	6.36	33	0.4%	-1.00 [-4.20, 2.20]	<del></del>
Pedro 2008	5.19	4.82	34	8.54	6.99	35	0.5%	-3.35 [-6.18, -0.52]	<del></del>
Su 2016	6.16	2.57	94	9.79	2.7	49	4.4%	-3.63 [-4.55, -2.71]	<del></del>
Ye 2018	6.18	2.63	1642	10.36	3.19	1654	93.5%	-4.18 [-4.38, -3.98]	
Total (95% CI)			1852			1817	100.0%	-4.13 [-4.32, -3.94]	<b>•</b>
Heterogeneity: Chi <sup>2</sup> =	,	,	,,			-10 -5 0 5 10			
Test for overall effect:	∠ = 41.9	6 (P <	0.0000	Favours Alpha blocker Favours Placebo					

Figure 3: Time to stone passage

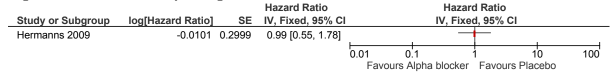


Figure 4: Stone passage

•	Alpha blo	ocker	Place	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI
Abdel-Meguid 2010	61	75	42	75	7.0%	1.45 [1.16, 1.82]	
Agrawal 2009	52	68	12	34	2.4%	2.17 [1.35, 3.48]	
Ahmad 2015	42	49	26	48	5.3%	1.58 [1.19, 2.10]	<del></del>
Al-Ansari 2010	41	50	28	46	5.8%	1.35 [1.03, 1.76]	<del></del>
Furyk 2016	140	161	127	155	13.1%	1.06 [0.97, 1.17]	<del>-</del>
Hermanns 2009	39	45	40	45	10.1%	0.97 [0.84, 1.14]	+
Ochoa-Gomez 2011	22	32	23	33	4.5%	0.99 [0.71, 1.36]	<del></del>
Pedro 2008	25	34	27	35	5.7%	0.95 [0.73, 1.25]	<del></del>
Pickard 2015	216	249	202	246	14.0%	1.06 [0.98, 1.14]	<del> -</del>
Su 2016	78	95	29	49	6.3%	1.39 [1.08, 1.78]	<del></del>
Sur 2015	36	52	27	59	4.3%	1.51 [1.09, 2.11]	<del></del>
Wang 2016	48	62	33	61	5.8%	1.43 [1.10, 1.87]	_ <del>-</del>
Ye 2018	1419	1642	1300	1654	15.6%	1.10 [1.07, 1.13]	•
Total (95% CI)		2614		2540	100.0%	1.19 [1.09, 1.29]	<b>♦</b>
Total events	2219		1916				
Heterogeneity: Tau <sup>2</sup> = 0 Test for overall effect: 2			•	P < 0.00	001); I <sup>2</sup> = 7	71% <del> </del>	1.1 0.2 0.5 1 2 5 10  Favours placebo Favours alpha blocker

Figure 5: Hospitalisation

	Alpha blo	ocker	Placel	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	I M-H, Fixed, 95% CI
Ahmad 2015	0	49	1	48	5.7%	0.33 [0.01, 7.83]	· _
Furyk 2016	20	198	23	195	86.8%	0.86 [0.49, 1.51]	<del></del>
Hermanns 2009	6	45	2	45	7.5%	3.00 [0.64, 14.08]	<del>-  </del>
Total (95% CI)		292		288	100.0%	0.99 [0.59, 1.64]	
Total events	26		26				
Heterogeneity: Chi <sup>2</sup> =	2.69, df = 2	(P = 0.2)	26); I <sup>2</sup> = 26	6%			
Test for overall effect:	Z = 0.05 (P	= 0.96)					0.1 0.2 0.5 1 2 5 10 Favours Alpha blocker Favours Placebo

Figure 6: Use of healthcare services (representation to ED)

	Alpha blocker		Placel	00	Risk Ratio	Risk Ratio					
Study or Subgroup	Events	Total	<b>Events</b>	Total	M-H, Fixed, 95% CI	M-H, I	Fixed, 95% CI	<u> </u>			
Furyk 2016	31	198	35	195	0.87 [0.56, 1.36]	.     .   .   .   .   .   .   .   .   .	<del></del>				
						0.1 0.2 0.5	1 2	5	10		
						Favours Alpha block	er Favours F	Placebo			

Figure 7: Adverse events (unspecified)

	Alpha blo	ocker	Placebo		Placebo Risk Ratio		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	I M-H, Fixed, 95% CI	
Ahmad 2015	0	49	0	48		Not estimable		
Su 2016	7	94	0	49	24.5%	7.89 [0.46, 135.42]	<del></del>	
Wang 2016	10	62	2	61	75.5%	4.92 [1.12, 21.53]		
Total (95% CI)		205		158	100.0%	5.65 [1.50, 21.29]		
Total events	17		2					
Heterogeneity: Chi2 =	0.09, df = 1	(P = 0.7)	$(7); I^2 = 0$					
Test for overall effect:	Z = 2.56 (P	= 0.01)		0.1 0.2 0.5 1 2 5 10 Favours Alpha blocker Favours Placebo				

Figure 8: Adverse events (retrograde ejaculation)

Alpha blocker		ocker	Place	bo		Risk Ratio	Risk Ratio		
Study or Subgroup	Events Total		Events	Total	Weight	M-H, Fixed, 95% C	I M-H, Fix	ed, 95% CI	
Agrawal 2009	3	68	0	34	1.3%	3.55 [0.19, 66.84]			
Al-Ansari 2010	1	32	0	35	0.9%	3.27 [0.14, 77.57]		•	
Hermanns 2009	2	39	0	36	1.0%	4.63 [0.23, 93.20]		-	
Ochoa-Gomez 2011	2	15	0	21	0.8%	6.88 [0.35, 133.64]		-	$\longrightarrow$
Sur 2015	11	72	1	80	1.9%	12.22 [1.62, 92.34]			
Ye 2018	67	1642	48	1654	94.0%	1.41 [0.98, 2.02]		<b>—</b>	
Total (95% CI)		1868		1860	100.0%	1.73 [1.23, 2.43]		•	
Total events	86		49						
Heterogeneity: Chi <sup>2</sup> =	6.47, df = 5	(P = 0.2)	$(26); I^2 = 23$			1 10	400		
Test for overall effect:	Z = 3.17 (P	= 0.002	2)	0.01 0.1 Favours Alpha blocker	1 10 Favours Placebo	100			

Figure 9: Adverse events (dizziness)

_	Alpha bl	ocker	Place	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	I M-H, Fixed, 95% CI
Agrawal 2009	9	68	2	34	4.5%	2.25 [0.51, 9.84]	-
Al-Ansari 2010	2	50	2	46	3.5%	0.92 [0.14, 6.27]	
Hermanns 2009	0	45	1	45	2.5%	0.33 [0.01, 7.97]	<del>-</del>
Ochoa-Gomez 2011	2	32	0	33	0.8%	5.15 [0.26, 103.30]	<del></del>
Pedro 2008	4	34	0	35	0.8%	9.26 [0.52, 165.65]	<del></del>
Sur 2015	8	119	2	120	3.4%	4.03 [0.87, 18.60]	<u>+</u>
Ye 2018	52	1642	50	1654	84.4%	1.05 [0.71, 1.54]	<del>-</del>
Total (95% CI)		1990		1967	100.0%	1.28 [0.92, 1.79]	•
Total events	77		57				
Heterogeneity: Chi <sup>2</sup> =	7.23, df = 6	(P = 0.3)	30); I <sup>2</sup> = 1	7%			
Test for overall effect:							0.1 0.2 0.5 1 2 5 10 Favours Alpha blocker Favours Placebo

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### Figure 10: Adverse events (headache)

	Alpha bl	ocker	Place	bo		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	I M-H, Fixed, 95% CI
Agrawal 2009	8	68	1	34	2.7%	4.00 [0.52, 30.69]	<del>-  </del>
Al-Ansari 2010	2	50	2	46	4.2%	0.92 [0.14, 6.27]	<del></del>
Sur 2015	4	119	0	120	1.0%	9.07 [0.49, 166.72]	<del></del>
Ye 2018	41	1642	46	1654	92.1%	0.90 [0.59, 1.36]	
Total (95% CI)		1879		1854	100.0%	1.06 [0.72, 1.56]	
Total events	55		49				
Heterogeneity: Chi <sup>2</sup> =	4.37, df = 3	P = 0.2	$(22); I^2 = 3$	1%			0.1 0.2 0.5 1 2 5 10
Test for overall effect:	Z = 0.32 (P	9 = 0.75)					Favours Alpha blocker Favours Placebo

Figure 11: Adverse events (hypotension)

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	Favours Alpha bl	ocker	Place	bo		Peto Odds Ratio	Peto Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	Peto, Fixed, 95% CI	Peto, Fixed, 95% CI
Agrawal 2009	0	68	0	34		Not estimable	
Al-Ansari 2010	1	50	0	46	100.0%	6.82 [0.13, 344.93]	
Total (95% CI)		118		80	100.0%	6.82 [0.13, 344.93]	
Total events	1		0				
Heterogeneity: Not app	olicable						0.01 0.1 1 10 100
Test for overall effect: 2	Z = 0.96 (P = 0.34)						Favours Alpha blocker Favours Placebo

Figure 12: Pain intensity (people experiencing pain episodes)

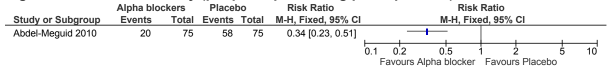


Figure 13: Pain intensity (pain episodes)

	Alpha	block	ers	Pl	acebo	)		Mean Difference		fference			
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI		IV, Fixed	d, 95% CI		
Al-Ansari 2010	1.6	1.3	50	2.3	1.4	46	43.4%	-0.70 [-1.24, -0.16]		-			
Wang 2016	2.39	1.3	62	2.75	1.38	61	56.6%	-0.36 [-0.83, 0.11]		-			
Total (95% CI)			112			107	100.0%	-0.51 [-0.86, -0.15]		<b>♦</b>			
Heterogeneity: Chi <sup>2</sup> = Test for overall effect:				I <sup>2</sup> = 0%					-10 -	1 5 (Alpha blocker	) Favours Pla	5 cebo	10

Figure 14: Pain intensity (pain score >0) at 1 week

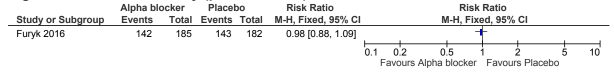


Figure 15: Pain intensity (pain score >0) at 2 weeks

	Alpha blo	ocker	Placel	bo	Risk Ratio		Ris	k Ratio		
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI		M-H, Fi	xed, 95%	CI	
Furyk 2016	60	176	58	177	1.04 [0.77, 1.40]			+		
						0.1 0.2	0.5	1	2 5	10
						Favours	Alpha blocker	r Favou	rs Placebo	

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Figure 16: Pain intensity (pain score >0) at 3 weeks

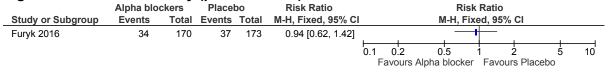


Figure 17: Pain intensity (pain score >0) at 4 weeks

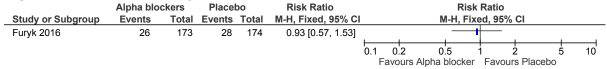


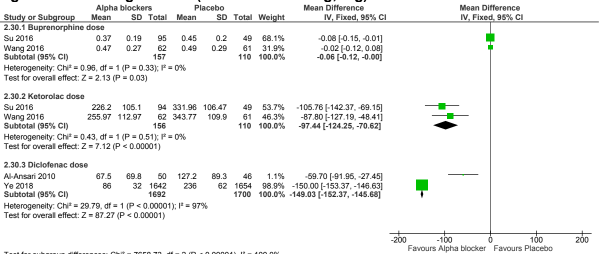
Figure 18: Analgesic use (number of people)

	Alpha blo	ckers	Place	bo		Risk Ratio	Risk	Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Rand	dom, 95% CI		
Ahmad 2015	9	49	19	48	44.1%	0.46 [0.23, 0.92]				
Ye 2018	31	1642	155	1654	55.9%	0.20 [0.14, 0.29]	_			
Total (95% CI)		1691		1702	100.0%	0.29 [0.13, 0.66]				
Total events	40		174							
Heterogeneity: Tau <sup>2</sup> = Test for overall effect:				0.03); I²	= 78%		0.1 0.2 0.5 Favours Alpha blocker	1 2 Favours Place	5 ebo	10

Figure 19: Analgesic use (number of times)

	Alpha	block	ers	PI	acebo			Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
Al-Ansari 2010	0.9	0.93	50	1.8	1.3	46	98.8%	-0.90 [-1.36, -0.44]	
Pedro 2008	8.63	8.58	34	9.41	9.08	35	1.2%	-0.78 [-4.95, 3.39]	
Total (95% CI)			84			81	100.0%	-0.90 [-1.35, -0.45]	<b>◆</b>
Heterogeneity: Chi <sup>2</sup> = Test for overall effect:				I <sup>2</sup> = 0%					-10 -5 0 5 10 Favours Alpha blocker Favours Placebo

Figure 20: Analgesic use (mean dose of drug; mg)



Test for subgroup differences: Chi² = 7658.73, df = 2 (P < 0.00001),  $I^2$  = 100.0%

# 2 E.1.2 Alpha blockers versus no treatment (pain management only)

Figure 21: Time to stone passage (days)

Bajwa 2013		Alpha	block			treatme			Mean Difference	Mean Difference
Alizadeh 2014  3.7 5.07 50 4.7 8.03 46 5.0% -1.00 [-3.71, 1.71]  Bajwa 2013  15.7 3.72 30 20.93 3.43 30 6.0% -5.23 [-7.04, -3.42]  Bayraktar 2017  9.3 5.8 60 8.7 6.4 64 5.6% 0.60 [-1.55, 2.75]  De Sio 2006  4.4 2.1 50 7.5 1.8 46 6.9% -3.10 [-3.88, -2.32]  Itoh 2011  9.29 5.91 55 13.4 5.9 56 5.6% -4.11 [-6.31, -1.91]  Lojanapiwat 2008  10.03 6.87 50 23 0.0001  25 5.9% -12.97 [-14.87, -11.07]  Lv 2014  7.85 2.21 70 10.65 2.92 33 6.7% -2.80 [-3.92, -1.68]  Mohseni 2006  3.18 2.5 32 5.86 2.67 32 6.5% -2.68 [-3.92, -1.68]  Rahim 2012  13.3 6.31 45 19.18 4.66 45 5.5% -2.68 [-3.95, -1.41]  Rahim 2012  13.3 6.31 45 19.18 4.66 45 5.5% -2.68 [-3.95, -1.41]  Sayed 2008  7.32 0.78 45 12.59 9.46 35 3.8% -0.29 [-4.12, 3.54]  Sayed 2008  7.32 0.78 45 12.53 2.12 45 7.0% -5.21 [-5.87, -4.55]  Sen 2017  13.51 4.09 47 19.6 4.2 19 5.5% -6.09 [-8.31, -3.87]  Wang 2008  6.3 2.26 64 10.1 3 31 6.6% -3.80 [-4.99, -2.61]  Yilmaz 2005  6 0.83 86 10.54 2.12 28 6.9% -4.54 [-5.34, -3.74]  Yuksel 2015  8.03 4.99 35 12.91 6.14 35 5.1% -4.88 [-7.50, -2.26]  To 2011  7.65 2.1 88 9.4 2.48 43 6.9% -1.75 [-2.61, -0.89]	Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Bajwa 2013	Ahmed 2010	7.896	7.21	59	13.9	6.99	28	4.5%	-6.00 [-9.18, -2.83]	<del></del>
Bayraktar 2017 9.3 5.8 60 8.7 6.4 64 5.6% 0.60 [-1.55, 2.75] De Sio 2006 4.4 2.1 50 7.5 1.8 46 6.9% -3.10 [-3.88, -2.32]	Alizadeh 2014	3.7	5.07	50	4.7	8.03	46	5.0%	-1.00 [-3.71, 1.71]	<del>+</del>
De Sio 2006	Bajwa 2013	15.7	3.72	30	20.93	3.43	30	6.0%	-5.23 [-7.04, -3.42]	<del></del>
Itoh 2011 9.29 5.91 55 13.4 5.9 56 5.6% -4.11 [-6.31, -1.91]	Bayraktar 2017	9.3	5.8	60	8.7	6.4	64	5.6%	0.60 [-1.55, 2.75]	<del></del>
Lojanapiwat 2008 10.03 6.87 50 23 0.0001 25 5.9% -12.97 [-14.87, -11.07]	De Sio 2006	4.4	2.1	50	7.5	1.8	46	6.9%	-3.10 [-3.88, -2.32]	<del>-</del>
Lv 2014 7.85 2.21 70 10.65 2.92 33 6.7% -2.80 [-3.92, -1.68] ——  Mohseni 2006 3.18 2.5 32 5.86 2.67 32 6.5% -2.68 [-3.95, -1.41] ——  Rahim 2012 13.3 6.31 45 19.18 4.66 45 5.5% -5.88 [-8.17, -3.59] ——  Sameer 2014 12 6.67 35 12.29 9.46 35 3.8% -0.29 [-4.12, 3.54] ——  Sayed 2008 7.32 0.78 45 12.53 2.12 45 7.0% -5.21 [-5.87, -4.55] ——  Sen 2017 13.51 4.09 47 19.6 4.2 19 5.5% -6.09 [-8.31, -3.87] ——  Wang 2008 6.3 2.26 64 10.1 3 31 6.6% -3.80 [-4.99, -2.61] ——  Yilmaz 2005 6 0.83 86 10.54 2.12 28 6.9% -4.54 [-5.34, -3.74] ——  Yuksel 2015 8.03 4.99 35 12.91 6.14 35 5.1% -4.88 [-7.50, -2.26] ——  Zhou 2011 7.65 2.1 88 9.4 2.48 43 6.9% -1.75 [-2.61, -0.89] ——	toh 2011	9.29	5.91	55	13.4	5.9	56	5.6%	-4.11 [-6.31, -1.91]	<del></del>
Mohseni 2006 3.18 2.5 32 5.86 2.67 32 6.5% -2.68 [-3.95, -1.41] ————————————————————————————————————	ojanapiwat 2008	10.03	6.87	50	23	0.0001	25	5.9%	-12.97 [-14.87, -11.07]	<del></del>
Rahim 2012 13.3 6.31 45 19.18 4.66 45 5.5% -5.88 [-8.17, -3.59]  Sameer 2014 12 6.67 35 12.29 9.46 35 3.8% -0.29 [-4.12, 3.54]  Sayed 2008 7.32 0.78 45 12.53 2.12 45 7.0% -5.21 [-5.87, -4.55]  Sen 2017 13.51 4.09 47 19.6 4.2 19 5.5% -6.09 [-8.31, -3.87]  Wang 2008 6.3 2.26 64 10.1 3 31 6.6% -3.80 [-4.99, -2.61]  Yilmaz 2005 6 0.83 86 10.54 2.12 28 6.9% -4.54 [-5.34, -3.74]  Yuksel 2015 8.03 4.99 35 12.91 6.14 35 5.1% -4.88 [-7.50, -2.26]  Zhou 2011 7.65 2.1 88 9.4 2.48 43 6.9% -1.75 [-2.61, -0.89]	_v 2014	7.85	2.21	70	10.65	2.92	33	6.7%	-2.80 [-3.92, -1.68]	<del>-</del>
Sameer 2014 12 6.67 35 12.29 9.46 35 3.8% -0.29 [-4.12, 3.54] Sayed 2008 7.32 0.78 45 12.53 2.12 45 7.0% -5.21 [-5.87, -4.55] Sen 2017 13.51 4.09 47 19.6 4.2 19 5.5% -6.09 [-8.31, -3.87] Wang 2008 6.3 2.26 64 10.1 3 31 6.6% -3.80 [-4.99, -2.61] Yilmaz 2005 6 0.83 86 10.54 2.12 28 6.9% -4.54 [-5.34, -3.74] Yuksel 2015 8.03 4.99 35 12.91 6.14 35 5.1% -4.88 [-7.50, -2.26] Zhou 2011 7.65 2.1 88 9.4 2.48 43 6.9% -1.75 [-2.61, -0.89]	Mohseni 2006	3.18	2.5	32	5.86	2.67	32	6.5%	-2.68 [-3.95, -1.41]	<del>-</del>
Sayed 2008 7.32 0.78 45 12.53 2.12 45 7.0% -5.21 [-5.87, -4.55] ———————————————————————————————————	Rahim 2012	13.3	6.31	45	19.18	4.66	45	5.5%	-5.88 [-8.17, -3.59]	<del></del>
Sen 2017     13.51     4.09     47     19.6     4.2     19     5.5%     -6.09 [-8.31, -3.87]       Wang 2008     6.3     2.26     64     10.1     3     31     6.6%     -3.80 [-4.99, -2.61]        Yilmaz 2005     6     0.83     86     10.54     2.12     28     6.9%     -4.54 [-5.34, -3.74]        Yuksel 2015     8.03     4.99     35     12.91     6.14     35     5.1%     -4.88 [-7.50, -2.26]       Zhou 2011     7.65     2.1     88     9.4     2.48     43     6.9%     -1.75 [-2.61, -0.89]	Sameer 2014	12	6.67	35	12.29	9.46	35	3.8%	-0.29 [-4.12, 3.54]	<del></del>
Wang 2008 6.3 2.26 64 10.1 3 31 6.6% -3.80 [-4.99, -2.61] Think 2005 6 0.83 86 10.54 2.12 28 6.9% -4.54 [-5.34, -3.74] Think 2005 8.03 4.99 35 12.91 6.14 35 5.1% -4.88 [-7.50, -2.26] Think 2011 7.65 2.1 88 9.4 2.48 43 6.9% -1.75 [-2.61, -0.89]	Sayed 2008	7.32	0.78	45	12.53	2.12	45	7.0%	-5.21 [-5.87, -4.55]	<del>-</del>
Yilmaz 2005 6 0.83 86 10.54 2.12 28 6.9% -4.54 [-5.34, -3.74] ————————————————————————————————————	Sen 2017	13.51	4.09	47	19.6	4.2	19	5.5%	-6.09 [-8.31, -3.87]	<del></del>
Yilmaz 2005 6 0.83 86 10.54 2.12 28 6.9% -4.54 [-5.34, -3.74] ————————————————————————————————————	Nang 2008	6.3	2.26	64	10.1	3	31	6.6%	-3.80 [-4.99, -2.61]	-
Zhou 2011 7.65 2.1 88 9.4 2.48 43 6.9% -1.75 [-2.61, -0.89]	Yilmaz 2005	6	0.83	86	10.54	2.12	28	6.9%	-4.54 [-5.34, -3.74]	<del>-</del>
	Yuksel 2015	8.03	4.99	35	12.91	6.14	35	5.1%	-4.88 [-7.50, -2.26]	
Total (95% CI) 901 641 100.0% -4.14 [-5.23, -3.04]	Zhou 2011	7.65	2.1	88	9.4	2.48	43	6.9%	-1.75 [-2.61, -0.89]	~
	Γotal (95% CI)			901			641	100.0%	-4.14 [-5.23, -3.04]	•

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Figure 22: Stone passage

i igaic zz.	Otonic	pus	Juge				
	Alpha blo	ckers	No treat	ment		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% CI
Ahmed 2010	48	59	14	28	3.3%	1.63 [1.10, 2.40]	_ <del></del>
Aldemir 2011	25	31	11	29	2.6%	2.13 [1.29, 3.49]	
Alizadeh 2014	41	50	30	46	4.7%	1.26 [0.98, 1.61]	<del></del>
Arrabal-Martin 2010	30	35	19	35	3.8%	1.58 [1.13, 2.20]	<del></del>
Bajwa 2013	23	30	11	30	2.5%	2.09 [1.26, 3.48]	
Balci 2014	19	25	9	25	2.2%	2.11 [1.20, 3.72]	
Bayraktar 2017	49	60	18	64	3.2%	2.90 [1.93, 4.37]	<del></del>
Chau 2011	19	22	14	20	3.8%	1.23 [0.89, 1.72]	+-
De Sio 2006	45	50	27	46	4.5%	1.53 [1.18, 1.99]	<del></del>
El Said 2015	15	28	7	26	1.6%	1.99 [0.97, 4.09]	<del></del>
Erturhan 2007	22	30	12	30	2.6%	1.83 [1.12, 2.99]	<del></del>
Ferre 2009	27	38	24	39	3.9%	1.15 [0.84, 1.59]	+-
Ibrahim 2013	46	52	12	23	3.2%	1.70 [1.13, 2.54]	<del></del>
Islam 2012	27	32	13	28	3.1%	1.82 [1.19, 2.78]	<del></del>
Itoh 2011	40	55	31	56	4.3%	1.31 [0.99, 1.75]	<del></del>
Kupeli 2004	8	15	3	15	0.8%	2.67 [0.87, 8.15]	<del>                                     </del>
Lojanapiwat 2008	27	50	1	25	0.3%	13.50 [1.95, 93.69]	
Lv 2014	62	70	20	33	4.3%	1.46 [1.10, 1.95]	<del></del>
Mohseni 2006	29	32	20	32	4.2%	1.45 [1.08, 1.94]	<del></del>
Mustafa 2016	51	60	32	60	4.5%	1.59 [1.23, 2.07]	_ <del>-</del>
Rahim 2012	37	45	22	45	3.9%	1.68 [1.21, 2.34]	<del></del>
Resim 2005	26	30	22	30	4.6%	1.18 [0.91, 1.53]	+-
Sameer 2014	30	35	7	35	1.7%	4.29 [2.18, 8.43]	-
Sayed 2008	40	45	23	45	4.1%	1.74 [1.28, 2.36]	
Sen 2017	33	47	5	19	1.4%	2.67 [1.23, 5.79]	
Sun 2009	27	30	8	30	2.0%	3.38 [1.84, 6.18]	<del></del>
Thapa 2014	28	35	21	35	4.0%	1.33 [0.97, 1.83]	<del></del>
Wang 2008	51	64	17	31	3.7%	1.45 [1.03, 2.05]	-
Yilmaz 2005	67	86	15	28	3.6%	1.45 [1.01, 2.09]	•
Yuksel 2015	32	35	25	35	4.8%	1.28 [1.01, 1.62]	-
Zhou 2011	68	88	13	43	2.8%	2.56 [1.60, 4.08]	
Total (95% CI)		1364		1066	100.0%	1.64 [1.48, 1.82]	•
Total events	1092		506				
Heterogeneity: Tau <sup>2</sup> = Test for overall effect:				< 0.0001	); I <sup>2</sup> = 57%	b	0.1 0.2 0.5 1 2 5 10  Favours no treatment Favours Alpha blocker

Figure 23: Hospitalisation

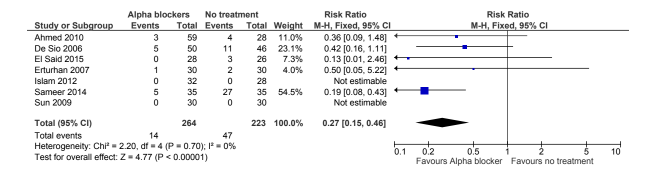


Figure 24: Use of healthcare services (return to ED/primary care visit)

	Alpha blo	ckers	No treat	ment	Risk Ratio			Risk	Ratio		
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fix	ed, 95% (	CI	
Ferre 2009	6	38	8	39	0.77 [0.29, 2.01]			<del> </del>		-	
						0.1	0.2	0.5	1 2	2 5	10
							Favours	Alpha blocker	Favours	no treatment	

Figure 25: Adverse events (unspecified)

	Favours Alpha b	locker	No treati	ment		Peto Odds Ratio	Peto Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	Peto, Fixed, 95% CI	Peto, Fixed, 95% CI
Aldemir 2011	0	31	0	29		Not estimable	
Alizadeh 2014	0	50	0	46		Not estimable	
Arrabal-Martin 2010	0	35	0	35		Not estimable	
El Said 2015	4	28	0	26	43.0%	7.72 [1.03, 58.18]	
Erturhan 2007	0	30	0	30		Not estimable	
Ferre 2009	0	38	0	39		Not estimable	
Sayed 2008	0	45	0	45		Not estimable	
Wang 2008	6	64	0	31	57.0%	4.80 [0.83, 27.71]	<del></del>
Yilmaz 2005	0	86	0	28		Not estimable	
Total (95% CI)		407		309	100.0%	5.89 [1.57, 22.13]	
Total events	10		0				
Heterogeneity: Chi <sup>2</sup> = 0	.12, df = 1 (P = 0.7	3); $I^2 = 0^9$	%				
Test for overall effect: 2	Z = 2.62 (P = 0.009	)					0.01 0.1 1 10 100  Favours Alpha blocker Favours no treatment

Figure 26: Adverse events (dizziness)

_			,	•		•		
	Alpha blo	ckers	No treat	ment		Risk Ratio	Risk Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	CI M-H, Fixed, 95% CI	
Balci 2014	2	25	0	25	3.1%	5.00 [0.25, 99.16]	· ·	$\rightarrow$
Chau 2011	2	33	0	34	3.0%	5.15 [0.26, 103.33]	j	$\rightarrow$
De Sio 2006	1	50	0	46	3.2%	2.76 [0.12, 66.22]	•	$\rightarrow$
Kupeli 2004	1	39	0	39	3.1%	3.00 [0.13, 71.46]	· · · · · · · · · · · · · · · · · · ·	$\rightarrow$
Lv 2014	10	70	8	33	66.4%	0.59 [0.26, 1.35]	<del></del> _	
Resim 2005	5	30	3	30	18.3%	1.67 [0.44, 6.36]	<del></del>	
Sun 2009	2	30	0	30	3.1%	5.00 [0.25, 99.95]	· ·	<b>→</b>
Total (95% CI)		277		237	100.0%	1.34 [0.74, 2.40]		
Total events	23		11					
Heterogeneity: Chi2 =	6.54, df = 6 (	P = 0.37	); I <sup>2</sup> = 8%				1 1 1 1	<del></del>
Test for overall effect:	Z = 0.96 (P	= 0.33)					0.1 0.2 0.5 1 2 5 Favours Alpha blocker Favours no treatment	10

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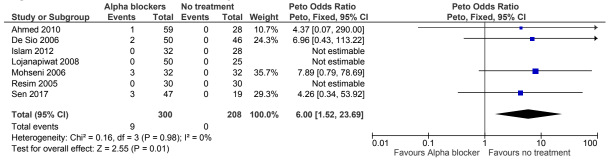


Figure 28: Adverse events (retrograde ejaculation)

					J	,	<i>)</i>
_	Alpha blo	ckers	No treat	ment	_	Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	I M-H, Fixed, 95% CI
Ahmed 2010	2	59	0	28	23.8%	2.42 [0.12, 48.73]	<del></del>
Lojanapiwat 2008	0	35	0	20		Not estimable	
Lv 2014	1	41	0	18	24.3%	1.36 [0.06, 31.81]	<del></del>
Resim 2005	0	22	1	23	51.9%	0.35 [0.01, 8.11]	-
Total (95% CI)		157		89	100.0%	1.09 [0.21, 5.67]	
Total events	3		1				
Heterogeneity: Chi <sup>2</sup> =	0.79, df = 2	P = 0.67	'); I <sup>2</sup> = 0%				
Test for overall effect:	Z = 0.10 (P	= 0.92)	•				0.01 0.1 1 10 100 Favours Alpha blocker Favours no treatment

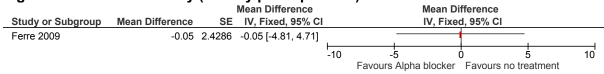
Figure 29: Adverse events (headache)

	Alpha blo	ckers	No treat	ment		Risk Ratio		Ris	k Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	<u> </u>	M-H, Fiz	ced, 95% CI		
Lv 2014	4	70	0	33	14.5%	4.31 [0.24, 77.78]				-	<b>→</b>
Resim 2005	4	30	4	30	85.5%	1.00 [0.28, 3.63]		-			
Total (95% CI)		100		63	100.0%	1.48 [0.47, 4.69]					
Total events	8		4								
Heterogeneity: Chi2 =	0.88, df = 1	P = 0.35	$(1)$ ; $I^2 = 0\%$				<u> </u>	0.2 0.5	<del>                                     </del>	<u> </u>	10
Test for overall effect:	Z = 0.66 (P	= 0.51)					0.1	0.2 0.5 Favours Alpha blocker	Favours no t	reatment	10

Figure 30: Pain intensity (people experiencing pain episodes)

ga. 0 00.			'''' \ P'	oop.	o onp	51.151.1511.19 PG		opioodoo,
	Alpha blo	ckers	No treat	ment		Risk Ratio		Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	I	M-H, Fixed, 95% CI
Aldemir 2011	20	31	23	29	33.1%	0.81 [0.59, 1.12]		<del></del>
Mustafa 2016	36	60	48	60	66.9%	0.75 [0.59, 0.96]		-
Sun 2009	0	30	0	30		Not estimable		
Total (95% CI)		121		119	100.0%	0.77 [0.64, 0.94]		•
Total events	56		71					
Heterogeneity: Chi2 =	0.16, $df = 1$ (	P = 0.69	); $I^2 = 0\%$				-	0.2 0.5 1 2 5 10
Test for overall effect:	Z = 2.63 (P =	= 0.008)					0.1	0.2 0.5 1 2 5 10 Favours Alpha blocker Favours no treatment

Figure 31: Pain intensity (colicky pain episodes)



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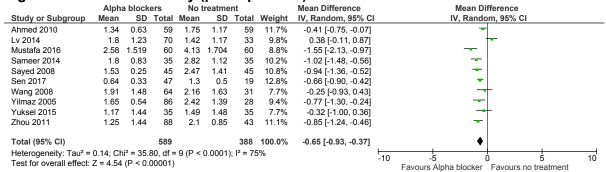


Figure 33: Pain intensity (VAS score) at 3 days

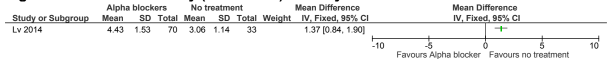


Figure 34: Pain intensity (VAS score) at 7 days

	Alpha blockers		No treatment			Mean Difference	Mean Difference						
Study or Subgroup			Total	IV, Fixed, 95% CI			IV, Fixed	d, 95% CI					
Lv 2014	3.2	1.7	70	1.57	0.5	33	1.63 [1.20, 2.06]				+		
								-10	-5	5 (	)	5	10
									Favours A	Alpha blocker	Favours no tre	eatment	

Figure 35: Analgesic use (number of people using analgesia)

	Alpha blo	ckers	No treat	ment		Risk Ratio		Ris	k Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C		M-H, Fi	xed, 95% CI		
Aldemir 2011	10	31	18	29	32.1%	0.52 [0.29, 0.93]			-		
Arrabal-Martin 2010	9	35	21	35	36.2%	0.43 [0.23, 0.80]					
De Sio 2006	5	50	17	46	30.5%	0.27 [0.11, 0.67]	_				
Lojanapiwat 2008	1	50	0	25	1.1%	1.53 [0.06, 36.25]	<b>←</b>		<u> </u>		$\longrightarrow$
Total (95% CI)		166		135	100.0%	0.42 [0.29, 0.62]					
Total events	25		56								
Heterogeneity: Chi <sup>2</sup> = 2	2.04, df = 3 (	P = 0.57	); I <sup>2</sup> = 0%				<u> </u>	000	<del>                                     </del>	<del></del> _	
Test for overall effect:	Z = 4.37 (P	< 0.0001	)				0.1	0.2 0.5 Favours Alpha blocke	r Favours no treatn	nent	10

Figure 36: Analgesic use (number of times)

_	Alpha blockers Mean SD Total 1.48 2.15 50		No t	reatme	ent		Mean Difference	Mean Difference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Alizadeh 2014	1.48	2.15	50	2.3	4.31	46	21.4%	-0.82 [-2.20, 0.56]	<del></del>
Bayraktar 2017	1.3	0.4	60	1.4	0.4	64	28.0%	-0.10 [-0.24, 0.04]	•
Itoh 2011	0.3	0.9	55	1.5	3.1	56	25.2%	-1.20 [-2.05, -0.35]	
Sayed 2008	0.14	0.5	45	2.78	2.7	45	25.4%	-2.64 [-3.44, -1.84]	
Total (95% CI)			210			211	100.0%	-1.18 [-2.49, 0.13]	•
0 ,	erogeneity: $Tau^2 = 1.59$ ; $Chi^2 = 43.59$ , $df = 3$ (P < 0.00001); for overall effect: $Z = 1.76$ (P = 0.08)				01); I² =	93%	-10	0 -5 0 5 10 Favours Alpha blocker Favours no treatment	

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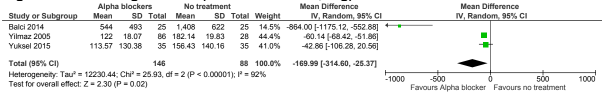


Figure 38: Analgesic use (days)

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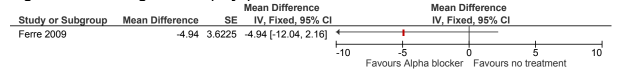


Figure 39: Analgesic use (Pethidine dose, mg)

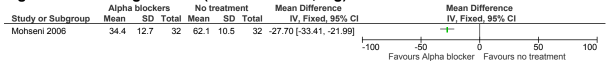


Figure 40: Analgesic use (Ketorolac dose, mg)

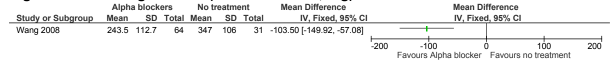


Figure 41: Analgesic use (Buprenorphine dose, mg)

		Alpha	block	ers	No to	reatme	ent	Mean Difference			Mean Di	fference		
	Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI			IV, Fixed	d, 95% CI		
Ī	Wang 2008	0.38	0.3	34	0.39	0.33	31	-0.01 [-0.16, 0.14]						
									-10	-	5 (	)	5	10
										Favours	Alpha blocker	Favours no t	reatment	

# 5 E.1.3 Calcium channel blockers versus placebo

Figure 42: Stone passage

J	Calcium channel blo	ckers	Place	bo	Risk Ratio			Risk	Ratio		
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fixe	ed, 95%	CI	
Pickard 2015	214	247	202	246	1.06 [0.98, 1.14]				+		
						0.1	0.2	0.5	1 2	2 5	10
							Favo	urs Placebo	Favour	rs CC blocke	er

# 8 E.1.4 Calcium channel blockers versus no treatment (pain management only)

#### Figure 43: Time to stone passage (days)

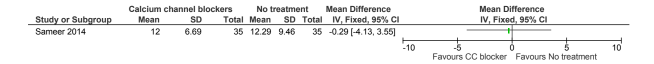


Figure 44: Stone passage

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Figure 45: Hospitalisation

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_	Calcium channel blo	ckers	No treat	ment		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Islam 2012	0	31	0	28		Not estimable	
Sameer 2014	11	35	27	35	100.0%	0.41 [0.24, 0.69]	
Total (95% CI)		66		63	100.0%	0.41 [0.24, 0.69]	
Total events	11		27				
Heterogeneity: Not applic	able					ŀ	0.1 0.2 0.5 1 2 5 10
Test for overall effect: Z =	= 3.37 (P = 0.0007)					'	Favours CC blocker Favours No treatment

Figure 46: Adverse events (hypotension)

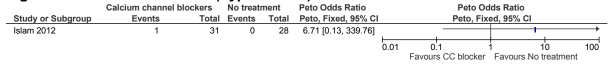


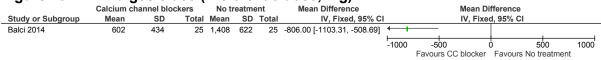
Figure 47: Adverse events (dizziness)

9		,,,,,,,		,						
	Calcium channel b	lockers	No treat	ment	Risk Difference		Risk	Difference	ce	
Study or Subgroup	Events	Events         Total           0         25			M-H, Fixed, 95% CI		M-H,	Fixed, 95°	% CI	
Balci 2014	0	0	25	0.00 [-0.07, 0.07]			+			
						-1	-0.5	or Favo	0.5	1

Figure 48: Pain intensity (pain episodes)

J	-		<i>)</i> (1: :				,					
	Calcium cl	hannel blo	ckers	No ti	reatme	ent	Mean Difference		Mea	n Differen	ce	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV,	Fixed, 95%	CI	
Sameer 2014	2.91	1.01	35	2.82	1.12	35	0.09 [-0.41, 0.59]			+		
								-10	-5	0	5	10
									Favours CC bloc	ker Favo	urs No treatmen	nt

Figure 49: Analgesic use (Diclofenac dose, mg)



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#### Alpha blockers versus Calcium channel blockers 2 E.1.5

Figure 50: Time to stone passage (days)

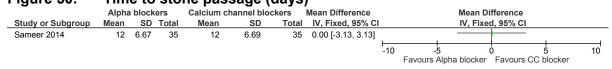


Figure 51: Stone passage

	Alpha blo	ckers	Calcium channel b	lockers		Risk Ratio		Risk Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl		M-H, Random, 9	5% CI	
Balci 2014	19	25	16	25	8.7%	1.19 [0.82, 1.71]		<del> </del>	_	
Gandhi 2013	51	64	32	58	12.2%	1.44 [1.11, 1.88]			_	
Islam 2012	27	32	22	31	11.9%	1.19 [0.91, 1.56]		+		
Pickard 2015	216	249	214	247	20.0%	1.00 [0.93, 1.07]		+		
Sameer 2014	30	35	21	35	10.7%	1.43 [1.06, 1.93]		<del></del>	_	
Ye 2011	1530	1596	1171	1593	20.8%	1.30 [1.26, 1.35]				
Zhang 2009	75	102	66	97	15.7%	1.08 [0.90, 1.29]		<del> -</del>		
Total (95% CI)		2103		2086	100.0%	1.20 [1.05, 1.39]		•		
Total events	1948		1542							
Heterogeneity: Tau <sup>2</sup> =	0.02; Chi <sup>2</sup> =	51.71, d	$f = 6 (P < 0.00001); I^2$	<sup>2</sup> = 88%			0.1 0.2	0.5	<del></del>	10
Test for overall effect:	Z = 2.58 (P =	= 0.010)	. ,						ours Alpha block	

Figure 52: Hospitalisation

	Alpha blo	ckers	Calcium channel	blockers		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Islam 2012	0	32	0	31		Not estimable	<u></u>
Sameer 2014	5	35	11	35	100.0%	0.45 [0.18, 1.17]	
Total (95% CI)		67		66	100.0%	0.45 [0.18, 1.17]	
Total events	5		11				
Heterogeneity: Not ap	plicable					<u> </u>	4 00 05 4 0 5 40
Test for overall effect:	Z = 1.63 (P =	= 0.10)				0.	1 0.2 0.5 1 2 5 10 Favours Alpha blocker Favours CC blocker

Figure 53: Adverse events (headache)

			(	,								
	Alpha blo	ckers	Calcium channel bl	ockers	Risk Ratio			Risk	Ratio			
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fix	ed, 95%	CI		
Gandhi 2013	32	64	25	58	1.16 [0.79, 1.70]				+-			
						0.1	0.2	0.5	1	2	5	10
							Favours	Alpha blocker	Favour	s CC b	locker	

Figure 54: Adverse events (dizziness)

_	Alpha blo	ckers	Calcium channel b	lockers		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Balci 2014	2	25	0	25	13.7%	5.00 [0.25, 99.16]	-
Gandhi 2013	16	64	3	58	86.3%	4.83 [1.48, 15.74]	
Total (95% CI)		89		83	100.0%	4.86 [1.62, 14.56]	
Total events	18		3				
Heterogeneity: Chi <sup>2</sup> = 0 Test for overall effect:			); I <sup>2</sup> = 0%				0.1 0.2 0.5 1 2 5 10  Favours Alpha blocker Favours CC blocker



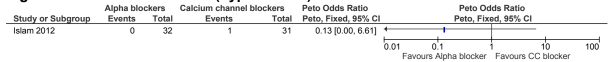


Figure 56: Adverse events (unspecified)

			- d		-,							
	Alpha blo	ckers	Calcium channel b	lockers	Risk Ratio			Ris	k Ratio			
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fix	ked, 95%	CI		
Ye 2011	90	1596	98	1593	0.92 [0.69, 1.21]				+			
						0.1	0.2	0.5	1	2	5	10
							Favours A	Alpha blocker	Favou	rs CC bloc	ker	

Figure 57: Adverse events (flushing)

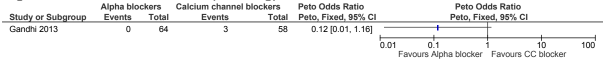


Figure 58: Pain intensity (pain episodes)

_	Alpha	a block	ers	Calcium ch	nannel bloc	kers	Mean Difference			Mean Di	fference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI			IV, Fixed	d, 95% CI	
Sameer 2014	1.8	0.83	35	2.91	1.01	35	-1.11 [-1.54, -0.68]			+		
								-10	-	5 (	) !	5 10
									Favours	Alpha blocker	Favours CC b	locker

Figure 59: Analgesic use (mg)

9 a. 0 00.	,	~.ອ`	,0.0	400 (	່ອ <i>າ</i>							
	Alpha	block	ers	Calcium ch	annel bloc	kers	Mean Difference		IV	lean Dif	ference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		ľ	V, Fixed	I, 95% CI	
Balci 2014	544	493	25	602	434	25	-58.00 [-315.47, 199.47]					
								-500	-250 Favours Alpha b	locker	250 Favours CC bloc	500
									i avouis Aipiia b	JIOCKEI	I avours CC bloc	KCI

Figure 60: Analgesic use (number of people using analgesics)

	Alpha blo	ckers	Calcium channel ble	ockers	Risk Ratio			Risk	Ratio			
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fix	ed, 95%	CI		
Ye 2011	24	1596	77	1593	0.31 [0.20, 0.49]							
						0.1	0.2	0.5	1_ 2	2	5	10
							Favoure 4	Alpha blocker	Favour	CC blocks	ar .	

Figure 61: Analgesic use (number of diclofenac injections)

	Alpha	block	ers	Calcium c	hannel bloo	ckers	Mean Difference			Mean Di	fference		
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI			IV, Fixe	d, 95% CI		
Gandhi 2013	0.42	0.14	64	1.19	0.59	58	-0.77 [-0.93, -0.61]			†			
								-10	-: Favours	5 Alpha blocker	) { Favours CC b	5 Jocker	10

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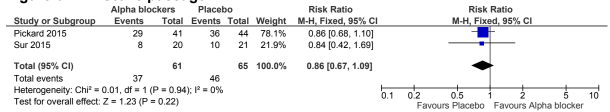
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# 1 E.2 Mid ureteric stones <10mm in adults

# 2 E.2.1 Alpha blockers versus placebo

Figure 62: Stone passage



# 3 E.2.2 Alpha blockers versus no treatment (pain management only)

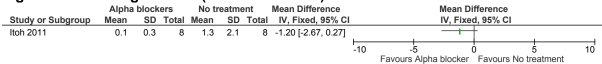
Figure 63: Time to stone passage

_	Favours	Alpha blo	cker	No to	reatme	ent	Mean Difference		Mea	n D	ifference		
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, I	ixe	d, 95% CI		
Itoh 2011	8.67	5.03	8	21	5.03	8	-12.33 [-17.26, -7.40]	+					
								-100	-50		Ó	50	100
									Favours Alpha bloc	ker	Favours No	treatment	

Figure 64: Stone passage



Figure 65: Analgesic use (number of times)



### 6 E.2.3 Calcium channel blockers versus placebo

## Figure 66: Stone passage

	Calcium channel b	locker	Place	bo	Risk Ratio			Ris	k Ratio	1		
Study or Subgroup	Events	Total	<b>Events</b>	Total	M-H, Fixed, 95% CI			M-H, Fi	xed, 95	% CI		
Pickard 2015	32	40	36	44	0.98 [0.79, 1.20]							
						$\vdash$	-		+	$-\!\!\!\!\!-$		
						0.1	0.2	0.5	1	2	5	10
							F	avours placebo	o Favo	ours calc	cium chann	el b

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# 1 E.2.4 Alpha blockers versus Calcium channel blockers

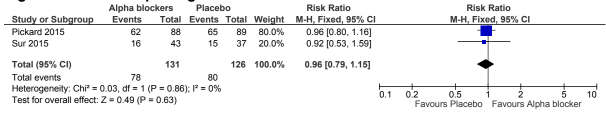
#### Figure 67: Stone passage

	Alpha blo	ckers	Calcium channel b	lockers	Risk Ratio			Ris	k Rati	0		
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, F	xed, 9	5% CI		
Pickard 2015	29	41	32	40	0.88 [0.69, 1.14]				+			
						0.1	0.2	0.5	1	2	5 blooker	10

# 2 E.3 Proximal ureteric stones <10mm in adults

# 3 E.3.1 Alpha blockers versus placebo

# Figure 68: Stone passage



# 5 E.3.2 Alpha blockers versus no treatment (pain management only)

Figure 69: Time to stone passage (days)

	Alpha blockers Mean SD Total		No to	reatme	ent		Mean Difference		Mean Di	fference			
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI		IV, Fixed	d, 95% CI		
Itoh 2011	13.45	13.48	26	18.73	8.66	28	26.5%	-5.28 [-11.37, 0.81]		-			
Lee 2014	14.3	7.9	44	19.6	8.5	35	73.5%	-5.30 [-8.96, -1.64]					
Total (95% CI)			70			63	100.0%	-5.29 [-8.43, -2.16]		<b>*</b>			
Heterogeneity: Chi <sup>2</sup> = Test for overall effect:	,	•	,,	I <sup>2</sup> = 0%					-100	-50 ( Favours Alpha blocker	) Favours No	50 treatment	100

Figure 70: Stone passage

	Alpha blo	ckers	No treatr	nent		Risk Ratio		Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C		M-H, Fixed, 95% CI
Chau 2011	8	11	3	14	6.0%	3.39 [1.17, 9.86]		
Ibrahim 2013	13	22	1	4	3.9%	2.36 [0.42, 13.37]		<del></del>
Itoh 2011	15	26	15	28	33.0%	1.08 [0.67, 1.73]		<del></del>
Lee 2014	40	54	25	54	57.1%	1.60 [1.15, 2.22]		<del>-</del>
Total (95% CI)		113		100	100.0%	1.57 [1.20, 2.03]		•
Total events	76		44					
Heterogeneity: Chi <sup>2</sup> =	4.62, df = 3 (	P = 0.20	); I <sup>2</sup> = 35%					
Test for overall effect:	Z = 3.35 (P	= 0.0008	)				0.1	0.2 0.5 1 2 5 10 Favours No treatment Favours Alpha blocker

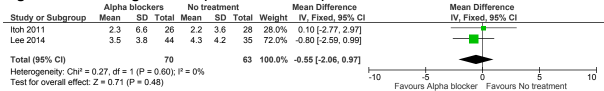
Figure 71: Quality of life (EuroQoL)

•	Alpha	blook	0.00	No 6	reatme	nné ,	Mean Difference		Moon F	Difference	
	Aipiia	DIOCK	613	NOU	eaume	FIIL	Weall Dillerence		IVICALI L	Milerence	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fix	ed, 95% CI	
Lee 2014	5.4	0.6	44	5.5	8.0	35	-0.10 [-0.42, 0.22]			<del>†</del> .	
								-10	-5	0 5	10
									Favours No treatment	Favours Alpha blocker	

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# 2 E.3.3 Calcium channel blockers versus placebo

Figure 73: Stone passage

_	Calcium channel b	locker	Placel	bo	Risk Ratio			Risl	k Ratio		
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fix	ced, 95% C	CI	
Pickard 2015	58	92	65	89	0.86 [0.71, 1.06]			-	+		
						0.1	0.2	0.5	1 2	<u>2</u> 5	10
							Fa	vours placebo	Favours	calcium chai	nnel b

# 3 E.3.4 Alpha blockers versus Calcium channel blockers

Figure 74: Stone passage

	Alpha blo	ckers	Calcium channel b	lockers	Risk Ratio			Risl	Ratio			
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fix	ed, 95%	CI		
Pickard 2015	62	88	58	92	1.12 [0.91, 1.37]				+-			
						0.1	0.2	0.5	1 Eavour	2 s Δlnha hl	5 ookor	10

# 4 E.4 Distal ureteric stones <10mm in children

## 5 E.4.1 Alpha blockers versus placebo

Figure 75: Time to stone passage (days)

	Alpha	block	ers	Pla	aceb	0		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	I IV, Random, 95% CI
Elgalaly 2017	7	1.3	18	10.4	4.7	19	48.7%	-3.40 [-5.60, -1.20]	
Mokhless 2012	8.2	3.2	33	14.5	4.5	28	51.3%	-6.30 [-8.29, -4.31]	<del></del>
Total (95% CI)			51			47	100.0%	-4.89 [-7.73, -2.05]	
Heterogeneity: Tau <sup>2</sup> = Test for overall effect:				1 (P = 0	).06);	I <sup>2</sup> = 73	%		-10 -5 0 5 10
			,						Favours Alpha blocker Favours placebo

Figure 76: Stone passage (4 weeks)

			-3 - ( -		,			
	Alpha blo	ckers	Place	bo		Risk Ratio		Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI		M-H, Fixed, 95% CI
Elgalaly 2017	16	18	14	19	41.2%	1.21 [0.88, 1.65]		+
Mokhless 2012	29	33	18	28	58.8%	1.37 [1.01, 1.85]		<b></b> -
Total (95% CI)		51		47	100.0%	1.30 [1.04, 1.62]		•
Total events	45		32					
Heterogeneity: Chi2 =	0.32, df = 1	P = 0.57	); I <sup>2</sup> = 0%				<u> </u>	02 05 1 2 5 10
Test for overall effect:	Z = 2.33 (P =	= 0.02)					0.1	0.2 0.5 1 2 5 10  Favours placebo Favours Alpha blocker

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	Alpha blo	ckers	No treat	ment	Peto Odds Ratio		Peto O	dds Ratio	
Study or Subgroup	Events	Total	Events	Total	Peto, Fixed, 95% CI		Peto, Fi	xed, 95% CI	
Elgalaly 2017	3	18	0	19	8.82 [0.86, 90.57]			<u> </u>	
						0.01	0.1	1 10	100

Figure 78: Adverse events (headache)

	Alpha blo	ckers	No treat	ment	Risk Ratio			Risk	Ratio	
Study or Subgroup	Events	Total	<b>Events</b>	Total	M-H, Fixed, 95% CI			M-H, Fixe	ed, 95% CI	
Mokhless 2012	1	33	1	28	0.85 [0.06, 12.95]			-		
						0.01	0.	1	1 10	100
							Favours A	Alpha blocker	Favours No treatment	

Figure 79: Adverse events (hypotension)

	Alpha blo	ckers	No treat	ment	Risk Difference			Risk D	ifference	9	
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fix	ed, 95%	CI	
Mokhless 2012	0	33	0	28	0.00 [-0.06, 0.06]				+	1	
						-1	-0	).5	Ó	0.5	1
							Favours	Alpha blocker	Favoui	rs No treatment	

Figure 80: Pain intensity (pain episodes)

	Alpha	block	ers	Pla	aceb	0		Mean Difference	Mean Difference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI	
Elgalaly 2017	2.3	1.4	18	4.7	2.6	19	43.1%	-2.40 [-3.74, -1.06]	<del></del>	
Mokhless 2012	1.4	1.2	33	2.2	1.4	28	56.9%	-0.80 [-1.46, -0.14]	-	
Total (95% CI)			51			47	100.0%	-1.49 [-3.04, 0.06]	•	
Heterogeneity: Tau <sup>2</sup> = Test for overall effect:				1 (P = 0	).04);	l <sup>2</sup> = 77	%		-10 -5 0 5 Favours Alpha blocker Favours placebo	10

# E.4.2 Alpha blockers versus no treatment (pain management only)

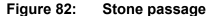
Figure 81: Time to stone passage (days)

	Alpha	block	ers	No ti	reatme	ent		Mean Difference		Mean D	ifference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% C		IV, Rando	om, 95% CI	
Aldaqadossi 2015	7.7	1.9	31	18	1.73	32	50.1%	-10.30 [-11.20, -9.40]				
Aydogdu 2009	5.9	2.1	19	6.1	2.3	20	49.9%	-0.20 [-1.58, 1.18]		ı	•	
Total (95% CI)			50			52	100.0%	-5.26 [-15.16, 4.63]		•	<b>\</b>	
Heterogeneity: Tau <sup>2</sup> = Test for overall effect:				df = 1 (P	0.0	0001);	l <sup>2</sup> = 99%		-100	-50	050	100
rest for overall effect.	2 - 1.04	(1 – 0	.50)							Favours Alpha blocker	Favours No treatment	

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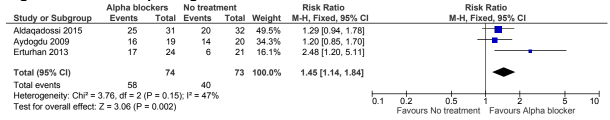


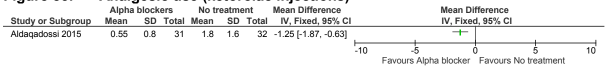
Figure 83: Adverse events (unspecified)

_				•		,	
	Alpha bloo	ckers	No treat	ment		Risk Difference	Risk Difference
Study or Subgroup	Events	Total	<b>Events</b>	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Aldaqadossi 2015	0	31	0	32	61.8%	0.00 [-0.06, 0.06]	-
Aydogdu 2009	0	19	0	20	38.2%	0.00 [-0.09, 0.09]	+
Total (95% CI)		50		52	100.0%	0.00 [-0.05, 0.05]	<b>•</b>
Total events	0		0				
Heterogeneity: Chi2 =	0.00, df = 1 (I)	P = 1.00	); $I^2 = 0\%$				-1 -0.5 0 0.5
Test for overall effect:	Z = 0.00 (P =	= 1.00)					Favours Alpha blocker Favours No treatment

Figure 84: Pain intensity (daily pain episodes)

	Alpha blockers			No to	reatme	ent	Mean Difference	Mean Difference					
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI				d, 95% CI		
Aldaqadossi 2015	1.6	1.6	31	2.5	1.9	32	-0.90 [-1.77, -0.03]	+				1	1
								-10	{	5	Ó	5	10
									Favours	Alpha blocker	Favours No	treatment	

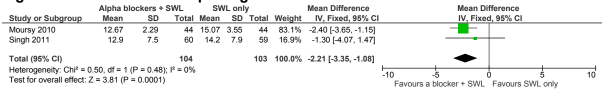
Figure 85: Analgesic use (ketorolac injections)



# 4 E.5 Adjunctive therapy: distal ureteric stones <10mm in adults

# E.5.1 Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only

Figure 86: Time to stone passage



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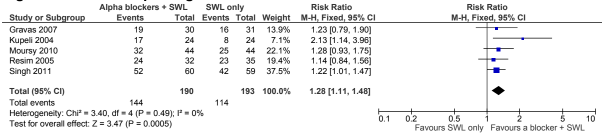


Figure 88: Hospitalisation

	Alpha blockers	+ SWL	SWL o	nly	Risk Ratio			Risk	Ratio		
Study or Subgroup	Events	Total	<b>Events</b>	Total	M-H, Fixed, 95% CI			M-H, Fixe	ed, 95% CI		
Moursy 2010	12	44	19	44	0.63 [0.35, 1.14]				_		
							72	0,5	<del>                                     </del>	<u> </u>	10
						U. I Fav	ours a	blocker + SWL	Favours SW	L only	10

Figure 89: Adverse events (abnormal ejaculation)

_	Alpha blockers -	SWL o	nly		Peto Odds Ratio	Peto Odds Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	Peto, Fixed, 95% CI	Peto, Fixed, 95% CI
Moursy 2010	6	28	0	27	84.5%	8.71 [1.62, 46.76]	
Resim 2005	1	21	0	22	15.5%	7.75 [0.15, 390.96]	-
Total (95% CI)		49		49	100.0%	8.56 [1.83, 40.08]	
Total events	7		0				
Heterogeneity: Chi <sup>2</sup> = 0	0.00, df = 1 (P = 0.9)	$96$ ); $I^2 = 0$	%				0.01 0.1 1 10 100
Test for overall effect:	Z = 2.72 (P = 0.006)	5)					Favours a blocker + SWL Favours SWL only

Figure 90: Adverse events (dizziness)

	Alpha blockers	+ SWL	SWL o	nly		Peto Odds Ratio	Peto Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	Peto, Fixed, 95% Cl	Peto, Fixed, 95% CI
Gravas 2007	2	30	0	31	29.0%	7.91 [0.48, 129.46]	
Kupeli 2004	1	39	0	39	14.8%	7.39 [0.15, 372.38]	
Resim 2005	4	32	0	35	56.2%	8.97 [1.20, 66.79]	
Total (95% CI)		101		105	100.0%	8.40 [1.86, 37.87]	
Total events	7		0				
Heterogeneity: Chi <sup>2</sup> = 0	0.01, df = 2 (P = 1.0	$(0)$ ; $I^2 = 0$	%				0.01 0.1 1 10 100
Test for overall effect:	Z = 2.77 (P = 0.006)	5)					Favours a blocker + SWL Favours SWL only

Figure 91: Adverse events (headache)

9	,	• • • • • •					
_	Alpha blockers	+ SWL	SWL only			Peto Odds Ratio	Peto Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	Peto, Fixed, 95% CI	Peto, Fixed, 95% CI
Moursy 2010	4	44	0	44	37.8%	7.94 [1.08, 58.33]	
Resim 2005	5	32	2	35	62.2%	2.84 [0.60, 13.45]	
Total (95% CI)		76		79	100.0%	4.19 [1.23, 14.28]	
Total events	9		2				
Heterogeneity: Chi <sup>2</sup> = 0	0.63, df = 1 (P = 0.4	43); $I^2 = 0$	%				0.01 0.1 1 10 100
Test for overall effect: 2	Z = 2.29 (P = 0.02)						Favours a blocker + SWL Favours SWL only

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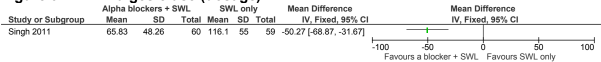




Figure 93: Analgesic use (number of analgesics)

	Alpha blockers + SWL			SW	L on	ly	Mean Difference Mean Difference					
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fixe	d, 95% CI		
Moursy 2010	4.39	2.42	44	6.11	3.1	44	-1.72 [-2.88, -0.56]		_			
								-10	<del> </del> -5	0	5	10
									blocker + SWL	Favours SW	only	

Figure 94: Analgesic use (dosage)



# 3 E.5.2 Alpha blockers as adjunctive therapy to ureteroscopy versus ureteroscopy only

Figure 95: Stone passage

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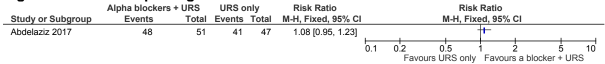


Figure 96: Use of healthcare services (length of hospital stay)

J	Alpha blockers + URS			UR	S on	ly	Mean Difference	Mean Difference				
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fixe	d, 95% CI		
Abdelaziz 2017	1.2	0.6	51	1.7	0.9	47	-0.50 [-0.81, -0.19]		+			
									1			
								<sup>'</sup> -10 -	5 (	b s	5	10
								Favours a b	locker + URS	Favours URS	only	

# 7 E.5.3 Alpha blockers as adjunctive therapy to ureteroscopy versus placebo and ureteroscopy

Figure 97: Stone passage

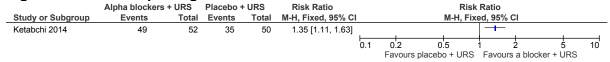


Figure 98: Pain intensity (colic episodes)

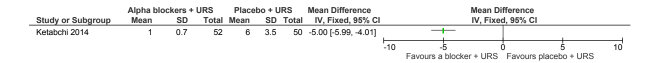
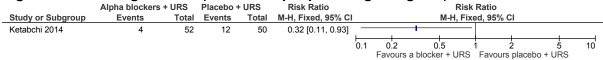


Figure 99: Analgesic use (number of people using analgesia)



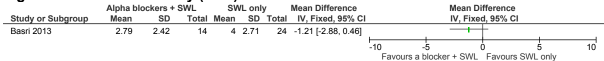
# E.6 Adjunctive therapy: distal ureteric stones 10-20mm in adults

# 4 E.6.1 Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only

Figure 100: Time to stone passage (days)

	Alpha blockers + SWL			SV	/L onl	y	Mean Difference	Mean Difference						
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI			V, Fixed, 95%	Fixed, 95% CI			
Basri 2013	9.86	6.94	14	12.42	9.38	24	-2.56 [-7.78, 2.66]				_			
								-10	-5	<u> </u>	+	10		
									vours a blocker	+ SWL Favo	urs SWL only	10		

Figure 101: Pain intensity (VAS)



# 7 E.7 Adjunctive therapy: mid ureteric stones 10-20mm in adults

# 8 E.7.1 Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only

Figure 102: Time to stone passage (days)

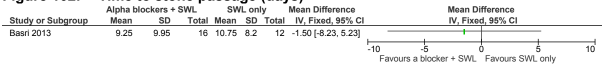


Figure 103: Pain intensity (VAS)

94.0 .00.			, , .	,,,,,,							
	Alpha blo	ockers +	SWL	SW	L only	у	Mean Difference		Mean Di	fference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fixe	d, 95% CI	
Basri 2013	2.38	2.42	16	3	3.91	12	-0.62 [-3.13, 1.89]				
								-10 -	5	5	10
								Favours a h	Nocker + SWI	Favours SWI	only

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# E.8 Adjunctive therapy: proximal ureteric stones <10mm in adults

# E.8.1 Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only

Figure 104: Time to stone passage (days)

	Alpha blockers + SWL		SWL only				Mean Difference	Mean Difference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Agarwal 2009	30.7	19.6	20	39	19.9	20	13.1%	-8.30 [-20.54, 3.94]	<del></del>
Ates 2012	4.14	1.78	35	3.61	2.7	44	35.9%	0.53 [-0.46, 1.52]	•
Cho 2013	9.5	4.8	41	18.6	20.6	43	24.6%	-9.10 [-15.43, -2.77]	<del></del>
Singh 2011	26.78	11.96	59	31.28	18.31	58	26.4%	-4.50 [-10.11, 1.11]	-
Total (95% CI)			155			165	100.0%	-4.32 [-9.85, 1.21]	•
Heterogeneity: Tau <sup>2</sup> = 1 Test for overall effect: 2			if = 3 (P	= 0.004)	); I <sup>2</sup> = 77	7%			-50 -25 0 25 50 Favours a blocker + SWL Favours SWL only

Figure 105: Stone passage

			_									
_	Alpha blockers	Alpha blockers + SWL			Risk Ratio			Risk Ratio				
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI			M-H, Fixed,	95% CI		
Agarwal 2009	19	20	18	20	11.1%	1.06 [0.88, 1.26]			+-			
Ates 2012	33	35	35	44	19.2%	1.19 [1.00, 1.41]			-	-		
Cho 2013	39	41	40	43	24.1%	1.02 [0.92, 1.14]			+			
Eryildirim 2016	20	28	17	26	10.9%	1.09 [0.76, 1.57]			<del></del>	_		
Park 2013	37	44	29	44	17.9%	1.28 [1.00, 1.64]			-	<del>-</del>		
Singh 2011	28	30	27	30	16.7%	1.04 [0.89, 1.21]			+			
Total (95% CI)		198		207	100.0%	1.11 [1.03, 1.21]			<b>•</b>			
Total events	176		166									
Heterogeneity: Chi <sup>2</sup> =	5.25, df = 5 (P = 0.	39); I <sup>2</sup> = 5	%					0.2	<del>,  </del>	<del></del>	<u></u> _	
Test for overall effect:	Z = 2.59 (P = 0.01	0)					0.1		0.5 1 SWL only Fa	vours a blo	ocker + SV	10 //L

Figure 106: Hospitalisation

_	Alpha blockers + SWL			SV	/L only	y	Mean Difference	Mean Difference				
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fixe	d, 95% CI		
Ates 2012	0.51 0.7 35			0.52	0.62	44	-0.01 [-0.31, 0.29]		-	+		
								-10	-5	0 !	5	10
							Favours a	blocker + SWL	Favours SWL	only		

Figure 107: Use of healthcare services (ED visits)

	Alpha b	Alpha blockers + SWL			/L onl	y	Mean Difference		Mean Dif	ference		
Study or Subgrou	up Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fixed	l, 95% CI		
Eryildirim 2016	0.82	0.9	28	1.42	1.07	26	-0.60 [-1.13, -0.07]		+			
								-10 -5	d		5	10
								Favours a blo	ocker + SWL	Favours SWL	only	

Figure 108: Quality of life (EQ5D)

	-,	,	,-		,							
	Alpha blo	ckers +	SWL	SW	/L onl	y	Mean Difference		Me	an Difference	e	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV	, Fixed, 95%	CI	
Eryildirim 2016	0.82	0.11	28	0.78	0.09	26	0.04 [-0.01, 0.09]		ı	+		
								-1	-0.5	Ó	0.5	
									Favours SWL	only Favou	rs a blocker + S	WL

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# Figure 109: Quality of life (EQ5D VAS)

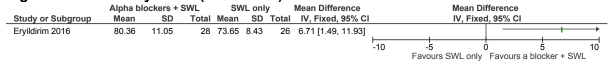


Figure 110: Adverse events (dizziness)

_	Alpha blockers -	+ SWL	SWL o	nly		Peto Odds Ratio	Peto Odds Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	Peto, Fixed, 95% CI	Peto, Fixed, 95% CI	
Cho 2013	2	41	0	43	66.4%	7.95 [0.49, 129.44]		<b>→</b>
Park 2013	1	44	0	44	33.6%	7.39 [0.15, 372.38]	-	<b>→</b>
Total (95% CI)		85		87	100.0%	7.76 [0.80, 75.32]		_
Total events	3		0					
Heterogeneity: Chi <sup>2</sup> = 0	0.00, $df = 1$ ( $P = 0.9$	$(8); I^2 = 0$	%				0.01 0.1 1 10	100
Test for overall effect:	Z = 1.77 (P = 0.08)						Favours a blocker + SWL Favours SWL only	100

Figure 111: Adverse events (retrograde ejaculation)

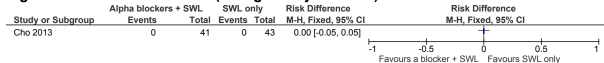


Figure 112: Pain intensity (VAS)

	Alpha blockers + SWL			SI	WL only	1		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Agarwal 2009	2.53	1.79	20	3.83	2.8	20	13.5%	-1.30 [-2.76, 0.16]	<del></del>
Ates 2012	6.89	1.02	35	6.59	1.58	44	21.9%	0.30 [-0.28, 0.88]	+-
Cho 2013	5.33	1.22	41	6.43	1.36	43	22.1%	-1.10 [-1.65, -0.55]	<b>-</b>
Eryildirim 2016	5.86	1.41	28	6.65	1.57	26	19.8%	-0.79 [-1.59, 0.01]	<del></del>
Singh 2011	2.492	0.757	59	4.181	1.724	58	22.7%	-1.69 [-2.17, -1.21]	*
Total (95% CI)			183			191	100.0%	-0.89 [-1.68, -0.10]	•
Heterogeneity: Tau <sup>2</sup> = 0 Test for overall effect: 2			= 4 (P <	0.0001	); I <sup>2</sup> = 86	6%			-10 -5 0 5 10 Favours a blocker + SWL Favours SWL only

Figure 113: Pain intensity (colic episodes)

•			• •				,				
	Alpha blo	ckers +	SWL	SV	/L onl	y	Mean Difference		Mean Di	fference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fixed	d, 95% CI	
Eryildirim 2016	2.54	2.55	28	4.92	3.08	26	-2.38 [-3.89, -0.87]				
								-10 -	5	<u> </u>	10
									olocker + SWL	Favours SWL	only

Figure 114: Analgesic use (dosage)

_	Alpha blockers + SWL			SI	NL only	/	Mean Difference		Mean Di	fference	
Study or Subgroup			Total	Mean	SD	Total	IV, Fixed, 95% CI		IV, Fixed	d, 95% CI	
Eryildirim 2016	242	196.6	28	431.7	246.5	26	-189.70 [-309.20, -70.20]	<del></del>			
								-500 -250		250	500
								Favours a blocker	+ SWL	Favours SWL only	

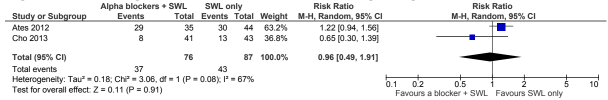
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Figure 115: Analgesic use (number of people using analgesia)



# E.8.2 Alpha blockers as adjunctive therapy to shock wave lithotripsy versus placebo and shock wave lithotripsy

Figure 116: Time to stone passage (days)

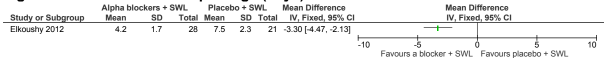


Figure 117: Stone passage

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# E.9 Adjunctive therapy: proximal ureteric stones 10-20mm in adults

# 6 E.9.1 Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only

Figure 118: Time to stone passage (days)

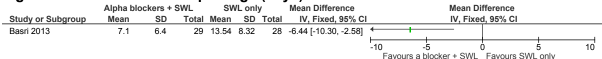
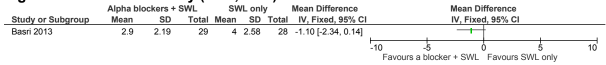


Figure 119: Stone passage

<b>J</b>	Alpha blockers	_	SWL o	nly	Risk Ratio			R	isk Rati	0		
Study or Subgroup	Events	Total	<b>Events</b>	Total	M-H, Fixed, 95% CI			M-H,	Fixed, 9	5% CI		
Singh 2011	26	29	23	28	1.09 [0.88, 1.35]		<del></del>					
						0.1	0.2	0.5	1	2	<del></del>	10
							Fave	oure SM/L o	nly Fa	oure a blo	ckar + SI	Λ/Ι

Figure 120: Pain intensity (VAS, 0-10)



# E.9.2 Alpha blockers as adjunctive therapy to ureteroscopy versus ureteroscopy only

#### Figure 121: Time to stone passage (days)

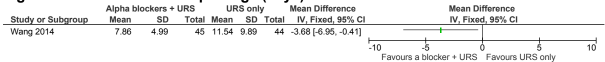


Figure 122: Stone passage

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_	Alpha blockers	+ URS	URS o	nly		Risk Ratio			Risk Ratio	0		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI		ľ	N-H, Fixed, 9	5% CI		
Ahmed 2017	74	81	67	84	61.3%	1.15 [1.01, 1.30]			-			
Wang 2014	44	45	41	44	38.7%	1.05 [0.96, 1.15]			+			
Total (95% CI)		126		128	100.0%	1.11 [1.02, 1.21]			<b>•</b>			
Total events	118		108									
Heterogeneity: Chi <sup>2</sup> =	, ,	,,	9%				0.1	0.2 0	.5 1	2		10
Test for overall effect:	Z = 2.34 (P = 0.02)	)						Favours U	RS only Fav	ours a bloc	cker + UF	RS

Figure 123: Use of healthcare services (Hospitalisation time)

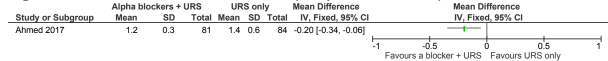


Figure 124: Hospitalisation (readmission)

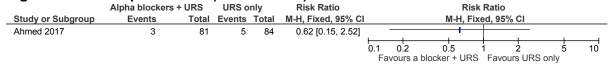


Figure 125: Adverse events (dizziness)

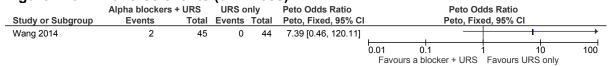


Figure 126: Pain intensity (colic episodes)

_	Alpha blockers	+ URS	URS o	nly	Risk Ratio			Risk	Ratio			
Study or Subgroup	Events	Total	Events	Total	M-H, Fixed, 95% CI			M-H, Fix	ed, 95%	CI		
Wang 2014	2	45	10	44	0.20 [0.05, 0.84]	+	<del></del>					
						0.1	0.2	0.5	1	2	5	10
						Fa	vours a b	locker + URS	Favou	rs URS	only	

# **Appendix F: GRADE tables**

Table 33: Clinical evidence profile: Alpha blockers versus placebo for distal ureteric stones <10mm in adults

			Quality as:	sessment		No of pat	ients		Effect	Quality	Importance	
No of studie	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other consideration s	Alpha blockers	Placebo (<10mm)	Relative (95% CI)	Absolute		
Stone p	assage (follow	-up 1-4 weeks;	assessed with	number of people	spontaneously p	assing stones	during follow up	)				
	randomised trials	no serious risk of bias	serious <sup>1</sup>	no serious indirectness	serious <sup>1</sup>	none	2219/2614 (84.9%)	60.9%	RR 1.19 (1.09 to 1.29)	116 more per 1000 (from 55 more to 177 more)	⊕⊕OO LOW	CRITICAL
Time to	stone passage	(follow-up 2-4	weeks; measu	red with: mean nu	mber of days for s	pontaneous st	one passage; Be	tter indicate	ed by lowe	r values)		
	randomised trials	very serious <sup>1</sup>	serious <sup>1</sup>	no serious indirectness	no serious imprecision <sup>1</sup>	none	1852	1817	-	MD 4.13 lower (4.32 to 3.94 lower)	⊕OOO VERY LOW	CRITICAL
Time to	stone passage	e (follow-up 3 w	reeks; assesse	d with: mean numb	per of hours for sp	ontaneous sto	ne passage )					
	randomised trials	serious <sup>3</sup>	no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	-	0%	HR 0.99 (0.55 to 1.78)	-	⊕000 VERY LOW	CRITICAL
Hospita	lisation (follow	v-up 3-4 weeks;	assessed with	: number of people	e hospitalized dur	ing follow up)						
	randomised trials	no serious risk of bias		no serious indirectness	very serious <sup>2</sup>	none	26/292 (8.9%)	4.4%	RR 0.99 (0.59 to 1.64)	0 fewer per 1000 (from 18 fewer to 28 more)	⊕⊕OO LOW	CRITICAL

	randomised	no serious risk		no serious	very serious <sup>2</sup>	none	31/198	18%	RR 0.87	23 fewer per	$\oplus \oplus OO$	CRITICAL
	trials	of bias	inconsistency	indirectness			(15.7%)		(0.56 to 1.36)	1000 (from 79 fewer to 65 more)	LOW	
dve	rse events (uns	pecified) (follow	up 2-4 weeks;	assessed with:	number of people	experiencing a	dverse events dur	ing follow ι	ıp)			
	randomised	very serious <sup>3</sup>	no serious	serious <sup>4</sup>	no serious	none	17/205	0%	RR 5.65	-	⊕ООО	CRITICAL
	trials		inconsistency		imprecision		(8.3%)		(1.5 to 21.29)		VERY LOW	
ve	rse events (retro	ograde ejaculatio	on) (follow-up 3	3-4 weeks; asse	ssed with: number	of people exp	eriencing retrograd	e ejaculatio	on during fo	ollow up )		
	randomised	serious <sup>3</sup>	serious <sup>1</sup>	no serious	no serious	none	86/1868	0%	Peto OR	20 more per 1000	⊕⊕ОО	CRITICAL
	trials			indirectness	imprecision		(4.6%)		1.78 (1.26 to 2.51)	·	LOW	
lve	randomised trials	very serious <sup>3</sup>	no serious inconsistency	no serious indirectness	serious <sup>1</sup>	none	77/1990 (3.9%)	w up)	RR 1.28 (0.92 to 1.79)	6 more per 1000 (from 2 fewer to 17 more)	⊕000 VERY LOW	CRITICA
lve	rse events (head	dache) (follow-u	p 4 weeks; ass	essed with: nun	nber of people exp	eriencing head	ache during follow	up)	1.73)	17 more)	LOW	
	randomised	no serious risk	no porious	no serious	very serious <sup>1</sup>	none	55/1879	2.9%	RR 1.06	2 more per 1000	⊕⊕OO	CRITICAL
	trials	of bias	inconsistency	indirectness	very serious	none	(2.9%)	2.970	(0.72 to 1.56)	(from 8 fewer to 16 more)	LOW	CRITICAL
dvo	rse events (hype	otension) (follow	v-up 4 weeks; a	ssessed with: r	number of people e	xperiencing h	/potension during	follow up)				
146	randomised	serious <sup>1</sup>	no serious	no serious	very serious <sup>1</sup>	none	1/118	0%	Peto OR	9 more per 1000	⊕ООО	CRITICAL
106			inconsistency	indirectness			(0.85%)		6.82 (0.13	(from 18 fewer to	VERY	

1	randomised	serious <sup>3</sup>	no serious	no serious	no serious	none	20/75	77.3%	RR 0.34	510 fewer per	⊕⊕⊕О	IMPORTAN
•	trials	Scrious	inconsistency	indirectness	imprecision	Horic	(26.7%)	77.070	(0.23 to	· ·	MODERA	
							(==::,-,		0.51)	fewer to 595	TE	
									,	fewer)	. –	
										,		
Pain	intensity (pain e	episodes) (follow	-up 2-4 weeks	measured with	: mean number of	pain episodes	; Better indicated b	y lower valu	ues)			
2	randomised	serious <sup>3</sup>	no serious	no serious	serious <sup>2</sup>	none	112	107	Ι -	MD 0.51 lower	⊕⊕00	IMPORTAN
_	trials	Scrious	inconsistency	indirectness	Scrious	none	112	107		(0.86 to 0.15	LOW	T
										lower)	LOW	
										1011017		
Pain	intensity (pain s	score >0) at 1 we	ek (follow-up 1	weeks: assess	ed with: verbal nur	neric pain sca	le)		1	ļ		<b>!</b>
			-	,			,					
1	randomised	no serious risk	no serious	no serious	no serious	none	142/185	78.6%	RR 0.98	16 fewer per	$\oplus \oplus \oplus \oplus$	IMPORTAN
	trials	of bias	inconsistency	indirectness	imprecision		(76.8%)		(0.88 to	1000 (from 94	HIGH	Т
							(		1.09)	fewer to 71 more)		
									,	,		
Pain	intensity (pain s	score >0) at 2 we	eks (follow-up	2 weeks: asses	sed with: verbal nu	ımeric pain sc	ale)		ļ	<u> </u>		
				,								
1	randomised	no serious risk	no serious	no serious	very serious <sup>2</sup>	none	60/176	32.8%	RR 1.04	13 more per 1000	⊕⊕00	IMPORTAN
	trials	of bias	inconsistency	indirectness			(34.1%)		(0.77 to	(from 75 fewer to	LOW	Т
							, ,		1.4)	131 more)		
Pain	intensity (pain s	score >0) at 3 we	eks (follow-up	3 weeks; asses	sed with: verbal nu	ımeric pain sc	ale)		II.		I.	
1	randomised	no serious risk	no serious	no serious	very serious <sup>2</sup>	none	34/170	21.4%	RR 0.94	13 fewer per	⊕⊕00	IMPORTAN
	trials	of bias	inconsistency	indirectness			(20%)		(0.62 to	1000 (from 81	LOW	Т
									1.42)	fewer to 90 more)		
Pain	intensity (pain s	score >0) at 4 we	eks (follow-up	4 weeks; asses	sed with: verbal nu	ımeric pain sc	ale)					
1	randomised	no serious risk	no serious	no serious	very serious <sup>2</sup>	none	26/173	16.1%	RR 0.93	11 fewer per	⊕⊕00	IMPORTAN
	trials	of bias	inconsistency	indirectness			(15%)		(0.57 to	1000 (from 69	LOW	Т
							, ,		1.53)	fewer to 85 more)		
										ĺ		
Anal	gesic use (follow	w-up 4 weeks; as	sessed with: n	umber of peopl	e using analgesics	during follow	up period )					
	-						,					

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2	randomised trials	no serious risk of bias	very serious <sup>1</sup>	no serious indirectness	no serious imprecision	none	40/1691 (2.4%)	24.5%	RR 0.29 (0.13 to 0.66)	174 fewer per 1000 (from 83 fewer to 213 fewer)	⊕⊕OO LOW	IMPORTAN T
Analge	esic use (numb	er of times) (foll	ow-up 4-12 we	eks; measured	with: mean numbe	r of times anal	gesics were used o	during follov	v up; Bette	r indicated by low	er values	)
2	randomised trials	serious <sup>3</sup>	no serious inconsistency	serious <sup>4</sup>	no serious imprecision	none	84	81	-	MD 0.9 lower (1.35 to 0.45 lower)	⊕⊕⊕O MODERA TE	IMPORTAN T
Analge	esic use (Bupre	enorphine dose)	(measured wit	h: mean dose o	f Buprenorphine u	sed during foll	ow up ; Better indi	cated by lov	ver values)			
2	randomised trials	very serious <sup>3</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	157	159	-	MD 0.07 lower (0.12 to 0.02 lower)	⊕⊕OO LOW	IMPORTAN T
Analge	esic use (Ketor	olac dose) (follo	w-up 2 weeks;	measured with:	mean dose of Ke	torolac used d	uring follow up; Be	tter indicate	d by lower	values)		
2	randomised trials	very serious <sup>3</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	156	159	-	MD 97.44 lower (124.25 to 70.62 lower)	⊕⊕OO LOW	IMPORTAN T
Analge	esic use (Diclo	fenac dose) (foll	ow-up 4 weeks	; measured with	n: mean dose of Di	clofenac used	during follow up; I	Better indica	ted by low	er values)		
2	randomised trials	no serious risk of bias	very serious <sup>1</sup>	no serious indirectness	no serious imprecision	none	1692	1700	-	MD 149.03 lower (152.37 to 145.68 lower)		IMPORTAN T

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Downgraded by 1 or 2 increments because the point estimate varies widely across studies, the confidence intervals across studies show minimal or no overlap, or heterogeneity, 12>50%, p<0.05, unexplained by subgroup analysis

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>&</sup>lt;sup>3</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>&</sup>lt;sup>4</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)

Table 34: Clinical evidence profile: Alpha blockers versus no treatment (pain management only) for distal ureteric stones <10mm in adults

			Quality as:	sessment			Ne	o of patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers	No treatment (pain management only) (<10mm)	Relative (95% CI)	Absolute		
Stone pa	ssage (follow	v-up 10 d	ays - 8 weeks ; as	ssessed with: n	umber of peop	ole spontaneously	/ passing st	ones during follow up	)			
	trials	serious <sup>1</sup>	serious <sup>1</sup> (follow-up 2-8 we		no serious imprecision with: mean nu	none mber of days for	1092/1364 (80.1%) spontaneou	51.1% s stone passage ; Bet	(1.48 to 1.82)	327 more per 1000 (from 245 more to 419 more) by lower values)	⊕000 VERY LOW	CRITICAL
17		very serious <sup>1</sup>	very serious <sup>1</sup>	no serious indirectness	no serious imprecision	none	901	641	-	MD 4.14 lower (5.23 to 3.04 lower)	⊕OOO VERY LOW	CRITICAL
Adverse	events (unsp	ecified) (	follow-up 10 day	s - 4 weeks ; as	sessed with: r	number of people	experiencin	g adverse events duri	ng follow up	)		
9	randomised trials	very serious <sup>1</sup>		no serious indirectness	no serious imprecision	none	10/407 (2.5%)	0%	Peto OR 5.89 (1.57 to 22.13)	-	⊕⊕OO LOW	CRITICAL
Adverse	events (dizzi	ness) (fo	llow-up 2-6 week	s; assessed wit	th: number of p	l people experienci	ng dizzines	s during follow up )				
7	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious <sup>3</sup>	none	23/277 (8.3%)	0%	RR 1.34 (0.74 to 2.4)	-	⊕OOO VERY LOW	CRITICAL
Adverse	events (hypo	otension)	(assessed with:	number of peop	ole experiencir	ng hypotension di	ıring follow	up )				

	randomised	very	no serious	no serious	no serious	none	9/300	0%	Peto OR 6		$\oplus \oplus OO$	CRITICAL
	trials	serious <sup>1</sup>	inconsistency	indirectness	imprecision		(3%)		(1.52 to 23.69)		LOW	
lvers	e events (retro	grade eja	culation) (follov	v-up 2-8 weeks	; assessed with	n: number of p	eople experiencir	ng retrograde eja	culation during f	ollow up)		
	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious <sup>3</sup>	none	3/157 (1.9%)	0%	RR 1.09 (0.21 to 5.67)	-	⊕000 VERY LOW	CRITICA
lvers	e events (head	lache) (fo	llow-up 2-6 wee	ks; assessed w	rith: number of	people experi	encing headache	during follow up	)			
		very serious <sup>1</sup>	no serious inconsistency	serious <sup>2</sup>	very serious <sup>3</sup>	none	8/100 (8%)	6.7%	RR 1.48 (0.47 to 4.69)	32 more per 1000 (from 36 fewer to 247 more)	⊕000 VERY LOW	CRITICA
spita	alisation (follow	w-up 2-4 \	weeks; assessed	d with: number	of people adm	itted to hospit	al during follow u	p)				
		very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	14/264 (5.3%)	11.5%	RR 0.27 (0.15 to 0.46)	84 fewer per 1000 (from 62 fewer to 98 fewer)	⊕⊕OO LOW	CRITICA
se of	healthcare ser	vices (ret	urn to ED/prima	ry care visit) (fo	ollow-up 2 wee	ks; assessed	with: number of p	eople returning t	o ED or having a	n unscheduled pri	mary car	e visit)
		very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious <sup>3</sup>	none	6/38 (15.8%)	20.5%	RR 0.77 (0.29 to 2.01)	47 fewer per 1000 (from 146 fewer to 207 more)	⊕000 VERY LOW	CRITICA
					mbar of needle	experiencing	pain during follow	w up)				
ain in	tensity (follow	-up 10 da	ys-4 weeks; ass	essed with: nu	inber of people	experiencing	pain daining rono.					

	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious <sup>3</sup>	none	0	-	-	MD 4.94 lower (12.04 lower to	⊕000 VERY	IMPORTAN
			,							2.16 higher)	LOW	
nalge	esic use (Pethic	dine dose	l ) (follow-up 4 w	eeks; measure	d with: mean do	se of Pethidine	used during f	ollow up ; Better indi	icated by lowe	r values)		
	randomised	very	no serious	no serious	no serious	none	32	32	_	MD 27.7 lower	⊕⊕OO	IMPORTAI
	trials	serious1	inconsistency	indirectness	imprecision					(33.41 to 21.99	LOW	
	trials		Ţ		·					lower)	LOW	
nalge	trials esic use (Ketor	olac dose	e) (follow-up 2 w	reeks; measure	d with: mean do	T		follow up; Better indi	icated by lower	lower)		IMPORTA
nalge	trials esic use (Ketor	olac dose	no serious	no serious	d with: mean do	ose of Ketorolac	used during	follow up; Better indi	cated by lower	r values)  MD 103.5 lower	⊕⊕OO	IMPORTA
nalge	trials esic use (Ketor	olac dose	e) (follow-up 2 w	reeks; measure	d with: mean do	T			icated by lower	lower)		IMPORTA
	trials  esic use (Ketoro randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	64		-	ND 103.5 lower (141.57 to 65.43 lower)	⊕⊕OO	IMPORTA
	trials  esic use (Ketore randomised trials  esic use (Bupre	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	64	62	-	ND 103.5 lower (141.57 to 65.43 lower)	⊕⊕OO LOW	IMPORTA

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 35: Clinical evidence profile: Calcium channel blockers versus placebo for distal ureteric stones <10mm in adults

			Quality asse	essment			No of pa	itients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Calcium channel blockers	placebo (<10mm)	Relative (95% CI)	Absolute		

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population or the majority of the evidence had indirect outcomes

<sup>&</sup>lt;sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>&</sup>lt;sup>4</sup> Downgraded by 1 or 2 increments because the point estimate varies widely across studies, the confidence intervals across studies show minimal or no overlap or heterogeneity, I2>50%, p<0.04, unexplained by subgroup analysis

Stone passage (follow-up 28-45 days; assessed with: number of people spontaneously passing stones during follow up )														
	more per 1000 $\oplus \oplus \oplus \oplus$ n 16 fewer to 115 HIGH more)													

Table 36: Clinical evidence profile: Calcium channel blockers versus no treatment (pain management only) for distal ureteric stones <10mm in adults

			Quality as	sessment			No	of patients	E	ffect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Calcium channel blockers	no treatment (pain management only) (<10mm)	Relative (95% CI)	Absolute		
Stone pa	ssage (follo	w-up 4 we	eeks; assessed v	vith: number o	f people spont	aneously passing	g stones dur	ing follow up )	-			
3	randomised trials			no serious indirectness	no serious imprecision	none	59/91 (64.8%)	36%	RR 1.95 (1.4 to 2.71)	342 more per 1000 (from 144 more to 616 more)	⊕⊕⊕O MODERATE	CRITICAL
Time to	l stone passaç	je (follow	-up 4 weeks; me	asured with: m	lean number o	f days for sponta	neous stone	passage ; Better inc	licated by low	er values)		
1	randomised trials				no serious imprecision	none	35	35	-	MD 0.29 lower (4.13 lower to 3.55 higher)	⊕⊕⊕O MODERATE	CRITICAL
Hospital	isation (follo	w-up 4 w	eeks; assessed v	with: number o	f people admit	tted to hospital du	uring follow	up)				
2	randomised trials			no serious indirectness	no serious imprecision	none	11/66 (16.7%)	38.6%	RR 0.41 (0.24 to 0.69)	228 fewer per 1000 (from 120 fewer to 293 fewer)	⊕⊕⊕O MODERATE	CRITICAL
Pain inte	nsity (pain e	pisodes)	(follow-up 4 wee	eks; measured	with: mean nu	mber of pain epis	odes during	   follow up; Better ind	dicated by low	ver values)		

	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	35	35	-	MD 0.09 higher (0.41 lower to 0.59 higher)	⊕⊕OO LOW	IMPORTAN
vers	e events (hype	otension)	(follow-up 4 we	eeks; assessed	with: number	of people exper	iencing hypote	ension during follow	up)			
	randomised trials		no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	1/31 (3.2%)	0%	Peto OR 6.71 (0.13 to 339.76)	-	⊕OOO VERY LOW	CRITICAL
dvers	e events (dizz	iness) (fo	ollow-up 4 week	s; assessed wi	th: number of p	people experien	cing dizziness	during follow up)	<u>'</u>			
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	0/25 (0%)	0%	see comment	0 fewer per 1000 (7 fewer to 7 more) <sup>3</sup>	⊕⊕OO LOW	CRITICAL
nalge	sic use (Diclo	fenac dos	se) (follow-up 4	weeks; measu	red with: mean	Diclofenac dos	e during follow	up ; Better indicate	ed by lower va	lues)		

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 37: Clinical evidence profile: Alpha blockers versus Calcium channel blockers for distal ureteric stones <10mm in adults

			Quality as:	sessment			No o	f patients		Effect	0	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers	Calcium channel blockers (<10mm)	Relative (95% CI)	Absolute	Quality	Importance
Stone pas	ne passage (follow-up 4 weeks; assessed with: number of people spontaneously passin						tones during	g follow up)	<del>'</del>			

<sup>&</sup>lt;sup>3</sup> Risk difference calculated in Review Manager

7	randomised trials	serious <sup>1</sup>	serious <sup>2</sup>	no serious indirectness	serious <sup>3</sup>	none	1948/2103 (92.6%)	68%	RR 1.2 (1.05 to 1.39)	136 more per 1000 (from 34 more to 265 more)	⊕000 VERY LOW	CRITICAL
Time to s	tone passag	e (follow-	-up 4 weeks; mea	sured with: mea	an number of d	lays for spontane	ous stone pa	assage ; Better i	ndicated by lo	ower values)		
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	35	35	-	MD 0 higher (3.13 lower to 3.13 higher)	⊕⊕⊕O MODERATE	CRITICAL
Hospitali	sation (follow	v-up 4 we	eeks; assessed w	ith: number of p	eople requirin	g hospitalisation	during follow	v up)				
2	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>3</sup>	none	5/67 (7.5%)	15.7%	RR 0.45 (0.18 to 1.17)	86 fewer per 1000 (from 129 fewer to 27 more)	⊕⊕OO LOW	CRITICAL
Adverse	events (head	ache) (fo	llow-up 4 weeks;	assessed with:	number of peo	ople experiencing	headache d	uring follow up	)			
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious <sup>3</sup>	none	32/64 (50%)	43.1%	RR 1.16 (0.79 to 1.7)	69 more per 1000 (from 91 fewer to 302 more)	⊕OOO VERY LOW	CRITICAL
Adverse	events (dizzi	ness) (fol	llow-up 4 weeks;	assessed with:	number of pec	pple experiencing	dizziness du	ıring follow up )				
2	randomised trials	very serious <sup>1</sup>	no serious inconsistency	serious <sup>4</sup>	no serious imprecision	none	18/89 (20.2%)	2.6%	RR 4.86 (1.62 to 14.56)	100 more per 1000 (from 16 more to 353 more)	⊕OOO VERY LOW	CRITICAL
Adverse	events (hypo	tension)	(follow-up 4 weel	ks; assessed wi	th: number of	people experienci	ng hypotens	sion during follo	w up)			
1	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious <sup>3</sup>	none	0/32 (0%)	3.2%	Peto OR 0.13 (0 to 6.61)	28 fewer per 1000 (from 32 fewer to 147 more)	⊕OOO VERY LOW	CRITICAL
Adverse	events (not s	pecified)	(follow-up 4 wee	ks; assessed w	ith: number of	people experienc	ing adverse	events during fo	ollow up)			
1	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>3</sup>	none	90/1596 (5.6%)	6.2%	RR 0.92 (0.69 to 1.21)	5 fewer per 1000 (from 19 fewer to 13 more)	⊕OOO VERY LOW	CRITICAL

	randomised trials	very serious <sup>1</sup>	no serious inconsistency	serious <sup>4</sup>	serious <sup>3</sup>	none	0/64 (0%)	5.2%		45 fewer per 1000 (from 51 fewer to 8	⊕000 VERY LOW	CRITICAL
			,						( ,	more)		
nalge	sic use (mg) (f	ollow-up	4 weeks; measu	red with: mean	Diclofenac mg	used during	follow up; Better ir	idicated by I	ower values)			
	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>3</sup>	none	25	25	-	MD 58 lower (315.47 lower to 199.47 higher)	⊕000 VERY LOW	IMPORTAN
nalge	sic use (follow	up 4 wee	eks; assessed w	ith: number of	people using a	nalgesics dur	ing follow up )					
	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	24/1596 (1.5%)	4.8%	RR 0.31 (0.2 to 0.49)	33 fewer per 1000 (from 24 fewer to 38 fewer)	⊕⊕⊕O MODERATE	IMPORTAN
nalge	sic use (follow	-up 4-12 v	weeks; measure	d with: mean a	nalgesic use ; I	Better indicate	ed by lower values)					
	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	64	58	-	MD 0.77 lower (0.93 to 0.61 lower)		IMPORTAI
ain in	tensity (pain e	pisodes)	(follow-up 4 wee	eks; measured v	vith: mean nun	nber of pain e	pisodes; Better ind	icated by lov	wer values)			
	randomised	serious <sup>1</sup>	no serious	no serious	no serious	none	35	35	-	MD 1.11 lower	⊕⊕⊕О	IMPORTA

Table 38: Clinical evidence profile: Alpha blockers versus placebo for mid ureteric stones <10mm in adults

Quality assessment	No of patients	Effect	Quality	Importance

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 or 2 increments because the point estimate varies widely across studies, the confidence intervals across studies show minimal or no overlap, or heterogeneity, 12>75%, p<0.05, unexplained by subgroup analysis.

<sup>&</sup>lt;sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>&</sup>lt;sup>4</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments) or the majority of the evidence had indirect outcomes

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No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	considerations	Alpha blockers	Placebo (<10mm)	Relative (95% CI)	Absolute		
Stone pas	sage (follow-i	ıp 4 weeks	s; assessed with:	number of peopl	e spontaneoi	usly passing stone	es during fol	low up )				
	randomised trials			no serious indirectness	serious <sup>2</sup>	none	37/61 (60.7%)	64.7%	RR 0.86 (0.67 to 1.09)	91 fewer per 1000 (from 214 fewer to 58 more)	⊕⊕OO LOW	CRITICAL

Renal and ureteric stones: Medical expulsive therapy

CONSULTATION

Table 39: Clinical evidence profile: Alpha blockers versus no treatment (pain management only) for mid ureteric stones <10mm in adults

			Quality as	sessment			No of	f patients			Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers	No treatment (<10mm)	Relative (95% CI)	Absolute		
Stone pa	ssage (follow	up 8 wee	ks; assessed with	l n: number of pec	pple spontaneou	l usly passing ston	es during fo	ollow up )				
2	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	9/14 (64.3%)	16.3%	RR 4.09 (1.09 to 15.33)	504 more per 1000 (from 15 more to 1000 more)	⊕OOO VERY LOW	CRITICAL
Time to s	tone passage	e (follow-u	ip 8 weeks; measi	red with: mean	number of days	for spontaneous	stone pass	sage ; Better in	dicated by lo	wer values)		
1	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	8	8	-	MD 12.33 lower (17.26 to 7.4 lower)	⊕⊕OO LOW	CRITICAL
Analgesi	use (follow-	up 8 week	s; measured with	: mean number	of times analge	sics were used du	iring follow	up ; Better ind	licated by lov	ver values)		
1	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	8	8	-	MD 1.2 lower (2.67 lower to 0.27 higher)	⊕OOO VERY LOW	IMPORTANT

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

1 Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 40: Clinical evidence profile: Calcium channel blockers versus placebo for mid ureteric stones <10mm in adults

			Quality asse	essment			No of patients	5		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Calcium channel blockers versus placebo	Control	Relative (95% CI)	Absolute	Quanty	importance
Stone pas	ssage (follow-	up 4 week	(s)									
	randomised trials			no serious indirectness	serious²	none	32/40 (80%)	81.8%	RR 0.98 (0.79 to 1.2)	16 fewer per 1000 (from 172 fewer to 164 more)		CRITICAL

Renal and ureteric stones: Medical expulsive therapy

Table 41: Clinical evidence profile: Alpha blockers versus Calcium channel blockers for mid ureteric stones <10mm in adults

			Quality asse	essment			No	of patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers	Calcium channel blockers (<10mm)	Relative (95% CI)	Absolute		
Stone pas	sage (follow-	up 4 week	ks; assessed with	: number of peo	ple spontane	ously passing sto	nes during	follow up )				
	randomised trials			no serious indirectness	serious <sup>2</sup>	none	29/41 (70.7%)	80%	RR 0.88 (0.69 to 1.14)	96 fewer per 1000 (from 248 fewer to 112 more)	0000	CRITICAL

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 42: Clinical evidence profile: Alpha blockers versus placebo for proximal ureteric stones <10mm in adults

(from ⊕⊕OO CR nore) LOW	CRITICAL
	00 (from ⊕⊕OO

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 43: Clinical evidence profile: Alpha blockers versus no treatment (pain management only) for proximal ureteric stones <10mm in adults

			Quality as	sessment			N	o of patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers	No treatment (pain management only) (<10mm)	Relative (95% CI)	Absolute		
Stone pa	ssage (follow	-up 4-8 w	reeks; assessed v	with: number of	f people sponta	neously passing	stones dur	ing follow up)				
4	randomised trials			no serious indirectness	serious <sup>2</sup>	none	76/113 (67.3%)	35.7%	RR 1.57 (1.2 to 2.03)	203 more per 1000 (from 71 more to 368 more)	⊕⊕OO LOW	CRITICAL
Time to s	tone passage	e (follow-	up 4-8 weeks; me	easured with: m	ean number of	days for spontar	eous stone	passage ; Better indic	cated by low	ver values)		
2				no serious indirectness	serious <sup>2</sup>	none	70	63	-	MD 5.29 lower (8.43 to 2.16 lower)	⊕OOO VERY LOW	CRITICAL

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

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rand	domised v	very	no serious	no serious	no serious	none	70	63	-	MD 0.55 lower	$\oplus \oplus OO$	IMPORTAI
trials	s s	serious1	inconsistency	indirectness	imprecision					(2.06 lower to 0.97	LOW	
										higher)		
ality of life	e (EuroQol	L) (follov	 v-up 4 weeks; m	easured with: n	lean score on E	L EuroQol ; Bette	er indicated by	lower values)				
ality of life	e (EuroQol	L) (follov	 v-up 4 weeks; m	easured with: n	nean score on E	EuroQoI ; Bette	er indicated by	lower values)				
	· ·	, , <u> </u>	v-up 4 weeks; m	easured with: n	nean score on E	EuroQol ; Bette	er indicated by	lower values)	-	MD 0.1 lower (0.42	⊕000	CRITICA
	domised v	very				·		,	-	MD 0.1 lower (0.42 lower to 0.22	⊕OOO VERY	CRITICA

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 44: Clinical evidence profile: Calcium channel blockers versus placebo for proximal ureteric stones <10mm in adults

			Quality asse	essment			No of patients	s		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Calcium channel blockers versus placebo	Control	Relative (95% CI)	Absolute	Quanty	importance
Stone pas	ssage (follow-	-up 4 wee	ks)									
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	58/92 (63%)	73%	RR 0.86 (0.71 to 1.06)	102 fewer per 1000 (from 212 fewer to 44 more)	⊕⊕OO LOW	CRITICAL

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

#### Table 45: Clinical evidence profile: Alpha blockers versus Calcium channel blockers for proximal ureteric stones <10mm in adults

Quality assessment	No of patients	Effect	Quality	Importance

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations		Calcium channel blockers (<10mm)	Relative (95% CI)	Absolute		
Stone pas	ssage (follow	up 4 wee	ks; assessed with	number of peo	ple spontane	ously passing sto	nes during	follow up )			ļ	
1	randomised	serious <sup>1</sup>	no serious	no serious	serious <sup>2</sup>	none	62/88	63%	RR 1.12	76 more per 1000	$\oplus \oplus \mathrm{OO}$	CRITICAL
	trials		inconsistency	indirectness			(70.5%)		(0.91 to 1.37)	(from 57 fewer to 233	LOW	ĺ
										more)		i
<sup>1</sup> Downgra	ded by 1 incre	ement if the	e majority of the evi	dence was at higl	h risk of bias,	and downgraded b	y 2 increme	nts if the majority of t	the evidence v	vas at very high risk of b	oias	
<sup>2</sup> Downgra	ded by 1 incre	ement if the	e confidence interva	al crossed one MI	D or by 2 incr	ements if the confid	dence interv	al crossed both MIDs	3			

Table 46: Clinical evidence profile: Alpha blockers versus placebo for distal ureteric stones <10mm in children

			Quality asse	essment			No of pa	tients		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers	Placebo	Relative (95% CI)	Absolute	Quanty	Importance	
Stone pas	Stone passage (follow-up 4 weeks)												
2	randomised trials			no serious indirectness	serious <sup>2</sup>	none	45/51 (88.2%)	69%	RR 1.3 (1.04 to 1.62)	207 more per 1000 (from 28 more to 428 more)	⊕⊕OO LOW	CRITICAL	
Time to st	tone passage	(days) (fo	llow-up (4 weeks);	Better indicated	by lower va	lues)							
	randomised trials	serious <sup>1</sup>	serious <sup>3</sup>	no serious indirectness	serious <sup>2</sup>	none	51	47	-	MD 4.89 lower (7.73 to 2.05 lower)	⊕OOO VERY LOW	CRITICAL	
Adverse e	Adverse events (headaches/dizziness) (follow-up 4 weeks)												
	randomised trials			no serious indirectness	serious²	none	3/18 (16.7%)	0%	Peto OR 8.82 (0.86 to 90.57)	167 more per 1000 (from 21 fewer to 354 more) <sup>4</sup>	⊕⊕OO LOW	CRITICAL	

Adverse events (headaches) (follow-up 4 weeks)														
	randomised trials	serious <sup>1</sup>	no serious inconsistency		very serious <sup>2</sup>	none	1/33 (3%)	3.6%	RR 0.85 (0.06 to 12.95)	5 fewer per 1000 (from 34 fewer to 430 more)	⊕OOO VERY LOW	CRITICAL		
Adverse e	Adverse events (hypotension)													
	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious	none	0/33 (0%)	0%	See comment	0 fewer per 1000 (from 62 fewer to 62 more) <sup>4</sup>	⊕OOO VERY LOW	CRITICAL		
Pain inter	Pain intensity (number of pain episodes) (follow-up 4 weeks; Better indicated by lower values)													
	randomised trials	serious <sup>1</sup>	very serious <sup>5</sup>	no serious indirectness	serious <sup>2</sup>	none	51	47	-	MD 1.49 lower (3.04 lower to 0.06 higher)	⊕OOO VERY LOW	IMPORTANT		

Renal and ureteric stones: Medical expulsive therapy

Table 47: Clinical evidence profile: Alpha blockers versus no treatment (pain management only) for distal ureteric stones <10mm in children

	Quality assessment  No of Pisk of Other							patients		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers	No treatment (<10mm)	Relative (95% CI)	Absolute			
Stone pas	Stone passage (follow-up 3-4 weeks; assessed with: number of people spontaneously passing stones)												

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs.

<sup>3</sup> Downgraded by 1 or 2 increments because heterogeneity, I2= 73%, p= > 0.1, unexplained by subgroup analysis

<sup>4</sup> Risk difference calculated in Review Manager

<sup>5</sup> Downgraded by 1 or 2 increments because heterogeneity, I2= 77%, p= > 0.1, unexplained by subgroup analysis

3	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	58/74 (78.4%)	62.5%	RR 1.45 (1.14 to 1.84)	281 more per 1000 (from 87 more to 525 more)	⊕OOO VERY LOW	CRITICAL
ime to	stone passage	e (follow-u	ıp 3-4 weeks; me	easured with: me	an number of d	ays for spontan	eous stone pas	ssage ; Better	indicated by	lower values)		
	randomised trials	very serious <sup>1</sup>	very serious <sup>3</sup>	no serious indirectness	very serious <sup>2</sup>	none	50	52	-	MD 5.26 lower (15.16 lower to 4.63 higher)	⊕OOO VERY LOW	CRITICAL
ain int	ensity (daily pa	ain episod	les) (follow-up 4	weeks; measure	d with: mean n	umber of daily p	ain episodes d	luring follow	up ; Better ind	icated by lower values	s)	
	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	31	32	-	MD 0.9 lower (1.77 to 0.03 lower)	⊕000 VERY LOW	IMPORTAN
nalges	sic use (follow-	up 4 weel	ks; measured wit	th: mean number	r of times analg	esics were used	during follow	up ; Better in	dicated by low	ver values)		
	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	31	32	-	MD 1.25 lower (1.87 to 0.63 lower)	⊕OOO VERY LOW	IMPORTAI
dverse	e events (follow	v-up 3-4 w	eeks; assessed	with: number of	people experie	ncing adverse e	vents (unspeci	ified))	1			1
	randomised	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	0	0%	see comment	MD 0 more per 1000 (50 fewer to 50 more) <sup>4</sup>	⊕⊕OO LOW	CRITICA

Table 48: Clinical evidence profile: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only for distal ureteric stones <10mm in adults

Quality assessment	No of patients	Effect	Quality	Importance

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>&</sup>lt;sup>3</sup> Downgraded by 1 or 2 increments because the point estimate varies widely across studies, the confidence intervals across studies show minimal or no overlap, or heterogeneity, I2>50%, p<0.04, unexplained by subgroup analysis.

<sup>&</sup>lt;sup>4</sup> Risk difference calculated in Review Manager

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers + SWL	SWL (<10mm)	Relative (95% CI)	Absolute		
Stone pas	ssage (follow	-up 15 da	ys - 6 weeks; asso	essed with: num	ber of people s	tone free at the er	d of follow up	o)				
5 Time to s	randomised trials	serious <sup>1</sup>	no serious inconsistency	serious <sup>2</sup>	serious <sup>3</sup>	none	144/190 (75.8%) Better indicate	56.8%	RR 1.28 (1.11 to 1.48)	159 more per 1000 (from 62 more to 273 more)	⊕OOO VERY LOW	CRITICAL
2	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>3</sup>	none	104	103	-	MD 2.21 lower (3.35 to 1.08 lower)	⊕OOO VERY LOW	CRITICAL
Adverse	events (dizzir	ness) (follo	ow-up 15 days - 6	weeks; assesse	ed with: number	of people experie	encing dizzine	ess during t	follow up )			1
3	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	7/101 (6.9%)	0%	Peto OR 8.4 (1.86 to 37.87)	69 more per 1000 (from 17 more to 122 more) <sup>4</sup>	⊕⊕OO LOW	CRITICAL
Analgesi	use (follow-	up 4 weel	 ks; measured with	i: mean number	of times analge	esics were used d	uring follow u	p ; Better ii	ndicated by low	er values)		
1	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>3</sup>	none	44	44	-	MD 1.72 lower (2.88 to 0.56 lower)	⊕OOO VERY LOW	IMPORTANT
Hospitali	sation (follow	v-up 4 wee	ks; assessed with	h: number of pe	ople hospitalize	l ed during follow u	o)					
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>3</sup>	none	12/44 (27.3%)	43.2%	RR 0.63 (0.35 to 1.14)	160 fewer per 1000 (from 281 fewer to 60 more)	⊕⊕OO LOW	CRITICAL
Adverse	 events (abnor	rmal ejacı	l ılation) (follow-up	4-6 weeks; ass	essed with: nun	 nber of people exp	periencing ab	l normal ejad	ulation during	follow up )		
2	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	7/49 (14.3%)	0%	Peto OR 8.56 (1.83 to 40.08)	142 more per 1000 (from 40 more to 246 more) <sup>4</sup>	⊕⊕OO LOW	CRITICAL

	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>3</sup>	none	9/76 (11.8%)	2.9%	Peto OR 4.19 (1.23 to 14.28)	88 more per 1000 (from 1 more to 427 more)	⊕OOO VERY LOW	CRITICA
dverse	events (hypo	tension) (	follow-up 6 weeks	s; assessed with	n: number of pe	ople experiencing	hypotension	during follo	ow up)			
	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	0/32 (0%)	0%	See comment	0 more per 1000 (from 60 fewer to 60 more) <sup>4</sup>	⊕⊕OO LOW	CRITICA
nalges	ic use (dosag	e) (follow-	-up 4 weeks; mea	sured with: mea	n dosage (mg) (	of Diclofenac duri	ng follow up;	l Better indic	cated by lower v	ralues)		
	randomised trials	very serious <sup>1</sup>	no serious inconsistency	serious <sup>2</sup>	no serious imprecision	none	60	59	-	MD 50.27 lower (68.87 to 31.67 lower)	⊕000 VERY	IMPORTA

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 49: Clinical evidence profile: Alpha blockers as adjunctive therapy to ureteroscopy versus ureteroscopy only for distal ureteric stones <10mm in adults

			Quality as:	sessment			No of pa	atients		Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers + URS	URS (<10mm)	Relative (95% CI)	Absolute			
Stone pas	Stone passage (follow-up 2 weeks; assessed with: number of people stone-free at the end of follow up)												

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)

<sup>&</sup>lt;sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>&</sup>lt;sup>4</sup> Risk difference calculated in Review Manager

1	randomised trials	serious <sup>1</sup>			no serious imprecision	none	48/51 (94.1%)	87.2%	RR 1.08 (0.95 to 1.23)	70 more per 1000 (from 44 fewer to 201 more)	⊕⊕⊕O MODERATE	CRITICAL		
Use of he	Jse of healthcare services (measured with: length of hospital stay; Better indicated by lower values)													
1	randomised trials	serious <sup>1</sup>		no serious indirectness	serious <sup>2</sup>	none	51	47	-	MD 0.5 lower (0.81 to 0.19 lower)	⊕⊕OO LOW	CRITICAL		

<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 50: Clinical evidence profile: Alpha blockers as adjunctive therapy to ureteroscopy versus placebo and ureteroscopy for distal ureteric stones <10mm in adults

			Quality as	sessment			No of p	oatients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers + URS	placebo + URS (<10mm)	Relative (95% CI)	Absolute		
Stone pas	ssage (follow	-up 2 wee	ks; assessed witl	h: number of pe	ople stone free	at the end of follo	w up )					
	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	49/52 (94.2%)	70%	RR 1.35 (1.11 to 1.63)	245 more per 1000 (from 77 more to 441 more)	⊕000 VERY LOW	CRITICAL
Analgesio	use (follow-	up 2 weel	ks; assessed with	: number of peo	ple using analo	jesia during follov	up)					
	randomised trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious²	none	4/52 (7.7%)	24%	RR 0.32 (0.11 to 0.93)	163 fewer per 1000 (from 17 fewer to 214 fewer)	⊕OOO VERY LOW	IMPORTANT
Pain inter	nsity (colic ep	oisodes) (	follow-up 2 weeks	s; measured with	n: mean numbe	r of colic episodes	s during follo	w up ; Better ir	ndicated by I	ower values)		
	randomised trials	very serious <sup>1</sup>	no serious inconsistency		no serious imprecision	none	52	50	-	MD 5 lower (5.99 to 4.01 lower)	⊕⊕OO LOW	IMPORTANT

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 51: Clinical evidence profile: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only for distal ureteric stones 10-20mm in adults

Renal and ureteric stones: Medical expulsive therapy

	•		Quality assess	ment			No of pat	ients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers + SWL	SWL (10- 20mm)	Relative (95% CI)	Absolute		
Time to sto	one passage (	follow-up (	unclear; measured	with: number	of days for	stone passage; Be	tter indicated by	lower value	es)			
1	randomised trials	l , .	no serious inconsistency	serious <sup>2</sup>	serious <sup>3</sup>	none	14	24	-	MD 2.56 lower (7.78 lower to 2.66 higher)	⊕000 VERY LOW	CRITICAL
Pain intens	sity (VAS) (foll	ow-up und	clear; measured wit	h: visual ana	logue scale;	range of scores: 0	-10; Better indic	ated by low	er values		L	
1	randomised trials		no serious inconsistency	serious <sup>2</sup>	serious <sup>3</sup>	none	14	24	-	MD 1.21 lower (2.88 lower to 0.46 higher)	⊕000 VERY LOW	IMPORTANT

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 52: Clinical evidence profile: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only for mid ureteric stones 10-20mm in adults

Quality assessment	No of patients	Effect	Quality	Importance
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<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)

<sup>&</sup>lt;sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers + SWL	SWL (10- 20mm)	Relative (95% CI)	Absolute		
ne passage (f	follow-up	unclear; measured	with: number	r of days for	stone passage; Be	tter indicated by	lower value	es)			
andomised rials	serious <sup>1</sup>	no serious inconsistency	serious <sup>2</sup>	very serious <sup>3</sup>	none	16	12	-	MD 1.5 lower (8.23 lower to 5.23 higher)	⊕OOO VERY LOW	CRITICAL
ty (VAS) (foll	ow-up und	clear; measured wit	th: visual ana	logue scale;	range of scores: 0	-10; Better indic	ated by low	er values	)		
andomised rials	serious <sup>1</sup>	no serious inconsistency	serious <sup>2</sup>	serious <sup>3</sup>	none	16	12	-	MD 0.62 lower (3.13 lower to 1.89 higher)	VERY	IMPORTANT
a ri:	e passage (findomised als	bias  e passage (follow-up indomised serious)  y (VAS) (follow-up undomised serious)	bias Inconsistency  e passage (follow-up unclear; measured ndomised als inconsistency  y (VAS) (follow-up unclear; measured with ndomised serious no serious	bias Inconsistency Indirectness e passage (follow-up unclear; measured with: number ndomised als inconsistency serious²  y (VAS) (follow-up unclear; measured with: visual analondomised serious¹ no serious serious²	bias Inconsistency Indirectness Imprecision  e passage (follow-up unclear; measured with: number of days for sendomised als inconsistency serious seri	bias Inconsistency Indirectness Imprecision considerations  e passage (follow-up unclear; measured with: number of days for stone passage; Be  ndomised serious¹ no serious serious² very serious³  y (VAS) (follow-up unclear; measured with: visual analogue scale; range of scores: 0	bias Inconsistency Indirectness Imprecision considerations + SWL  e passage (follow-up unclear; measured with: number of days for stone passage; Better indicated by ndomised als serious¹ no serious serious² very serious³ none 16  y (VAS) (follow-up unclear; measured with: visual analogue scale; range of scores: 0-10; Better indicated by serious² serious² serious² none 16	Design Risk of bias Inconsistency Indirectness Imprecision Considerations Alpha blockers SWL (10-20mm)  e passage (follow-up unclear; measured with: number of days for stone passage; Better indicated by lower value als serious none serious serious serious none 16 12  y (VAS) (follow-up unclear; measured with: visual analogue scale; range of scores: 0-10; Better indicated by low none serious none serious serious serious serious none 16 12	Design Risk of bias Inconsistency Indirectness Imprecision Considerations Alpha blockers + SWL (10-20mm) (95% CI)  The passage (follow-up unclear; measured with: number of days for stone passage; Better indicated by lower values)  The passage (follow-up unclear; measured with: number of days for stone passage; Better indicated by lower values)  The passage (follow-up unclear; measured with: viewal analogue scale; range of scores: 0-10; Better indicated by lower values)  The passage (follow-up unclear; measured with: visual analogue scale; range of scores: 0-10; Better indicated by lower values)  The passage (follow-up unclear; measured with: visual analogue scale; range of scores: 0-10; Better indicated by lower values)	bias inconsistency indirectness imprecision considerations + SWL 20mm) (95% CI) Absolute  e passage (follow-up unclear; measured with: number of days for stone passage; Better indicated by lower values)  ndomised serious¹ no serious serious² very none 16 12 - MD 1.5 lower (8.23 lower to 5.23 higher)  y (VAS) (follow-up unclear; measured with: visual analogue scale; range of scores: 0-10; Better indicated by lower values)  ndomised serious¹ no serious serious² serious³ none 16 12 - MD 0.62 lower (3.13)	Design Risk of bias Inconsistency Indirectness Imprecision Considerations Alpha blockers + SWL (10-20mm) (95% CI) Absolute  e passage (follow-up unclear; measured with: number of days for stone passage; Better indicated by lower values)  Indomised als serious no serious inconsistency serious serious serious serious serious none 16 12 - MD 1.5 lower (8.23 lower to 5.23 higher) VERY LOW  Yellow-up unclear; measured with: visual analogue scale; range of scores: 0-10; Better indicated by lower values)

Renal and ureteric stones: Medical expulsive therapy

CONSULTATION

Table 53: Clinical evidence profile: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only for proximal ureteric stones <10mm in adults

	Quality assessment  Other							No of patients Effect			Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers + SWL	SWL (<10mm)	Relative (95% CI)	Absolute		
Stone pas	ssage (follow	-up 2-12 v	weeks; assessed	with: number of	people stone t	ree at the end of	follow up)					
	randomised trials			no serious indirectness	no serious imprecision	none	176/198 (88.9%)	84.8%	RR 1.11 (1.03 to 1.21)	93 more per 1000 (from 25 more to 178 more)	⊕⊕⊕O MODERATE	CRITICAL
Time to s	tone passage	e (follow-ι	ıp 2-12 weeks; m	easured with: no	umber of days t	for stone passage	; Better indic	ated by lov	wer values)			

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)

<sup>&</sup>lt;sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

	andomised	very	serious <sup>2</sup>	no serious	no serious	none	155	165	-	MD 4.32 lower (9.85		CRITICAL
ľ	rials	serious <sup>1</sup>		indirectness	imprecision					lower to 1.21 higher)	VERY LOW	
ain inten	sity (VAS) (f	ollow-up	2-12 weeks; mea	asured with: vis	ual analogue so	ale ; range of s	scores: 0-10; Bett	ter indicate	ed by lower value	ies)	l	
r	andomised	very	serious <sup>2</sup>	no serious	serious <sup>3</sup>	none	183	191	-	MD 0.89 lower (1.68	⊕OOO	IMPORTAN
t	rials	serious <sup>1</sup>		indirectness						to 0.1 lower)	VERY LOW	
ospitalis	ation (follow	/-up 2 we	eks; measured v	vith: mean num	ber of Hospitalis	sations; Better	indicated by low	er values)				
r	andomised	very	no serious	serious <sup>4</sup>	no serious	none	35	44	-	MD 0.01 lower (0.31	⊕OOO	CRITICAL
t	rials	serious <sup>1</sup>	inconsistency		imprecision					lower to 0.29 higher)	VERY LOW	
nalgesic	use (follow-	up 2-6 w	eeks; assessed v	with: number of	people using a	nalgesia durinç	g follow up)					
r	andomised	very	serious <sup>2</sup>	serious <sup>4</sup>	very serious <sup>3</sup>	none	37/76	49.2%	RR 0.96 (0.49	20 fewer per 1000	⊕OOO	IMPORTAN
t	rials	serious <sup>1</sup>					(48.7%)		to 1.91)	(from 251 fewer to 448 more)	VERY LOW	
dverse e	vents (dizzii	l ness) (foll	ow-up 3-6 week	s; assessed wit	h: number of pe	ople experienc	ing dizziness du	ring follow	/ up )			
r	andomised	very	no serious	no serious	serious <sup>3</sup>	none	3/85	0%	Peto OR 7.76	35 more per 1000	⊕ООО	CRITICAL
t	rials	serious <sup>1</sup>	inconsistency	indirectness			(3.5%)		(0.8 to 75.32)	(from 9 fewer to 80 more) <sup>5</sup>	VERY LOW	
dverse e	vents (retro	grade eja	culation) (follow	up 6 weeks; as	sessed with: nu	mber of people	e experiencing re	trograde e	 ejaculation duri	ng follow up )		
r	andomised	very	no serious	no serious	no serious	none	0/41	0%	See comment	0 more per 1000	⊕⊕00	CRITICAL
t	rials	serious <sup>1</sup>	inconsistency	indirectness	imprecision		(0%)			(from 45 fewer to 45 more) <sup>5</sup>	LOW	
nalgesic	use (dosage	e) (follow	l -up 4 weeks; me	asured with: me	ean dosage (mg	) of Diclofenac	used during follo	ow up; Bet	ter indicated by	v lower values)		
r	andomised	very	no serious	no serious	serious <sup>3</sup>	none	28	26	-	MD 189.7 lower	⊕OOO	IMPORTAN
t	rials	serious <sup>1</sup>	inconsistency	indirectness						(309.2 to 70.2 lower)	VERY LOW	
		1	1	1	1			1	1		1	I

1	trials	very serious <sup>1</sup>	no serious inconsistency	no serious indirectness		none	28	26	-	MD 0.6 lower (1.13 to 0.07 lower)	⊕000 VERY LOW	CRITICAL
Pain inter	nsity (renal co	olic episo	des) (follow-up 4	weeks; measur	ed with: mean i	number of renal co	olic episodes	during foll	ow up ; Better	indicated by lower v	alues)	
1		very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>3</sup>	none	28	26	-	MD 2.38 lower (3.89 to 0.87 lower)	⊕000 VERY LOW	IMPORTANT
Quality o	f life (EQ5D) (	follow-up	4 weeks; measu	red with: mean	score on EQ5D	; range of scores:	0-1; Better in	dicated by	higher values	)		
1		very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>3</sup>	none	28	26	-	MD 0.04 higher (0.01 lower to 0.09 higher)		CRITICAL
Quality o	f life (EQ5D V	AS) (folio	ow-up 4 weeks; m	easured with: n	nean score on E	Q5D visual analog	gue scale ; ra	nge of sco	res: 0-100; Bet	ter indicated by high	ner values)	
1		very serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>3</sup>	none	28	26	-	MD 6.71 higher (1.49 to 11.93 higher)	⊕000 VERY LOW	CRITICAL

Table 54: Clinical evidence profile: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus placebo and shock wave lithotripsy for proximal ureteric stones <10mm in adults

			Quality asses	ssment			No of	patients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers + SWL	Placebo + SWL (<10mm)	Relative (95% CI)	Absolute		
Stone pas	sage (follow	-up 3 months	; assessed with:	number of pe	ople stone free	at the end of follo	ow up )				L	

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 or 2 increments because the point estimate varies widely across studies, the confidence intervals across studies show minimal or no overlap or heterogeneity, I2=50%,

<sup>&</sup>lt;sup>3</sup> Risk difference calculated in Review Manager

1			no serious inconsistency	serious <sup>1</sup>	serious <sup>2</sup>	none	27/28 (96.4%)	66.7%	RR 1.45 (1.06 to 1.97)	300 more per 1000 (from 40 more to 647 more)	⊕⊕OO LOW	CRITICAL
Time to s	tone passage	(follow-up 3	months; measur	ed with: num	ber of days for s	stone passage ; B	etter indicated	l by lower valu	es)			
1	randomised trials		no serious inconsistency		no serious imprecision	none	28	21	-	MD 3.3 lower (4.47 to 2.13 lower)	⊕⊕OO LOW	CRITICAL

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)

Table 55: Clinical evidence profile: Alpha blockers as adjunctive therapy to shock wave lithotripsy versus shock wave lithotripsy only for proximal ureteric stones 10-20mm in adults

			Quality asses	sment			No of pa	tients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers + SWL	SWL (10- 20mm)	Relative (95% CI)	Absolute		
Time to st	one passage	(follow-up	unclear; measure	ed with: numb	er of days to	stone passage ; E	Better indicated	d by lower v	ralues)			
1	randomised trials	very serious <sup>1</sup>	no serious inconsistency	serious <sup>2</sup>	serious <sup>3</sup>	none	29	28	-	MD 6.44 lower (10.3 to 2.58 lower)	⊕000 VERY LOW	CRITICAL
Pain inten	sity (VAS) (fo	llow-up ur	l nclear; measured v	with: visual a	nalogue scal	e; range of scores	: 0-10; Better ii	ndicated by	lower values)			
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	serious <sup>2</sup>	serious <sup>3</sup>	none	29	28	-	MD 1.1 lower (2.34 lower to 0.14 higher)	⊕000 VERY LOW	IMPORTANT
Stone pas	sage (follow-	up 3 mont	hs; assessed with	: number of p	eople stone	free at the end of f	follow up )	Į.		!		

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>&</sup>lt;sup>3</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

	randomised	very	no serious	serious <sup>2</sup>	serious <sup>3</sup>	none	26/29	82.1%	RR 1.09 (0.88	74 more per 1000 (from	$\oplus$ OOO	CRITICAL
	trials	serious1	inconsistency				(89.7%)		to 1.35)	99 fewer to 287 more)	VERY	
											LOW	

Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias

Table 56: Clinical evidence profile: Alpha blockers as adjunctive therapy to ureteroscopy versus ureteroscopy only for proximal ureteric stones 10-20mm in adults

			Quality as	sessment			No of pa	tients	Effect		Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Alpha blockers + URS	URS (10- 20mm)	Relative (95% CI)	Absolute	,	
Stone pa	ssage (follow	/-up 4-6 w	eeks; assessed v	 vith: number of	people stone fr	ee at the end of fo	llow up )					
2	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	118/126 (93.7%)	86.5%	RR 1.11 (1.02 to 1.21)	95 more per 1000 (from 17 more to 182 more)	⊕⊕⊕O MODERATE	CRITICAL
Use of he	ealthcare serv	vices (Hos	spitalisation time)	(follow-up adm	ission; measur	ed with: length of	hospital stay	for proce	dure; Better ind	dicated by lower valu	ies)	
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	serious <sup>2</sup>	none	81	84	-	MD 0.2 lower (0.34 to 0.06 lower)	⊕⊕OO LOW	CRITICAL
Hospitali	sation (readn	nission) (f	l follow-up 8 weeks	; assessed with	: number of pe	ople readmitted to	hospital dur	ing follow	up)			
1	randomised trials	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	3/81 (3.7%)	6%	RR 0.62 (0.15 to 2.52)	23 fewer per 1000 (from 51 fewer to 91 more)	⊕OOO VERY LOW	CRITICAL
Time to s	tone passage	e (follow-u	up 6 weeks; meas	sured with: num	ber of days for	stone passage; B	etter indicate	d by lower	values)			

<sup>&</sup>lt;sup>2</sup> Downgraded by 1 or 2 increments because the majority of the evidence included an indirect population (downgrade by one increment) or a very indirect population (downgrade by two increments)

<sup>&</sup>lt;sup>3</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

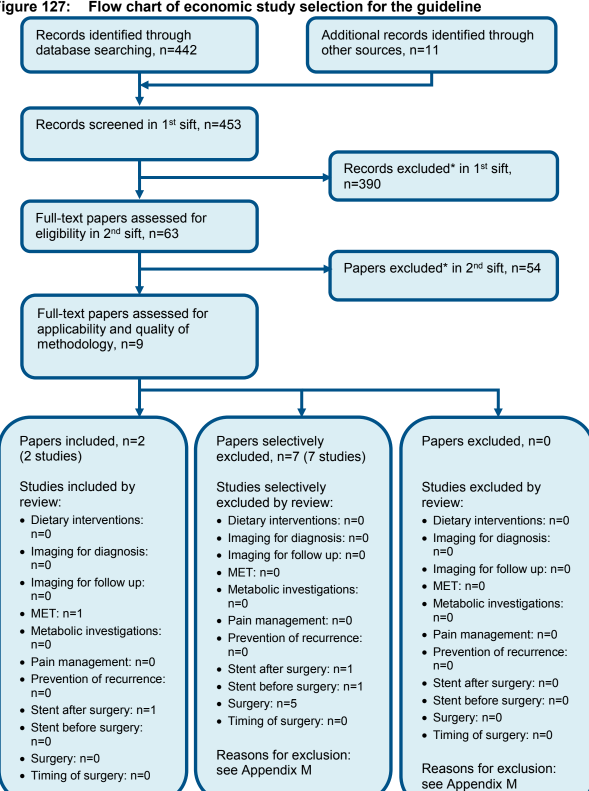
1	randomised	very	no serious	no serious	serious <sup>2</sup>	none	45	44	-	MD 3.68 lower (6.95	$\oplus$ OOO	CRITICAL
	trials	serious <sup>1</sup>	inconsistency	indirectness						to 0.41 lower)	VERY LOW	
Pain inter	nsity (uretera	l colic rat	e) (follow-up 6 we	eeks; assessed	with: number o	f people experienc	cing ureteral of	colic durin	g follow up )			
					-					<u> </u>		
1	randomised	very	no serious	no serious	serious <sup>2</sup>	none	2/45	22.7%	RR 0.2 (0.05	182 fewer per 1000	$\oplus$ OOO	IMPORTANT
	trials	serious <sup>1</sup>	inconsistency	indirectness			(4.4%)		to 0.84)	(from 36 fewer to 216	VERY LOW	
										fewer)		
Adverse e	events (dizzir	ness) (foll	ow-up 6 weeks; a	ssessed with: n	umber of peop	le experiencing di	zziness durin	g follow u	<b>p</b> )			
1	randomised	very	no serious	no serious	very serious <sup>2</sup>	none	2/45	0%	Peto OR 7.39	44 more per 1000	$\oplus$ OOO	CRITICAL
	trials	serious1	inconsistency	indirectness			(4.4%)		(0.46 to	(from 28 fewer to 117	<b>VERY LOW</b>	
									120.11)	more) <sup>3</sup>		
					1							

<sup>&</sup>lt;sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias <sup>2</sup> Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

<sup>&</sup>lt;sup>3</sup> Risk difference calculated in Review Manager

## Appendix G: Health economic evidence selection

Figure 127: Flow chart of economic study selection for the guideline



<sup>\*</sup> Non-relevant population, intervention, comparison, design or setting; non-English language

1

2

# Appendix H: Health economic evidence tables

[Please note, only cite studies using the Main Endnote library for the guideline. This can be found at N:\NCGC Guidelines\[guideline]\5-Development\Searches\[Guideline]\ main database. **Under no circumstances should you cite from the search results library.**]

Renal and ureteric stones: Medical expulsive therapy

Study	Pickard 2015 <sup>145</sup>			
Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALYs)  Study design: CUA (health outcome: QALYs)  Approach to analysis: This within trial analysis comparing the cost effectiveness of MET (nifedipine or tamsulosin) vs placebo and MET drugs to each other (tamsulosin vs nifedipine). The data were taken from 24 UK hospitals from 1167 participants and data on resource use and quality of life data was collected in all patients at baseline, 4 and 12 weeks after	Population: Patients presented as an emergency with a diagnosis of ureteric colic at UK NHS hospitals and diagnosed with a symptomatic ureteric stone of ≤ 10 mm in maximum dimension  Patient characteristics: N: unclear as only complete data was used for the economic analysis Mean age: 43.1 (tamsulosin group), 42.3 (nifedipine group), 42.8 (placebo) Male (a):82.2% (tamsulosin group), 82.8% (nifedipine group), 77.9% (placebo)  Intervention 1: Placebo	Total costs (mean per patient): Intervention 1: NR Intervention 2: NR Intervention 3: NR Intervention 4: NR Incremental (2–1): -£42 Incremental (4-3):-£87  Currency & cost year: 2012-13 British Pounds  Cost components incorporated: Drugs costs (interventions, analgesics, antibiotics), Resource use costs (GP appointment, outpatient appointment and admissions), diagnostic tests costs, unit costs of further active intervention like stents use or cost of	QALYs (mean per patient): Intervention 1: 0.20 Intervention 2: 0.19 Intervention 3: 0.20 Intervention 4: 0.19 Incremental (2-1): -0.001 (adjusted) Incremental (4-3): -0.002 (adjusted)	Intervention 2 versus Intervention 1): Intervention 2 is less expensive and less effective than intervention 1, so the ICER of 1 versus 2 is presented for ease of interpretation; Placebo vs MET = £42,000 (c) Probability Intervention 2 cost effective (£20K/30K threshold):56%/51%  ICER (Intervention 3 versus Intervention 4): Intervention 4 is less expensive and less effective than intervention 3, so the ICER of 3 versus 4 is presented for ease of interpretation; Nifedipine vs Tamsulosin = £43,500 Probability intervention 4 cost effective(£20K/30K threshold):61%/55%  Analysis of uncertainty: Non-parametric bootstrapping was used to generate 1000 estimates of mean costs and QALYs for each treatment

Perspective: UK NHS

**Time horizon/Followup**: the period of the clinical study (12 weeks)

Treatment effect duration:4 weeks

Discounting: N/A

Intervention 2: Medical Expulsive Therapy consisted of Nifedipine, 30mg-MR capsules, or Tamsulosin hydrochloride 0.4 mg, for a maximum of 28 days

**Intervention 3**: Nifedipine (MR capsules), 30mg-once daily, for a maximum of 28 days

#### Intervention 4:

Tamsulosin hydrochloride (MR capsules) 0.4 mg once daily, for a maximum of 28 days (b)

lithotripsy, participants costs (self-purchased health care such as prescription costs, over the counter medications, visits to non NHS health care providers)

group. Various one way sensitivity analyses were undertaken;

#### Using SF-6D instead of EQ-5D (d)

There was uncertainty around the QALY estimates derived using the EQ-5D that its sensitivity to capture the loss in QoL particularly in reference to acute pain was questioned. Therefore SF-36 responses were mapped onto the SF-6D measure.

- MET versus placebo: MET was again less expensive and less effective, and so comparing placebo to MET gave an ICER of £12,333 (placebo cost effective).
- Tamsulosin versus nifedipine:, Tamsulosin was again less expensive and less effective, and so comparing nifedipine to tamsulosin gave an ICER of £23,000 (nifedipine borderline cost effective).

# Multiple imputation for EQ-5D – replacing all missing EQ-5D data with highest EQ-5D score

- MET versus placebo: MET again less expensive and less effective, so comparing placebo to MET gave an ICER of £6,000 (placebo cost effective). Incremental cost only £6 so explains low ICER but incremental QALY still 0.001.
- Tamsulosin versus nifedipine: Tamsulosin is more expensive and more effective (both only slightly), giving an ICER of £24,677. So tamsulosin is above the cost effectiveness threshold slightly.

#### **Data sources**

CONSULTATION

Health outcomes: Results of the large RCT informing resource use for the cost effectiveness analysis. Questionnaires were designed to obtain information on stone passage or further intervention, pain, HRQoL and resource use, including NHS and personal costs. Participants were asked to complete trial questionnaires at baseline, 4 weeks post randomisation and 12 weeks post randomisation. The baseline questionnaire was completed in hospital before randomisation Quality-of-life weights: Health-related quality-of-life measures were collected at baseline, 4 weeks and 12 weeks by participant completion of the EQ-5D and the SF-36 questionnaires. Responses from the SF-36 questionnaire were also used as the basis of QALYs as a sensitivity analysis to validate the EQ-5D scores. They were mapped onto the existing Short Form questionnare-6 Dimensions (SF-6D) measure using a standard algorithm to allow utility values to be estimated for each time point. These utility scores were transformed to QALYs using the methods described above to provide an alternative measure of QALYs for each participant Cost sources: Unit costs (drug costs) were obtained from published sources such as the British National Formulary (BNF) and NHS reference costs (cost of diagnostic tests, outpatient costs for urology department for a consultant outpatient appointment, cost of interventions like lithotripsies, stents insertion and removal, cost of admission with no intervention, cost of any extra admission days using the long stay excess days tariff) Cost of a GP appointment were obtained from the Personal Social Services Research Unit costs of primary services. The unit cost data source year was 2012–13 and the currency was British pounds.

#### Comments

Source of funding: National Institute for Health Research Limitations: A cost utility analysis that is a within trial analysis based on a UK RCT, using an NHS perspective and the EQ-5D that reports changes in quality of life and costs coming from the use of MET (tamsulosin and nifedipine) and placebo. Study Included some participants costs that are not NHS costs related, and these were reported as part of NHS costs that they account for significant % of total costs of intervention; so it is difficult to separate participants' costs from the NHS costs in order to determine whether their magnitude is significant compared to the total costs of interventions. The categories where the patient reported outcomes fall include costs that are of similar amount in both interventions (MET, placebo), so unlikely changing the cost effectiveness results. Study used a time horizon of 12 weeks and not longer. That was justified by the authors as there weren't many people who still needed interventions at the end of the trial. However there were no extrapolation and therefore assumptions made about what this treatment would be which could impact incremental costs and effects because different numbers of people are stone free in each arm, and that is a potentially serious limitation detracting from overall study quality Other:

#### **Overall applicability:** Directly applicable **Overall quality:** Potentially serious limitations

Abbreviations: CUA: cost—utility analysis; da: deterministic analysis; EQ-5D: Euroqol 5 dimensions (scale: 0.0 [death] to 1.0 [full health], negative values mean worse than death); ICER: incremental cost-effectiveness ratio; MET: medical expulsive therapy; NR: not reported; pa: probabilistic analysis; SF-36: Short Form (36) Health Survey (scale: 0.0 (maximum disability to 100 no disability)QALYs: quality-adjusted life years WTP: Willingness to pay

- (a) Study reported % female participants for each intervention group and % male participants was worked out using the data from female
- (b) Interventions administered for up to 4 weeks until the stone passage
- (c) Note that the ICER reported in the table for MET vs placebo is reported as £4,355. Taking the incremental cost and dividing by the incremental QALY (-42/-0.001) is £42,000 which is much larger than the ICER reported. Therefore there must be a reporting error. Additionally this is reported as cost saving per QALY lost for MET versus placebo because MET is an intervention appearing in the bottom left quadrant of the cost effectiveness plane. However for ease of interpretation in cases like this the intervention should be switched around i.e. to compare placebo versus MET so that the less effective intervention is used as the comparator and so the ICER can be interpreted as it normally would (if less than £20,000 then intervention is cost effective versus the comparison).
- (d) At the different sensitivity analysis scenario where the uncertainty around the QALY estimates derived using EQ-5D is further investigated, costs also changed not just QALYs because they are using a different subset of people from the base case, because these are people who responded to the SF-36. Same applies to the multiple imputation analysis as well.

# Appendix I: Excluded studies

### I.1 Excluded clinical studies

1

3 Table 57: Studies excluded from the clinical review

Study	Exclusion reason
Abdelaziz 2017 <sup>2</sup>	Inappropriate comparison
Afridi 2017 <sup>4</sup>	Stone location not reported
Ahmed 2016 <sup>10</sup>	·
Ahmed 2014 12	Not review population Incorrect comparison
	Not available
Ahmed Pechuho 2012 <sup>11</sup>	
Amar 2017 <sup>19</sup>	Article not in English
Amer 2017 <sup>19</sup>	Systematic review is not relevant to review question or unclear PICO
Amer 2017 <sup>18</sup>	Systematic review is not relevant to review question or unclear PICO
Aydin 2017 <sup>23</sup>	Unclear stone size and location
Ayubov 2007 <sup>25</sup>	abstract
Bahn Zobbe 1986 <sup>26</sup>	Incorrect interventions
Bai 2017 <sup>27</sup>	Inappropriate comparison
Beach 2006 <sup>32</sup>	Systematic review is not relevant to review question or unclear PICO
Berger 2015 <sup>33</sup>	Mixed stone location
Bhagat 2007 <sup>34</sup>	Not review population
Borghi 1994 <sup>35</sup>	Mixed stone location
Campschroer 2014 <sup>36</sup>	Systematic review is not relevant to review question or unclear PICO
Cao 2014 <sup>37</sup>	Systematic review is not relevant to review question or unclear PICO
Caravati 198938	Crossover study
Cervenakov 2002 <sup>39</sup>	Inappropriate comparison
Cha 2012 <sup>40</sup>	Inappropriate comparison
Cho 2017 <sup>43</sup>	Mixed stone location
Clayman 2002 <sup>44</sup>	editorial comment
Cooper 2000 <sup>45</sup>	Inappropriate comparison
Daga 2016 <sup>46</sup>	Inappropriate comparison
De Nunzio 2016 <sup>47</sup>	Not review population
Dellabella 2003 <sup>50</sup>	Inappropriate comparison
Dellabella 2005 <sup>51</sup>	Inappropriate comparison
Dell'atti 2015 <sup>49</sup>	Inappropriate comparison
Dellis 2017 <sup>52</sup>	Stone location not reported
Ding 2016 <sup>53</sup>	Systematic review is not relevant to review question or unclear PICO
Doluoglu 2015 <sup>54</sup>	Inappropriate comparison
Elgalaly 2016 <sup>58</sup>	Inappropriate comparison
El-Gamal 2012 <sup>55</sup>	Inappropriate comparison
	, , ,

Study	Exclusion reason
Eryildirim 2015 <sup>62</sup>	Incorrect study design
Falahatkar 2011 <sup>64</sup>	Not review population
Fan 2013 <sup>65</sup>	Systematic review is not relevant to review question or unclear PICO
Georgescu 2015 <sup>69</sup>	Mixed stone location
Georgiev 2011 <sup>70</sup>	Incorrect study design
Glina 2015 <sup>71</sup>	Systematic review is not relevant to review question or unclear PICO
Gottlieb 2017 <sup>72</sup>	review of Pikard 2015
Goyal 2018 <sup>73</sup>	Incorrect interventions
Gravina 2005 <sup>75</sup>	Not review population
Griwan 2010 <sup>76</sup>	Inappropriate comparison
Gupta 2008 <sup>78</sup>	Comment
Gupta 2013 <sup>79</sup>	Inappropriate comparison
Gupta 2014 <sup>77</sup>	Review protocol
Gurbuz 2011 <sup>80</sup>	Inappropriate comparison
Hamidi Madani 201181	Incorrect interventions
Han 200682	Article not in English
Hollingsworth 2016 <sup>84</sup>	Systematic review is not relevant to review question or unclear PICO
Huang 2016 <sup>85</sup>	Systematic review is not relevant to review question or unclear PICO
Hussein 2010 <sup>86</sup>	Not review population
Hwang 201287	Incorrect study design
Itoh 2013 <sup>90</sup>	Appears to be a sub-analysis of Itoh 2011
Janane 201492	Not review population
Jayant 2014 93	Incorrect comparison
John 2010 <sup>94</sup>	Not review population
Kang 2009 97	Not in English
Kaneko 2010 <sup>96</sup>	Mixed stone location
Kc 2016 98	Incorrect comparison
Kim 2008 <sup>100</sup>	Article not in English
Kiraç 2013 <sup>101</sup>	Inappropriate comparison
Kobayashi 2008 <sup>102</sup>	Mixed stone location
Kohjimoto 2015 <sup>103</sup>	Inappropriate comparison
Koski 2018 <sup>104</sup>	Systematic review: references checked
Kroczak 2017 <sup>105</sup>	Literature review
Kumar 2013 <sup>108</sup>	Inappropriate comparison
Kumar 2014 <sup>107</sup>	Incorrect comparison
Kumar 2015 <sup>106</sup>	Inappropriate comparison
Lee 2012 <sup>110</sup>	Systematic review is not relevant to review question or unclear PICO
Li 1995 <sup>112</sup>	Not review population
Li 2015 <sup>114</sup>	Systematic review is not relevant to review question or unclear PICO
Li 2017 <sup>113</sup>	Systematic review is not relevant to review question or unclear PICO

Litasikos 2007 <sup>115</sup> Liu 2012 <sup>116</sup> Systematic review is not relevant to review question or unclear PICO Liu 2015 <sup>116</sup> Systematic review is not relevant to review question or unclear PICO Liu 2017 <sup>119</sup> Systematic review is not relevant to review question or unclear PICO Liu 2018 <sup>117</sup> Incorrect interventions Losek 2008 <sup>121</sup> Systematic review is not relevant to review question or unclear PICO Liu 2012 <sup>123</sup> Systematic review is not relevant to review question or unclear PICO Lu 2012 <sup>124</sup> Incorrect comparison Lu 2012 <sup>125</sup> Inappropriate comparison Lu 2014 <sup>126</sup> Systematic review is not relevant to review question or unclear PICO McClinton 2014 <sup>127</sup> Study protocol Micali 2006 <sup>129</sup> Not review population Micali 2007 <sup>128</sup> Inappropriate comparison Montiel-Jarquin Á 2017 <sup>133</sup> Inappropriate comparison Montiel-Jarquin Á 2017 <sup>133</sup> Not in English Mukhtarov 2007 <sup>135</sup> abstract Naja 2008 <sup>137</sup> Not review population Inappropriate comparison Inappropriate comparison Organi 2010 <sup>140</sup> Inappropriate comparison Inappropriate comparison Pico Picozzi 2011 <sup>147</sup> Systematic review is not relevant to review question or unclear PICO Picozzi 2011 <sup>147</sup> Systematic review is not relevant to review question or unclear PICO Picozzi 2011 <sup>148</sup> Not review population Porpiglia 2000 <sup>149</sup> Inappropriate comparison Inappropriate comparison Porpiglia 2000 <sup>149</sup> Inappropriate comparison Porpiglia 2000 <sup>150</sup> Inappropriate comparison Porpiglia 2000 <sup>151</sup> Inappropriate comparison Porpiglia 2000 <sup>152</sup> Incorrect study design Porpiglia 2009 <sup>151</sup> Second line therapy Portis 2018 <sup>154</sup> Incorrect study design Puvvada 2016 <sup>155</sup> Incorrect omparison Rahman 2017 <sup>158</sup> Inappropriate comparison Reddy 2016 <sup>161</sup> Incorrect study design Reddy 2016 <sup>161</sup> Incorrect study design Reddy 2016 <sup>161</sup> Incorrect interventions Saita 2004 <sup>165</sup> Not review population Not review population	Study	Exclusion reason
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, ,	Romics 2011 <sup>164</sup>	Incorrect interventions
Sarica 2006 <sup>167</sup> Not review population	Saita 2004 <sup>165</sup>	Incorrect study design
• •	Sarica 2006 <sup>167</sup>	Not review population
Schuler 2009 <sup>169</sup> Systematic review is not relevant to review question or unclear PICO	Schuler 2009 <sup>169</sup>	

Study	Exclusion reason
Seitz 2009 <sup>170</sup>	Systematic review is not relevant to review question or unclear PICO
Seungok 2009 <sup>172</sup>	Abstract
Shaaban 2008 <sup>173</sup>	Abstract
Shabana 2016 <sup>174</sup>	Inappropriate comparison
Shahat 2016 <sup>175</sup>	Not review population
Shokeir 2016 <sup>176</sup>	Incorrect comparison
Singh 2007 <sup>177</sup>	Systematic review is not relevant to review question or unclear PICO
Skolarikos 2015 <sup>181</sup>	Systematic review is not relevant to review question or unclear PICO
Skolarikos 2017 <sup>180</sup>	Systematic review: references checked
Skrekas 2003 <sup>182</sup>	abstract
Sridharan 2017 <sup>183</sup>	Systematic review: references checked
Sridharan 2018 <sup>184</sup>	Not available
Strohmaier 1994 <sup>185</sup>	Not review population
Sumer 2012 <sup>187</sup>	Not guideline condition. Not review population
Tasian 2014 <sup>190</sup>	Incorrect study design
Tian 2017 <sup>192</sup>	Systematic review is not relevant to review question or unclear PICO
Tsuzaka 2011 <sup>193</sup>	Incorrect comparison
Tuerxun 2017 <sup>194</sup>	Incorrect study design
Velazquez 2015 <sup>195</sup>	Systematic review is not relevant to review question or unclear PICO
Vicentini 2011 <sup>196</sup>	Not review population
Vincendeau 2010 <sup>197</sup>	Inappropriate comparison
Wang 2008 <sup>203</sup>	Article not in English
Wang 2009 <sup>199</sup>	Incorrect study design
Wang 2010 <sup>201</sup>	Inappropriate comparison
Wang 2016 <sup>202</sup>	Systematic review is not relevant to review question or unclear PICO
Wang 2017 <sup>204</sup>	Systematic review is not relevant to review question or unclear PICO
Yang 2016 <sup>206</sup>	Systematic review is not relevant to review question or unclear PICO
Yencilek 2010 <sup>209</sup>	Inappropriate comparison
Zaytoun 2012 <sup>212</sup>	Not review population
Zehri 2010 <sup>213</sup>	Incorrect study design
Zheng 2010 <sup>215</sup>	Systematic review is not relevant to review question or unclear PICO
Zhu 2010 <sup>217</sup>	Systematic review is not relevant to review question or unclear PICO

### I.2 Excluded health economic studies

3 None

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## Appendix J: Research recommendations

### J.1 Alpha blockers and ureteroscopy

Research Question: What is the clinical and cost effectiveness of tamsulosin as an adjunct to ureteroscopy?

#### Why this is important:

Kidney and ureteric stones affect about 15% of the male population and 5% of the female population at some point in their lives. The incidence of kidney stones has been increasing because of their link to poor diet, obesity and diabetes. Kidney and ureteric stones can cause severe pain and morbidity. Ureteroscopy is a commonly used method of treating stones in the kidney or ureter, whereby a narrow telescope is advanced up the ureter and laser energy is applied to the stone through a small fibre. Fragments may be left to wash out or removed with a basket.

The ureteric is intrinsically narrow but its wall contains muscle which is known to relax when the patient is given a medication called tamsulosin, which is in common use for prostatic problems. Tamsulosin has been shown to improve the spontaneous passage rate of small ureteric stones and also fragment-clearance following shockwave lithotripsy and NICE guidelines have recommended alpha blockers such as tamsulosin are considered for such purposes.

The success of ureteroscopic stone clearance can be inhibited by the ureter being excessively tight. This might prevent the insertion of the ureteroscope into the ureter (failed access) or reduce the effectiveness of laser fragmentation or the spontaneous clearance of fragments. Ureteric stents are often used as a post-treatment safety measure if the ureter is felt to be tight or swollen up during a procedure. Ureteric stents are known to cause significant irritation symptoms due to mechanical rubbing in the urinary tract. There is also some evidence that these symptoms might be reduced by tamsulosin. Nevertheless, the studies that these finding are based on are small and the evidence quality is low so tamsulosin is not in widespread routine use for these purposes. A definitive RCT is required to determine if such a recommendation would be appropriate.

o determine il such a recommendation would be appropriate.	
PICO question	<b>Population:</b> Adults with ureteric or renal stones up to 20mm in size undergoing ureteroscopic treatment and no stent <b>Intervention(s):</b> Tamsulosin 400mcg od for 1 week prior to ureteroscopy and for 4 weeks after
	Comparison: Double-blind placebo controlled
	Outcomes:
	Primary outcome: Stone free rate as assessed by CT KUB at 4 weeks Secondary outcomes: failed access rate, operation time, stenting rate, needs for repeat ureteroscopy or adjunctive procedures, hospitalisation/ED attendance?, pain scores, quality of life (EQ-5D- 3L), stent symptoms, side effects, failed insertion of access sheath, cost per QALY.
Importance to patients or the population	Kidney stones are extremely common and cause significant morbidity. Ureteroscopy is a commonly used and effective method of treating kidney stones. The success of stone clearance can be inhibited by the tightness of the ureter. Simple measures to relax the ureter peri-operatively might improve the success of the procedure, reduce the need for secondary procedures and improve the procedure related morbidity and quality of life.

Research recommendations

Relevance to NICE	The NICE guidelines panel felt that the current evidence was of too low
guidance	quality to make a current recommendation on the use of tamsulosin for this purpose.
Relevance to the NHS	Tamsulosin is inexpensive and widely used by urologists. Ureteroscopic stone treatments are very common and improvements in its success rate will reduce the need for expensive secondary procedures and may reduce the cost of treatment related morbidity
National priorities	There is a strong link between diabetes, obesity and kidney stones and limiting the impact of these conditions is one of the top research priorities of the NHS. It is also a priority to test interventions and maximize effectiveness and cost-effectiveness.
Current evidence base	The current evidence is restricted to one or two studies with small numbers of participants for most outcome measures.
Equality	The recommendation is unlikely to impact on equality issues.
Study design	Double-blind placebo controlled RCT with health economic analysis
Feasibility	The trial is feasible and should be straightforward to carry out. There are a large number of such patients and a UK kidney stone trial network has already been established. The SUSPEND and TISU trials demonstrate this.
Other comments	The length of pre-treatment tamsulosin might be reviewed.
Importance	<ul> <li>Medium: the research is relevant to the recommendations in the guideline, but the research recommendations are not key to future updates.</li> </ul>