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OPEN Percutaneous nephrostomy versus retrograde ureteral stenting for acute upper obstructive uropathy: a systematic review and meta-analysis

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Acute obstructive uropathy is associated with significant morbidity among patients with any condition that leads to urinary tract obstruction. Immediate urinary diversion is necessary to prevent further damage to the kidneys. In many centres, the two main treatment options include percutaneous nephrostomy (PCN) and retrograde ureteral stenting (RUS). The purpose of this study if to compare the efficacy and safety of PCN and RUS for the treatment of acute obstructive uropathy. We searched the Cochrane Central Register of Controlled Trials (CENTRAL), MEDLINE, CINAHL, EMBASE, the World Health Organisation International Clinical Trials Registry Platform and Clinical Trials.gov. We also searched the reference lists of included studies to identify any additional trials. We included randomised controlled trials and controlled clinical trials comparing the outcomes of clinical improvement (septic parameters), hospitalisation duration, guality of life, urinary-related symptoms, failure rates, post-procedural pain [measured using a visual analogue scale (VAS)] and analgesics use. We conducted statistical analyses using random effects models and expressed the results as risk ratio (RR) and risk difference (RD) for dichotomous outcomes and mean difference (MD) for continuous outcomes, with 95% confidence intervals (CIs). Seven trials were identified that included 667 patients. Meta-analysis of the data revealed no difference in the two methods in improvement of septic parameters, quality of life, failure rates, post-procedural pain (VAS), or analgesics use. Patients receiving PCN had lower rates of haematuria and dysuria post-operatively and longer hospitalisation duration than those receiving RUS. PCN and RUS are effective for the decompression of an obstructed urinary system, with no significant difference in most outcomes. However, PCN is preferable to RUS because of its reduced impact on the patient's post-operative quality of life due to haematuria and dysuria, although it is associated with slightly longer hospitalisation duration.

Obstructive uropathy is one of the most common conditions affecting the urinary system and is a significant cause of renal impairment, leading to end-stage renal failure. It is a condition wherein impedance of urinary flow causes dilatation of the pelvicalyceal system, resulting in damage to the renal parenchyma¹; 9.2% of chronic kidney disease cases are caused by obstruction of the urinary tract². No or suboptimal treatment will lead to inevitable permanent chronic kidney disease through a combination of ischaemic or disuse-induced tubular injury, inflammation and interstitial renal fibrosis^{3,4}.

Urgent decompression is warranted in cases of acute obstructive uropathy, either percutaneously via a nephrostomy tube or retrogradely via ureteral stent placement. This decompression prevents further worsening of renal function, inflammation and ischaemia to renal parenchyma that can eventually progress to irreversible chronic kidney disease.

The choice of technique depends on operator preference, availability of interventional radiologists or urologists and the centre's standard protocol. At our institution, PCN is the preferred approach of urinary diversion

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for acute obstructive uropathy due to better availability of interventional radiologists and it is a much cheaper procedure as most of our patients are from lower income group.

Currently, there are no standard guidelines on the preferred method of urgent urinary tract decompression in the setting of acute obstructive uropathy. There is conflicting evidence in the literature regarding these two methods. PCN is associated with higher rates of sepsis and mortality compared to RUS⁵. Other outcomes such as time to definitive stone management, rates of spontaneous stone passage and initiation of stone metabolic workup were not found to be statistically different⁶. Therefore, an evidence-based comparison based on pooled data from available clinical trials comparing these two approaches is warranted to provide a potential basis for establishing a standard guideline in the management of acute upper obstructive uropathy. The objective of this study was to compare the efficacy and safety of PCN and RUS in treating this condition.

Results

Search results. We retrieved 3786 records from the search databases and four additional records from other sources (Fig. 1). In total, 563 records were screened after removing duplicates. We reviewed 30 full-text articles and excluded 23 articles, of which seven were non-comparative trials and six investigated empirical treatment. Empirical treatment refers to the use of PCN or RUS as preventive measures in conditions wherein obstructive uropathy is anticipated but has not yet occurred.

Included studies. We included seven trials (N = 667)⁷⁻¹⁴. These trials are summarized in Table 1.

Participants. All seven trials were conducted in healthcare settings. The total number of participants was 667. Four of the trials were conducted in the urology departments from referral cases^{9,10,12,14}, two trials included patients who presented to the emergency department^{7,13} and one trial included patients referred to a stone management unit⁸. One multicentre trial was conducted in urology units in Germany and Syria⁹. Two trials were RCTs^{9,10}, whereas five were CCTs^{7,8,12-14}.

Intervention. All trials compared PCN and RUS as the intervention and control groups, respectively. In all trials, the procedures were conducted by urologists. Percutaneous nephrostomies were conducted under ultrasound guidance with local anaesthesia. Retrograde ureteral stents were inserted using a cystoscope after patients were put under general anaesthesia. All stents were double-J stents.

Outcomes. Two trials analysed parameters of clinical success, which included septic parameters and hospitalisation duration in patients with obstructive uropathy^{7,12}. Three trials reported patients' quality of life and urinary-related symptoms^{8,13,14}. For the secondary outcomes, three trials reported failure rates^{7,9,10}, two trials reported post-procedural pain^{7,14} and four trials reported analgesics use^{8,9,13,14}.

All trials that reported quality of life used a similar questionnaire, that is, the EuroQol EQ-5D-3L^{8,14,15}, which is the three-level version of the EQ-5D. This measure includes a descriptive system that contains five dimensions: mobility, self-care, usual activities, pain/discomfort and anxiety/depression. Each dimension has three levels: no problems, some problems and extreme problems. Participants were asked to express their health state for the most appropriate statement in each of the five dimensions.

Excluded studies. We excluded 13 trials. Seven trials had no control group^{16–22}. Six trials analysed PCN and RUS as empirical treatment^{23–28}.

Risk of bias in included studies. The risk of bias assessment is shown in Figs. 2 and 3. The proportion of the assessment of included studies based on the risk of bias for each 'risk of bias' indicator is shown in Fig. 2. The risk of bias analysis for the individual studies is shown in Fig. 3.

Allocation. Three trials reported the use of the random number tables method of randomisation^{7,10,12}. One trial used quasi-randomisation based on odd- or even-numbered birth year²⁹. The remaining three trials did not use any randomisation; patients were allocated based on the preference of the performing surgeon, who was not aware of the trials^{8,14,15}.

Blinding. Blinding of participants and personnel was not feasible because all trials involved a procedure.

Incomplete outcome data. In one trial, RUS insertion failure occurred in three patients who were then subjected to PCN as the urinary diversion method. These patients were still included in the analysis in the RUS group¹⁰. In another trial, four of 20 patients in the RUS group had an unsuccessful procedure and thus were subjected to PCN⁹. The four patients were analysed as a separate group; the outcomes in the RUS group were analysed based on the remaining 16 patients. In one trial, two patients were missing from both the PCN and RUS groups because of loss to follow-up¹². These patients were not included in the final analysis. In one trial, four patients were missing for various reasons in each group¹¹. They were included in the analysis of outcomes.

Selective reporting. All seven trials reported outcomes based on the objectives and measured in the methods $^{7-14}$.





Other potential sources of bias. There were no other potential sources of bias in the included studies.

Effects of interventions. *Primary outcomes.* All trials reported the primary outcomes of this meta-analysis⁷⁻¹⁴:

1. Clinical success of PCN: Three trials reported the clinical success outcomes regarding improvement in septic parameters^{7,12,14}.

Author, year	Location	Age (years)	Settings	Recruitment	Study design	Number of participants, n	Interventions	Outcomes	
Ahmad 2012	Bahawalpur,	PCN: 43±9.65	Single contro	Referral to Uralogy	RCT	300	PCN, n = 200	Failure rates	
Allillad, 2015	Pakistan	RUS: 40±10.35		Referrar to Crology	KC1	300	RUS, n = 100	Haematuria	
		PCN: 63.1					PCN, n = 18	Mobility	
de Sousa, 2018	Braga, Portugal	RUS: 54.5	Single centre	Presented at ED	ССТ	50	RUS, n = 32	Self-care Usual activity Pain/discomfort Anxiety/depression Haematuria Dysuria Urgency Frequency Use of analgesics	
		PCN: 56±9					PCN, n = 13	Mobility	
Joshi, 2001	Bristol, UK	DJS: 55±14	Single centre	Referral to stone management unit	ССТ	34	DJS, n = 21	Self-care Usual activity Pain/discomfort Anxiety/depression Haematuria Dysuria Urgency Frequency Use of analgesics	
Makhmalii 2001	Aleppo, Syria	PCN: 49	Multinantes	Defensel to Unale an	DOT	40	PCN, n = 24	Failure rates	
Mokhinaji, 2001	Germany	RUS: 55	Multicentre	Referrat to Urology	KC1	40	RUS, n = 16	Use of analgesics	
		PCN: 41.3±13.0					PCN, n = 21	Failure rates	
Pearle, 1998	Texas, USA	RUS: 41.3±14.5	Single centre	Presented at ED	ССТ	42	RUS, n = 21	Use of analgesics Normalization for WBC normalization Duration to defer- vescence Duration of hospi- talization Pain post procedure	
		PCN: 54 (46.5-61)					PCN, n = 30	Mobility	
Shoshany, 2019	Tel Aviv, Israel	DJS: 55 (39.5–70.5)	Multicentre	Referral to Urology	ССТ	75	DJS, n = 45	Self-care Usual activity Pain/discomfort Anxiety/depression Haematuria Use of analgesics Pain post procedure	
		PCN: 58.21 ± 10.89					PCN, n = 53	Duration to defer-	
Wang, 2015 Taiwan		RUS: 57.52±11.93	Single centre	Referral to Urology	CCT	107	RUS, n = 54	vescence Duration for WBC normalization Duration of hospi- talization	

Table 1. Summary of included studies in meta-analysis.



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





	PCN		RUS				Mean Difference	Mean Difference	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI	IV, Random, 95% Cl
Pearle 1998	2	1.2	26	1.7	0.8	26	52.7%	0.30 [-0.25, 0.85]	
Wang 2016	4.74	1.62	53	4.37	1.46	54	47.3%	0.37 [-0.21, 0.95]	
Total (95% CI)			79			80	100.0%	0.33 [-0.07, 0.74]	
Heterogeneity: Tau² = Test for overall effect:	: 0.00; C Z = 1.62	hi ² = 0 (P = (_	-0.5 -0.25 0 0.25 0.5 Favours PCN Favours RUS					

Figure 4. Comparison between PCN and RUS for outcome duration to WBC normalisation.

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- a. Improvement in septic parameters:
 - i. WBC normalisation duration: Two trials compared the WBC normalisation duration (days) post-operatively^{7,12} (two trials; 149 participants; MD [95% C1] 0.33 [-0.07 to 0.74]; I²=0%; *P*=0.100; moderate-quality evidence; Fig. 4; Table 2).
 - ii. Duration to defervescence. Three trials reported the duration to defervescence^{7,12,14}; however, one trial reported the median duration (interquartile range [IQR]) and thus was not included in the meta-analysis¹⁴ (two trials; 142 participants; MD [95% CI] 0.33 [-0.46 to 0.53]; $I^2 = 0\%$; P = 0.890; moderate-quality evidence; Fig. 5; Table 2).

Patient or population: acute obstructive uropathy, Setting: emergency and day care, Intervention: PCN, Comparison: RUS											
	Anticipated absolute eff	ects* (95% CI)	Relative effect (95%	No. of participants	Certainty of the						
Outcomes	Risk with RUS	Risk with PCN	CI)	(studies)	evidence (GRADE)	Comments					
Duration to WBC normalisation	The mean duration to WBC normalisation was 0	MD 0.33 higher (0.07 lower to 0.74 higher)	-	159 (2 RCTs)	$ \begin{array}{c} \bigoplus \bigoplus \bigoplus \bigcirc MODER-\\ ATE^{a} \end{array} $	Assumed risk calculated from the mean risk across the RUS group of the two included trials					
Duration to deferves- cence	The mean duration to defervescence was 0	MD 0.03 higher (0.46 lower to 0.53 higher)	-	142 (2 RCTs)	⊕⊕⊖ MODER- ATE ª	Assumed risk calculated from the mean risk across the RUS group of the two included trials					
Hospitalisation duration	The mean hospitalisa- tion duration was 0	MD 1.82 higher (0.79 higher to 2.85 higher)	-	149 (2 RCTs)	$ \begin{array}{c} \bigoplus \bigoplus \bigoplus \bigcirc MODER-\\ ATE a \end{array} $	Assumed risk calculated from the mean risk across the RUS group of the two included trials					
QoL—mobility	31 per 100	24 per 100 (8 to 76)	RR 0.78 (0.25 to 2.48)	159 (3 RCTs)	$ \bigoplus_{LOW} \bigcirc_{a,b,c} VERY $	Assumed risk calculated from the mean risk across the RUS group of the three included trials					
QoL—usual activity	34 per 100	53 per 100 (19 to 100)	RR 1.57 (0.55 to 4.53)	159 (3 RCTs)	$ \bigoplus_{LOW} \bigcirc_{a,b,c} VERY $	Assumed risk calculated from the mean risk across the RUS group of the three included trials					
QoL—pain/discomfort	47 per 100	46 per 100 (35 to 59)	RR 0.97 (0.75 to 1.26)	159 (3 RCTs)	⊕⊕⊖⊖ LOW a,b	Assumed risk calculated from the mean risk across the RUS group of the three included trials					
Haematuria	38 per 100	21 per 100 (14 to 32)	RR 0.56 (0.37 to 0.85)	459 (4 RCTs)	$\bigoplus_{LOW} \bigcirc_{a,b,c} VERY$	Assumed risk calculated from the mean risk across the RUS group of the four included trials					

Table 2. Summary of findings for comparison between percutaneous nephrostomy and retrograde ureteral stenting for acute obstructive uropathy. ***The risk in the intervention group** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). CI: Confidence interval; MD: Mean difference; RR: Risk ratio. GRADE Working Group grades of evidence. High certainty: We are very confident that the true effect lies close to that of the estimate of the effect, but there is a possibility that it is substantially different. Low certainty: Our confidence in the effect, but there is a possibility that it is substantially different from the estimate of the effect. Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect.

	PCN RUS							Mean Difference		Mean Di	fference		
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI		IV, Rando	m, 95% Cl		
Pearle 1998	2.3	1.5	18	2.6	2.1	17	16.3%	-0.30 [-1.52, 0.92]					
Wang 2016	2.6	1.35	53	2.5	1.48	54	83.7%	0.10 [-0.44, 0.64]					
Total (95% CI)			71			71	100.0%	0.03 [-0.46, 0.53]					
Heterogeneity: Tau² = Test for overall effect:	-2	-1 (Favours [PCN]) Favours (I	1 RUS]	2								

Figure 5. Comparison between PCN and RUS for outcome duration to defervescence.

	DCN RUS						Mean Difference	Mean Difference	
Study or Subgroup	Mean	SD	Total	Mean SD Total			Weight	IV, Random, 95% CI	IV, Random, 95% Cl
Pearle 1998	4.5	3.7	21	3.2	2.8	21	26.9%	1.30 [-0.68, 3.28]	
Wang 2016	10.25	3.53	53	8.24	2.77	54	73.1%	2.01 [0.81, 3.21]	_
Total (95% CI) Heterogeneity: Tau² = Test for overall effect:	0.00; C Z = 3.46	hi² = 0 5 (P = 0	74 .36, df 0.0005)	= 1 (P =	0.55);	75 ² = 0%	100.0%	1.82 [0.79, 2.85]	-2 -1 0 1 2 Favours PCN Favours RUS

Figure 6. Comparison between PCN and RUS for outcome hospitalisation duration.

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- 2. Hospitalisation duration: Three trials reported the hospitalisation duration^{7,12,14}; however, one trial reported the findings as median (IQR) and thus was not included in the meta-analysis¹⁴ (two trials; 149 participants; MD [95% CI] 1.82 [0.79 to 2.85]; $I^2 = 0\%$; P < 0.001; moderate-quality evidence; Fig. 6; Table 2).
- 3. Quality of life: Three trials reported patient quality of life^{8,14,15}. All trials had a high risk of random sequence generation and allocation concealment.



Figure 7. Comparison between PCN and RUS for outcome QoL - mobility.



Figure 8. Comparison between PCN and RUS for outcome QoL - self-care.

	PC	1	RUS	6		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
de Sousa 2018	5	18	5	32	27.7%	1.78 [0.59, 5.32]	
Joshi 2001	10	13	19	21	38.1%	0.85 [0.61, 1.18]	
Shoshany 2019	17	30	9	45	34.2%	2.83 [1.46, 5.50]	_
Total (95% CI)		61		98	100.0%	1.57 [0.55, 4.53]	
Total events	32		33				
Heterogeneity: Tau ² = Test for overall effect:	0.73; Chi Z = 0.84 (i ² = 15.9 (P = 0.4	99, df = 2 10)	(P = 0.	0003); I ř :	= 87%	0.1 0.2 0.5 1 2 5 10 Favours PCN Favours RUS

Figure 9. Comparison between PCN and RUS for outcome QoL - usual activity.

	PCN RUS					Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	I M-H, Random, 95% CI
de Sousa 2018	4	18	5	32	4.6%	1.42 [0.44, 4.63]]
Joshi 2001	11	13	20	21	68.4%	0.89 [0.69, 1.14]] 📕
Shoshany 2019	16	30	21	45	27.0%	1.14 [0.72, 1.81]	1
Total (95% CI)		61		98	100.0%	0.97 [0.75, 1.26]	⊥ ♦
Total events	31		46				
Heterogeneity: Tau ² =	= 0.01; Ch	i ^z = 2.2	8, df = 2 (P = 0.3	2); I ² = 12	%	
Test for overall effect:	Z=0.22	(P = 0.8	(3)				Favours PCN Favours RUS

Figure 10. Comparison between PCN and RUS for outcome QoL - pain/discomfort.

- a) Mobility: Three trials reported quality of life regarding mobility^{8,14,15} (RR [95% CI] 0.78 [0.25 to 2.48]; RD [95% CI] −0.10 [−0.42 to 0.22]; I²=73%; *P*=0.670; very low-quality evidence; Fig. 7; Table 2).
- b) Self-care: Three trials reported quality of life regarding self-care^{8,14,15} (RR [95% CI] 2.76 [0.55 to 13.85]; RD [95% CI] 0.12 (-0.18 to 0.43]; I²=50%; P=0.220; Fig. 8).
- c) Usual activity: Three trials reported quality of life regarding usual activity^{8,14,15}. (RR [95% CI] 1.57 [0.55 to 4.53]; RD [95% CI] 0.13 [-0.16 to 0.41]; I² = 87%; P = 0.400; very low-quality evidence; Fig. 9; Table 2).
- d) Pain/discomfort: Three trials reported quality of life regarding pain or discomfort^{8,14,15} (RR [95% CI] 0.97 [0.75 to 1.26]; RD [95% CI] 0.00 [-0.13 to 0.13]; I² = 12%; P = 0.830; low-quality evidence; Fig. 10; Table 2).
- e) Anxiety/depression: Three trials reported quality of life regarding anxiety or depression^{8,14,15} (RR [95% CI] 0.81 [0.56 to 1.16]; RD [95% CI] 0.12 [– 0.26 to 0.03]; $I^2 = 0\%$; P = 0.250; Fig. 11).
- 4. Urinary-related symptoms:

	PCM	1	RUS	;		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% CI
de Sousa 2018	9	18	19	32	45.5%	0.84 [0.49, 1.45]	
Joshi 2001	7	13	12	21	34.4%	0.94 [0.50, 1.76]	
Shoshany 2019	6	30	16	45	20.1%	0.56 [0.25, 1.27]	
Total (95% CI)		61		98	100.0%	0.81 [0.56, 1.16]	•
Total events	22		47				
Heterogeneity: Tau ² = Test for overall effect:	0.00; Chi Z = 1.15 (i ^z = 1.00 (P = 0.2	3, df = 2 (:5)	P = 0.5	8); I² = 0%	b	0.01 0.1 1 10 100 Favours PCN Favours RUS

Figure 11. Comparison between PCN and RUS for outcome QoL - anxiety/depression.

	PCN		RUS	5		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% Cl
Ahmad 2013	9	200	10	100	17.3%	0.45 [0.19, 1.07]	
de Sousa 2018	3	18	22	32	12.6%	0.24 [0.08, 0.70]	_
Joshi 2001	8	13	18	21	37.7%	0.72 [0.45, 1.14]	
Shoshany 2019	11	30	25	45	32.4%	0.66 [0.39, 1.13]	
Total (95% CI)		261		198	100.0%	0.56 [0.37, 0.85]	•
Total events	31		75				
Heterogeneity: Tau ² = Test for overall effect:	: 0.06; Chi Z = 2.72 (i ² = 4.6: (P = 0.0	5, df = 3 (106)	P = 0.2	0); I² = 35	%	0.01 0.1 1 10 100 Favours PCN Favours RUS

Figure 12. Comparison between PCN and RUS for outcome haematuria.



Figure 13. Comparison between PCN and RUS for outcome dysuria.

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	PCN RUS					Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% CI	M-H, Random, 95% Cl
de Sousa 2018	8	18	24	32	66.3%	0.59 [0.34, 1.03]	
Joshi 2001	3	13	17	21	33.7%	0.29 [0.10, 0.79]	
Total (95% CI)		31		53	100.0%	0.46 [0.23, 0.94]	-
Total events	11		41				
Heterogeneity: Tau ² = Test for overall effect:	0.11; Chi Z = 2.15 (i ² = 1.6 (P = 0.0	%	0.1 0.2 0.5 1 2 5 10 Favours PCN Favours RUS			

Figure 14. Comparison between PCN and RUS for outcome frequency.

- a) Haematuria: Four trials reported haematuria post-procedure^{8,10,14,15}. All trials had high risk of random sequence generation and allocation concealment (RR [95% CI] 0.56 [0.37 to 0.85]; RD [95% CI] -0.24
 - [-0.47 to -0.01]; I² = 35%; P < 0.001; very low-quality evidence; Fig. 12; Table 2).
- b) Dysuria: Two trials reported dysuria^{8,15} (RR 0.28, 95% CI 0.15 to 0.54; RD − 0.61, 95% CI − 0.78 to − 0.43; I² = 0%; P < 0.001; moderate-quality evidence; Fig. 13; Table 2).
- c) Frequency: Two trials reported urinary frequency post-procedure^{8,15} (RR 0.46, 95% CI 0.23 to 0.94; RD − 0.44, 95% CI − 0.71 to − 0.17; I² = 47%; P = 0.030; Fig. 14).
- d) Urgency: Two trials reported urinary urgency^{8,15} (RR 0.56, 95% CI 0.15 to 2.13; RD − 0.37, 95% CI − 0.90 to 0.17; I² = 91%; P = 0.400; Fig. 15).



Figure 15. Comparison between PCN and RUS for outcome urgency.

	PCN		RUS			Risk Ratio	Risk Ratio		
Study or Subgroup	Events Total Even			Total	Weight	M-H, Random, 95% Cl		M-H, Random, 95% Cl	
Ahmad 2013	5	200	3	200	51.8%	1.67 [0.40, 6.88]			
Mokhmalji 2001	0	20	4	20	25.7%	0.11 [0.01, 1.94]			
Pearle 1998	1	21	0	21	22.5%	3.00 [0.13, 69.70]			
Total (95% CI)		241		241	100.0%	0.95 [0.16, 5.58]			
Total events	6		7						
Heterogeneity: Tau ² =	: 1.05; Chi	i ^z = 3.3!	9, df = 2 (P = 0.1	8); l ² = 41	%			
Test for overall effect:	Z=0.06	(P = 0.9	15)				0.005	Favours PCN Favours RUS	21

Figure 16. Comparison between PCN and RUS for outcome failure rate.

	F	/CN		R	US			Mean Difference	Mean Difference		
Study or Subgroup	Mean [VAS]	SD [VAS]	Total	Mean [VAS]	SD [VAS]	Total	Weight	IV, Random, 95% CI [VAS]	IV, Random, 95% CI [VAS]		
Pearle 1998	3.72	7.7605	21	0.65	1.356	21	33.4%	3.07 [-0.30, 6.44]			
Shoshany 2019	1.19	1.52	30	1.02	2.04	45	66.6%	0.17 [-0.64, 0.98]	-#-		
Total (95% CI)			51			66	100.0%	1.14 [-1.54, 3.82]			
Heterogeneity: Tau ² = 2.64; Chi ² = 2.69, df = 1 (P = 0.10); l ² = 63%											
Test for overall effect:	Z = 0.83 (P = 0	Favours PCN Favours RUS									

Figure 17. Comparison between PCN and RUS for outcome post-procedural pain (as measured by VAS).

	PCN		RUS		Risk Difference		Risk Difference
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	M-H, Random, 95% CI
de Sousa 2018	5	18	22	32	24.5%	-0.41 [-0.67, -0.15]	_
Mokhmalji 2001	2	20	7	20	24.8%	-0.25 [-0.50, -0.00]	
Pearle 1998	8	21	1	21	25.3%	0.33 [0.11, 0.56]	
Shoshany 2019	19	30	18	45	25.4%	0.23 [0.01, 0.46]	
Total (95% CI)		89		118	100.0%	-0.02 [-0.37, 0.33]	
Total events	34		48				
Heterogeneity: Tau ² = 0.11; Chi ² = 25.75, df = 3 (P < 0.0001); l ² = 88%							
Test for overall effect:	Z=0.10	(P = 0.9	12)		,,		-1 -0.5 U 0.5 Favours PCN Favours RUS

Figure 18. Comparison between PCN and RUS for outcome of analgesics use.

Secondary outcomes.

1. Failure rates

Three trials reported the number of failure rates for each procedure^{10,29,30}. Two of these trials had a high risk of random sequence generation and allocation concealment^{10,29} (RR [95% CI] 0.95 [0.16 to 5.58]; RD [95% CI] -0.02 [-0.13 to 0.09]; $I^2 = 41\%$; P = 0.950; Fig. 16).

2. Post-procedural pain (VAS)

Two trials reported quantitative assessment of post-procedural pain using VAS^{14,30} (two trials; 117 participants; MD [95% C1] 1.14 (-1.54 to 3.82); $I^2 = 63\%$; P = 0.400; Fig. 17).

3. Analgesics use

Four trials reported analgesics use to alleviate post-procedural pain^{14,15,29,30}. Three trials had a high risk of random sequence generation and allocation concealment^{14,15,29} (RR [95% CI] 0.95 [0.31 to 2.93]; RD [95% CI] -0.02 [-0.37 to 0.33]; I^2 = 81%; P = 0.920; Fig. 18).

Discussion

This review aimed to compare the efficacy and safety of PCN and RUS for the treatment of acute obstructive uropathy based on available RCTs and CCTs. Consequently, only a few trials could be included in this metaanalysis for each outcome. PCN was associated with an increase in the hospitalisation duration compared with RUS. However, for outcomes of urinary-related symptoms, the findings were in favour of PCN, whereby more patients in the RUS group experienced haematuria and dysuria. These findings may affect the decision-making process of choosing these treatment options because although PCN may lead to longer hospitalisation duration, it has a reduced risk of causing a deleterious effect on the patient's quality of life because of haematuria or dysuria after discharge to home. There were no differences regarding WBC normalisation duration, duration to deferves-cence, all five components of quality of life, urinary urgency and frequency, failure rates, quantitative assessment of pain post-procedure and analgesics use. Only one trial reported the comparison of the radiation dose received by patients undergoing these procedures; thus, the evidence is not conclusive. However, this is an important outcome that should be considered because exposure to radiation has substantial adverse effects on patients.

We conducted a thorough and extensive search of the databases and included seven trials that compared PCN and RUS. Stone disease constitutes the largest proportion of the obstruction cause in the included trials. Hence, the findings may not be applicable to other causes of obstructive uropathy, such as extrinsic obstruction due to malignancy or bladder abnormalities secondary to neurogenic bladder.

All included trials had a high risk of performance bias because of the surgical nature of the interventions, wherein patients are required to receive a detailed explanation and provide consent before the procedures are initiated, rendering the blinding of participants and personnel infeasible. However, this situation reflects the real-life scenario encountered in daily practice. Furthermore, most of the included trials also had a high risk of selection bias. Of these trials, five had a high risk of bias regarding random sequence generation, of which four also had a high risk of allocation concealment. In these trials, the decision to conduct either PCN or RUS was made solely on the preference of the performing surgeon based on experience or local protocol. A high risk of attrition bias was present in one trial, wherein patients who had an unsuccessful RUS were subjected to PCN.

We determined that the quality of evidence for the main comparisons ranged from moderate to very low. The quality of evidence was judged to be very low for quality of life outcomes regarding the impact of the procedures on mobility and usual activity of the patients and any occurrence of haematuria. This judgement was due to the high risk of performance, detection and selection bias of the relevant trials assessing these outcomes. We also detected a high heterogeneity among trials in these outcomes, which contributed to the reduced certainty of evidence in the findings.

We conducted an extensive literature search using multiple databases and by checking the reference lists of all related trials to minimise publication bias. For one trial, we were unable to analyse the data for the outcomes of duration to defervescence and hospitalisation duration because the data were presented in a non-usable format.

One systematic review involving six trials comparing the use of PCN with RUS for managing ureteral obstruction in benign and malignant aetiologies reported no difference regarding the time to clinical improvement in both PCN and RUS³¹. However, PCN was associated with a higher risk of post-procedure bacterial colonisation. Additionally, both procedures were also reported to have a high success rate. RUS insertion failure was reported to be amenable with PCN, but the contrary was not true. Regarding the quality of life assessment, this review reported that both procedures were associated with worsening quality of life. Nevertheless, no significant difference was reported.

In comparison with our review, most of these findings had an overall agreement with our data. However, a few exceptions were noted. The hospitalisation duration was significantly shorter in patients receiving RUS than in those receiving PCN, but PCN was superior regarding reported post-procedural haematuria and dysuria. Of the six trials in the previous review, four similar trials were included in our review^{8,10,29,30}. The other two trials were excluded because of different reported outcomes of interest.

We recognised several limitations of our review. Firstly, there are limited number of trials conducted to compare the efficacy and safety of PCN versus RUS in the treatment of acute upper obstructive uropathy. Some of the outcomes could only be analysed based on data extracted from two studies. These include improvement in septic parameters, hospitalisation duration, urinary-related symptoms i.e. dysuria, frequency and urgency, and post-procedural pain. Secondly, there is a lack of RCTs conducted to compare these two types of intervention. We only found two RCTs to be included in our study. This is probably attributable to the urgent nature of the procedures rendering randomization difficult. Thirdly, all included trials contained high risk of performance bias. This was due to the surgical nature of the interventions which require written informed consent prior to the procedures. Hence, blinding of participants and personnel was not feasible.

In conclusion, we found no significant difference between PCN and RUS regarding improvement in septic parameters or quality of life post-procedure. Both methods are safe with high success rates. In clinical practice, we recommend the use of PCN rather than RUS, because the reduced quality of life due to haematuria and dysuria is much more devastating to patients over the long term. If further research is undertaken to compare PCN and RUS for the treatment of acute obstructive uropathy, we suggest the inclusion of radiation exposure and improvement of renal function as the main comparison.

Methods

Eligibility criteria. *Types of studies.* We included randomised controlled trials (RCTs) and controlled clinical trials (CCTs) comparing PCN and RUS.

Types of participants. We included patients with acute obstructive uropathy secondary to any cause who received PCN or RUS.

Types of interventions. We included studies on PCN conducted by a physician, using any size of nephrostomy tube, under ultrasound or fluoroscopy guidance with RUS as the comparison.

Types of outcome. We reviewed data on clinical success rates, described as improvement of septic parameters regarding duration to defervescence and normalisation of white blood cell count and hospitalisation duration. We evaluated information on quality of life and analysed data on urinary-related symptoms including haematuria, dysuria, frequency and urgency. For secondary outcomes, we reviewed information regarding failure rates of each procedure, post-procedural pain and analgesics use.

Search strategies. We searched the Cochrane Central Register of Controlled Trials CENTRAL (latest issue), MEDLINE, EMBASE and CINAHL on 16 October 2019. A repeat search for trials that was performed on the 16 February 2021 yielded no additional trials that fulfil our inclusion criteria. The search strategy is described in Appendix 1. We adopted the search strategy for other databases. We restricted the publications to English language only.

We checked the reference list of identified RCTs, CCTs and review articles to find unpublished trials or trials not identified by electronic searches. We searched for ongoing trials through the World Health Organisation International Clinical Trials Registry Platform (http://www.who.int/ictrp/en/ and www.clinicaltrials.gov).

Trial selection. We screened the titles and abstracts from the searches and obtained the full text of articles that appeared to meet the eligibility criteria or in instances wherein information was insufficient to determine study eligibility. We evaluated the eligibility of the trials independently and documented the reasons for exclusion. We resolved any disagreements between the review authors by discussion.

Data extraction. Using a data extraction form, from each of the selected trials we extracted study settings, participant characteristics (age, sex and ethnicity), methodology (number of participants randomised and analysed), white blood cell count (WBC) normalisation duration, duration to defervescence, hospitalisation duration, quality of life based on questionnaires, urinary-related symptoms (e.g. haematuria, dysuria, urgency and frequency), technical success of each procedure to evaluate failure rate, pain score based on a visual analogue scale (VAS) and analgesics use.

Risk of bias assessment. We assessed the risk of bias based on random sequence generation, allocation concealment, blinding of participants and personnel, blinding of outcome assessors, completeness of outcome data, selectivity of outcome reporting and other bias³². We resolved any disagreements by discussion.

Grading quality of evidence. We assessed the quality of evidence for primary and secondary outcomes on the basis of the GRADE methodology for risk of bias, inconsistency, indirectness, imprecision and publication bias. The risk of bias was classified as very low, low, moderate or high³³.

Statistical analyses. We performed meta-analyses using Review Manager 5.3 software (RevMan 2014) and used the random effects model to pool data. We assessed the presence of heterogeneity in two steps. First, we assessed apparent heterogeneity at face value by comparing populations, settings, interventions and outcomes. Second, we assessed statistical heterogeneity using the I^2 statistic³².

Thresholds for the interpretation of the I² statistic can be misleading because the importance of inconsistency depends on several factors. We used the guide to the interpretation of heterogeneity as outlined: 0%-40%might not be important, 30%-60% may represent moderate heterogeneity, 50%-90% may represent substantial heterogeneity and 75%-100% would be considerable heterogeneity³².

We measured the treatment effect for dichotomous outcomes using risk ratios (RRs) and risk difference (RD) and for continuous outcomes using mean differences (MDs), both with 95% confidence intervals (CIs). No subgroup analysis was conducted in this meta-analysis.

We checked the included trials for the unit of analysis errors. These errors can occur when trials randomise participants to intervention or control groups in clusters but analyse the results using the total number of individual participants. We adjusted the results from trials showing the unit of analysis errors on the basis of the mean cluster size and intra-cluster correlation coefficient³².

We contacted the original trial authors to request missing or inadequately reported data. We conducted analyses of the available data when missing data were not available.

We conducted a sensitivity analysis to investigate the impact of the risk of bias for sequence generation and allocation concealment of included studies.

Ethical standard. The manuscript does not contain original clinical studies.

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Author contributions

Z.K.A.I.: protocol development, data collection, manuscript writing/editing. N.M.N.: protocol development, manuscript writing/editing. M.SA.: manuscript writing/editing.

Competing interests

The authors declare no competing interests.

Additional information

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