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Sepsis

Sepsis: recognition, assessment and early management

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Disclaimer

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Appendices

Appendix I: Economic evidence tables

I.1 Scoring systems

None.

I.2 Signs and symptoms

None.

I.3 Blood tests

None.

I.4 Lactate

None.

I.5 Serum creatinine

None.

I.6 Disseminated intravascular coagulation

None.

I.7 Antimicrobial treatment

None.

I.8 IV fluid administration

None.

I.9 Escalation of care

None.

I.10 Inotropic agents and vasopressors

None.

I.11 Supplemental oxygen

None.

I.12 Use of bicarbonate

None.

I.13 Early goal-directed therapy (EGDT)

Study	Mouncey 2015 ⁸²⁷			
Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
Economic analysis: CUA (health outcome: QALYs)	Population: Patients with early signs of	Total costs (mean per patient):	QALYs (mean per patient): Intervention 1: 0.054	ICER (Intervention 2 versus Intervention 1): Intervention 2 dominated (more expensive

Study design: Within trial analysis (RCT)

Approach to analysis: Analysis of individual level data for mortality and EQ-5D. Unit costs were applied to resource use.

Perspective: UK NHS

Time horizon/Follow-up 90 days QoL follow up Treatment effect duration: Resuscitation protocol was followed for 6 hours

Discounting: Costs: NR; Outcomes: NR

septic shock

N = 1251

Patient characteristics:

Mean age: invtn 1 = 64.3 (15.5), intvn 2 = 66.4 (14.6) Male: invtn 1 = 58.6%, intvn 2 = 57%

Intervention 1:

Usual care
The usual care group
continued to receive
monitoring, investigation
and treatment as

determined by the clinician.

Intervention 2:

Early Goal Directed Therapy (EGDT).

Following a resuscitation protocol involving central venous catheter insertion with central venous oxygen saturation monitoring capability and intensive therapy of other interventions

Intervention 1: £11,424 Intervention 2: £12,414 Incremental (2–1): £989 (95% CI: -726 to 2,705; p=NR)

Currency & cost year: 2012 UK pounds

Cost components incorporated:

- Equipment and consumables 2 monitors capable of oxygen saturation monitoring assumed to be needed per hospital. Costs of consumables including the catheter capable of monitoring, pressure transducers.
- Blood products and dobutamine
- Staff time to deliver the protocol; time for vascular catheter insertion and time for monitoring patients (assumed 10 minutes of nurse time per hr of the resus protocol). Staff time for training, assumed to be 20 minutes per ED staff member every 5 years (5 years assumed to be the life

Intervention 2: 0.054 Incremental (2–1): -0.001 (95% CI: -0.006 to 0.005); p=0.85) and less benefit)

Probability Intervention 2 cost-effective (£20K/30K threshold): 12%/12% (read from graph)

Analysis of uncertainty:

Some form of PSA undertaken ^(a) to generate cost effectiveness plane and cost effectiveness acceptability curve. 500 estimates obtained.

Sensitivity analyses undertaken include:

- Manufacturer list price used for monitoring machines instead of discounted price used in base case
- Staff monitoring time varied from 10 minutes per hour in the base case to 5 and 15 minutes.
- Location of protocol implementation; if protocol is implemented in the ED, staffneed to be trained but in critical care they do not. Sensitivity analysis assumed that the protocol was implemented either exclusively in the ED or critical care.
- Re-admission data in the base case was gathered both from the health services questionnaire sent out and the Intensive Care National Audit & Research Centre Case Mix Programme Database. In a sensitivity analysis only the database was used to avoid any potential double counting.
- Baseline covariates were adjusted for components of the Mortality in Emergency Department Sepsis (MEDS) score

of the protocol) - Hospital stay/ICU stay - Re-admissions	- Costs and QALYs were assumed to be gamma distributed, compared to normally distributed in the base case.
	EGDT remained cost-ineffective in all sensitivity analyses.

Data sources

Health outcomes: Mortality data taken from the RCT (proMISe trial) alongside the economic evaluation.

Quality-of-life weights: EQ-5D scores were elicited at 90 days, assuming an EQ-5D score of zero at randomisation, and a linear interpolation between randomisation and 90 days. Zero QALYs were assumed for people who died before 90 days.

Cost sources: Costs of monitor and central venous catheter with monitoring capability was derived from the manufacturer. These costs are over 50% discount on list prices. It was assumed each site would require 2 monitors which would have a lifespan on average of 5 years. Monitor costs per patient were calculated by dividing the total costs of the monitors (£4000) by the expected number of eligible patients over 5 years. Annual number of eligible patients calculated by taking average number of potentially eligible patients per site per year from the trial screening log data (23 patients per site per year). Some consumables sourced from hospital finance departments. Training costs per patient per hour derived from total training costs per site divided by eligible patients over 5 years. Blood products from NHS blood and transplant price list 2012. Drugs from BNF 2012. Staff costs and outpatient and community health service costs from PSSRU 2012. Hospital stay costs from NHS reference costs 2012.

Comments

Source of funding: NR **Limitations:** Adverse events not taken account of in cost effectiveness analysis (either their treatment costs or impact on QoL). Methodology behind probabilistic analysis unclear. Short time horizon.

Overall applicability(d): Directly applicable Overall quality: potentially serious limitations

Abbreviations: 95% CI: 95% confidence interval; 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; NR: not reported; pa: probabilistic analysis; PSSRU: Personal Social Services Research Unit; QALYs: quality-adjusted life years (a) The paper states incremental costs and QALYs were estimated using 'a seemingly unrelated regression model', and they used 'the estimates of the means, variances and the covariance from the regression model to generate 500 estimates of incremental costs and QALYs from the joint distribution of these endpoints'. By generating a cost effectiveness plane and cost effectiveness acceptability curve this implies some kind of probabilistic analysis was done but the methodology quoted isn't clear.

I.14 Monitoring

None.

I.15 Patient education, information and support

None.

I.16 Training and education

Study	Suarez 2011 ¹⁰⁷⁶			
Study details	Population & interventions	Costs	Health outcomes	Cost-effectiveness
Economic analysis: CEA/CUA (health outcome: Life Years Gained and QALYs) Study design: Within trial analysis Approach to analysis: Preeducation program cohort (2 months before program) was compared to a post education program cohort (4 months after program). Program consisted of a 2 month educational program of training physicians and nursing staff from the emergency department, medical, and surgical wards, and ICU in early recognition of severe sepsis and the treatments in the Surviving Sepsis Campaign (SSC) protocol. Unit costs applied to prospective study data. Multivariable regression models were used to adjust for baseline	Population: Patients with severe sepsis Patient characteristics: N = 2319 (b) Mean age = 62.2 (SD: 16.3) Male = 60.8% Intervention 1: Pre-intervention cohort, the 2 months prior to the educational program Intervention 2: Post intervention cohort, the 4 months following educational program.	Total costs (mean per patient): Intervention 1: £14,427 Intervention 2: £15,906 Incremental (2–1): £1,479 (95% CI: NR; p=NR) Currency & cost year: 2006 Spanish Euros presented here as 2006 UK pounds ^(c) Cost components incorporated: Unit costs for emergency visits, surgical and medical ward daily stays, and ICU daily stays. Cost associated with the pharmacological and non- pharmacological interventions of the SSC protocol. One of the goals of the SSC protocol is	QALYs (mean per patient): Intervention 1: 3.75 Intervention 2: 4.12 Incremental (2–1): 0.37 (95% CI: 0.02-0.73; p=NR) Life Years Gained (mean per patient): Intervention 1: 5.44 Intervention 2: 5.98 Incremental (2–1): 0.54 (95% CI: 0.02-1.05; p=NR)	ICER (Intervention 2 versus Intervention 1): £5,476 per QALY gained (the 'adjusted' ICER) (pa) 95% CI: NR Probability Intervention 2 cost-effective (£20K threshold): 94% (read off graph) Probabilistic analysis was undertaken using non parametric bootstrapping with 2000 replications. Analysis of uncertainty: One way sensitivity analyses undertaken include: - Changing the rate for sepsis survivors from 0.51 to 0.39. Making this value even more restrictive Quality of life weight was changed from 0.69 to 0.75 The ICER was also calculated for different utility values. Only for very low utility values (lower than 0.2) was the ICER more than £20,000 (read off graph) Discounting of Life Years Gained and

Economic evidence tables

differences of costs, QALYs, and Life Years Gained. Perspective: Spanish healthcare system perspective. Time horizon/Follow-up: Post intervention cohort was a 4 month period after intervention introduced. Costs were only considered up until hospital discharge. Lifetime horizon for life years. Treatment effect duration: Discounting: Costs: NA; Health maintaining glucose control; the average cost per patient reported in a cost effectiveness analysis of insulin therapy was used. Patients who achieved the goal were applied the cost of the intensive therapy group, and patients who did not meet the goal were applied the cost of the conventional therapy group.	QALYs was changed from 3% to 0%. - Discounting of Life Years Gained and QALYs was changed from 3% to 5%. - The cost of the education and training program and cost of staff time spent attending the sessions was not included in base case. These costs were included in a sensitivity analysis. All sensitivity analyses generated results similar to that of the base case.
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outcomes: 3% Data sources

Health outcomes: Mortality and resource use data derived from a cohort before and after study (Ferrer 2008 ⁴⁵⁰). Age and gender specific life expectancy for each survivor taken from the 2006 Spanish like expectancy tables. These were adjusted using the estimated reduction rate for sepsis survivors of 0.51 ⁹⁴⁵.

Quality-of-life weights: The quality of lie weight used was 0.69. This utility weight was obtained from a study of 6 month survivors of severe sepsis using the EQ-5D. 395

Cost sources: unit costs for emergency visits, surgical and medical ward daily stays, and ICU daily stays were from the Spanish National Health Institute. Pharmacological intervention costs from the SSC protocol were from the Spanish physician's desk reference. Non-pharmacological intervention costs were obtained from their suppliers. Insulin therapy cost was the average cost per patient from a cost effectiveness study on insulin therapy¹¹²² (€144 for intensive therapy and €72 for conventional therapy). All prices in the study were adjusted to 2006 values using the Spanish consumer price index. Long term costs after discharge were not included. The costs of the training program were not included in the base case, but were included in a sensitivity analysis (€54,270).

Comments

Source of funding: Supported by a grant from the Instituto de Salud Carlos III. Limitations: Only includes short term costs. Data on effectiveness from a cohort study, not RCT. Base case did not include cost if the intervention itself. Methodology not always clear; particularly around where adjusted ICER comes from. Other: The paper states that both the incremental costs and incremental QALYs/Life Years Gained were 'obtained by adjusting multivariable regression models to take into account possible baseline imbalances'. The ICER that is reported in the study is stated to be the 'adjusted ICER' (6,428 Euros or £5,476). It is unclear whether the 'adjusted' ICER reported is the deterministic or probabilistic ICER, however the paper states the ICER in the text (as well as a table) then immediately in the next sentence states that

nearly all the bootstrap replications were below the threshold used of 30,000 euros. Thus implying this is likely to be the probabilistic ICER.

Overall applicability: Partially applicable^(d) Overall quality^(e) Potentially serious limitations

Abbreviations: CEA: cost-effectiveness analysis; 95% CI: 95% confidence interval; CUA: cost-utility 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; NR: not reported; pa: probabilistic analysis; QALYs: quality-adjusted life years

- (a) The post intervention cohort are those that would benefit from the 'treatment effect' of the education program. This cohort included patients during the 4 month period after the intervention. The time horizon for health outcome was lifetime so life expectancy was applied to the survivors. Therefore there is an assumption being made about the continuation of the study effect because life years will continue to vary between arms as different numbers of people will be alive in the pre and post intervention cohorts. The utility being applied to the groups is the same because the utility is the utility of sepsis survivors and is not impacted by the intervention except by the impact on mortality.
- (b) Note that the study this economic evaluation is based on is included in the clinical review (Ferrer2008) and the number of patients included in the study is higher than that reported here because there was also a third observation period (one year after the pre intervention group, to test the longevity of the education program) included in the clinical paper that is separate to the pre and post intervention cohorts.
- (c) Converted using 2006 purchasing power parities 883
- (d) Directly applicable / Partially applicable / Not applicable
- (e) Minor limitations / Potentially serious limitations / Very serious limitations

Appendix J: GRADE tables

J.1 Scoring systems

None.

J.2 Signs and symptoms

None.

J.3 Blood tests

None.

J.4 Lactate

None.

J.5 Serum creatinine

None.

J.6 Disseminated intravascular coagulation (DIC)

Table 1: Disseminated intravascular coagulation (DIC) and all-cause mortality

			Quality asse	essment			No of patients		Effect		Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	DIC	Control	OR (95% CI)	Abcoluto		
28-day morta	8-day mortality - Gando 2008											
		,	no serious inconsistency	very serious ²	no serious imprecision	none	65	264	1.22 (1.00 to 1.49)	_4	VERY LOW	CRITICAL
28-day morta	ality - Gando 2013						•					
		,	no serious inconsistency	no serious indirectness	no serious imprecision	none	292	332	1.28 (1.14 to 1.44)	_4	VERY LOW	CRITICAL
28-day morta	ality - Ogura 2014											
			no serious inconsistency	no serious indirectness	no serious imprecision	none	292	332	1.73 (1.09 to 2.75)	_4	VERY LOW	CRITICAL
In-hospital n	nortality - Gando 20	007										
1		,	no serious inconsistency	no serious indirectness	serious ³	none	11	34	4.22 (1.42 to 12.59)	_4	VERY LOW	CRITICAL
n-hospital mortality - Gando 2007A												

		very serious ¹		no serious indirectness	serious ³	none	20	28	40.50 (4.54 to 360.98)	_4	VERY LOW	CRITICAL
In-hospital mortality - Ogura 2014												
		very serious ¹			no serious imprecision	none	292	332	1.55 (1.01 to 2.37)	_4	VERY LOW	CRITICAL

¹ Risk of bias mainly due to the lack of evidence that physicians treating patients were blinded to the DIC status. The assumed lack of blinding means that knowledge of DIC could affect treatment, which would possibly affect outcome.

J.7 Antimicrobial treatment

Table 2: <1 hour versus >1 hour (adult population)

			Quality asse	No of patients	Effect		Quality	Importance					
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	<1h versus >1h (multivariable analysis)	Control	OR (95% CI)	Absolute		•	
Mortality													
	observational studies				no serious imprecision	none	-	-	OR 0.87 (0.81 to 0.94)	_2	VERY LOW	CRITICAL	
Mortality -	lortality - ICU setting												

² The majority of the evidence included an indirect population (downgraded by one increment) or a very indirect population (downgraded by two increments)

³ Downgraded by 1increment due to a very imprecise result expressed by a very wide confidence interval

⁴ N/A as only adjusted or unadjusted OR was provided

5	observational studies				no serious imprecision	none	-	-	Not estimable	_2	VERY LOW	CRITICAL		
Mortality -	Mortality - ED setting													
3	observational studies			no serious indirectness	serious ³	none	-	-	Not estimable	_2	VERY LOW	CRITICAL		

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 Absolute effect not estimable as the crude event rate for the control group was not provided ³ Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 3: <2 hours versus >2 hours (adult population)

			No of patients	Effec	et	Quality	Importance					
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	<2h versus >2h (multivariable analysis)	Control	OR (95% CI)	Absolute		Importance
Mortality												
	observational studies			no serious indirectness	serious ²	none	-	-	OR 0.73 (0.51 to 1.04)	_3	VERY LOW	CRITICAL
Mortality -	Mortality - ICU setting											

	observational studies			no serious indirectness	serious ²	none	-	-	OR 0.14 (0.02 to 0.88)	_3	VERY LOW	CRITICAL		
Mortality -	Mortality - ED setting													
	observational studies				no serious imprecision	none	-	-	OR 0.78 (0.54 to 1.12)	_3	VERY LOW	CRITICAL		

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs ³ Absolute effect not estimable as the crude event rate for the control group was not provided

Table 4: <3 hours versus >3 hours (adult population)

		Quality assess		No of patients		Effect		Quality	Importance			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	<3h versus >3h (multivariable analysis)	Control	OR (95% CI)	Absolute		importance
Mortality	Mortality											
	observational studies			no serious indirectness	serious ²	none	-	-	OR 0.7 (0.57 to 0.86)	_3	VERY LOW	CRITICAL
Mortality -	Mortality - ICU setting											
1	observational	serious ¹	no serious	no serious	serious ²	none	-	-	OR 0.8 (0.6 to	_3	VERY	CRITICAL

		studies	inconsistency	indirectness					1.07)		LOW	
M	lortality -	ED setting										
5		observational studies		no serious indirectness	serious ²	none	-	-	OR 0.62 (0.47 to 0.82)	_3	VERY LOW	CRITICAL

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs ³ Absolute effect not estimable as the crude event rate for the control group was not provided

Table 5: <4 hours versus >4 hours (adult population)

			No of patients		Effec	t	Quality	Importance				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	<4h versus >4h (multivariable analysis)	Control	OR (95% CI)	Absolute		mportance
Mortality	Nortality											
	observational studies	serious ¹			very serious ²	none	3/25 (12%)	-	OR 0.86 (0.49 to 1.53)	_3	VERY LOW	CRITICAL
Mortality -	Mortality - ED setting											
	observational studies	serious ¹			very serious²	none	-	-	OR 0.86 (0.49 to 1.53)	_3	VERY LOW	CRITICAL

Table 6: <5 hours versus >5 hours (adult population)

Tuble 0.	13 110013 VC1		No of patients		Effec	t	Quality	Importance				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	<5h versus >5h (multivariable analysis)	Control	OR (95% CI)	Absolute		importance
Mortality												
	observational studies		no serious inconsistency		very serious²	none	-	-	OR 0.65 (0.26 to 1.62)	_3	VERY LOW	CRITICAL
Mortality -	ED setting											
	observational studies	serious ¹	no serious inconsistency		very serious²	none	-	-	OR 0.65 (0.26 to 1.62)	_3	VERY LOW	CRITICAL

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

Table 7: <6 hours versus >6 hours (adult population)

Quality assessment	No of patients	Effect	Quality Importan	ıce
--------------------	----------------	--------	------------------	-----

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs ³ Absolute effect not estimable as the crude event rate for the control group was not provided

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

³ Absolute effect not estimable as the crude event rate for the control group was not provided

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	<6h versus >6h (multivariable analysis)	Control	OR (95% CI)	Absolute		
Mortality												
3	observational studies			no serious indirectness	serious²	none	-	-	OR 0.72 (0.58 to 0.9)	_3	VERY LOW	CRITICAL
Mortality -	ICU setting											
2	observational studies	serious ¹	serious ⁴	no serious indirectness	serious ²	none	-	-	OR 0.79 (0.57 to 1.08)	_3	VERY LOW	CRITICAL
Mortality -	ED setting											
1	observational studies			no serious indirectness	serious²	none	-	-	OR 0.67 (0.5 to 0.9)	_3	VERY LOW	CRITICAL

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs ³ Absolute effect not estimable as the crude event rate for the control group was not provided

Table 8: Hourly treatment delay (ICU, adult population)

Quality assessment	No of patients	Effect	Quality	Importance

⁴ I2=60% (p=0.11)

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Hourly treatment delay (ICU)	Control	OR (95% CI)	Absolute		
In-hospital	mortality											
	observational studies				no serious imprecision	none	-	-	OR 1.12 (1.1 to 1.14)	_2	⊕000 VERY LOW	CRITICAL

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 Absolute effect not estimable as the crude event rate for the control group was not provided

Table 9: Parenteral antibiotics prior to admission to hospital

			Quality assess		No of patients		Effec	ct	Quality	Importance		
No of studies	Design Inconsistency Indirectness Imprecision					Other considerations	Parenteral antibiotics prior to admission to hospital (GP)	Control	OR (95% CI)	Absolute	•	mportance
Mortality												
4	observational studies	,			very serious ²	none	-	-	OR 0.58 (0.21 to 1.58)	_3	VERY LOW	CRITICAL

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

³ Absolute effect not estimable as the crude event rate for the control group was not provided

Table 10: <1 hour versus >1 hour (PICU, paediatric population)

			Quality asses		No of patients Effect				Quality	Importance		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Parenteral antibiotics prior to admission to hospital (GP)	Control	OR (95% CI)	Absolute		•
Mortality												
4				no serious indirectness	very serious²	none	-	-	OR 0.6 (0.13 to 2.86)	_3	VERY LOW	CRITICAL

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs ³ Absolute effect not estimable as the crude event rate for the control group was not provided

Table 11: <2 hours versus >2 hours (PICU, paediatric population)

			Quality assess		No of patients Effect			et	Quality	Importance		
No of studies	Design Inconsistency Indirectness Imprecision					Other considerations	Parenteral antibiotics prior to admission to hospital (GP)	Control	OR (95% CI)	Absolute	•	
Mortality												
la					very serious²	none	-	-	OR 0.41 (0.13 to 1.35)	_3	VERY LOW	CRITICAL

Table 12: <3 hours versus >3 hours (PICU, paediatric population)

			Quality asses		No of patients Effect				Quality	Importance		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Parenteral antibiotics prior to admission to hospital (GP)	Control	OR (95% CI)	Absolute		
Mortality	ortality											
4				no serious indirectness	serious ²	none	-	-	OR 0.25 (0.08 to 0.79)	_3	VERY LOW	CRITICAL

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

Table 13: <4 hours versus >4 hours (PICU, paediatric population)

			Quality asses	sment			No of patients		Effe	ct	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Parenteral antibiotics prior to admission to hospital (GP)	Control	OR (95% CI)	Absolute		importance
Mortality												
1		very serious¹		no serious indirectness	serious ²	none	-	-	OR 0.28 (0.1	_3	VERY	CRITICAL

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

³ Absolute effect not estimable as the crude event rate for the control group was not provided

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

³ Absolute effect not estimable as the crude event rate for the control group was not provided

				to 0.81)	LOW	
					1 1	

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

J.8 IV fluid administration

Table 14: Clinical evidence profile: 6% HES versus 0.9% saline in adults with sepsis

		·	Quality asse	ssment		·	No of patier	nts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	6% HES versus 0.9% saline	Control	Relative (95% CI)	Absoluta		
90-day mortality												
1	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	serious²	none	248/976 (25.4%)	224/945 (23.7%)	`	17 more per 1000 (from 19 fewer to 59 more)	LOW	CRITICAL

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

Table 15: Clinical evidence profile: Crystalloid versus colloid plus crystalloid in adults with severe sepsis

Quality assessment	No of patients	Quality	Importance

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

³ Absolute effect not estimable as the crude event rate for the control group was not provided

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

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Crystalloid versus colloid + crystalloid	Control	Relative (95% CI)	Absolute			
Hospital r	ospital mortality												
		very serious ¹	no serious inconsistency	no serious indirectness	serious ²	none	101/235 (43%)	121/258 (46.9%)		38 fewer per 1000 (from 117 fewer to 56 more)	VERY LOW	CRITICAL	
ICU morta	ICU mortality												
		very serious ¹	no serious inconsistency	no serious indirectness	very serious²	none	72/235 (30.6%)	99/258 (38.4%)		77 fewer per 1000 (from 146 fewer to 8 more)	VERY LOW	CRITICAL	

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 16: Clinical evidence profile: 20% albumin versus 6% HES in adults with severe sepsis

			Quality asse	ssment			No of patier	nts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	20% albumin versus 6% HES	Control	Relative Absolute		Quanty	Importance
28-day mortality												

	randomised trials				very serious ²	none	4/30 (13.3%)	6/26 (23.1%)	RR 0.58 (0.18 to 1.83)	97 fewer per 1000 (from 189 fewer to 192 more)	VERY LOW	CRITICAL
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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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 17: Clinical evidence profile: 4% albumin versus 0.9% Sodium Chloride RP in adults with severe sensis

Table 17	able 17: Clinical evidence profile: 4% albumin versus 0.9% Sodium Chloride BP in adults with severe sepsis													
			Quality asse	essment			No of patients			Effect				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	4% albumin versus 0.9% Sodium Chloride BP	Control	Relative (95% CI)	Absolute	Quality	Importance		
28-day me	ortality (univa	riate analysis	\$)											
1	randomised trials		no serious inconsistency	no serious indirectness	serious ²	none		217/615 (35.3%)	RR 0.87 (0.74 to 1.02)	46 fewer per 1000 (from 92 fewer to 7 more)	LOW	CRITICAL		
28-day me	ortality (multi	variate analys	sis)											
1	randomised trials		no serious inconsistency		no serious imprecision	none		166/467 (35.5%)	OR 0.71 (0.52 to 0.97)	_3	HIGH	CRITICAL		

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

³ Adjusted odds ratio

Table 18: Clinical evidence profile: Albumin versus crystalloids in adults with sepsis

			Quality asse	essment			No of patie	nts		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Albumin versus crystalloids	Control	Relative (95% CI)	Absolute	quanty	
All-cause	mortality											
1			no serious inconsistency		no serious imprecision	none	710/1937 (36.7%)	763/1941 (39.3%)	RR 0.93 (0.86 to 1.01)	28 fewer per 1000 (from 55 fewer to 4 more)	⊕⊕⊕O MODERATE	CRITICAL
90-day me	ortality							<u>'</u>				
1	randomised trials	very serious ²			no serious imprecision	none	115/283 (40.6%)	116/286 (40.6%)	RR 1 (0.82 to 1.22)	0 fewer per 1000 (from 73 fewer to 89 more)	⊕⊕OO LOW	CRITICAL

Table 19: Clinical evidence profile: Albumin versus colloids in adults with sepsis

			Quality assess	ment			No of patier	nts		Effect	Quality	Importance
No of	Design	Risk of	Inconsistency	Indirectness	Imprecision	Other	Albumin versus	Control Absolute				

¹ Downgraded by 1 increment because of inconsistencies regarding the study population ² 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

studies		bias				considerations	colloids		(95% CI)			
Mortality	1	1										
1	randomised trials		no serious inconsistency	serious ²	serious ³	none		58/156 (37.2%)	RR 1.02 (0.76 to 1.36)	7 more per 1000 (from 89 fewer to 134 more)	VERY LOW	CRITICAL

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

Table 20: Clinical evidence profile: Packed red blood cells (PRBC) plus EGDT versus EGDT only in adults with septic shock

			Quality assess	ment		No of patie	nts		Effect	Quality	Importance		
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	PRBC + EGDT versus EGDT	Control	Relative (95% CI)	Absolute	<u> </u>		
Hospital n	Hospital mortality												
			no serious inconsistency	no serious indirectness	very serious¹	none	14/34 (41.2%)	20/59 (33.9%)		71 more per 1000 (from 98 fewer to 366 more)	VERY LOW	CRITICAL	

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

Table 21: Clinical evidence profile: Red blood cells (RBC) for low threshold (≤7g/dl) versus high threshold (≤9g/dl) in adults with septic shock

Quality assessment	No of patients	Effect	Quality	Importance

Downgraded by 1 increment because of differences regarding the study population
 Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	RBC at low versus high threshold	Control	Relative (95% CI)	Absolute		
90-day mo	ortality											
1			no serious inconsistency		no serious imprecision	none	216/502 (43%)	223/496 (45%)	RR 0.97 (0.84 to 1.11)	13 fewer per 1000 (from 72 fewer to 49 more)	MODERATE	CRITICAL
90-day mo	ortality - >70 y	ears of age										
1	randomised trials		no serious inconsistency		no serious imprecision	none	93/173 (53.8%)	98/185 (53%)	RR 1.01 (0.84 to 1.23)	5 more per 1000 (from 85 fewer to 122 more)	MODERATE	CRITICAL
90-day mo	ortality - 70 ye	ars or young	er									
1	randomised trials		no serious inconsistency		no serious imprecision	none	123/329 (37.4%)	125/311 (40.2%)	RR 0.93 (0.77 to 1.13)	28 fewer per 1000 (from 92 fewer to 52 more)	MODERATE	CRITICAL

¹ Intervention does not fall within the 6-hour time frame

Table 22: Clinical evidence profile: 0-2 litres versus 2-4 litres of fluids in adults with severe sepsis

			Quality assess	sment			No of pat	ients		Effect	Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	0-2L versus 2-4L	Control	Relative (95% CI)	Absolute		

Hospital n	nortality										
1	observational studies	very serious ¹	no serious indirectness	serious ²	none	97/210 (46.2%)	82/186 (44.1%)	•	22 more per 1000 (from 71 fewer to 132 more)	VERY LOW	CRITICAL
ICU morta	lity										
1	observational studies	very serious ¹	no serious indirectness	serious ²	none	66/210 (31.4%)	66/186 (35.5%)		39 fewer per 1000 (from 117 fewer to 60 more)	VERY LOW	CRITICAL

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 23: Clinical evidence profile: 0-2 litres versus >4 litres of fluids in adults with severe sepsis

	Quality assessment									Quality	Importance	
No of studies	studies Design bias Inconsistency Indirectness Imprecision consideration								Relative (95% CI)	Absolute	Quanty	importance
Hospital m	nortality											
	observational studies	very serious ¹		no serious indirectness	serious²	none	97/210 (46.2%)	45/100 (45%)	RR 1.03 (0.79 to 1.33)	13 more per 1000 (from 94 fewer to 149 more)	VERY LOW	CRITICAL
ICU morta	lity											
1	observational	very	no serious	no serious	serious ²	none	66/210	41/100	RR 0.77 (0.56	94 fewer per 1000 (from 180	VERY	CRITICAL

S	studies	serious ¹	inconsistency	indirectness		(31.4%)	(41%)	to 1.04)	fewer to 16 more)	LOW	

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 24: Clinical evidence profile: 2-4 litres versus >4 litres of fluids in adults with severe sepsis

	Quality assessment									Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	2-4L versus >4L	Control	Relative (95% CI)	Absolute	Quanty	importanio
Hospital m	ortality											
				no serious indirectness	very serious ²	none	82/186 (44.1%)	45/100 (45%)	RR 0.98 (0.75 to 1.28)	9 fewer per 1000 (from 112 fewer to 126 more)	VERY LOW	CRITICAL
ICU mortal	ity											
				no serious indirectness	serious²	none	66/186 (35.5%)	45/100 (45%)	RR 0.79 (0.59 to 1.05)	94 fewer per 1000 (from 185 fewer to 22 more)	VERY LOW	CRITICAL

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 25: Clinical evidence profile: High volume (20-40ml Ringer lactate/kg) versus low volume (20ml Ringer lactate/kg) in children with septic shock

Quality assessment	No of patients	Effect	Quality	Importance
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No of studies	Dasian	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	High volume versus low volume	Control	Relative (95% CI)	Absolute	
Cumula	ive 72-hour su	rvival									
1	randomised trials				no serious imprecision	none	52/74 (70.3%)	55/73 (75.3%)		53 fewer per 1000 (from 173 fewer to 105 more)	CRITICAL

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

J.9 Escalation of care

None.

J.10 Inotropic agents and vasopressors

Table 26: Clinical evidence profile: Norepinephrine versus vasopressin for adults with septic shock

	Quality assessment							No of patients Effect			Quality	Importance
No of studies	Design Inconsistency Indirectness Imprecision				Other considerations	Norepinephrine versus vasopressin	Control	Relative (95% CI)	Absolute	quanty	portunes	
28-day m	28-day mortality											
1				no serious indirectness	serious ¹	none		140/396 (35.4%)	RR 1.11 (0.93 to	39 more per 1000 (from 25 fewer to 117	MODERATE	CRITICAL

									1.33)	more)		
90-day m	ortality											
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	188/379 (49.6%)	172/392 (43.9%)		57 more per 1000 (from 13 fewer to 136 more)	MODERATE	CRITICAL
ICU mort	ality											
2	randomised trials	serious ²	no serious inconsistency	no serious indirectness	very serious ¹	none	13/25 (52%)	11/28 (39.3%)	RR 1.26 (0.72 to 2.21)	102 more per 1000 (from 110 fewer to 475 more)	VERY LOW	CRITICAL
Requiring	g renal replac	ement thera	py at 48 hours									
1	randomised trials	serious ²	no serious inconsistency	no serious indirectness	very serious ¹	none	8/15 (53.3%)	5/15 (33.3%)	RR 1.6 (0.68 to 3.77)	200 more per 1000 (from 107 fewer to 923 more)	VERY LOW	NOT IMPORTANT
New onse	et of tachyarr	hythmias										
1	randomised trials	serious ²	no serious inconsistency	no serious indirectness	very serious ¹	none	4/15 (26.7%)	1/15 (6.7%)	RR 4 (0.5 to 31.74)	200 more per 1000 (from 33 fewer to 1000 more)	VERY LOW	NOT IMPORTANT

¹ Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs ² 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 27: Norepinephrine versus dopamine for adults with septic shock

	·	Quality as		·		No of patients			Effect				
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Norepinephrine versus dopamine	Control	Relative (95% CI)	Absolute	Quality	Importance	
28-day mo	ortality												
	randomised trials	,	no serious inconsistency	no serious indirectness	serious²	none	51/118 (43.2%)	67/134 (50%)	RR 0.86 (0.66 to 1.13)	70 fewer per 1000 (from 170 fewer to 65 more)	VERY LOW	CRITICAL	
Mortality	ortality												
	randomised trials	,		no serious indirectness	serious ²	none	23/40 (57.5%)	28/40 (70%)	RR 0.82 (0.59 to 1.15)	126 fewer per 1000 (from 287 fewer to 105 more)	VERY LOW	CRITICAL	
Hospital r	nortality												
	randomised trials	,		no serious indirectness	very serious ²	none	7/16 (43.8%)	10/16 (62.5%)	RR 0.7 (0.36 to 1.37)	188 fewer per 1000 (from 400 fewer to 231 more)	VERY LOW	CRITICAL	
Incidence	of arrhythmi	as			!								
1	randomised trials	,	no serious inconsistency	no serious indirectness	no serious imprecision	none	14/118 (11.9%)	51/134 (38.1%)	RR 0.31 (0.18 to 0.53)	263 fewer per 1000 (from 179 fewer to 312	LOW	NOT IMPORTANT	

										fewer)		
Length of stay in the hospital (Better indicated by lower values)												
1				no serious indirectness	no serious imprecision	none	118	134	-	MD 0.7 lower (4.36 lower to 2.96 higher)	LOW	IMPORTANT
Length of stay on the ICU (Better indicated by lower values)												
1				no serious indirectness	no serious imprecision	none	118	134	-	MD 0.7 higher (1.15 lower to 2.55 higher)	LOW	IMPORTANT

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 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs

Table 28: Norepinephrine versus epinephrine for adults with septic shock

	Quality assessment									Effect	Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Norepinephrine versus epinephrine	Control	Relative (95% CI)	Absolute		·	
28-day m	28-day mortality												
1				no serious indirectness	serious ¹	none	24/82 (29.3%)	17/76 (22.4%)	RR 1.31 (0.76 to 2.24)	69 more per 1000 (from 54 fewer to 277 more)	MODERATE	CRITICAL	

90-day m	ortality										
1		no serious risk of bias	no serious indirectness	serious ¹	none	30/82 (36.6%)	23/74 (31.1%)	RR 1.18 (0.76 to 1.83)	56 more per 1000 (from 75 fewer to 258 more)	MODERATE	CRITICAL

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

Table 29: Dopexamine versus dopamine for adults with septic shock

	Quality assessment								Effect			Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Dopexamine versus dopamine	Control	Relative (95% CI)	Absolute		
28-day mo	ortality											
1		no serious risk of bias			very serious¹	none	5/20 (25%)	4/21 (19%)	RR 1.31 (0.41 to 4.2)	59 more per 1000 (from 112 fewer to 610 more)	LOW	CRITICAL

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

Table 30: Norepinephrine plus dobutamine versus epinephrine for adults with septic shock

Quanty assessment and a parisment and a parism		Quality assessment	No of patients	Effect	Quality	Importance
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No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Norepinephrine + dobutamine versus epinephrine	Control	Relative (95% CI)	Absolute			
28-day m	28-day mortality												
	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	58/169 (34.3%)	64/161 (39.8%)	RR 0.86 (0.65 to 1.14)	56 fewer per 1000 (from 139 fewer to 56 more)	MODERATE	CRITICAL	
90-day mortality													
	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	85/169 (50.3%)	84/161 (52.2%)	RR 0.96 (0.78 to 1.19)	21 fewer per 1000 (from 115 fewer to 99 more)	HIGH	CRITICAL	
7-day mo	7-day mortality												
1	randomised trials		no serious inconsistency	no serious indirectness	serious ¹	none	34/169 (20.1%)	40/161 (24.8%)	RR 0.81 (0.54 to 1.21)	47 fewer per 1000 (from 114 fewer to 52 more)	MODERATE	CRITICAL	
14-day m	14-day mortality												
	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	44/169 (26%)	56/161 (34.8%)	RR 0.75 (0.54 to 1.04)	87 fewer per 1000 (from 160 fewer to 14 more)	MODERATE	CRITICAL	
Mortality	Mortality												

2		very serious ²	no serious inconsistency	no serious indirectness	very serious ¹	none	13/26 (50%)	13/26 (50%)	RR 1 (0.58 to 1.71)	0 fewer per 1000 (from 210 fewer to 355 more)	VERY LOW	CRITICAL			
Mortality	Mortality at discharge from ICU														
1		no serious risk of bias	no serious inconsistency	no serious indirectness	no serious imprecision	none	75/169 (44.4%)	75/161 (46.6%)	RR 0.95 (0.75 to 1.21)	23 fewer per 1000 (from 116 fewer to 98 more)	HIGH	CRITICAL			
Mortality at discharge from hospital															
1			no serious inconsistency	no serious indirectness	no serious imprecision	none	82/169 (48.5%)	84/161 (52.2%)	RR 0.93 (0.75 to 1.15)	37 fewer per 1000 (from 130 fewer to 78 more)	HIGH	CRITICAL			
Number	of serious ad	verse event	s during catecho	lamine infusion							'				
1			no serious inconsistency	no serious indirectness	very serious ¹	none	41/169 (24.3%)	43/161 (26.7%)	RR 0.91 (0.63 to 1.31)	24 fewer per 1000 (from 99 fewer to 83 more)	LOW	NOT IMPORTANT			
Number of serious adverse events after catecholamine infusion															
1	randomised trials		no serious inconsistency	no serious indirectness	very serious ¹	none	13/169 (7.7%)	12/161 (7.5%)	RR 1.03 (0.49 to 2.19)	2 more per 1000 (from 38 fewer to 89 more)	LOW	NOT IMPORTANT			

¹ Downgraded by 1 increment if the confidence interval crossed one MID or by 2 increments if the confidence interval crossed both MIDs ² 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 31: Norepinephrine plus dopexamine versus norepinephrine plus epinephrine for adults with septic shock

			Quality asses	·		,	No of patients		Effect			
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	donovamino vorelle (Controll		Relative (95% CI)	Absolute	Quality	Importance
28-day me	ortality											
	randomised no serious no serious no serious no serious very none indirectness serious¹		none	2/12 (16.7%)	3/10 (30%)	RR 0.56 (0.11 to 2.7)	132 fewer per 1000 (from 267 fewer to 510 more)	LOW	CRITICAL			
90-day me	ortality											
			no serious inconsistency		very serious ¹	none	3/12 (25%)	4/10 (40%)	RR 0.62 (0.18 to 2.16)	152 fewer per 1000 (from 328 fewer to 464 more)	LOW	CRITICAL

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

Table 32: Norepinephrine plus epinephrine versus norepinephrine plus dobutamine for adults with septic shock

Quality assessment	No of patients	Effect	Quality	Importance
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No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Norepinephrine + epinephrine versus norepinephrine + dobutamine	Control	Relative (95% CI)	Absolute		
28-day m	ortality											
	randomised trials		no serious inconsistency	no serious indirectness	very serious ¹	none	15/30 (50%)	16/30 (53.3%)	RR 0.94 (0.57 to 1.53)	32 fewer per 1000 (from 229 fewer to 283 more)	LOW	CRITICAL
SOFA sc	ore at start (E	Better indic	ated by lower va	lues)								
	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	30	30	-	MD 0.8 higher (2.31 lower to 3.91 higher)		IMPORTANT
SOFA sc	ore at 24 hou	ırs (Better i	ndicated by lowe	er values)								
	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	30	30	-	MD 0.7 higher (2.41 lower to 3.81 higher)	MODERATE	IMPORTANT
SOFA sc	ore at 48 hou	ırs (Better i	ndicated by lowe	er values)								
	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	30	30	-	MD 0.6 higher (2.49 lower to 3.69 higher)		IMPORTANT

SOFA so	core at 72 hou	ırs (Better i	ndicated by lowe	er values)										
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	30	30	-	MD 0.6 higher (2.72 lower to 3.92 higher)		IMPORTANT		
SOFA so	core at 96 hou	ırs (Better i	ndicated by lowe	er values)										
1		no serious risk of bias	no serious inconsistency	no serious indirectness	serious ¹	none	30	30	-	MD 0.8 higher (2.62 lower to 4.22 higher)	MODERATE	IMPORTANT		
Acute co	cute coronary syndrome													
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ¹	none	1/30 (3.3%)	1/30 (3.3%)	RR 1 (0.07 to 15.26)	0 fewer per 1000 (from 31 fewer to 475 more)	LOW	NOT IMPORTANT		
Arrhythr	nias													
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ¹	none	4/30 (13.3%)	6/30 (20%)	RR 0.67 (0.21 to 2.13)	66 fewer per 1000 (from 158 fewer to 226 more)	LOW	NOT IMPORTANT		
Cerebra	stroke													
1	randomised trials	no serious risk of bias	no serious inconsistency	no serious indirectness	very serious ²	none	0/30 (0%)	0/30 (0%)	-	-	LOW	NOT IMPORTANT		

Limb is	chaemia									
1	randomised trials		very serious ¹	none	2/30 (6.7%)	3/30 (10%)	RR 0.67 (0.12 to 3.71)	33 fewer per 1000 (from 88 fewer to 271 more)	LOW	NOT IMPORTANT

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

J.11 Supplemental oxygen

None.

J.12 Use of bicarbonate

Table 33: Clinical evidence profile: bicarbonate versus no bicarbonate (28-day mortality)

			Quality asses	ssment			No of patients			Effect	Quality	Importance
No of studies					Imprecision	Other considerations	Bicarbonate versus no bicarbonate	Control Relative (95% CI)		Absolute	Quanty	Importance
28-day mo	ortality			•								
1	observational studies	- ,		no serious indirectness	very serious²	none	10/36 (27.8%)	12/36 (33.3%)		57 fewer per 1000 (from 197 fewer to 227 more)	VERY LOW	CRITICAL

¹ Case-control study. Small sample size

Table 34: Clinical evidence profile: bicarbonate versus no bicarbonate (Duration of critical care stay; Time to reversal of shock)

			Quality ass	essment			Median	[95% CI]		Effect	Quality	Importance
No of studies	I DESIGN I HINCONSISTENCYI INGIFECTINES I IMPRECISION I							Control group	Relative (95% CI)	Absolute	Quanty	importance
Duration of	of critical care sta	ау										
1	observational studies	very serious ¹	not estimable ²	no serious indirectness	not estimable²	none	44.5 [34-54] Hours	55 [39-60] Hours	-	-	VERY LOW	IMPORTANT

² Confidence interval crossed both standard MIDs

Time	Time to reversal of shock													
1	observational studies	very serious ¹	not estimable ²	no serious indirectness	not estimable ²	none	11.5 [6.0-16.0] days	16.0 [13.5-19.0] days	-	-	VERY LOW	MPORTANT		

¹ Case-control study. Small sample size

J.13 Early goal-directed therapy (EGDT)

Table 9: Clinical evidence profile: EGDT versus Usual care

			Quality as	sessment			No of pa	atients	Effect		Quality	Importance	
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	EGDT versus Control	Control	Relative (95% CI)	Absolute	_		
Primary n	ortality outco	ome of eac	ch study										
5	randomised trials	serious ¹	serious ²	no serious indirectness	no serious imprecision	none	495/2134 (23.2%)	582/2601 (22.4%)	RR 1.01 (0.9 to 1.12)	2 more per 1000 (from 22 fewer to 27 more)	LOW	CRITICAL	
90-day mo	ortality												
3	randomised trials		no serious inconsistency	no serious indirectness	no serious imprecision	none	460/1820 (25.3%)	598/2243 (26.7%)	RR 0.99 (0.89 to 1.11)	3 fewer per 1000 (from 29 fewer to 29 more)	MODERATE	CRITICAL	
ICU admis	ssion												
3	randomised trials		serious inconsistency	no serious indirectness	no serious imprecision	none	1677/1856 (90.4%)	1902/2324 (81.8%)	RR 1.11 (1.09 to 1.14)	91 more per 1000 (from 75 more to 116 more)	LOW	CRITICAL	
ICU lengt	length of stay for patient admitted to ICU (days) (Better indicated by lower values)												

² Non-parametric results

4	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	1825	2051	-	MD 0.02 lower (0.47 lower to 0.43 higher)	MODERATE	IMPORTANT			
ICU	ICU length of stay for patient admitted to ICU (days) - New Subgroup (Better indicated by lower values)														
4	randomised trials	serious ¹	no serious inconsistency	no serious indirectness	no serious imprecision	none	1825	2051	-	MD 0.02 lower (0.47 lower to 0.43 higher)	MODERATE	IMPORTANT			

¹Downgraded 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 ²Downgraded by 1 or 2 increments because:

J.14 Monitoring

None.

J.15 Patient education, information and support

None.

J.16 Training and education

None.

o The point estimate varies widely across studies, unexplained by subgroup analysis.

o The confidence intervals across studies show minimal or no overlap, unexplained by subgroup analysis

Heterogeneity, I²=50%, p=0.04, unexplained by subgroup analysis.

Appendix K: Forest plots

K.1 Scoring systems

None.

K.2 Signs and symptoms

K.2.1 Temperature

Figure 1: Sensitivity and specificity for temperature, adults

Temperature (adults): fever 38.5C and above to predict bacteriaemia in older patients FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) 0.27 [0.23, 0.31] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Pfitzenmeyer 1995 40 374 6 138 0.87 [0.74, 0.95] Temperature (adults): >38C to predict bacteriaemia in elderly patients TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Study Lindvig 2014 158 361 87 1518 0.64 [0.58, 0.70] 0.81 [0.79, 0.83] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Temperature (adults): fever to predict bacteriaemia in elderly patients Sensitivity (95% CI) Specificity (95% CI) TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Study Chassagne 1996 0 0 0 0 Not estimable Not estimable 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Temperature (adults): fever spike to predict bacteriaemia in elderly patients TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Chassagne 1996 0 0 0 0 Not estimable Not estimable 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Temperature (adults): >39C to predict septic complications in adult patients undergoing cardiac surgery TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Study 0.88 [0.62, 0.98] 14 53 2 41 0.44 [0.33, 0.54] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Temperature (adults): abnormal temperature (hypothermia or fever) in shock patients admitted to tertiary care centre via ED TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Study Seigel 2012 193 0 96 0 0.67 [0.61, 0.72] Not estimable 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Figure 2: Sensitivity and specificity for temperature, children

Temperature (children): predicting EOS / pneumonia in term new-borns >37 weeks Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Hofer 2012A 0 Not estimable Not estimable 0 0.2 0.4 0.6 0.8 o 0.2 0.4 0.6 0.8 1 Temperature (children): >38C for predicting post-operative infectious complications FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Angel 1994 2 125 0.67 [0.09, 0.99] 0.27 [0.20, 0.34] 0 0.2 0.4 0.2 0.4 0.6 0.8 0.6 0.8 Temperature (children): >39C for predicting post-operative infectious complications Specificity (95% CI) TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Study Angel 1994 1 16 2 155 0.33 [0.01, 0.91] 0.91 [0.85, 0.95] 0 0.2 0.4 0.6 0.8 0.2 0.4 0.6 0.8 1 Temperature (children): <40C or >40C for predicting SBI in febrile infants 8-12 weeks Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) TP FP FN TN Specificity (95% CI) Study 0.96 [0.93, 0.98] 0 0.2 0.4 0.6 0.8 1 Bonadio 1994 5 12 21 298 0.19 [0.07, 0.39] 0 0.2 0.4 0.6 0.8 1 Temperature (children): age-specific temperature-pulse centiles >97th centile for prediciting SBI in 3 months - 10 year olds Sensitivity (95% CI) Specificity (95% CI) TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Study **Brent 2011** 13 6 112 0.14 [0.00, 0.58] 0.90 [0.83, 0.94] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Temperature (children): age-specific temperature-pulse centiles >90th centile for prediciting SBI in 3 months - 10 year olds Sensitivity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Specificity (95% CI) Study TP FP FN TN 0.80 [0.71, 0.87] Brent 2011 3 88 0.25 [0.01, 0.81] 1 22 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Temperature (children): age-specific temperature-pulse centiles >75th centile for prediciting SBI in 3 months - 10 year olds Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Brent 2011 0 0 0 0 Not estimable Not estimable 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Temperature (children): age-specific temperature-pulse centiles >50th centile for prediciting SBI in 3 months - 10 year olds Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) ___ **Brent 2011** 5 83 6 133 0.45 [0.17, 0.77] 0.62 [0.55, 0.68] 0 0.2 0.4 0.6 0.8 1 0.2 0.4 0.6 0.8 1 Temperature (children): age-specific temperature-pulse centiles >97th centile for prediciting SBI (meningococcal) FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Brent 2011 12 191 4 109 0.75 [0.48, 0.93] 0.36 [0.31, 0.42] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Temperature (children): age-specific temperature-pulse centiles 90th-97th centile for prediciting SBI (meningococcal) TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Study Brent 2011 0 0 Not estimable Not estimable 0 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Temperature (children): age-specific temperature-pulse centiles 75th-90th centile for prediciting SBI (meningococcal) Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Brent 2011 Not estimable Not estimable 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 Temperature (children): age-specific temperature-pulse centiles 50th-75th centile for prediciting SBI (meningococcal) Study FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Brent 2011 Not estimable Not estimable 0 0.2 0.4 0.6 0.8 1 0.2 0.4 0.6 0.8 1 Temperature (children): <36C to predict bacteriaemia in neonates Sensitivity (95% CI) Specificity (95% CI) Specificity (95% CI) Study TP FP FN TN Sensitivity (95% CI) Hofer 2012 2 36 21 417 0.09 [0.01, 0.28] 0.92 [0.89, 0.94] 0.2 0.4 0.6 0.8 1 Temperature (children): >38.5C to predict bacteriaemia in neonates Specificity (95% CI) Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Hofer 2012 2 27 19 428 0.10 [0.01, 0.30] 0.94 [0.91, 0.96] 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

K.2.2 Heart rate

Figure 3: Sensitivity and specificity for heart rate, adults

TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Chassagne 1996 Not estimable Not estimable 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Figure 4: Sensitivity and specificity for heart rate, children Heart rate (children): tachycardia >180/min or bradycardia <100/min to predict culture-proven EOS in term neonates TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Study Specificity (95% CI) 14 81 37 344 0.27 [0.16, 0.42] 0.81 [0.77, 0.85] Hofer 2012 0.2 0.4 0.6 0.8 1 Heart rate (children): age-specific pulse centiles above 97th centile for SBI in 3 months - 10 year olds TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) 0 1 1 26 0.00 [0.00, 0.97] 0.96 [0.81, 1.00] Brent 2011 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Heart rate (children): age-specific pulse centiles above 90th centile for SBI in 3 months - 10 year olds TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Brent 2011 2 7 8 74 0.20 [0.03, 0.56] 0.91 [0.83, 0.96] 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Heart rate (children): age-specific pulse centiles above 75th centile for SBI in 3 months - 10 year olds TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Brent 2011 0.42 [0.15, 0.72] 0.76 [0.69, 0.82] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Heart rate (children): age-specific pulse centiles above 50th centile for SBI in 3 months - 10 year olds TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) 0.49 [0.43, 0.54] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Brent 2011 10 159 4 151 0.71 [0.42, 0.92] Heart rate (children): tachycardia for SBI in 3 months - 10 year olds TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) 23 196 11 284 0.59 [0.55, 0.64] 0.68 [0.49, 0.83] Brent 2011 Heart rate (children): age-specific pulse centiles above 97th centile for predicting meningococcal sepsis in children TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Study Brent 2011 0 0 0 0 Not estimable Not estimable 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Heart rate (children): age-specific pulse centiles above 75th centile for predicting meningococcal sepsis in children TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Study Brent 2011 0 0 0 Not estimable 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Heart rate (children): age-specific pulse centiles above 50th centile for predicting meningococcal sepsis in children Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Brent 2011 0 0 Not estimable Not estimable 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Heart rate (children): age-specific pulse centiles under 50th centile for predicting meningococcal sepsis in children Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Brent 2011 0 0 0 0 Not estimable Not estimable 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Heart rate (children): tachycardia for predicting meningococcal sepsis in children Study Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Brent 2011 Not estimable Not estimable 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

K.2.3 Blood pressure

Figure 5: Sensitivity and specificity for blood pressure, adults

Blood pressure (adults): HTI of ABP drops <95 mmHg SAP to predict 28-day mortality in adults TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Study Specificity (95% CI) 0.29 [0.13, 0.49] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 0.93 [0.90, 0.96] Dunser 2009A 230 20 16 8 Blood pressure (adults): HTI of ABP drops <75 mmHg MAP to predict 28-day mortality in adults TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Study Specificity (95% CI) 0.38 [0.14, 0.68] Dunser 2009A 244 8 17 5 0.93 [0.90, 0.96] 6 0.2 0.4 0.6 0.8 1 Blood pressure (adults): HTI of ABP drops <65 mmHg SAP to predict 28-day mortality in adults FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Study Specificity (95% CI) 0.26 [0.20, 0.33] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 4 52 0.95 [0.87, 0.99] Dunser 2009A 73 145 Blood pressure (adults): HTI of ABP drops <45 mmHg MAP to predict 28-day mortality in adults Study FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) 0.29 [0.23, 0.36] Dunser 2009A 80 134 5 55 0.94 [0.87, 0.98] ۲ 0 0.2 0.4 0.6 0.8 1 Blood pressure (adults): MAP 70 mmHg and under to predict onset of organ failure at 24h in adults Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) 0.70 [0.53, 0.83] Slotman 1997 19 12 0 28 1.00 [0.82, 1.00] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1 Blood pressure (adults): MAP 70 mmHg and under to predict onset of organ failure at 48h in adults TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Study Slotman 1997 1 45 0.93 [0.66, 1.00] 1.00 [0.92, 1.00] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 Blood pressure (adults): MAP 70 mmHg and under to predict onset of organ failure at 72h in adults TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI)

0.00 [0.00, 0.09]

0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

K.2.4 Respiratory rate

Study Slotman 1997

None.

K.2.5 Altered mental state

Figure 6: Sensitivity and specificity for altered mental state, adults

1.00 [0.82, 1.00]

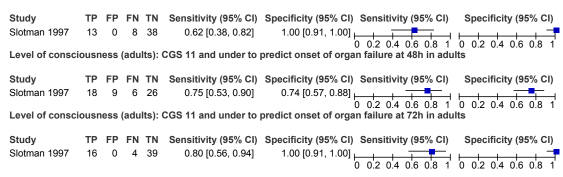
Altered mental state (adults): to predict bacteriaemia in elderly patients

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

K.2.6 Level of consciousness

Figure 7: Sensitivity and specificity for level of consciousness, adults

Level of consciousness (adults): CGS 11 and under to predict onset of organ failure at 24h in adults



K.2.7 Oxygen saturation

None.

K.2.8 Urine output

None.

K.2.9 Diarrhoea

None.

K.3 Blood tests

Note: studies for coupled sensitivity/specificity are listed in alphabetical order. Setting, target condition, and actual cut-off value reported by each study are included in the study name.

K.3.1 CRP, adults

Figure 8: Sensitivity and specificity for CRP (cut-off ≥5 mg/l), adults

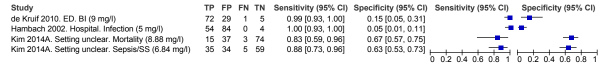


Figure 9: Sensitivity and specificity for CRP (cut-off ≥10 mg/l), adults

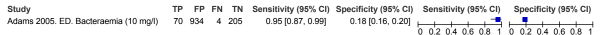


Figure 10: Sensitivity and specificity for CRP (cut-off ≥20 mg/l), adults

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Gaini 2006A. Hospital. Sepsis/SS (38 mg/l)	110	15	28	21	0.80 [0.72, 0.86]	0.58 [0.41, 0.74]	-	
Muller 2010. Hospital. Bacteraemia (20 mg/l)	70	775	3	77	0.96 [0.88, 0.99]	0.09 [0.07, 0.11]	-	
Nakamura 2009. Hospital. Bacteraemia (35 mg/l)	49	30	16	21	0.75 [0.63, 0.85]	0.41 [0.28, 0.56]		
Stucker 2005. Hospital. Infection (30 mg/l)	46	107	4	60	0.92 [0.81, 0.98]	0.36 [0.29, 0.44]	-	-
Yonemori 2001. Neutropenia. Infection (30.8 mg/l)	0	0	0	0	Not estimable	Not estimable		
							0 02 04 06 08 1	0 02 04 06 08 1

Figure 11: Sensitivity and specificity for CRP (cut-off ≥50 mg/l), adults

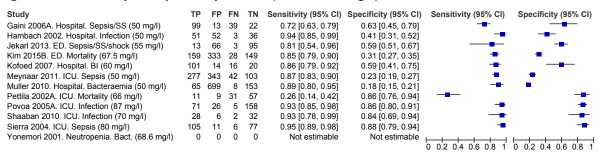


Figure 12: Sensitivity and specificity for CRP (cut-off ≥100 mg/l), adults

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Aalto 2004. ED. BSI (125 mg/l)	11	15	2	64	0.85 [0.55, 0.98]	0.81 [0.71, 0.89]		
Castelli 2004. ICU. Sepsis/SS (128 mg/l)	22	21	11	96	0.67 [0.48, 0.82]	0.82 [0.74, 0.89]		-
Castelli 2006. ICU. Sepsis/SS/shock (128 mg/l)	68	19	43	125	0.61 [0.52, 0.70]	0.87 [0.80, 0.92]	-	-
Cheval 2000. ICU. Sepsis (100 mg/l)	30	17	2	11	0.94 [0.79, 0.99]	0.39 [0.22, 0.59]	-	
Gaini 2006A. Hospital. Sepsis/SS (100 mg/l)	87	2	50	33	0.64 [0.55, 0.72]	0.94 [0.81, 0.99]	-	-
Hambach 2002. Hospital. Infection (100 mg/l)	45	34	9	54	0.83 [0.71, 0.92]	0.61 [0.50, 0.72]	-	-
Kim 2011. ED. Bacteraemia (100 mg/l)	22	81	16	167	0.58 [0.41, 0.74]	0.67 [0.61, 0.73]		-
Moreira 2010. Hospital. Sepsis (110 mg/l)	44	13	6	47	0.88 [0.76, 0.95]	0.78 [0.66, 0.88]	-	
Muller 2010. Hospital. Bacteraemia (100 mg/l)	59	571	14	281	0.81 [0.70, 0.89]	0.33 [0.30, 0.36]	0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 13: Sensitivity and specificity for CRP (cut-off ≥150 mg/l), adults

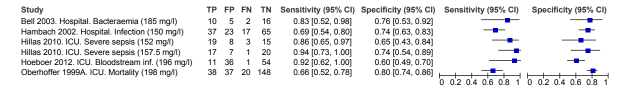
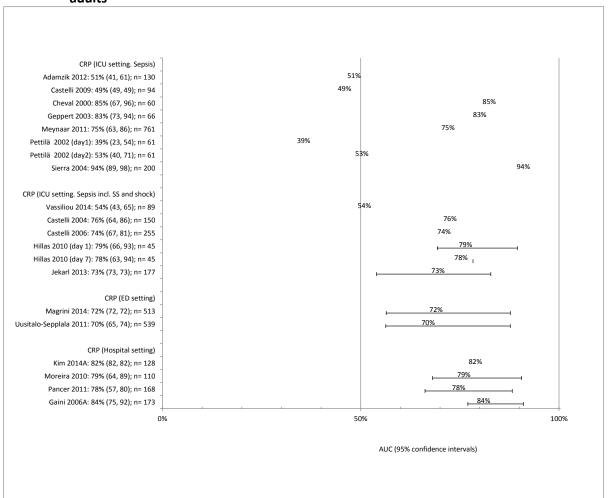
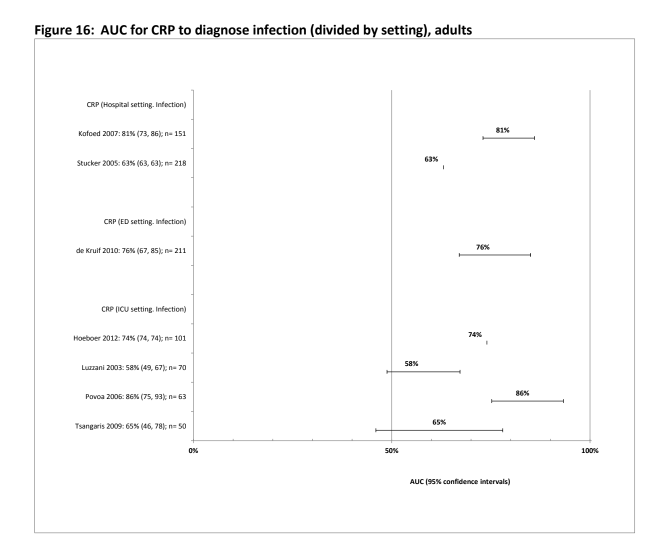


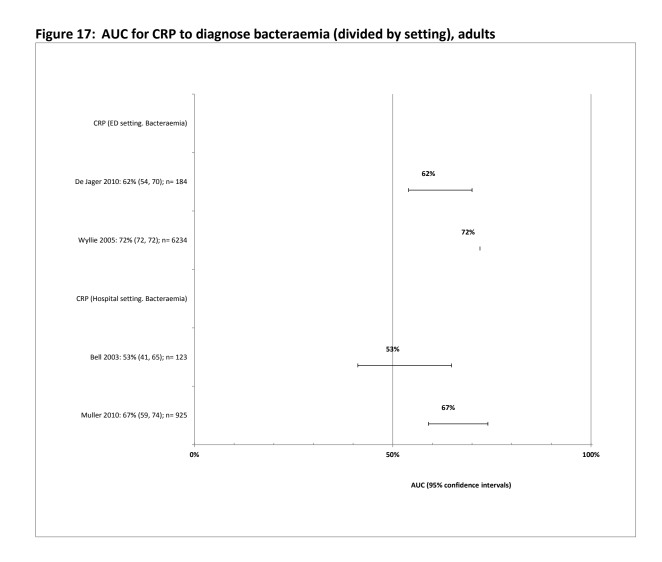
Figure 14: Sensitivity and specificity for CRP (cut-off ≥200 mg/l), adults

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Hoeboer 2012. ICU. Septic shock (208 mg/l)	24	15	10	52		0.78 [0.66, 0.87]		-
Muller 2010. Hospital. Bacteraemia (200 mg/l)	45	307	28	545	0.62 [0.50, 0.73]	0.64 [0.61, 0.67]		<u> </u>
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 15: AUC for CRP to diagnose sepsis, severe sepsis and septic shock (divided by setting), adults







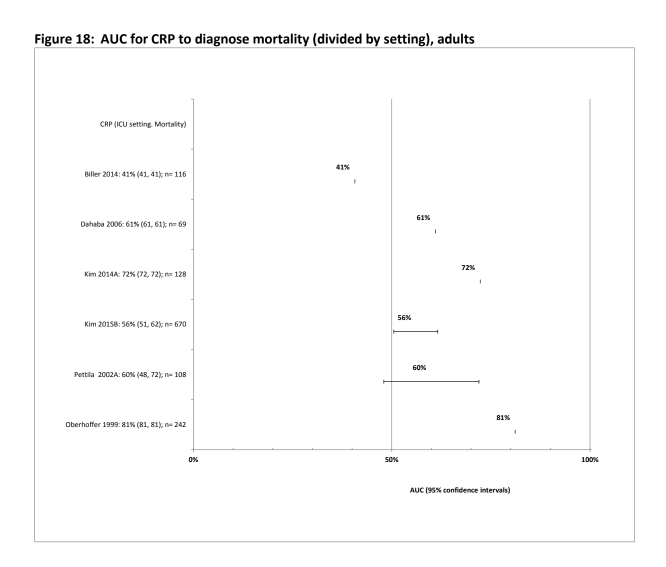


Figure 19: AUC for CRP to diagnose infection or bacteraemia in the immunocompromised subgroup (divided by setting), adults

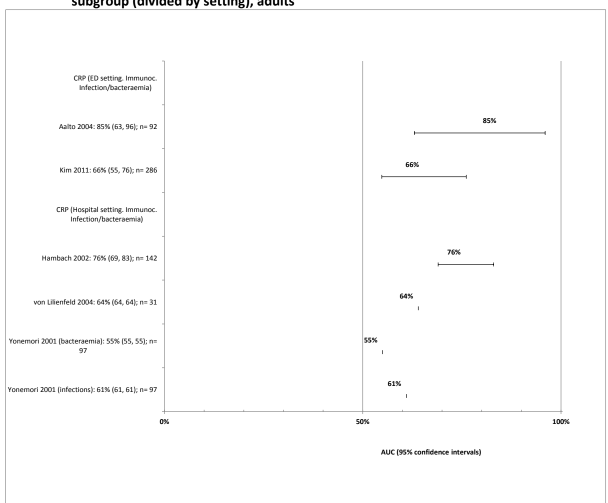


Figure 20: Odds ratio for CRP ratio (follow-up/initial level), adults



Figure 21: Odds ratio for CRP >8 mg/l versus CRP ≤8 mg/l, adults

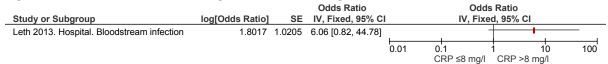


Figure 22: Odds ratio for CRP for diagnosing sepsis, adults

			Odds Ratio		(odds Ratio		
Study or Subgroup	log[Odds Ratio]	SE Weig	ght IV, Fixed, 95% CI		IV, I	Fixed, 95%	CI	
Uusitalo-Sepplala 2011. ED. Sepsis/SS	0.2852	0.0975	1.33 [1.10, 1.61]			+		
				0.01	0.1	1	10	100
					Protective fa	ctor Risk	factor	

Figure 23: Odds ratio for CRP for diagnosing severe sepsis, adults

			Ouus Italio		,	ouus italio		
Study or Subgroup	log[Odds Ratio]	SE Weight	IV, Fixed, 95% CI		IV,	Fixed, 95%	CI	
Uusitalo-Sepplala 2011. ED. Sepsis/SS	0.0198 (0.1505	1.02 [0.76, 1.37]			+		
				0.01	0.1	1	10	100
				Р	rotective fa	actor Risk	factor	

K.3.2 CRP, children

Figure 24: Sensitivity and specificity for CRP (cut-off ≥20mg/I), children

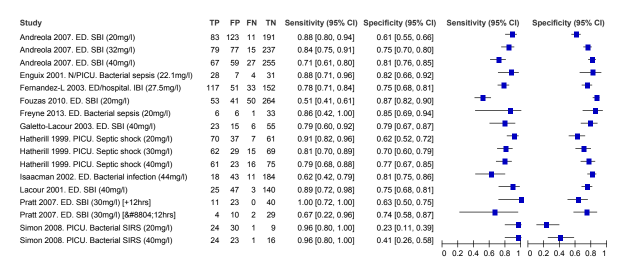


Figure 25: Sensitivity and specificity for CRP (cut-off ≥50mg/I), children

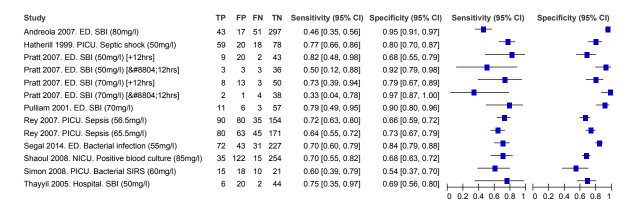


Figure 26: Sensitivity and specificity for CRP (cut-off ≥100mg/l), children

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Baez 2011. ICU. Post-op sepsis (100mg/l) [24hrs]	34	16	7	46	0.83 [0.68, 0.93]	0.74 [0.62, 0.84]	-	-
Baez 2011. ICU. Post-op sepsis (100mg/l) [48hrs]	37	19	4	43	0.90 [0.77, 0.97]	0.69 [0.56, 0.80]		
Baez 2011. ICU. Post-op sepsis (110mg/l) [24hrs]	38	24	3	38	0.93 [0.80, 0.98]	0.61 [0.48, 0.73]	-	
Baez 2011. ICU. Post-op sepsis (110mg/l) [48hrs]	36	7	5	55	0.88 [0.74, 0.96]	0.89 [0.78, 0.95]		
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 27: Sensitivity and specificity for CRP (cut-off ≥150mg/l), children

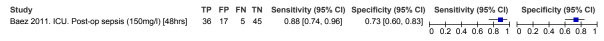


Figure 28: Sensitivity and specificity for CRP (cut-off ≥200mg/l), children

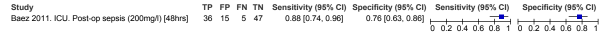


Figure 29: Sensitivity and specificity for change in CRP per day, children

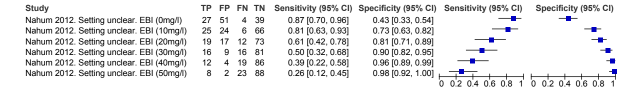
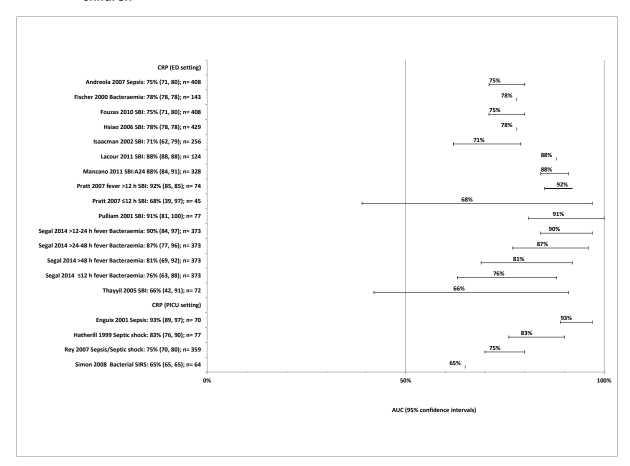


Figure 30: AUC for CRP to diagnose sepsis, severe sepsis and septic shock (divided by setting), children



K.3.3 CRP, neonates

Figure 31: Sensitivity and specificity for CRP (cut-off <5 mg/l), neonates

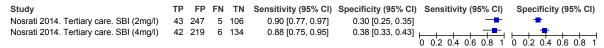


Figure 32: Sensitivity and specificity for CRP (cut-off ≥5mg/I), neonates

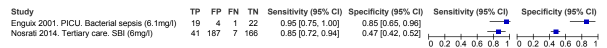


Figure 33: Sensitivity and specificity for CRP (cut-off ≥10mg/I), neonates

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Jacquot 2009. NICU. Late onset sepsis (10mg/l)	17	6	13	37	0.57 [0.37, 0.75]	0.86 [0.72, 0.95]		-
Nosrati 2014. Tertiary care. SBI (10mg/l)	40	138	8	215	0.83 [0.70, 0.93]	0.61 [0.56, 0.66]	-	-
Sherwin 2008. NICU. Late onset sepsis (18mg/l)	21	7	31	105	0.40 [0.27, 0.55]	0.94 [0.88, 0.97]	0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 34: Sensitivity and specificity for CRP (cut-off ≥20mg/I), neonates

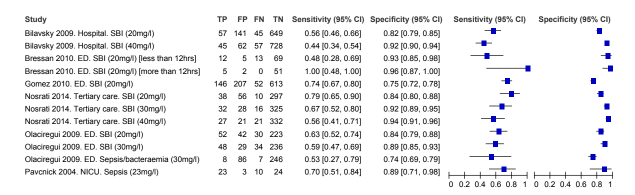
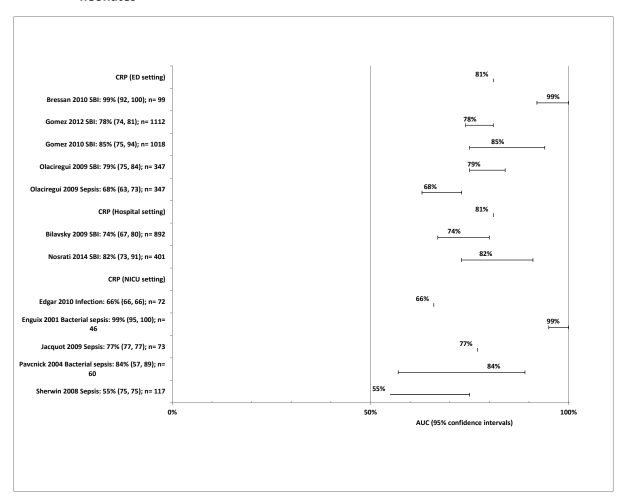


Figure 35: Sensitivity and specificity for CRP (cut-off ≥50mg/I), neonates

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Bilavsky 2009. Hospital. SBI (80mg/l)	24	14	78	776	0.24 [0.16, 0.33]	0.98 [0.97, 0.99]	-	
Gomez 2010. ED. SBI (70mg/l)	186	249	12	571	0.94 [0.90, 0.97]	0.70 [0.66, 0.73]		
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 36: AUC for CRP to diagnose sepsis, severe sepsis and septic shock (divided by setting), neonates



K.3.4 WBC, adults

Figure 37: Sensitivity and specificity for WBC (<1x109/I), adults

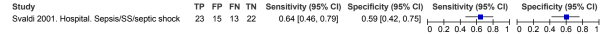


Figure 38: Sensitivity and specificity for WBC (>1x109/I), adults

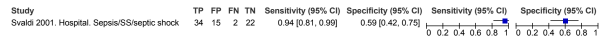


Figure 39: Sensitivity and specificity for WBC (<4x10⁹/l), adults

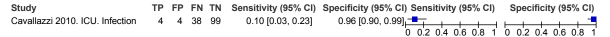


Figure 40: Sensitivity and specificity for WBC (>11x109/I), adults



Figure 41: Sensitivity and specificity for WBC (>12x109/I), adults

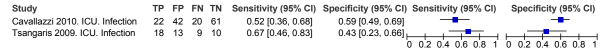


Figure 42: Sensitivity and specificity for WBC (>15x109/I), adults

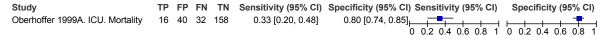


Figure 43: Sensitivity and specificity for WBC (>20.3x109/I), adults



Figure 44: Sensitivity and specificity for WBC (<4.3x109/l and >11x109/l), adults



Figure 45: Sensitivity and specificity for WBC (≤4x10⁹/l and ≥12x10⁹/l), adults

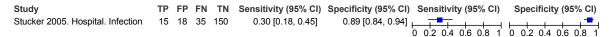


Figure 46: Sensitivity and specificity for WBC ($\leq 5 \times 10^9 / l$ and $\geq 20 \times 10^9 / l$), adults

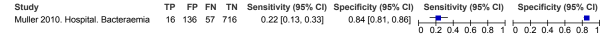


Figure 47: AUC for WBC to diagnose (1) sepsis, severe sepsis, septic shock; bacteraemia or infection; (3) mortality (divided by setting), adults

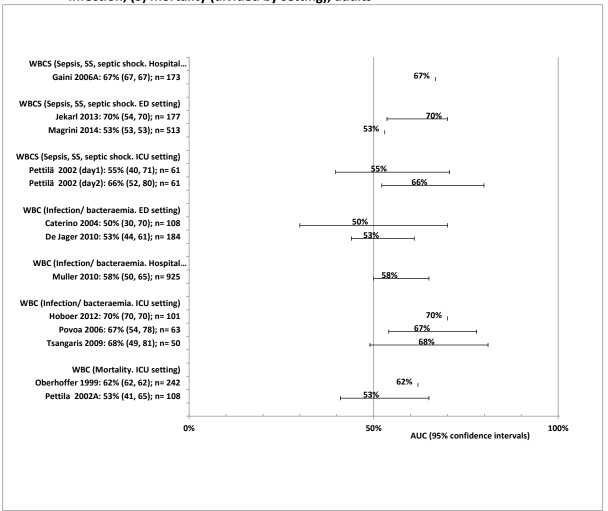


Figure 48: Odds ratio for WBC (>12x109/I), adults

				Odds Ratio		Odds	Ratio		
Study or Subgroup	log[Odds Ratio]	SE W	leight	IV, Fixed, 95% CI		IV, Fixe	d, 95% CI		
Freund 2012. ED. Sepsis/SS/septic shock	0.6043	0.2282		1.83 [1.17, 2.86]			-		
					0.01	0.1 Protective factor	1 Risk facto	10 or	100

Figure 49: Odds ratio for WCC (<4x10⁹/l or >20x10⁹/l), adults

_			Odds Ratio		(Odds Ra	tio	
Study or Subgroup	log[Odds Ratio] SE	Weight	IV, Fixed, 95% CI		IV,	Fixed, 9	5% CI	
Patterson 2012. ED. Bacteraemia	-0.4943 1.2573		0.61 [0.05, 7.17]			-		
				0.01	0.1	1	10	100
					Protective fa	actor Ri	sk factor	

Figure 50: Odds ratio for WBC (≤4x109/I or ≥12x109/I), adults

			Odds Ratio			Odds Ratio		
Study or Subgroup	log[Odds Ratio]	SE Weight	IV, Fixed, 95% CI		IV,	Fixed, 95%	CI	
Stucker 2005. Hospital. Infection	1.2528 0.4	1023	3.50 [1.59, 7.70]				т.	
				0.01	0.1	1	10	100
					Protective f	actor Risk	factor	

K.3.5 WBC, children

Figure 51: Sensitivity and specificity for WBC (<5x109/I), children



Figure 52: Sensitivity and specificity for WBC (>10x10⁹/I), children

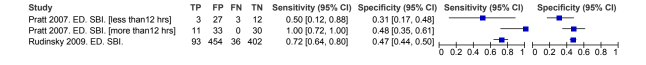


Figure 53: Sensitivity and specificity for WBC (>10.47x109/I), children



Figure 54: Sensitivity and specificity for WBC (>15x10⁹/l), children

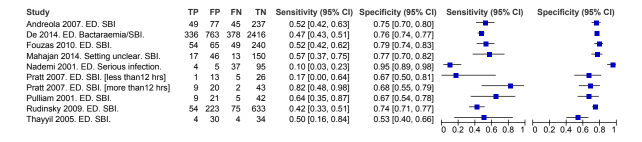


Figure 55: Sensitivity and specificity for WBC (>17.1x109/I), children



Figure 56: Sensitivity and specificity for WBC (>17.5x10⁹/I), children

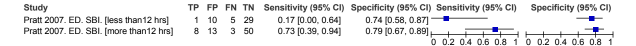


Figure 57: Sensitivity and specificity for WBC (>19x10⁹/l), children



Figure 58: Sensitivity and specificity for WBC (>20x109/I), children

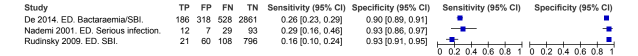


Figure 59: Sensitivity and specificity for WBC (>25x10⁹/I), children



Figure 60: Sensitivity and specificity for WBC (<5 or >15 x109/l), children

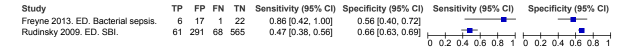
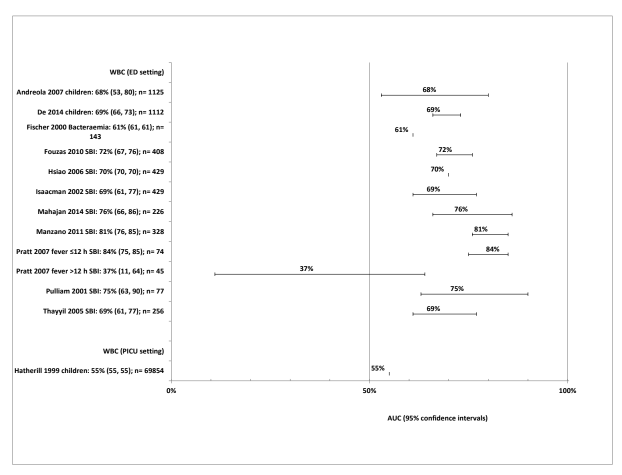


Figure 61: AUC for WBC to diagnose sepsis, severe sepsis and septic shock (divided by setting), children



K.3.6 WBC, neonates

Figure 62: Sensitivity and specificity for WBC (<1x109/I), neonates

Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Specificity

Figure 63: Sensitivity and specificity for WBC (<5x109/I), neonates

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)

Figure 64: Sensitivity and specificity for WBC (>5x109/I), neonates

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

Figure 65: Sensitivity and specificity for WBC (>10x109/I), neonates

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

Figure 66: Sensitivity and specificity for WBC (>15x109/I), neonates

FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Bilavsky 2009. Hospital. SBI. 49 126 53 664 0.48 [0.38, 0.58] 0.84 [0.81, 0.87] Bonsu 2003. ED. Bacteraemia. 8 834 9 2959 0.47 [0.23, 0.72] 0.78 [0.77, 0.79] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Figure 67: Sensitivity and specificity for WBC (>20x10⁹/l), neonates

TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Bilavsky 2009. Hospital. SBI. 22 38 80 752 0.22 [0.14, 0.31] 0.95 [0.93, 0.97] Bonsu 2003. ED. Bacteraemia. 266 3535 0.22 [0.03, 0.60] 0.93 [0.92, 0.94] 2182 12160 7474 48038 Hornik 2012 NICU Bacterial sensis 0.23 [0.22, 0.23] 0.80 [0.79, 0.80] 0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8

Figure 68: Sensitivity and specificity for WBC (>25x109/I), neonates

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

Figure 69: Sensitivity and specificity for WBC (>30x10⁹/l), neonates

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

Figure 70: Sensitivity or specificity for WBC (>50x109/I), neonates

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

Figure 71: Sensitivity and specificity for WBC (<4 or ≥20x10⁹/I), neonates

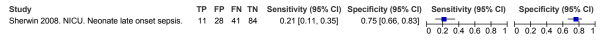


Figure 72: Sensitivity and specificity for WBC (>15 or <5 x109/I), neonates

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Bilavsky 2009. Hospital. SBI.	51	173	51	617	0.50 [0.40, 0.60]	0.78 [0.75, 0.81]	-	=
Bonsu 2003. ED. Bacteraemia.	17	1060	9	2725	0.65 [0.44, 0.83]	0.72 [0.71, 0.73]		
Bressan 2010. ED. SBI. [fever less than 12 hrs]	7	9	18	65	0.28 [0.12, 0.49]	0.88 [0.78, 0.94]		-
Bressan 2010. ED. SBI. [fever more than 12 hrs]	4	5	1	48	0.80 [0.28, 0.99]			
							n n 2 n 4 n 6 n 8 1	0 02 04 06 08 1

Figure 73: Sensitivity and specificity for WBC (>20 or <4.1 x109/l), neonates

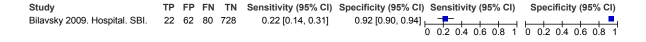
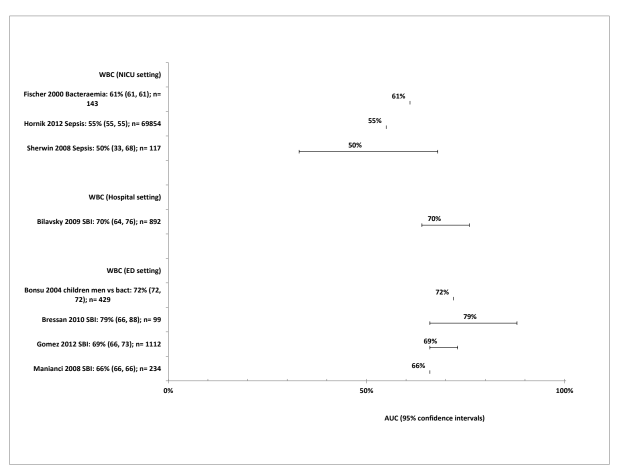


Figure 74: Sensitivity and specificity for WBC (>20 or <5 x109/l), neonates



Figure 75: AUC for WBC to diagnose sepsis, severe sepsis and septic shock (divided by setting), neonates



K.3.7 Leukocytes, adults

Figure 76: Multivariable odds ratio for leukocyte count, adults

			Odds Ratio			Od	lds Rat	io			
Study or Subgroup	log[Odds Ratio]	SE Weight	IV, Fixed, 95% CI			IV, Fi	xed, 95	5% CI			
de Kruif 2010. ED. Bacterial infection	0.1178 0	0.1178 0.0616 1			+						
				0.1	0.2	0.5	1	2	5	10	
					Prote	ective fact	or Ris	k factor			

Figure 77: AUC for leukocyte sedimentation rate, adults

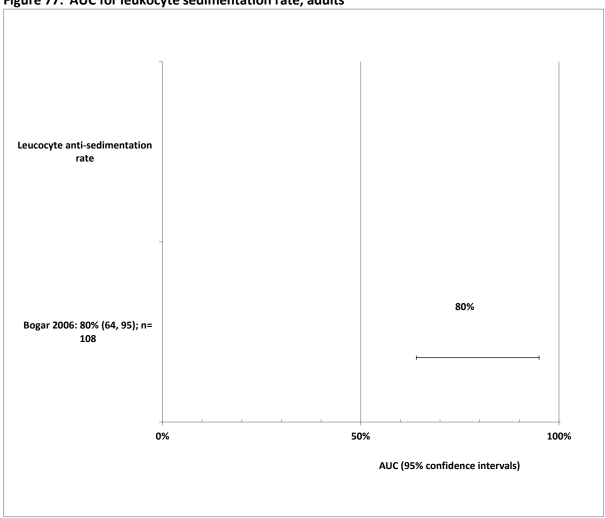


Figure 78: Odds ratio for leukocyte count ≥4x10⁹/l or ≤12x10⁹/l compared to <4x10⁹/l or >12x10⁹/l, adults

			Odds Ratio	lds Ratio Odds Ratio				
Study or Subgroup	log[Odds Ratio]	SE Weight	IV, Fixed, 95% CI		IV, Fixe	d, 95% CI		
Leth 2013. Hospital. Bloodstream infection	0.0677	0.2654	1.07 [0.64, 1.80]			 		
				0.01	0.1	1	10	100
					Protective factor	Risk factor		

K.3.8 Leukocytes, children

Figure 79: Sensitivity and specificity for leukocytes (>7.1x10⁹/l), children

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Fernandez-lopez 2003. ED/hospital. IBI.	81	48	69	154	0.54 [0.46, 0.62]	0.76 [0.70, 0.82] [<u> </u>	
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 80: Sensitivity and specificity for leukocytes (>15x109/l), children

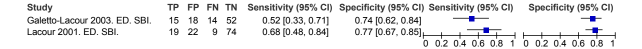
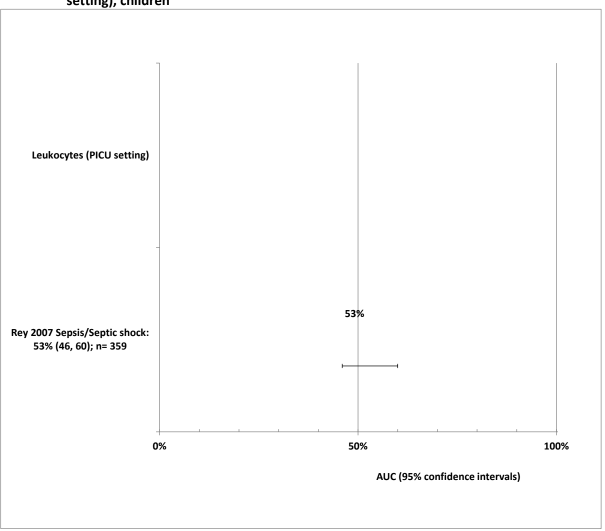


Figure 81: AUC for leukocytes to diagnose sepsis, severe sepsis and septic shock (divided by setting), children



K.3.9 Leukocytes, neonates

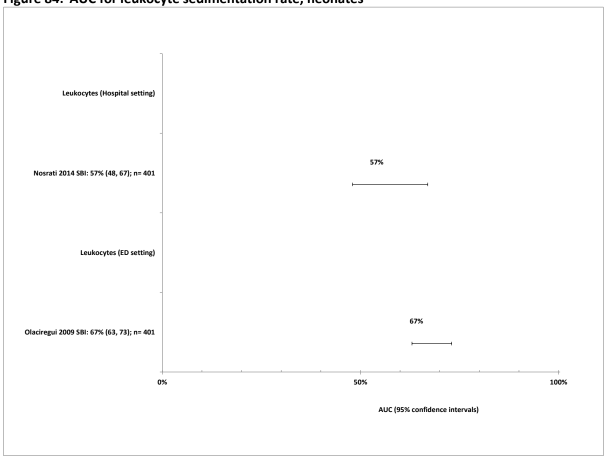
Figure 82: Sensitivity and specificity for leukocytes (>10x109/I), neonates



Figure 83: Sensitivity and specificity for leukocytes (>15x109/I), neonates







K.3.10 Neutrophils, adults

Figure 85: Sensitivity and specificity for neutrophil count (cut-off 7.5x109/I), adults



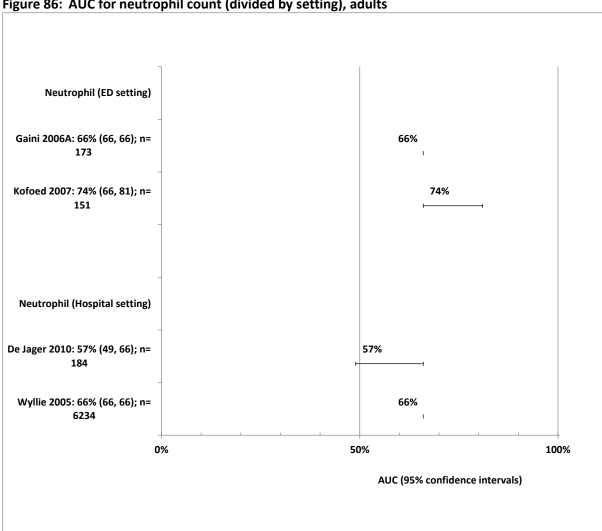


Figure 86: AUC for neutrophil count (divided by setting), adults

Figure 87: Odds ratio for neutrophil count (>80%), adults



Figure 88: Odds ratio for neutrophil count ≥2x109/I or ≤7x109/I compared to <2x109/I or >7x109/I, adults



K.3.11 Neutrophils, children

Figure 89: Sensitivity and specificity for neutrophil count (cut-off 6.45x10⁹/l), children

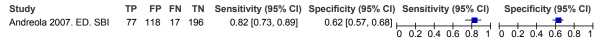


Figure 90: Sensitivity and specificity for neutrophil count (cut-off 10x109/I), children

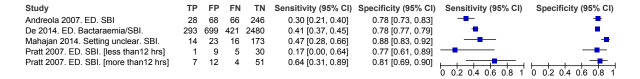


Figure 91: Sensitivity and specificity for neutrophil count (cut-off 10.2x10⁹/l), children

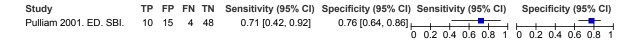


Figure 92: Sensitivity and specificity for neutrophil count (cut-off 10.6x10⁹/I), children

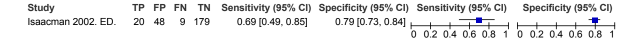


Figure 93: Sensitivity and specificity for neutrophil count (cut-off 11x109/I), children

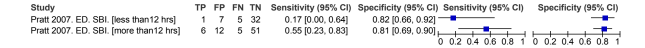


Figure 94: Sensitivity and specificity for neutrophil count (cut-off 12x109/I), children

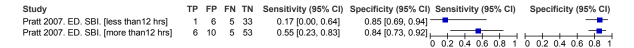


Figure 95: Sensitivity and specificity for neutrophil count (cut-off 13x10⁹/I), children

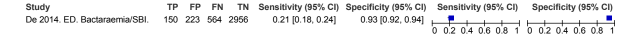
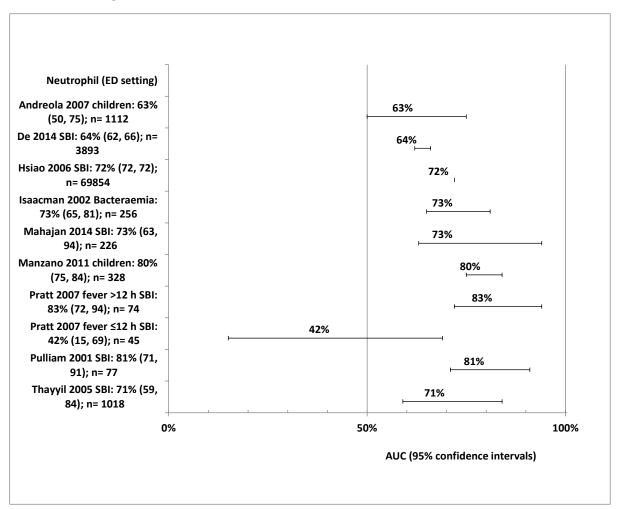


Figure 96: Sensitivity and specificity for neutrophil count (cut-off 15x10⁹/l), children



Figure 97: AUC for neutrophils to diagnose sepsis, severe sepsis and septic shock (divided by setting), children



K.3.12 Neutrophils, neonates

Figure 98: Sensitivity and specificity for neutrophil count (cut-off 1x109/I), neonates

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Hornik 2012. NICU. Bacterial sepsis.	232	1204	9424	58994	0.02 [0.02, 0.03]	0.98 [0.98, 0.98]	0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 99: Sensitivity and specificity for neutrophil count (cut-off 1.5x109/I), neonates

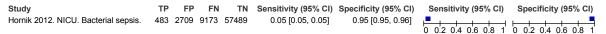
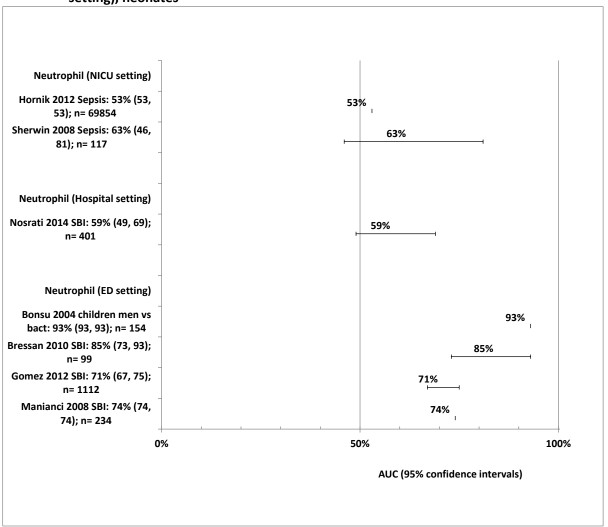


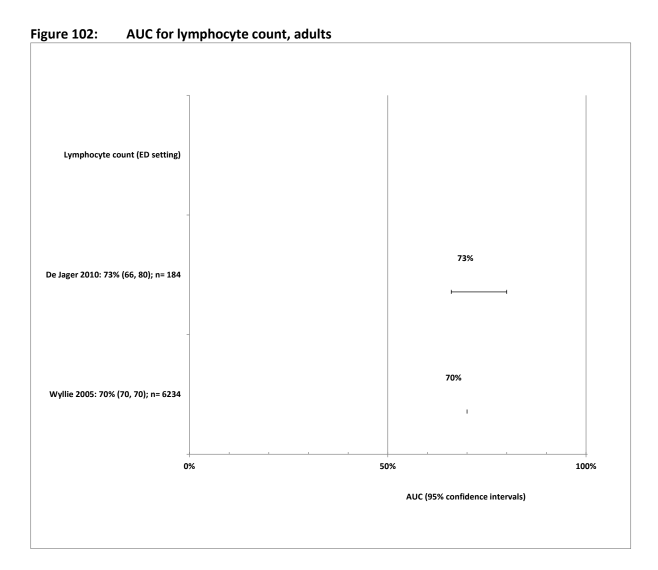
Figure 100: Sensitivity and specificity for neutrophil count (cut-off 10x10⁹/l), neonates

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Bressan 2010. ED. SBI. [fever less than 12 hrs]	5	2	20	72	0.20 [0.07, 0.41]	0.97 [0.91, 1.00]	_	-
Bressan 2010. ED. SBI. [fever more than 12 hrs]	4	0	1	53	0.80 [0.28, 0.99]	1.00 [0.93, 1.00]		-
Sherwin 2008. NICU. Neonate late onset sepsis.	17	8	35	104	0.33 [0.20, 0.47]	0.93 [0.86, 0.97]		
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 101: AUC for neutrophils to diagnose sepsis, severe sepsis and septic shock (divided by setting), neonates



K.3.13 Lymphocytes, adults



K.3.14 Lymphocytes, children

None.

K.3.15 Lymphocytes, neonates

None.

K.3.16 Lactate, adults

Figure 103: Sensitivity and specificity for lactate (>1.5 mmol/l), adults



Figure 104: Sensitivity and specificity for lactate (>1.7 mmol/l), adults



Figure 105: AUC for lactate to diagnose (1) sepsis, severe sepsis, septic shock; (2) bacteraemia or infection; (3) mortality, adults

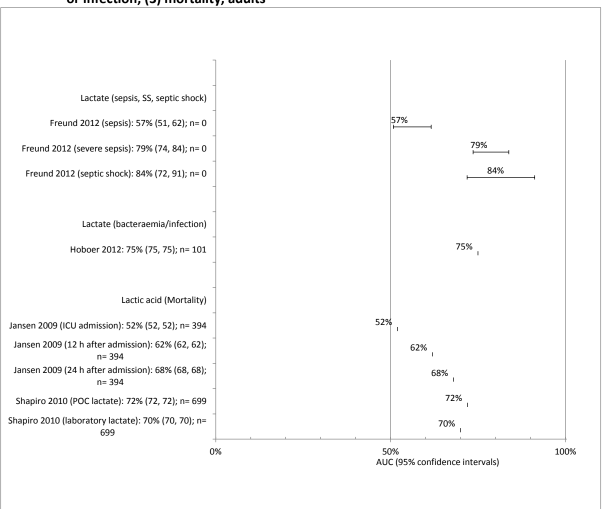


Figure 106: Odds ratio for lactate (>2 mmol/l) for the diagnosis of severe sepsis, adults

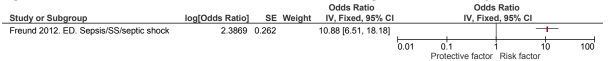
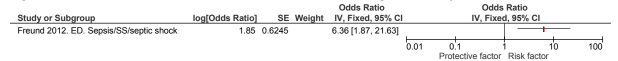


Figure 107: Odds ratio for lactate (>2 mmol/l) for the diagnosis of septic shock, adults



K.3.17 Lactate, children

None.

K.3.18 Lactate, neonates

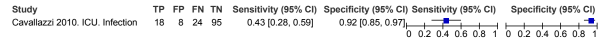
None.

K.3.19 Bands, adults

Figure 108: Sensitivity and specificity for bands (>8.5%), adults



Figure 109: Sensitivity and specificity for bands (>10%), adults



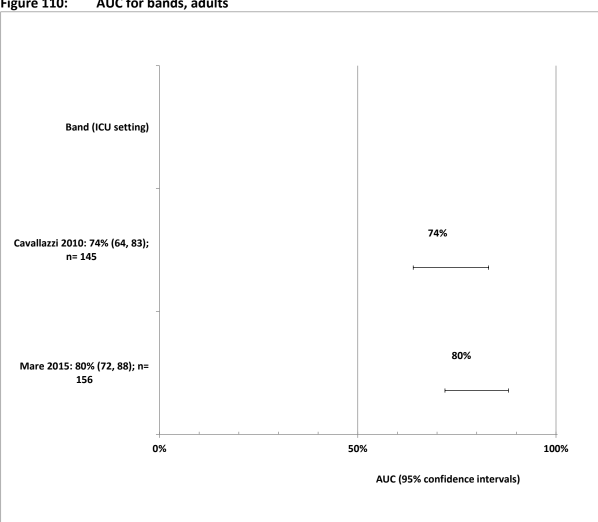


Figure 110: AUC for bands, adults

K.3.20 Bands, children

Sensitivity and specificity for bands (>1.5x109/I), children Figure 111:

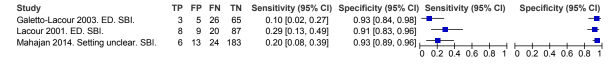
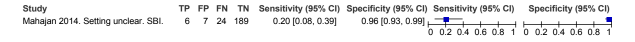


Figure 112: Sensitivity and specificity for bands (>1.8x10⁹/l), children



Setting), children

Band count (ED setting)

67%

Mahajan 2014 SBI: 67% (55, 78); n= 226

0%

50%

AUC (95% confidence intervals)

Figure 113: AUC for bands to diagnose sepsis, severe sepsis and septic shock (divided by setting), children

K.3.21 Bands, neonates

None.

K.3.22 Haemoglobin, adults

Figure 114: Odds ratio for haemoglobin (≤100 g/l), adults

			Odds Ratio		(Odds Ratio		
Study or Subgroup	log[Odds Ratio]	SE Weight	IV, Fixed, 95% CI		IV,	Fixed, 95%	CI	
Patterson 2012. ED. Bacteraemia	-0.3425 1.0	.0628	0.71 [0.09, 5.70]			-		
				0.01	0.1	1	10	100
					Protective fa	ctor Risk	actor	

K.3.23 Haemoglobin, children

None.

K.3.24 Haemoglobin, neonates

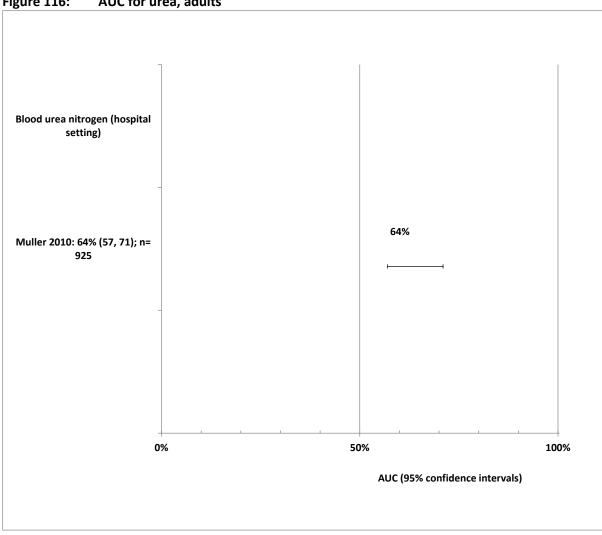
None.

K.3.25 Urea, adults

Figure 115: Sensitivity and specificity for urea (>11 mmol/I), adults







K.3.26 Urea, children

None.

K.3.27 Urea, neonates

None.

K.3.28 Creatinine, adults

None.

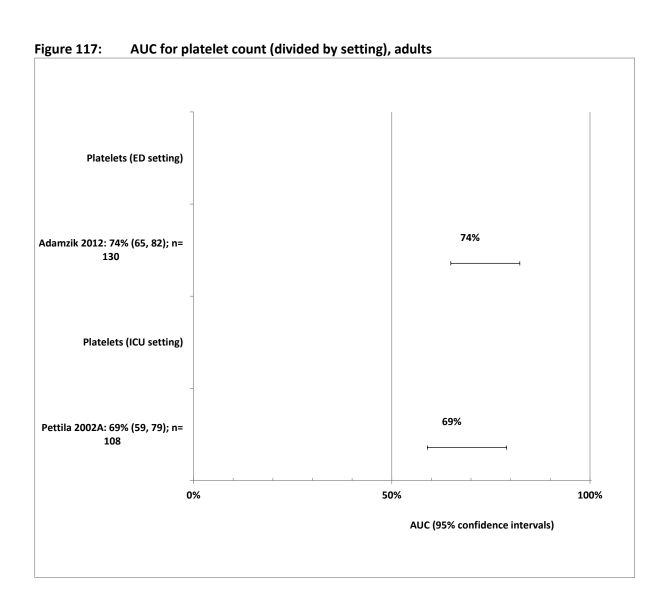
K.3.29 Creatinine, children

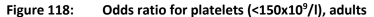
None.

K.3.30 Creatinine, neonates

None.

K.3.31 Platelets, adults







K.3.32 Platelets, children

Figure 119: Sensitivity and specificity for platelets (>68x109/l), children



Figure 120: Sensitivity and specificity for platelets (>400x109/l), children



Figure 121: Sensitivity and specificity for platelets (>450x109/l), children



Figure 122: Sensitivity and specificity for platelets (>500x10⁹/l), children



Figure 123: Sensitivity and specificity for platelets (>600x10⁹/l), children

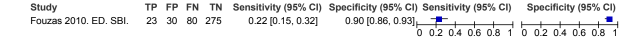


Figure 124: Sensitivity and specificity for platelets (20% increase), children

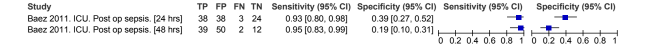
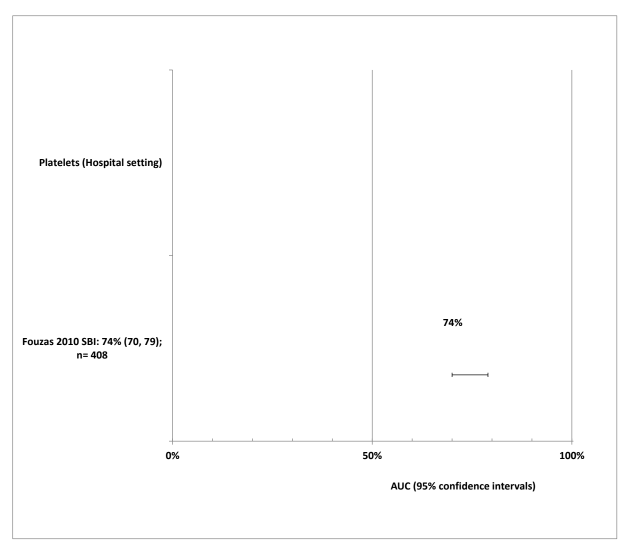


Figure 125: AUC for platelets to diagnose sepsis, severe sepsis and septic shock (divided by setting), children



K.3.33 Platelets, neonates

Figure 126: Sensitivity and specificity for platelets (>50x109/I), neonates

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Hornik 2012. NICU. Bacterial sepsis.	744	1324	8912	58874	0.08 [0.07, 0.08]	0.98 [0.98, 0.98]	0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Figure 127: Sensitivity and specificity for platelets (>100x10⁹/l), neonates

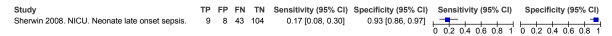


Figure 128: Sensitivity and specificity for platelets (<100x109/l), neonates

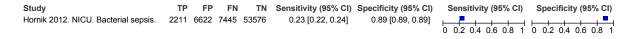
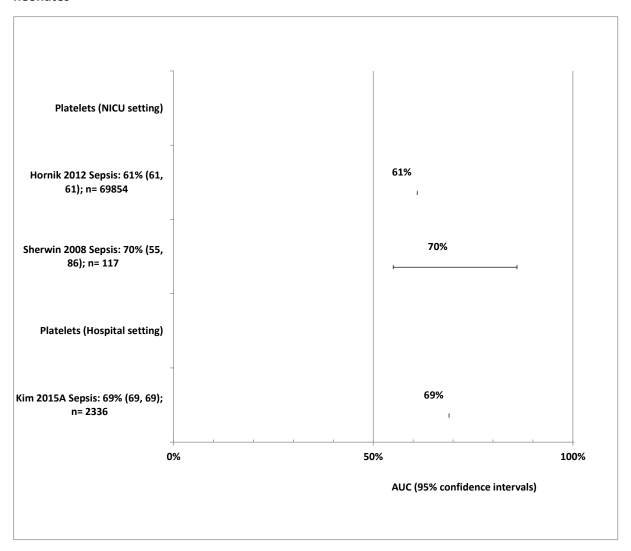
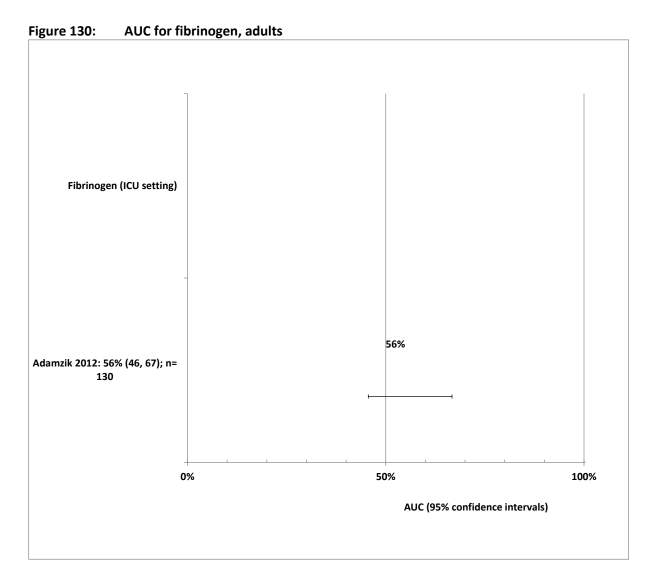


Figure 129: AUC for platelets to diagnose sepsis, severe sepsis and septic shock (divided by setting), neonates

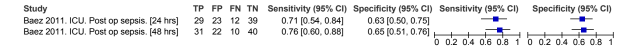


K.3.34 Fibrinogen, adults



K.3.35 Fibrinogen, children

Figure 131: Sensitivity and specificity for fibrinogen (20% increase), children



K.3.36 Fibrinogen, neonates

None.

K.3.37 Thrombin time, adults

Odds ratio for prothrombin time (≥18.4 seconds), adults Figure 132:

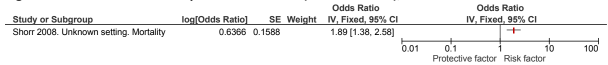


Figure 133: AUC for thrombin time (divided by setting), adults

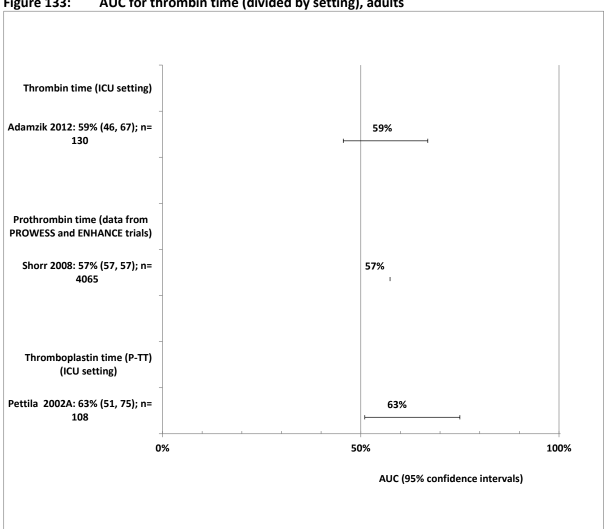


Figure 134: Odds ratio for anti-thrombin III (<53%), adults

			Odds Ratio		Odds	Ratio		
Study or Subgroup	log[Odds Ratio]	SE Weight	IV, Fixed, 95% CI		IV, Fixed	d, 95% CI		
Shorr 2008. Unknown setting. Mortality	0.8416	0.1609	2.32 [1.69, 3.18]			+ .		
				0.01	0.1 Protective factor	1 10 Risk factor	100	

K.3.38 Thrombin time, children

None.

K.3.39 Thrombin time, neonate	(.3.39	HIGHIOTHI	ume,	neonate
-------------------------------	---------------	-----------	------	---------

None.

K.3.40 Bilirubin, adults

None.

K.3.41 Bilirubin, children

None.

K.3.42 Bilirubin, neonates

None.

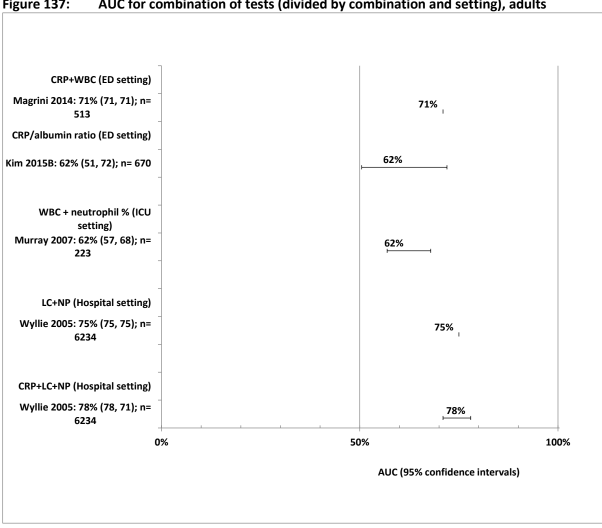
K.3.43 Combination of tests, adults

Figure 135: Sensitivity and specificity for bands (>10%) and WBC (>12x109/I), adults



Figure 136: Sensitivity and specificity for CRP/albumin ratio (>5.09), adults





AUC for combination of tests (divided by combination and setting), adults Figure 137:

Odds ratio for CRP >100 mg/l and lactate <4.0 mmol/l, adults **Figure 138:**

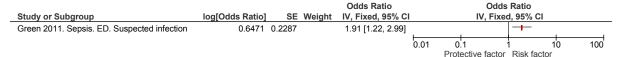


Figure 139: Odds ratio for CRP >100 mg/l and lactate ≥4.0 mmol/l, adults

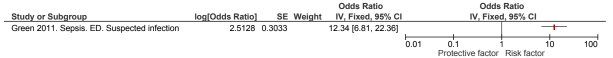
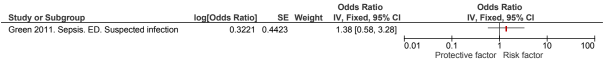


Figure 140: Odds ratio for CRP ≤100 mg/l and lactate ≥4.0 mmol/l, adults



K.3.44 Combination of tests, children

Figure 141: Sensitivity and specificity for CRP (>31mg/l) or WBC (>17.1x10⁹/l), children

Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Sensitivity (95% CI) Specificity (95% CI) Specificity

Figure 142: Sensitivity and specificity for CRP (>36mg/l) or ANC (>10.5x10⁹/l), children

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

 Isaacman 2002. ED.
 23
 114
 6
 114
 0.79 [0.60, 0.92]
 0.50 [0.43, 0.57]
 0.20 0.4 0.6 0.8 1
 0.20 0.4 0.6 0.8 1
 0.02 0.4 0.6 0.8 1

Figure 143: Sensitivity and specificity for CRP (>85mg/l) and ANC (>10x10⁹/l) or WBC (>15 x10⁹/l), children

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

Figure 144: Sensitivity and specificity for CRP (>85mg/l) and ANC (>10x10 9 /l) and WBC (>15 x10 9 /l), children

 Study
 TP
 FP
 FN
 TN
 Sensitivity (95% CI)
 Specificity (95% CI)
 Sensitivity (95% CI)
 Specificity (95% CI)

K.3.45 Combination of tests, neonates

None.

K.4 Lactate

None.

K.5 Serum creatinine

Figure 145: Serum creatinine level increase per 0.1 mg/dl: 28-day mortality

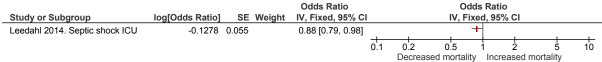


Figure 146: Initial serum creatinine >3.0 mg/dl: in-hospital mortality

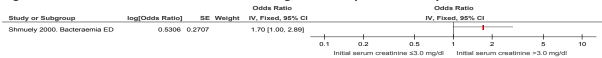


Figure 147: Initial serum creatinine >0.7 mg/dl: in-hospital mortality

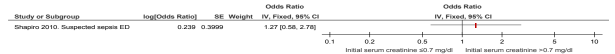
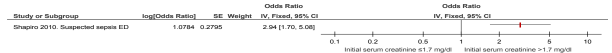


Figure 148: Initial serum creatinine >1.7 mg/dl: in-hospital mortality



K.6 Disseminated intravascular coagulation (DIC)

Figure 149: 28-day mortality (multivariable analysis)

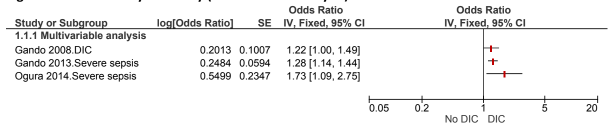


Figure 150: In-hospital mortality (multivariable and univariable analyses)

	-		Odds Ratio			Odds Ratio		
Study or Subgroup	log[Odds Ratio]	SE	IV, Fixed, 95% CI			IV, Fixed, 95% CI		
1.2.1 Multivariable analysis								
Gando 2007.SIRS/sepsis	1.441	0.557	4.22 [1.42, 12.59]				-	
Ogura 2014. Severe sepsis	0.4357	0.2182	1.55 [1.01, 2.37]			 1 		
1.2.2 Univariable analysis								
Gando 2007A.SIRS/sepsis	3.7013	1.1161	40.50 [4.54, 360.98]					
				L 05	——————————————————————————————————————		<u> </u>	
				0.05	0.2	No DIC DIC	5	20

K.7 Antimicrobial treatment

Figure 151: Mortality: <1 hour versus >1 hour, adult population

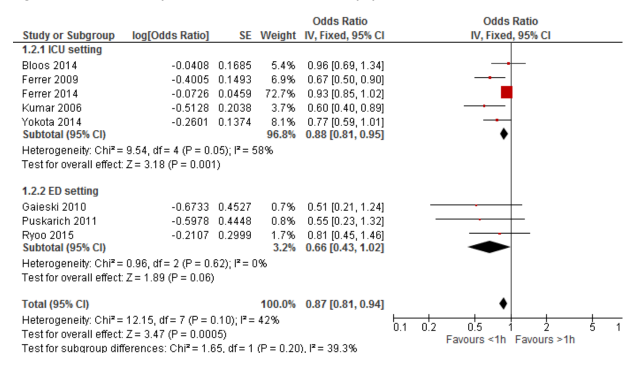


Figure 152: Mortality <2 hours versus >2 hours, adult population

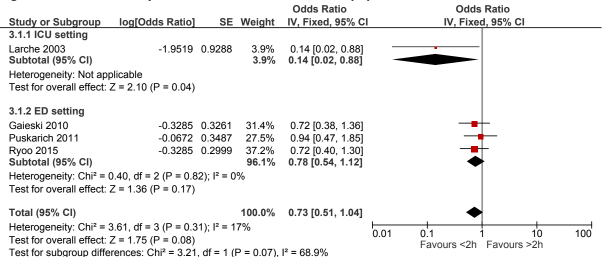


Figure 153: Mortality <3 hours versus >3 hours

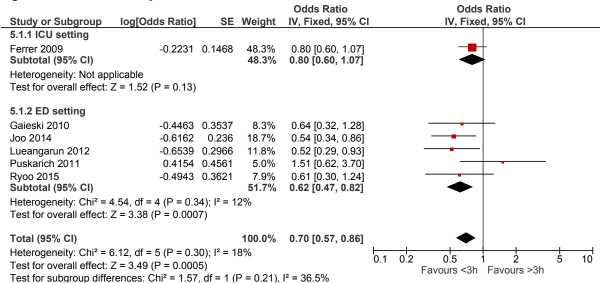


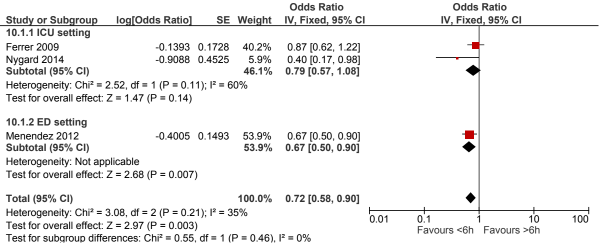
Figure 154: Mortality <4 hours versus >4 hours

		_			Odds Ratio		Odds I	Ratio	
	Study or Subgroup	log[Odds Ratio]	SE	Weight	IV, Fixed, 95% CI		IV, Fixed,	95% CI	
_	7.1.2 ED setting								
	Gaieski 2010	-0.2231	0.4218	47.6%	0.80 [0.35, 1.83]		-	_	
	Puskarich 2011	0.9416	0.8082	13.0%	2.56 [0.53, 12.50]		+	-	
	Ryoo 2015	-0.4155154	0.463304	39.4%	0.66 [0.27, 1.64]			_	
	Subtotal (95% CI)			100.0%	0.86 [0.49, 1.53]		•	•	
	Heterogeneity: Chi ² =	2.18, $df = 2$ ($P = 0$.	34); I² = 8%						
	Test for overall effect:	Z = 0.51 (P = 0.61)	ı						
	T-4-1 (05% CD			400.00	0.0010.40.4.501				
	Total (95% CI)			100.0%	0.86 [0.49, 1.53]		_	•	
	Heterogeneity: Chi ² =	2.18, $df = 2$ ($P = 0$.	34); I² = 8%			0.01 0.1		10	100
	Test for overall effect:	Z = 0.51 (P = 0.61)	ı			0.01 0.1	Favoure	Favours >4h	100
	Test for subgroup diffe	erences: Not appli	cable				i avouis	avouis 7411	

Figure 155: Mortality <5 hours versus >5 hours

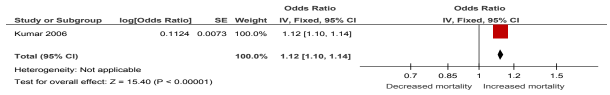
				Odds Ratio		Odds R	Ratio	
Study or Subgroup	log[Odds Ratio]	SE	Weight	IV, Fixed, 95% CI		IV, Fixed,	95% CI	
9.1.1 ED setting								
Gaieski 2010	-0.1508	1.0037	21.6%	0.86 [0.12, 6.15]		-		
Puskarich 2011	0.3716	1.1674	16.0%	1.45 [0.15, 14.29]				
Ryoo 2015	-0.7339692	0.590773	62.4%	0.48 [0.15, 1.53]			-	
Subtotal (95% CI)			100.0%	0.65 [0.26, 1.62]		◆	-	
Heterogeneity: Chi ² =	0.81, $df = 2$ ($P = 0$.	67); I² = 0%						
Test for overall effect:	Z = 0.92 (P = 0.36)							
Total (95% CI)			100.0%	0.65 [0.26, 1.62]			-	
Heterogeneity: Chi²=	0.81, $df = 2$ ($P = 0$.	67); I² = 0%			0.01 (1 1	10	100
Test for overall effect:	Z = 0.92 (P = 0.36)				0.01		Favours >5h	100
Test for subgroup diff	ferences: Not appli	cable				i avours i	avours - SII	





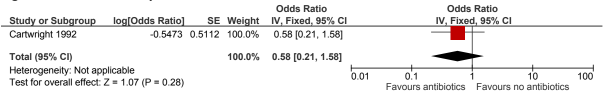
K.7.1 Hourly treatment delay

Figure 157: In-hospital mortality for hourly treatment delay



K.7.2 Parenteral antibiotics prior to admission to hospital

Figure 158: Mortality



K.7.3 PICU setting, paediatric population

Figure 159: PICU mortality: <1 hour versus >1 hour

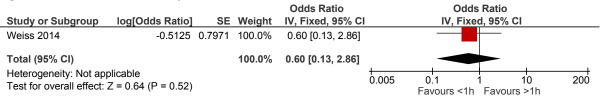


Figure 160: PICU mortality: <2 hours versus >2 hours

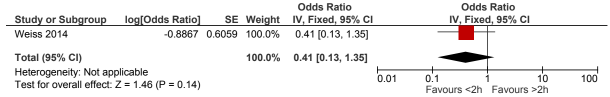


Figure 161: PICU mortality: <3 hours versus >3 hours

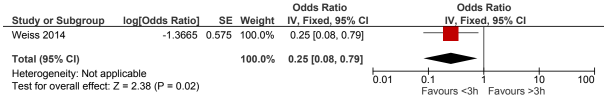


Figure 162: PICU mortality: <4 hours versus >4 hours

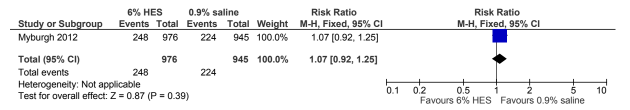


K.8 IV fluid administration

K.8.1 6% HES versus 0.9% saline in adults with sepsis

K.8.1.1 Mortality at 28 days

Figure 163: Mortality at 90 days



K.8.2 Crystalloid versus colloid plus crystalloid in adults with severe sepsis

K.8.2.1 Mortality at 28 days

Figure 164: Hospital mortality

	Crystal	loid	Colloid + crys	stalloid		Risk Ratio			Risk	Ratio			
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	l		M-H, Fixe	ed, 95% C	1		
McInthyre 2007A	101	235	121	258	100.0%	0.92 [0.75, 1.12]			-	-			
Total (95% CI)		235		258	100.0%	0.92 [0.75, 1.12]			•	-			
Total events	101		121										
Heterogeneity: Not ap Test for overall effect:		P = 0.38	3)				0.1	0.2 Favo	0.5 urs Crystalloid	1 2 Favours	Colloid + c	5 crysta	10

Figure 165: ICU mortality

	Crystal	lloid	Colloid + crys	stalloid		Risk Ratio			Risk	Ratio			
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	l		M-H, Fixe	d, 95% (CI		
McInthyre 2007A	72	235	99	258	100.0%	0.80 [0.62, 1.02]			-				
Total (95% CI)		235		258	100.0%	0.80 [0.62, 1.02]			•				
Total events	72		99										
Heterogeneity: Not ap Test for overall effect:		P = 0.07	")				0.1	0.2 Favo	0.5 urs Crystalloid	1 2 Favours	2 Colloid	5 + cryst	10 al

K.8.3 20% albumin versus 6% HES in adults with severe sepsis

K.8.3.1 Mortality at 28 days

Figure 166: 28-day mortality

•	•		•				
	Albun	nin	Colloid	- HES		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Dolecek 2009	4	30	6	26	100.0%	0.58 [0.18, 1.83]	
Total (95% CI)		30		26	100.0%	0.58 [0.18, 1.83]	
Total events	4		6				
Heterogeneity: Not app	plicable						0.1 0.2 0.5 1 2 5 10
Test for overall effect:	Z = 0.93 (P = 0.3	5)				0.1 0.2 0.5 1 2 5 10 Favours Albumin Favours HFS

K.8.4 4% albumin versus 0.9% Sodium Chloride BP in adults with severe sepsis

K.8.4.1 Mortality at 28 days

Figure 167: 28-day mortality (univariate analysis)

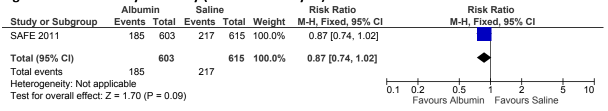
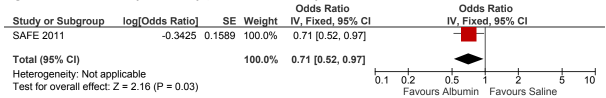


Figure 168: 28-day mortality (multivariate analysis)



Note: adjusted OR

K.8.5 Albumin versus crystalloids in adults with sepsis

K.8.5.1 Mortality at 28 days

Figure 169: Mortality

	Albun	nin	Crystall	oids		Risk Ratio			Risk	Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C			M-H, Fixe	ed, 95% CI		
Patel 2014	710	1937	763	1941	100.0%	0.93 [0.86, 1.01]						
Total (95% CI)		1937		1941	100.0%	0.93 [0.86, 1.01]			•			
Total events	710		763									
Heterogeneity: Not app Test for overall effect:		P = 0.0	9)				0.1	0.2 Favo	0.5 ours Albumin	1 2 Favours Cry	5 stalloids	10

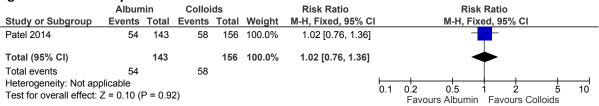
Figure 170: 90-day mortality

	Albun	nin	Crystall	loids		Risk Ratio			Risk	Ratio)		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI			M-H, Fix	ed, 95	% CI		
ALBIOS 2014	115	283	116	286	100.0%	1.00 [0.82, 1.22]			-	-			
Total (95% CI)		283		286	100.0%	1.00 [0.82, 1.22]			•				
Total events	115		116										
Heterogeneity: Not ap Test for overall effect:	•	P = 0.9	9)				0.1	0.2 Favours	0.5 20% albumin	1 Favo	2 ours crvs	5 stalloids	10

K.8.6 Albumin versus colloids in adults with sepsis

K.8.6.1 Mortality at 28 days

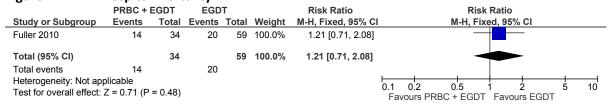
Figure 171: Mortality



K.8.7 Packed red blood cells (PRBC) plus EGDT versus EGDT only in adults with septic shock

K.8.7.1 Mortality at 28 days

Figure 172: Hospital mortality



K.8.8 Red blood cells (RBC) for low threshold (≤7 g/dl) versus high threshold (≤9 g/dl) in adults with septic shock

K.8.8.1 Mortality at 28 days

Figure 173: 90-day mortality

0	1	_									
	Low thre	shold	High thre	shold		Risk Ratio		R	sk Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C		M-H, F	Fixed, 95% CI	l	
11.1.1 > 70 years of a	ge										
Holst 2014	93	173	98	185	42.4%	1.01 [0.84, 1.23]			-		
Subtotal (95% CI)		173		185	42.4%	1.01 [0.84, 1.23]			*		
Total events	93		98								
Heterogeneity: Not ap	plicable										
Test for overall effect:	Z = 0.15 (P	= 0.88)									
11.1.2 70 years or yo	unger										
Holst 2014	123	329	125	311	57.6%	0.93 [0.77, 1.13]			-		
Subtotal (95% CI)		329		311	57.6%	0.93 [0.77, 1.13]			•		
Total events	123		125								
Heterogeneity: Not ap	plicable										
Test for overall effect:	Z = 0.73 (P	= 0.47)									
Total (95% CI)		502		496	100.0%	0.97 [0.84, 1.11]			•		
Total events	216		223								
Heterogeneity: Chi ² =	0.39, df = 1	(P = 0.5)	3); I ² = 0%					0.2 0.5	+ +	<u></u> _	10
Test for overall effect:	Z = 0.49 (P	= 0.63)					0.1	Favours low thresho	ı ∠ İd Favours İ	high threshold	10
Test for subgroup diffe	erences: Ch	$i^2 = 0.39$, df = 1 (P =	0.53), I	$^{2} = 0\%$			i avoaio iow tilicollo	ia ravours i	ngir an conoid	

K.8.9 0-2 litres versus 2-4 litres of fluid in adults with severe sepsis

K.8.9.1 Mortality at 28 days

Figure 174: Hospital mortality

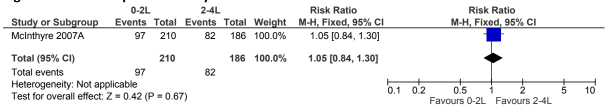


Figure 175: ICU mortality

	0-2L		2-4L			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
McInthyre 2007A	66	210	66	186	100.0%	0.89 [0.67, 1.17]	-
Total (95% CI)		210		186	100.0%	0.89 [0.67, 1.17]	•
Total events	66		66				
Heterogeneity: Not app Test for overall effect: 2		P = 0.3	9)				0.1 0.2 0.5 1 2 5 10 Favours 0-2L Favours 4L

K.8.10 0-2 litres versus >4 litres of fluids in adults with severe sepsis

K.8.10.1 Mortality at 28 days

Figure 176: Hospital mortality

•	•		•				
	0-2L		>4L			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
McInthyre 2007A	97	210	45	100	100.0%	1.03 [0.79, 1.33]	-
Total (95% CI)		210		100	100.0%	1.03 [0.79, 1.33]	•
Total events	97		45				
Heterogeneity: Not ap	plicable						0.1 0.2 0.5 1 2 5 10
Test for overall effect:	Z = 0.20 (P = 0.8	4)				Favours 0-2L Favours >4L

Figure 177: ICU mortality

	0-2L	_	>4L			Risk Ratio			Ris	sk Rati	io		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI			M-H, F	ixed, 9	5% CI		
McInthyre 2007A	66	210	41	100	100.0%	0.77 [0.56, 1.04]			-				
Total (95% CI)		210		100	100.0%	0.77 [0.56, 1.04]			<				
Total events	66		41										
Heterogeneity: Not ap Test for overall effect:		P = 0.0	9)				0.1	0.2 F	0.5 avours 0-2	1 L Fav	2 /ours >4L	5	10

K.8.11 2-4 litres versus >4 litres of fluids in adults with severe sepsis

K.8.11.1 Mortality at 28 days

Figure 178: Hospital mortality

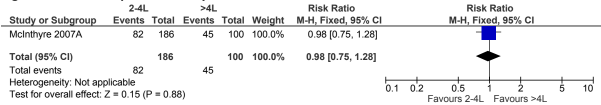


Figure 179: ICU mortality

	2-4L		>4L			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
McInthyre 2007A	66	186	45	100	100.0%	0.79 [0.59, 1.05]	-
Total (95% CI)		186		100	100.0%	0.79 [0.59, 1.05]	•
Total events	66		45				
Heterogeneity: Not app Test for overall effect:		P = 0.1	1)				0.1 0.2 0.5 1 2 5 10 Favours 2-4 Favours >4

K.8.12 High volume (20-40 ml Ringer lactate/kg) versus low volume (20 ml Ringer lactate/kg) in children with septic shock

K.8.12.1 Mortality at 28 days

Figure 180: Cumulative 72-hour survival

	20-40ml RL	per kg	20ml RL	per kg		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Santhanam 2008	52	74	55	73	100.0%	0.93 [0.77, 1.14]	-
Total (95% CI)		74		73	100.0%	0.93 [0.77, 1.14]	•
Total events	52		55				
Heterogeneity: Not ap	plicable						0.1 0.2 0.5 1 2 5 10
Test for overall effect:	Z = 0.69 (P = 0)	0.49)					Favours High Favours Low

K.9 Escalation of care

None.

K.10 Inotropic agents and vasopressors

K.10.1 Norepinephrine versus vasopressin for adults with septic shock

K.10.1.1 Mortality

Figure 181: 28-day mortality

	Norepinephrine		Vasopressin			Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Russell 2008	150	382	140	396	100.0%	1.11 [0.93, 1.33]	
Total (95% CI)		382		396	100.0%	1.11 [0.93, 1.33]	
Total events	150		140				
Heterogeneity: Not ap	•					-	0.5 0.7 1 1.5 2
Test for overall effect:	: Z = 1.13 (P =	= 0.26)					Favours Norepinephrine Favours Vasopressin

Figure 182: 90-day mortality

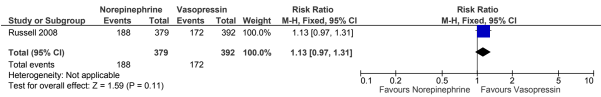


Figure 183: ICU mortality

	Norepinep	hrine	Vasopre	essin		Risk Ratio		Risk	Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI		M-H, Fixe	ed, 95% CI		
Lauzier 2006	3	10	3	13	24.6%	1.30 [0.33, 5.12]			-		
Morelli 2009 (TERLIVAP)	10	15	8	15	75.4%	1.25 [0.69, 2.26]					
Total (95% CI)		25		28	100.0%	1.26 [0.72, 2.21]		~			
Total events	13		11								
Heterogeneity: Chi ² = 0.00, Test for overall effect: Z = 0			0%				0.1	0.2 0.5 Favours Norepinephrine	1 2 Favours Va	5 sopressin	10

K.10.1.2 Adverse events

Figure 184: Requiring renal replacement therapy

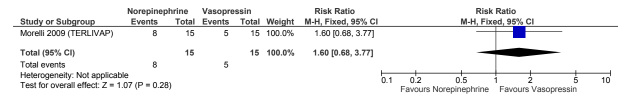
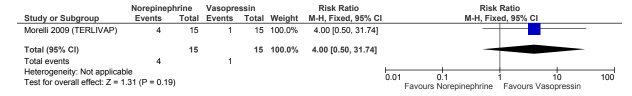


Figure 185: New onset of tachyarrhythmias



Note: this forest plot has a different scale

K.10.2 Norepinephrine versus dopamine for adults with septic shock

K.10.2.1 Mortality

Figure 186: 28-day mortality

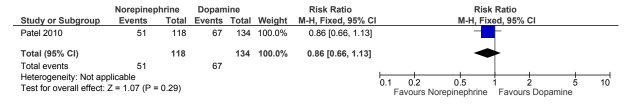


Figure 187: All-cause mortality

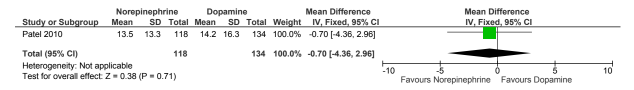
	Norepinep	hrine	Dopan	nine		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% C	M-H, Fixed, 95% CI
Marik 1994	5	10	6	10	21.4%	0.83 [0.37, 1.85]	
Mathur 2007	14	25	19	25	67.9%	0.74 [0.49, 1.11]	
Ruokonen 1993	4	5	3	5	10.7%	1.33 [0.58, 3.09]	-
Total (95% CI)		40		40	100.0%	0.82 [0.59, 1.15]	•
Total events	23		28				
Heterogeneity: Chi2 =	1.55, df = 2 (F	P = 0.46); $I^2 = 0\%$				
Test for overall effect:	Z = 1.15 (P =	0.25)					0.1 0.2 0.5 1 2 5 10 Favours Norepinephrine Favours Dopamine

Figure 188: Hospital mortality

	Norepinep	hrine	Dopan	nine		Risk Ratio	Risk Ratio		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95%	CI	
Martin 1993	7	16	10	16	100.0%	0.70 [0.36, 1.37]	-		
Total (95% CI)		16		16	100.0%	0.70 [0.36, 1.37]			
Total events	7		10						
Heterogeneity: Not ap Test for overall effect:		: 0.30)					0.01 0.1 1 Favours Norepinephrine Favours	10 s Dopamine	100

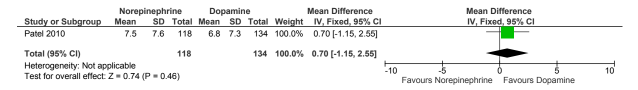
K.10.3 Duration of hospital stay

Figure 189: Length of stay in hospital



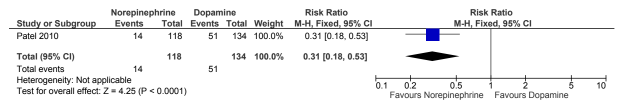
K.10.4 Duration of critical care stay

Figure 190: ICU length of stay



K.10.5 Adverse events

Figure 191: Incidence of arrhythmias



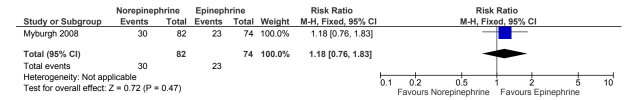
K.10.6 Norepinephrine versus epinephrine for adults with septic shock

K.10.6.1 Mortality

Figure 192: 28-day mortality

	Norepinep	hrine	Epinepl	nrine		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	M-H, Fixed, 95% CI
Myburgh 2008	24	82	17	76	100.0%	1.31 [0.76, 2.24]	
Total (95% CI)		82		76	100.0%	1.31 [0.76, 2.24]	
Total events	24		17				
Heterogeneity: Not ap Test for overall effect:		0.33)					0.1 0.2 0.5 1 2 5 10 Favours Norepinephrine Favours Epinephrine

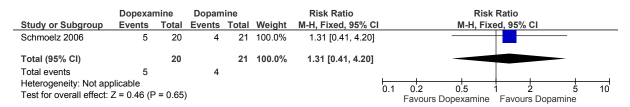
Figure 193: 90-day mortality



K.10.7 Dopexamine versus dopamine for adults with septic shock

K.10.7.1 Mortality at 28 days

Figure 194: 28-day mortality



K.10.8 Norepinephrine plus dobutamine versus epinephrine for adults with septic shock

K.10.8.1 Mortality

Figure 195: 7-day mortality



Figure 196: 14-day mortality

	Norepi + dobuta	amine	Epinepl	nrine		Risk Ratio			Risk	Ratio			
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI			M-H, Fix	ed, 95%	CI		
Annane 2007 (CATS)	44	169	56	161	100.0%	0.75 [0.54, 1.04]				t			
Total (95% CI)		169		161	100.0%	0.75 [0.54, 1.04]			•	+			
Total events	44		56										
Heterogeneity: Not appl Test for overall effect: Z							0.1 Fa	0.2 vours Nore	0.5 epi + dobutam.	1 Favou	2 irs Epiner	5 ohrine	10

Figure 197: 28-day mortality

	Norepi + dobuta	amine	Epinepl	nrine		Risk Ratio			Risk	Ratio)		
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI			M-H, Fix	ed, 95	% CI		
Annane 2007 (CATS)	58	169	64	161	100.0%	0.86 [0.65, 1.14]			_	-			
Total (95% CI)		169		161	100.0%	0.86 [0.65, 1.14]			•	-			
Total events	58		64										
Heterogeneity: Not app Test for overall effect: 2							0.1 Fa	0.2 vours Nore	0.5 epi + dobutam.	1 Favo	2 ours Epinep	5 ohrine	10

Figure 198: 90-day mortality

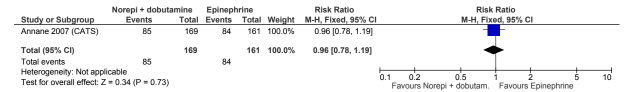


Figure 199: All-cause mortality

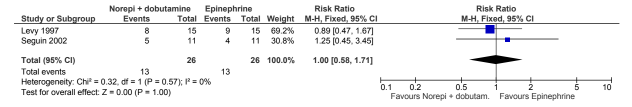


Figure 200: Mortality at discharge from the ICU

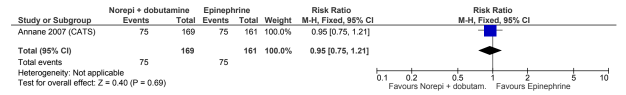


Figure 201: Mortality at discharge from the hospital

	Norepi + dobuta	amine	Epinepl	nrine		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Fixed, 95% CI	I M-H, Fixed, 95% CI
Annane 2007 (CATS)	82	169	84	161	100.0%	0.93 [0.75, 1.15]	-
Total (95% CI)		169		161	100.0%	0.93 [0.75, 1.15]	•
Total events	82		84				
Heterogeneity: Not appl Test for overall effect: Z							0.1 0.2 0.5 1 2 5 10 Favours Norepi + dobutam. Favours Epinephrine

K.10.8.2 Adverse events

Figure 202: Number of adverse events during catecholamine infusion



Figure 203: Number of adverse events after catecholamine infusion



K.10.9 Norepinephrine plus dopexamine versus epinephrine for adults with septic shock

K.10.9.1 Mortality

Figure 204: 28-day mortality

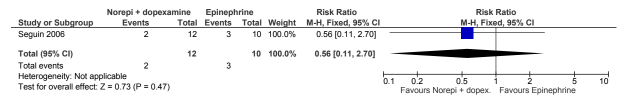
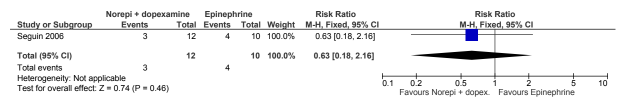


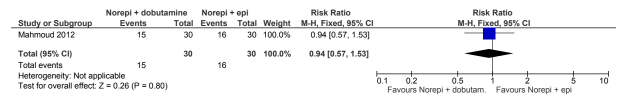
Figure 205: 90-day mortality



K.10.10 Norepinephrine plus epinephrine versus norepinephrine plus dobutamine for adults with septic shock

K.10.10.1 Mortality at 28 days

Figure 206: 28-day mortality



K.10.10.2 Number of organs supported

Figure 207: SOFA score at start

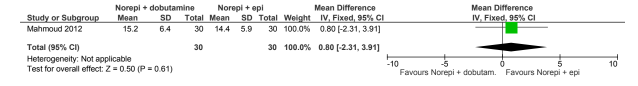


Figure 208: SOFA score at 24 hours

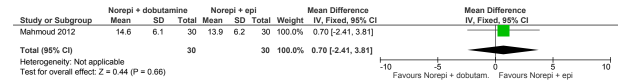


Figure 209: SOFA score at 48 hours

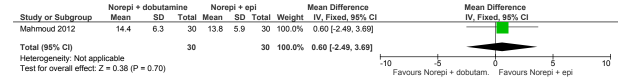


Figure 210: SOFA score at 72 hours

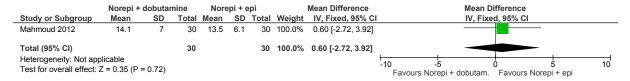
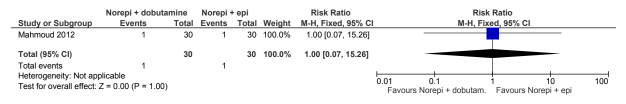


Figure 211: SOFA score at 96 hours

	Norepi +	dobutar	nine	Nore	pi + e	epi		Mean Difference		Mean Di	fference		
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI		IV, Fixe	d, 95% CI		
Mahmoud 2012	13.5	6.9	30	12.7	6.6	30	100.0%	0.80 [-2.62, 4.22]					
Total (95% CI)			30			30	100.0%	0.80 [-2.62, 4.22]					
Heterogeneity: Not appl Test for overall effect: Z		= 0.65)							-10 - Favours Nor	5 epi + dobutam.	Favours Norep	5 i + epi	10

K.10.10.3 Adverse events

Figure 212: Acute coronary syndrome



Note: this forest plot has a different scale

Figure 213: Arrhythmias



Figure 214: Cerebral stroke

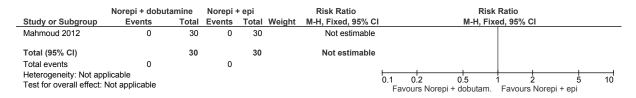
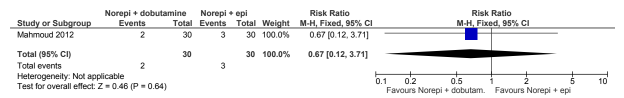


Figure 215: Limb ischaemia



K.11 Supplemental oxygen

None.

K.12 Use of bicarbonate

Figure 216: Bicarbonate versus no bicarbonate in sepsis. 28-day mortality



K.13 Early goal-directed therapy (EGDT)

K.13.1 The effect of EGDT versus a non-EGDT resuscitation strategy for people presenting to the ED with septic shock

K.13.1.1 Mortality

Figure 217: Primary mortality outcome of each study

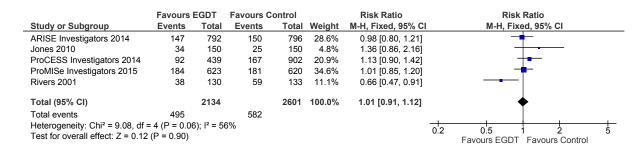


Figure 218: 90-day mortality

	Favours EG	DT Favours	Control		Risk Ratio	Risk Ratio
Study or Subgroup	Events T	otal Events	Total	Weight	M-H, Fixed, 95% CI Year	M-H, Fixed, 95% CI
ProMISe Investigators 2015	184	623 181	620	35.8%	1.01 [0.85, 1.20]	
ProCESS Investigators 2014	129	405 267	827	34.7%	0.99 [0.83, 1.17] 2014	
ARISE Investigators 2014	147	792 150	796	29.5%	0.98 [0.80, 1.21] 2014	
Total (95% CI)	1	820	2243	100.0%	1.00 [0.90, 1.11]	
Total events	460	598				
Heterogeneity: Chi ² = 0.05, df =	= 2 (P = 0.97); I	$ ^2 = 0\%$			-	0.7 0.85 1 1.2 1.5
Test for overall effect: Z = 0.09	(P = 0.93)					Favours EGDT Favours Control

K.13.1.2 ICU Utilisation

Figure 219: ICU admission^a



a: ICU admission refers to the rate of ICU admission from ED; 'favours EGDT means a lower ICU admission rate for the EGDT group and 'favours control' means a higher ICU admission rate for the EDGT group in the given trial.

Figure 220: ICU length of stay for patients admitted to ICU (days)

	Favo	urs E	GDT	Favou	urs Cor	ntrol		Mean Difference	Mean Difference
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Fixed, 95% CI	IV, Fixed, 95% CI
1.4.1 New Subgroup									
ARISE Investigators 2014	4.9	7.2	725	4.8	5.9	661	42.8%	0.10 [-0.59, 0.79]	-
Jones 2010	5.6	7.39	150	5.9	8.46	150	6.3%	-0.30 [-2.10, 1.50]	
ProCESS Investigators 2014	5.1	6.3	401	4.9	6.5	774	34.6%	0.20 [-0.57, 0.97]	
ProMISe Investigators 2015 Subtotal (95% CI)	5.7	8.1	549 1825	6.4	9.8	466 2051	16.3% 100.0%	-0.70 [-1.82, 0.42] -0.02 [-0.47, 0.43]	•
Heterogeneity: Chi ² = 1.94, df	= 3 (P =	0.58); I	2 = 0%						
Test for overall effect: Z = 0.09	9 (P = 0.9	3)							
Total (95% CI)			1825			2051	100.0%	-0.02 [-0.47, 0.43]	•
Heterogeneity: Chi ² = 1.94, df	= 3 (P =	0.58); I	2 = 0%					-	
Test for overall effect: Z = 0.09	P = 0.9	3)							Favours EGDT Favours Control
Test for subgroup differences:	Not appl	icable							1 avours LGD1 Favours Control

K.14 Monitoring

None.

K.15 Patient education, information and support

None.

K.16 Training and education

None.

Appendix L: Excluded clinical studies

L.1 Scoring systems

Table 35: Studies excluded from the clinical review

Table 33. Stut	dies excluded from the chilical review
Reference	Reason for exclusion
Adrie 2009 ²⁷	Setting (ICU)
Acharya 2007 ¹⁸	Setting (ICU)
Ait-Oufella 2011 ⁴²	Setting (ICU)
Alberti 2005 ⁵⁹	Setting (ICU)
Alsous 2000 ⁷⁰	Setting (ICU)
Anon 1999 ¹	Not scoring tool
Arnell 1996 ⁹⁶	Setting (ICU)
Arregui 1991 ⁹⁹	Setting (ICU)
Artero 2010 ¹⁰¹	Setting (ICU)
Ausania 2015 ¹⁰⁶	Not scoring tool
Bagshaw 2012 ¹²⁰	Not scoring tool (biomarkers)
Bains 2012 ¹³¹	Not scoring tool
Bang 2005 ¹³⁸	Not scoring tool
Barriere 1995 ¹⁴⁹	Systematic review including ICU setting
Baumgartner 1992 ¹⁵	Setting (ICU)
Bassetti 2014 ¹⁵³	Setting (ICU)
Bayer 2015 ¹⁶⁰	Development of a new scoring system, not externally validated
Beck 2014 ¹⁶³	Setting (ICU)
Behdad 2006 ¹⁶⁴	Population
Bencosme 1996 ¹⁶⁹	Setting (ICU)
Billeter 2009 ¹⁸⁹	Outcomes not analysed for scoring tool
Bleeker 2001 ¹⁹²	Not scoring tool
Boniatti 2011 ²¹²	Setting (ICU)
Bonig 2000 ²¹³	Setting (ICU)
Brunkhorst 2000 ²³⁴	Diagnostic accuracy of PCY, not a scoring system
Buist 2000 ²³⁸	Setting (ICU)
Byrne 1989 ²⁴⁴	Not scoring tool (theory behind the development of ASESPSIS)
Calle 2012 ²⁴⁸	Systematic review with different protocol
Calvano 1998 ²⁴⁹	Setting (surgical ICU)
Chan 2005 ²⁷⁹	Setting (ICU)
Charles 2008 ²⁸⁵	Setting (ICU)
Chawla 2007 ²⁸⁸	Setting (ICU)
Chen 2011 ²⁹⁸	Setting (ICU)
Chen 2006B ²⁹²	Setting (ICU)
Chen 1994 ²⁹¹	Setting (SICU)
Chen 2012 ²⁹⁶	Outcomes not analysed in relation to scoring tool
Close 2011 ³¹⁹	Not scoring tool

Reference	Reason for exclusion
Coslovsky 2015 ³³²	Development of a new scoring system, not externally validated
Cook 1992 ³²⁷	Setting (ICU)
Couto-Alves 2013 ³³⁴	Setting (PICU)
Croce 1992 ³³⁶	Setting (post-trauma). Outcomes not analysed in relation to scores at admission
Dabar 2015 ³⁴⁵	Comparison
Dabhi 2014 ³⁴⁶	Setting (ICU)
Das 2014 ³⁵³	Setting and when scores taken (post-surgical)
De Azevedo 2015 ³⁵⁷	Setting (ICU)
Deleon 2005 ³⁶³	Setting (PICU)
Dellinger 1988 ³⁷⁵	Setting (ICU)
Derkx 1996 ³⁷⁸	Setting (ICU)
Desai 2013 ³⁷⁹	Setting (MICU)
Eisen 2006 ⁴¹¹	Not scoring system
Elias 2015 ⁴¹⁶	Setting (ICU)
Emparanza 1988 ⁴¹⁹	Setting (PICU)
Escobar 2014 ⁴²⁹	Score immediately after birth (prior to hospital discharge)
Feng 2013 ⁴⁴⁶	Setting (ICU)
Flores 2001 ⁴⁶¹	Setting (ICU)
Furtado 2012 ⁴⁷²	Setting (ICU)
Garcia Paez 2008 ⁴⁸⁴	Not scoring system
Giamarellos-Bourboulis 2012 498	Setting (ICU)
Gogos 2003 ⁵⁰⁵	Not scoring system
Goitein 1985 ⁵⁰⁶	Setting (PICU)
Granja 2013 ⁵¹⁶	Setting (ICU)
Grozdanovski 2012 ⁵²²	Setting (ICU)
Hachimi-Idrissi 1998 ⁵⁴³	Setting (ICU)
Han 2006 ⁵⁵⁰	Narrative review
Henry 2015 ⁵⁶¹	Setting (ICU)
Hillas 2010 ⁵⁷²	Setting (ICU)
Hoen 1993 ⁵⁷⁸	Not scoring system
Holme 2013 ⁵⁷⁹	Setting (NICU), population (neonates)
Inal 2009 ⁵⁹⁷	Setting (ICU)
Jaimes 2005 ⁶⁰⁹	Outcomes not analysed
Jiang 2015 ⁶²²	Setting (ICU)
Jones 2008 ⁶²⁷	Incorrect study design
Kaur 2014 ⁶⁴³	Setting (PICU)
Kellner 2013 ⁶⁴⁷	Setting (ICU)
Khwannimit 2009 ⁶⁵⁷	Setting (ICU)
Kumar 2003 ⁶⁹¹	Setting (ICU included in outcome with ward)
Landesberg 2015 ⁷⁰⁶	No prognostic scores
Legall 1993 ⁷¹³	Setting (ICU)
Lee 1993 ⁷¹⁶	Setting (ICU)

Reference	Reason for exclusion
Maher 1989 ⁷⁵⁹	Setting (ICU)
Marra 2006a ⁷⁷⁴	Setting (ICU)
Marshall 2014 ⁷⁷⁶	Narrative review
McGillicuddy 2009 ⁷⁹¹	Not diagnostic accuracy of a scoring system
Mei 2007 ⁷⁹⁶	Not diagnostic accuracy of a scoring system
Mohan 2015 ⁸¹⁵	Setting (ICU)
Moreno 1999 ⁸²⁴	Setting (ICU)
Naved 2011 ⁸⁴²	Setting (ICU)
Oda 2000 ⁸⁷⁰	Setting (ICU)
Paul 2006 ⁹⁰⁴	Development of a new scoring system, not externally validated
Paul 2007A ⁹⁰⁶	Not mortality predictor
Pilz 1991 ⁹²¹	Not a study
Pollock 1991 ⁹²⁷	Setting (PICU)
Pollock 1997 ⁹²⁶	Setting (PICU)
Presterl 1997 ⁹³⁶	Setting (ICU)
Que 2015 ⁹⁴⁶	Setting (ICU)
Rhee 2009 ⁹⁶⁶	Setting (ICU)
Richards 2011 ⁹⁶⁷	Setting (ICU)
Rixen 1996 ⁹⁷⁴	Setting (ICU)
LeGall1993 ⁷¹³	Setting (ICU)
Rogy 1996 ⁹⁸²	Setting (surgical ICU)
Rosenberg 2002 ⁹⁸⁷	Setting (ICU)
Routsi 2007 ⁹⁸⁹	Setting (ICU)
Shapiro 2009 ¹⁰²⁵	Not scoring tool
Silva 2001a ¹⁰³⁸	Setting (PICU)
Smith 2008 ¹⁰⁴⁷	Systematic review with different protocol
Smith 2008B ¹⁰⁴⁸	Systematic review with different protocol
Tafelski 2015 ¹⁰⁸⁶	Setting (ICU)
Tsai 2014 ¹¹⁰⁶	Not a scoring tool
Ueda 2014 ¹¹¹⁶	Setting (ICU)
Umscheid 2015 ¹¹¹⁹	Development of a new scoring system, not externally validated
van de Voorde 2013 ¹¹²¹	Outcomes not analysed in relation to scoring tool
Vincent 2011 1144	Outcomes not analysed in relation to scores at admission
Vincent 2011A ¹¹⁴⁴	Changes in score not analysed in regards to admission
Vincent 1996 1143	Not a study
Vincent 2003 ¹¹⁴⁵	Not a study
Viallon 2008 ¹¹⁴¹	Not scoring tool
Wang 2010 ¹¹⁵⁸	Setting (ICU)
Wilson 1990 ¹¹⁷²	Setting (post-surgical). Outcomes not analysed in relation to scores at admission
Wong 2008 ¹¹⁷⁹	Setting (PICU)
Wong 2014 ¹¹⁸⁰	Setting (PICU)
Wunder 2004 ¹¹⁸¹	Setting (ICU)

L.2 Signs and symptoms

Table 36: Studies excluded from the clinical review

Reference	Reason for exclusion
Aalto 2004 ¹⁰	No relevant outcomes and does not match review question (blood test)
Abrahamsen 2013 ¹⁴	No relevant outcomes
Abudu 2002 ¹⁵	No relevant outcomes and does not match review question (no signs and symptoms considered)
Acosta 2012 19	Inappropriate study design (case control)
Adam 2013 ²⁰	Not a study
Adams 1993 ²²	No relevant outcomes and does not match review question (maternal risk factors on neonatal outcomes) Incorrect study design (case-control study)
Adejuyigbe 2001 ²³	No relevant outcomes and does not match review question (no signs and symptoms considered)
Aebi 1996 ²⁹	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Ahkee 1997 ³⁴	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Ahn 2013A ³⁷	No relevant outcomes and does not match review question (blood test)
Aina-Mumuney 2007 ⁴⁰	No relevant outcomes and does not match review question (foetal monitoring on neonatal outcomes) Incorrect study design (case-control study)
Akpede 1993 ⁴⁴	No relevant outcomes and does not match review question (no signs and symptoms considered)
Akpede 1994 ⁴³	No relevant outcomes and does not match review question (prediction of meningitis in children with fever and seizure)
Al Jarousha 2008 ⁴⁶	Incorrect study design (case-control study)
Alam 2014 ⁵²	No relevant outcomes and does not match review question (maternal risk factors on neonatal outcomes)
Alberti 2005 ⁵⁹	No relevant outcomes
Alexander 1998 ⁶¹	No relevant outcomes and does not match review question (maternal risk factors on neonatal outcomes)
Alexander 1999 ⁶²	No relevant outcomes and does not match review question (maternal risk factors on neonatal outcomes)
Aliberti 2008 ⁶⁵	No relevant outcomes and does not match review question (prediction of clinical failure related to CAP)
Aliberti 2015 ⁶⁴	No relevant outcomes and does not match review question
Almuneef 2000 ⁶⁷	No relevant outcomes and does not match review question (maternal risk factors on neonatal outcomes)
Altunhan 2011 ⁷¹	No relevant outcomes and does not match review question (diagnostic accuracy of PCT)
Alves 2010 ⁷³	No relevant outcomes and does not match review question (no signs and symptoms considered)
Alves 2011 ⁷²	No relevant outcomes and does not match review question (no signs and symptoms considered)

Reference	Reason for exclusion
Ammann 2013 75	Setting not relevant.
Andersen 2004 ⁷⁸	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Andrews 2012 ⁸²	Systematic review with different protocol
Angsuwat 2010 84	No analysis on relevant outcomes.
Anon 2007 ³	Abstract only
Antonow 1998 86	No relevant outcomes and does not match review question (inappropriate comparisons)
Ariffin 2002 ⁹²	No relevant outcomes
Arsura 1998 ¹⁰⁰	No relevant outcomes and does not match review question (RDS). Sample size
Asiimwe 2015 ¹⁰²	No relevant analysis (no predictor analysis)
Ayoola 2003 ¹¹²	No relevant analysis.
Babay 2005 ¹¹³	No relevant outcomes and does not match review question (not a prognostic study; 8% of patients had sepsis)
Bagshaw 2007 129	No analysis on relevant outcomes. No relevant outcomes and does not match review question
Bagshaw 2008 ¹²⁶	No relevant outcomes and does not match review question (sepsis as risk factor for acute kidney injury)
Bang 2005b ¹³⁷	No relevant analysis.
Barati 2013 ¹⁴¹	No relevant outcomes and does not match review question (diagnostic accuracy of brain natriuretic peptide)
Barie 2004 ¹⁴⁵	No relevant outcomes and does not match review question (identification of source of infection)
Barnaby 2002 ¹⁴⁶	No relevant outcomes
Bas 2011 ¹⁵¹	No relevant outcomes and does not match review question (no signs and symptoms considered)
Baskaran 2008 ¹⁵²	No relevant outcomes and does not match review question (no signs and symptoms considered)
Bastos 1993 ¹⁵⁴	Does not match review question (GCS as predictor of mortality in any non-traumatic ICU admission; 3% had sepsis)
Bayer 2015 ¹⁶⁰	No relevant analysis (no signs and symptoms analysed)
Bejan 2014A ¹⁶⁶	No relevant analysis.
Bekhof 2013 ¹⁶⁷	Population does not match protocol (preterm infants)
Benito 2013 ¹⁷²	No relevant outcomes and does not match review question (diagnostic accuracy of blood tests)
Bernstein 2007 ¹⁸¹	No relevant outcomes and does not match review question (diagnostic accuracy of PCT)
Bettiol 2012 ¹⁸²	Cochrane review
Bettiol 2012 ¹⁸³	Cochrane review
Beuchee 2009 ¹⁸⁴	Population does not match protocol (preterm infants)
Bilavsky 2009 ¹⁸⁷	No relevant outcomes and does not match review question (diagnostic accuracy of blood tests)
Bilbault 2004 ¹⁸⁸	Does not match review question (gene expression)
Bizzarro 2011 ¹⁹⁰	No relevant outcomes and does not match review question (RDS)
Bleeker 2007 ¹⁹¹	Does not match review question (diagnostic accuracy of a tool to predict bacteraemia)

Reference	Reason for exclusion
Bochicchio 2001 ¹⁹⁵	Does not match review question (SIRS score to predict risk of infection)
Bochud 1994 ¹⁹⁶	Systematic review with different protocol
Boersma 1999 ¹⁹⁷	Does not match review question (review on discriminant rules to predict mortality in patients with community acquired pneumonia)
Bogar 2006 ¹⁹⁸	Does not match review question (diagnostic accuracy of PCT and leucocyte anti-sedimentation rate to predict bacteraemia)
Boland 1994 ²⁰⁰	No relevant outcomes and does not match review question (no signs and symptoms considered)
Bonadio 1990 ²⁰⁷	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Bonadio 1992 ²⁰⁹	No relevant outcomes and does not match review question (diagnostic accuracy of blood tests to predict serious bacterial infection)
Bonadio 1993 ²⁰⁶	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Bonadio 1993B ²¹⁰	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered, identification of pathogen)
Bonadio 1993C ²⁰⁸	No relevant outcomes and does not match review question (diagnostic accuracy of Young Infant Observation Scale to predict infection)
Bonig 2000 ²¹³	Does not match review question (blood tests)
Bonsu 2003 ²¹⁴	Does not match review question (diagnostic accuracy of WBC to predict bacteraemia)
Boockvar 2013 ²¹⁵	No relevant outcomes and does not match review question (predictors of delirium)
Bossink 1998 ²²⁰	No relevant outcomes
Bossink 1999 ²¹⁷	No relevant outcomes and does not match review question (development of model)
Bossink 2001 ²¹⁸	No relevant outcomes
Bozzetti 1991 ²²³	No relevant outcomes and does not match review question (no signs and symptoms considered)
Bressan 2012 ²²⁸	Does not match review question (diagnostic accuracy of PCT, CRP, WBC to predict serious bacterial infection)
Bressan 2012A ²²⁷	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Breuling 2015 ²²⁹	No relevant analysis (no diagnostic accuracy data)
Brunkhorst 2000 ²³⁴	No relevant outcomes and does not match review question (diagnostic accuracy of PCT)
Byer 2006 ²⁴²	Does not match review question (prediction of hypotension or toxic shock syndrome in patients with fever and erythroderma)
Caksen 2000 ²⁴⁵	No relevant outcomes and does not match review question (distribution of patients according to symptoms for septic arthritis and osteomyelitis)
Caljouw 2011 ²⁴⁷	No relevant outcomes and does not match review question
Carbonell 2008 ²⁵¹	No relevant outcomes and does not match review question
Carrieri 2003 ²⁵⁵	No relevant outcomes and does not match review question
Chaboyer 2008 ²⁷³	Does not match review question (prediction of adverse events after discharge from ICU; sepsis: 22%)
Chan 2014 ²⁸⁰	No relevant outcomes and does not match review question (biomarker profiling for the prediction of neutropenic fever)

Reference	Reason for exclusion
Chassagne 1996 ²⁸⁶	Incorrect analysis (no data given to validate summary results)
Chen 1992 ²⁸⁹	No relevant outcomes and does not match review question
Chen 2002 ²⁹³	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Chen 2007 ³⁰⁰	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Chen 2012B ²⁹⁹	No relevant outcomes and does not match review question (no signs and symptoms considered)
Chen 2014 ²⁹⁰	No relevant outcomes and does not match review question
Chia 1991 ³⁰³	No relevant outcomes and does not match review question (no signs and symptoms considered)
Chisti 2010 ³⁰⁶	Population not relevant (those with diarrhoea only in Bangladesh)
Chiu 1997 ³⁰⁷	No relevant outcomes and does not match review question (no signs and symptoms considered)
Churgay 1994 ³¹¹	No relevant outcomes and does not match review question (maternal risk factors on neonatal outcomes)
Chwals 1994 ³¹³	No relevant outcomes and does not match review question (diagnostic accuracy of blood tests)
Clemmer 1992 ³¹⁸	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Coburn 2012 321	Systematic review with different protocol.
Comstedt 2009 ³²⁵	No relevant outcomes and does not match review question (no signs and symptoms considered)
Corona 2004 ³²⁹	No relevant outcomes and does not match review question
Craig 2010 335	Outcomes reported only in figure.
D'Orio 1990 ³⁴²	No relevant outcomes.
da Silvia 2007 ³⁴³	No relevant outcomes and does not match review question (diagnostic accuracy of PCT)
Dalegrave 2012 ³⁴⁷	No relevant outcomes and does not match review question (no signs and symptoms considered)
Damas 1997 ³⁵⁰	No relevant outcomes and does not match review question (no signs and symptoms considered)
Daoud 1995 352	No relevant outcomes and does not match review question
Day 1992 ³⁵⁶	No relevant outcomes and does not match review question (maternal risk factors on neonatal outcomes)
de Macedo 2003 ³⁶⁴	No relevant outcomes.
De 2013 ³⁷⁰	No relevant outcomes and does not match review question (review traffic light system for predicting serious bacterial infections)
De2014 ³⁷¹	No relevant outcomes and does not match review question (diagnostic accuracy of blood tests to predict serious bacterial infection)
Devaux 1992 ³⁸³	No relevant outcomes and does not match review question (no signs and symptoms considered)
Dewhurst 2008 ³⁸⁶	Population does not match protocol (preterm infants)
Dickinson 2010 ³⁸⁹	Incorrect study design (narrative review)
Diepold 2008 ³⁹⁰	No relevant outcomes and does not match review question (diagnostic accuracy of blood tests: IL-6 and IL-8)
Dior 2014 ³⁹³	No relevant outcomes and does not match review question (maternal risk

Reference	Reason for exclusion
	factors on neonatal outcomes)
Dorio 1990 ³⁴²	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Drewry 2013 ³⁹⁸	Incorrect study design (case-control study)
Drewry 2015 399	No relevant analysis (no predictor analysis)
Drvar 2013 ⁴⁰²	No relevant outcomes and does not match review question
Dunser 2009 ⁴⁰⁸	No relevant outcomes reported
Dwyer 2011 ⁴⁰⁹	No relevant outcomes and does not match review question (diagnostic accuracy of prediction rules)
Ebersoldt 2007 ⁴¹⁰	Systematic review with different protocol
Elbanks 1993 ⁴³³	No relevant outcomes and does not match review question
Elting 1992 ⁴¹⁸	No relevant outcomes and does not match review question
Escobar 2000 ⁴²⁸	No relevant outcomes and does not match review question (maternal risk factors on neonatal outcomes)
Fairchild 2010 ⁴³⁷	Incorrect study design (narrative paper)
Fairchild 2013A ⁴³⁶	Incorrect study design (narrative paper)
Falguera 2009 ⁴³⁹	No relevant outcomes
Farley 1993 ⁴⁴²	No relevant outcomes and does not match review question
Fernandez-Perez 2005 ⁴⁴⁷	Review with different protocol
Fialkow 2006 451	No relevant outcomes and does not match review question
Figueroa-Damian 1999 452	No relevant outcomes and does not match review question
Filbin 2014 ⁴⁵³	No relevant outcomes and does not match review question
Finfer 2004 ⁴⁵⁴	No relevant outcomes and does not match review question
Fleming 2011 ⁴⁶⁰	Does not match protocol (no relevant analysis or outcomes)
Fok 1998 ⁴⁶³	No relevant outcomes and does not match review question (RDS). Setting not relevant
Galanakis 2002 ⁴⁷⁴	No relevant outcomes and does not match review question (RDS)
Galetto-Lacour 2010 ⁴⁷⁵	No relevant outcomes and does not match review question (diagnostic accuracy of blood tests to predict serious bacterial infection)
Gallagher 1994 ⁴⁷⁶	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Garra 2005 ⁴⁹⁰	No relevant outcomes and does not match review question (no signs and symptoms considered)
Gavazzi 2005 ⁴⁹¹	No relevant outcomes and does not match review question (no signs and symptoms considered)
George 1997 ⁴⁹²	No relevant outcomes and does not match review question (predictors of delirium)
Ghiorghis 1992 ⁴⁹⁶	Incorrect study design (case-control study)
Gille-Johnson 2012 ⁵⁰⁰	No relevant outcome
Goerlich 2014 ⁵⁰⁴	No relevant outcomes and does not match review question (no signs and symptoms considered)
Gogos 2003 ⁵⁰⁵	Does not match protocol (no relevant analysis or outcomes)
Goulet 2014 ⁵¹³	No relevant outcomes and does not match review question (no signs and symptoms considered)
Grander 2013 ⁵¹⁴	Does not match review question (prediction of mortality from critical illness, 8% sepsis)

Reference	Reason for exclusion
Griffin 2005 ⁵¹⁸	No relevant outcomes (results from multivariable analysis available in graphic form only)
Griffin 2007 ⁵¹⁹	No relevant outcomes (results from multivariable analysis available in graphic form only)
Guo 2015 ⁵³⁴	No relevant population (not people with sepsis)
Haj-Hassan 2011 ⁵⁴⁵	No relevant outcome
Hashavya 2001 ⁵⁵⁶	No relevant outcomes and does not match review question (blood test)
Hazan 2014 ⁵⁵⁸	No relevant outcomes and does not match review question (diagnostic accuracy of blood tests to predict serious bacterial infection)
Herbst 1997 ⁵⁶⁴	No relevant outcomes and does not match review question (maternal risk factors on neonatal outcomes)
Hernandez 2012 ⁵⁶⁶	No relevant outcomes and does not match review question (predictors of resuscitation)
Horeczko 2013 ⁵⁸⁴	No relevant outcomes and does not match review question
Housinger 1993 588	No relevant outcomes and does not match review question (blood test)
Hsiao 2006 ⁵⁹⁰	Outcomes not relevant (no analysis)
Ireland 2014 ⁵⁹⁹	No relevant outcomes and does not match review question (maternal predictors). Inappropriate comparison
Isfandiaty 2012 ⁶⁰²	No relevant outcomes and does not match review question (sepsis as a predictor of delirium)
Ismail 1997 ⁶⁰⁴	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered; prediction of nosocomial bacteraemia)
Iwashyna 2012 ⁶⁰⁵	No relevant outcomes and does not match review question
Jacobs 1990A ⁶⁰⁷	No relevant outcomes and does not match review question (no signs and symptoms considered)
Jain 2003 ⁶¹⁰	No relevant outcomes and does not match review question
Jeddi 2010 ⁶¹⁸	No relevant outcomes and does not match review question (diagnostic accuracy of blood tests)
Juncal 2011 ⁶³¹	No relevant outcomes
Karambin 2011 638	No relevant outcomes and does not match review question
Katsimpardi 2006 ⁶⁴⁰	Does not match review question (assessment of infectious complications in paediatric patients with acute lymphoblastic leukaemia)
Kayange 2010 ⁶⁴⁴	No relevant outcomes and does not match review question (inappropriate comparison)
Khaskheli 2013 ⁶⁵²	No relevant outcomes and does not match review question (no signs and symptoms considered)
Khassawneh 2009 ⁶⁵⁴	No relevant outcomes and does not match review question (inappropriate comparison)
Khurana 2011 ⁶⁵⁶	No relevant outcomes and does not match review question
Kibuuka 2015 ⁶⁵⁸	Incorrect population (malaria population)
Kim 2011A ⁶⁶⁵	No relevant outcomes and does not match review question (no signs and symptoms considered)
Kimmoun 2013 ⁶⁶⁹	No relevant outcomes
Landesberg 2012 ⁷⁰⁵	No relevant outcomes
Lannergard 2009 ⁷⁰⁷	Does not match review question (evaluation of biomarkers as prognostic tools for the decision to stop antibiotic therapy or to investigate oral stepdown therapy after an initial course of empiric intravenous cefuroxime or

Reference	Reason for exclusion
Tierer en de	a combination of cefuroxime and tobramycin)
Laterre 2005 ⁷¹⁰	No relevant outcomes and does not match review question (no signs and symptoms considered)
Laupland 2012 ⁷¹¹	No relevant outcomes
LeDoux 2000 ⁷¹⁴	No relevant outcomes and does not match review question (effect of vasopressor therapy)
Lefrant 2010 719	No relevant outcomes and does not match review question (scoring tool)
Leichtle 2013 ⁷²⁰	No relevant outcomes and does not match review question (no signs and symptoms considered)
Levy 2005 ⁷²³	No relevant outcomes and does not match review question
Liaw 1997 ⁷²⁶	No relevant outcomes and does not match review question
Lim 2012 ⁷²⁸	Inappropriate population (pre-term infants)
Mann-Salinas 2013 ⁷⁶⁵	Incorrect study design (case-control study)
Mesquida 2012 ⁸⁰¹	No relevant outcomes and does not match review question (no signs and symptoms considered)
Metsvaht 2009 803	No relevant outcomes and does not match review question (antimicrobial)
Mikkelsen 2013 ⁸⁰⁸	No relevant outcomes and does not match review question (development of ARDS in patients with sepsis)
Mitra 1993 813	Setting not relevant
Mobin 2012 ⁸¹⁴	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Nimri 2001 ⁸⁶⁰	No relevant outcomes and does not match review question (no signs and symptoms considered)
O'Leary 2015 ⁸⁶⁵	Incorrect population
Oostenbrink 2012 ⁸⁷⁹	No relevant outcomes
Ozalay 2006 ⁸⁸⁹	No relevant analysis
Papaioannou 2012 ⁸⁹¹	No relevant outcomes reported
Piazza 2004 ⁹²⁰	No relevant outcomes and does not match review question
Pontet 2003 ⁹²⁸	No relevant outcomes reported
Pope 2010 ⁹²⁹	No relevant outcomes and does not match review question (no signs and symptoms considered)
Quach 2008 944	No relevant outcomes and does not match review question (scoring tool)
Rackoff 1996 ⁹⁴⁷	No relevant outcomes and does not match review question (no signs and symptoms considered)
Ranes 2006 ⁹⁵⁰	No relevant outcomes and does not match review question (no signs and symptoms considered)
Razzaq 2013 ⁹⁵⁷	No relevant outcomes and does not match review question (no signs and symptoms considered)
Rehman 2014 ⁹⁵⁹	Incorrect study design (narrative study)
Ronco 1994 ⁹⁸³	No analysis of relevant variables
Santolaya 2008 ¹⁰⁰⁰	No relevant outcomes and does not match review question (prognostic value of blood tests)
Schultz 2013 ¹⁰¹⁰	No relevant outcomes reported
Sevastos 2008 ¹⁰²¹	No relevant outcomes and does not match review question
Shani 2008 ¹⁰²²	No relevant outcomes and does not match review question (RDS)

Reference	Reason for exclusion
Shapiro 2009 ¹⁰²⁵	Does not match protocol (sepsis scores)
Singh 2003 ¹⁰⁴³	Population does not match protocol (preterm infants)
Sirvent 2013 ¹⁰⁴⁴	No relevant outcomes and does not match review question (scoring tool)
Smith 1997 ¹⁰⁵²	No relevant outcomes and does not match review question (review to determine the rate of bacteraemia in women with pyelonephritis)
Sole-vidan 2011 ¹⁰⁵⁴	No relevant outcomes and does not match review question
Somogyi-Zalud 2000 ¹⁰⁵⁷	No relevant outcomes and does not match review question
Spanos 2010 ¹⁰⁶²	No relevant outcomes
Spruijt 2013 ¹⁰⁶⁴	No relevant outcomes
Sprung 2006 ¹⁰⁶⁵	No relevant outcomes and does not match review question (no signs and symptoms considered)
Stathakis 2007 ¹⁰⁶⁶	No relevant outcomes and does not match review question (no signs and symptoms considered, only blood markers)
Struelens 1991 ¹⁰⁷¹	Incorrect study design (case-control study)
Suchyta 1997 ¹⁰⁷⁹	No relevant outcomes and does not match review question
Tayek 2012 ¹⁰⁹¹	Review with different protocol
Thai 2012 ¹⁰⁹⁵	No relevant outcomes and does not match review question
Thompson 2009 ¹⁰⁹⁷	Case study
Thompson 2010 ¹⁰⁹⁶	Editorial
Torres 1991 ¹¹⁰⁰	Review with different protocol.
Toweill 2000 ¹¹⁰²	No relevant outcomes and does not match review question (no signs and symptoms considered)
Tsering 2011 ¹¹¹⁰	No relevant outcomes and does not match review question
Van den Bruel 2010 ¹¹²³	Systematic review
Vandissel 2005 ¹¹²⁵	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Venugopal 2012 ¹¹³⁹	No relevant outcomes and does not match review question (no signs and symptoms considered)
Vyles 2014 ¹¹⁵⁰	No relevant outcomes and does not match review question (no uni- or multi-variable analysis for signs and symptoms considered)
Wang 2009 ¹¹⁵⁷	No relevant outcomes and does not match review question (predicting mortality in patients with bacteraemia)
Waskerwitz 1981 ¹¹⁶¹	No relevant outcomes and does not match review question (no signs and symptoms considered)
Wojkowskamach 2012 ¹¹⁷⁶	Inappropriate population (hospitalised LBW newborns)
Xi 2010 ¹¹⁸²	No relevant outcomes and does not match review question (inappropriate comparisons)
Yahav 2015 ¹¹⁸⁶	No relevant analysis (no analysis of predictors)
Yang 2013 ¹¹⁹²	No relevant outcomes and does not match review question (ARDS)
Yossuck 2002 ¹²⁰¹	Inappropriate population (newborn)
Yu 2011 ¹²⁰²	No relevant outcomes and does not match review question (blood test)
Zaidi 1999 ¹²⁰⁴	No relevant outcomes and does not match review question

L.3 Blood tests

Table 37: Studies excluded from the clinical review

Study	Exclusion reason
Abdollahi 2012 ¹²	Invalid country
Aboud 2010 ¹³	Case-control study
Adamik 2000 ²¹	Invalid diagnostic tests
Adhikari 1986 ²⁴	Invalid outcomes
Adib 2012 ²⁵	Invalid country
Agrawal 2008 ³²	Invalid country
Agyeman 2011 ³³	Invalid population
Ahmed 2005 ³⁵	Invalid country
Ahn 2012 ³⁶	Invalid diagnostic tests
Aikawa 2005 ³⁸	Invalid population
Aimoto 2014 ³⁹	Invalid population
Al 2011 ⁴⁷	Invalid diagnostic tests
Alamgir 2006 ⁵⁵	Invalid analysis
Albright 2015 ⁶⁰	Invalid diagnostic tests
Al-Majali 2004 ⁴⁸	Invalid country
Al-Nawas 1996 ⁴⁹	Invalid outcomes
Al-Nawas 1996A ⁵⁰	Procalcitonin
Altunhan 2011 ⁷¹	Invalid country
Alves 2010 ⁷³	Invalid diagnostic tests
Al-Zwaini 2009 ⁵¹	Invalid country
Ambalavanan 2005 ⁷⁴	Invalid country Invalid population
Anbar 1986 ⁷⁷	Invalid outcomes
Ando 2012 ⁷⁹	
Anwer 2000 ⁸⁸	Invalid analysis
Aquino 2012 ⁹⁰	Invalid country Invalid outcomes
Arkader 2006 ⁹³	
Arnalich 1999 ⁹⁵	Invalid country
Arnon 2007 ⁹⁸	No prognostic or diagnostic data
Aube 1992 ¹⁰⁵	Invalid analysis Published before 1999
Aydemir 2014 ¹⁰⁹	Invalid country
Aydin 2013 ¹¹¹	Invalid country
Aydin 2014 ¹¹⁰	Invalid country
Bakker 1996 ¹³²	No data given
Balci 2003 ¹³³	Invalid country
Ballot 2004 ¹³⁵	Procalcitonin
Baorto 2001 ¹⁴⁰	Invalid population
Barati 2008 ¹⁴³	Procalcitonin
Barati 2015 ¹⁴²	Invalid country
Baron 1989 ¹⁴⁸	Invalid outcomes
Bates 1990 ¹⁵⁶	Invalid outcomes
Becchi 2008 ¹⁶¹	Invalid outcomes
Bender 2008 ¹⁷¹	Procalcitonin
Benitz 1998 ¹⁷³	Invalid setting
Benuck 1983 ¹⁷⁵	Invalid outcomes
Berger 1995 ¹⁷⁶	Invalid setting
Berkman 2009 ¹⁷⁹	Invalid diagnostic tests
Bernstein 2007 ¹⁸¹	Invalid outcomes
Bhaandari 2014 ¹⁸⁵	Invalid diagnostic tests
Bianchi 2004 ¹⁸⁶	Invalid country

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Bleeker 2001 ¹⁹²	Invalid analysis
Blommendahl 2002 ¹⁹³	Invalid population
Bloos 2014 ¹⁹⁴	Narrative review
Bojic 2014 ¹⁹⁹	Invalid country
Boskabadi 2010 ²¹⁶	Case-control study
Bossink 1998 ²²⁰	Published before 1999
Bossink 1999A ²¹⁹	Invalid diagnostic tests
Bossink 2001 ²¹⁸	Invalid outcomes
Brierley 2009 ²³⁰	Narrative review
Brodska 2009 ²³¹	Procalcitonin
Broner 1990 ²³³	Invalid setting
Buck 1994 ²³⁶	Invalid population
Byl 1997 ²⁴³	Published before 1999
Caldas 2008 ²⁴⁶	Not English
Calvano 1998 ²⁴⁹	Invalid diagnostic tests
Carrol 2002 ²⁵⁶	Invalid population
Carrol 2002A ²⁵⁷	Procalcitonin
Carrol 2005 ²⁵⁸	Invalid population
Casado-Flores 2006 ²⁶¹	Invalid population
Cazalis 2013 ²⁶⁷	Invalid diagnostic tests
Cekmez 2011 ²⁶⁹	Invalid country
Celik 2010 ²⁷⁰	Invalid country
Chaaban 2009 ²⁷²	Invalid analysis
Chalupa 2011 ²⁷⁵	Invalid outcomes
Chan 1997 ²⁷⁸	Invalid population
Chan 2002 ²⁸²	Invalid country
Chan 2004 ²⁸³	Invalid country
Chan 2011 ²⁸¹	Narrative review
Charles 2008 ²⁸⁵	Invalid outcomes
Chen 2010 ²⁹⁷	Narrative review
Chen 2014 ²⁹⁴	Invalid country
Chen 2014E ³⁰¹	Invalid country
Chen 2014F ²⁹⁵	Invalid country
Chiesa 2000 ³⁰⁴	Procalcitonin
Chiesa 2003 ³⁰⁵	Invalid analysis
Claessens 2010 ³¹⁶	Invalid population
Clec'h 2004 ³¹⁷	Procalcitonin
Coggins 2013 ³²³	Invalid analysis
Collighan 2004 ³²⁴	Invalid diagnostic tests
Contenti 2015A ³²⁶	Invalid diagnostic tests
Cortegiani 2014 ³³¹	Invalid outcomes
Couto 2007 ³³³	
	Invalid diagnostic tests
Couto-Alves 2013 ³³⁴	Not relevant to review question
Craig 2010 ³³⁵	Invalid diagnostic tests
Da Silva 2007A ³⁴⁴	Invalid population
Dalton 2012 ³⁴⁹	Invalid analysis
Davis 2015 ³⁵⁴	Invalid population
De 1998 ³⁶⁹	Invalid country
de Azevedo 2015 ³⁵⁷	Invalid country
De Blasi 2013 ³⁵⁸	Invalid study design
De Jager 2010 ³⁶²	Invalid study design
Debiane 2014 ³⁷²	Invalid population
Degroot 2014 ³⁶¹	Invalid diagnostic tests

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Dettmer 2015 ³⁸¹	Invalid comparison
Devran 2012 ³⁸⁵	Invalid country
DeWerra 1997 ³⁶⁸	Published before 1999
Dhanalakshmi 2015 ³⁸⁷	Invalid country
Dierkes 2009 ³⁹¹	Invalid diagnostic tests
Diez-Padrisa 2012 ³⁹²	Invalid country
Dornbusch 2003 ³⁹⁴	Procalcitonin
Draz 2013 ³⁹⁶	Invalid diagnostic tests
Drees 2012 ³⁹⁷	Invalid diagnostic tests
Drumheller 2012 ⁴⁰¹	Invalid diagnostic tests
Du 2002 ⁴⁰³	Invalid outcomes
Du 2003 ⁴⁰⁴	Invalid country
Du 2014 ⁴⁰⁵	Invalid country
Elawady 2014 ⁴¹⁵	Invalid country
El-Maghraby 2007 ⁴¹⁴	Invalid country
Endo 2008 ⁴²⁰	Invalid analysis
Engel 1998 ⁴²¹	Invalid diagnostic tests
Ersoy 2007 ⁴²⁵	Invalid outcomes
Escobar 2015 ⁴²⁷	Animal study
Fan 1989 ⁴⁴⁰	Invalid outcomes
Feng 2012 ⁴⁴⁵	Invalid country
Fisher 2000 ⁴⁵⁶	Invalid study design
Fleischhack 2000 ⁴⁵⁸	Invalid population
Fleischhack 2000A ⁴⁵⁹	Invalid population
Galetto-Lacour 2010 ⁴⁷⁵	Invalid study design
Garcia 2007 ⁴⁸⁵	Invalid diagnostic tests
Garland 2003 ⁴⁸⁷	Invalid population
Gerdes 1987 ⁴⁹⁴	Invalid setting
Ghosh 2001 ⁴⁹⁷	Invalid country
Gille-Johnson 2012 ⁵⁰⁰	Invalid outcomes
Greenberg 1990 ⁵¹⁷	Invalid outcomes
Gu 2015 ⁵²³	Invalid analysis
Guclu 2013 ⁵²⁴	Invalid study design
Guibourdenche 2002 ⁵²⁶	Invalid analysis
Guido 2012 ⁵²⁹	Invalid outcomes
Guillois 1994 ⁵³⁰	Invalid population
Gutovitz 2011 ⁵³⁶	Invalid comparison
Guven 2002 ⁵³⁷	Procalcitonin
Hall 2011 ⁵⁴⁷	Narrative review
Hanson 1983 ⁵⁵¹ Hariharan 2011 ⁵⁵⁴	Invalid study design
	Invalid outcomes
Hegadi 2015 ⁵⁵⁹	Invalid country
Hengst 2003 ⁵⁶⁰	Invalid study design
Heper 2006 ⁵⁶²	Invalid outcomes
Hermans 2012 ⁵⁶⁵	Invalid outcomes
Hernandez-Bou 2015 ⁵⁶⁸	Invalid population
Herzum 2008 ⁵⁶⁹	Narrative review
Hisamuddin 2015 ⁵⁷³	Invalid country
Ho 2008 ⁵⁷⁶	Invalid population
Hoppensteadt 2014A ⁵⁸³	Invalid diagnostic tests
Hoppensteadt 2015 ⁵⁸²	Invalid diagnostic tests
Hui 2012 ⁵⁹²	Invalid study design
lba 2014 ⁵⁹⁴	Narrative review

Jain 2014 ⁶¹¹	Invalid country
James 1999 ⁶¹²	Narrative review
Jansen 2009 ⁶¹⁴	Invalid study design
Janum 2011 ⁶¹⁵	Invalid outcomes
Jat 2011 ⁶¹⁷	Invalid country
Jeschke 2013 ⁶²⁰	Invalid analysis
Jordan 2000 ⁶³⁰	Invalid diagnostic tests
Juutilainen 2011A ⁶³³	Invalid population
Kasem 2012 ⁶³⁹	Procalcitonin
Katz 1992 ⁶⁴¹	Invalid population
Keshet 2009 ⁶⁴⁹	Invalid population
Keßler 1994 ⁶⁵⁰	Invalid outcomes
Khassawneh 2007 ⁶⁵³	Invalid country
Kim 2013A ⁶⁶⁸	Invalid outcomes
Kirschenbaum 2006 ⁶⁷²	Invalid outcomes
Kite 1988 ⁶⁷³	Invalid population
Kobayashi 2001 ⁶⁷⁷	Invalid outcomes
Kocabas 2007 ⁶⁷⁸	Invalid country
Kocazeybek 2003 ⁶⁷⁹	Invalid population
Kohli 1993 ⁶⁸⁰	Invalid country
Kohn 2001 ⁶⁸¹	Invalid study design
Koksal 2007 ⁶⁸²	Invalid country
Kono 1999 ⁶⁸⁴	Invalid outcomes
Krediet 1992 ⁶⁸⁶	Invalid setting
Krishna 2000 ⁶⁸⁸	Invalid country
Kumar 2010 ⁶⁹²	Invalid country
Kushimoto 2007 ⁶⁹⁶	Invalid outcomes
Kyr 2007 ⁶⁹⁷	Invalid diagnostic tests
Laborada 2003 ⁶⁹⁹	Invalid diagnostic tests
Lacaze-Masmonteil 2014 ⁷⁰⁰	Invalid analysis
Laham 2014 ⁷⁰¹	Invalid population
Lam 2008 ⁷⁰²	Invalid study design
Larsen 2011 ⁷⁰⁹	Invalid outcomes
Lee 2012A ⁷¹⁵	Invalid analysis
Leli 2014 ⁷²¹	Procalcitonin
Lichtenstern 2012 ⁷²⁷	Narrative review
Luz Fiusa 2013 ⁷⁴⁹	Invalid country
Lyle 2013 ⁷⁵⁰	Narrative review
MacKay 2011A ⁷⁵³	Invalid outcomes
Magudumana 2000 ⁷⁵⁷	Invalid population
Malik 2003 ⁷⁶²	Systematic review
Mannan 2010 ⁷⁶⁷	Invalid country
Manucha 2002 ⁷⁶⁸	Invalid country
Manzano 2010 ⁷⁶⁹	Procalcitonin
Manzon 2015 ⁷⁷⁰	Invalid diagnostic tests
Marecaux 1996 ⁷⁷¹	Invalid outcomes
Martinez-Albarran 2009 ⁷⁸⁰	Invalid country
Marzouk 1993 ⁷⁸¹	Invalid population
Mathers 1987 ⁷⁸³	Invalid setting
Mazur 1994 ⁷⁸⁹	Invalid outcomes
McKenzie 2009 ⁷⁹²	Invalid study design
Meidani 2013 ⁷⁹⁷	Cross-sectional study
Meisner 1998A ⁷⁹⁸	Invalid population

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Mencacci 2012 ⁷⁹⁹	Invalid diagnostic tests
Menon 2015 ⁸⁰⁰	Invalid country
Mimoz 1998 ⁸¹⁰	Invalid outcomes
Mintegi 2009 ⁸¹¹	Invalid population
Mistry 2013 ⁸¹²	Invalid population
Mokart 2005 ⁸¹⁷	Inconsistencies regarding units of measurement
Montiel-Jarquin 2012 ⁸¹⁹	Invalid country
Munoz 2004 ⁸³²	Procalcitonin
Murphy 2012A ⁸³⁴	Invalid analysis
Mustafa 2005 ⁸³⁶	Invalid country
Mustard 1987 ⁸³⁷	Invalid population
Naher 2011 ⁸³⁸	Invalid country
Neely 1998 ⁸⁴³	Invalid setting
Neely 2004 ⁸⁴⁴	Invalid diagnostic tests
Ng 2004A ⁸⁴⁷	Narrative review
Ng 2006 ⁸⁴⁸	Narrative review
Nijman 2011 ⁸⁵⁹	Invalid outcomes
Nijman 2013 ⁸⁵⁸	Invalid analysis
Nuntnarumit 2002 ⁸⁶³	Invalid country
Oberhoffer 1999 ⁸⁶⁸	Invalid outcomes
Oliveira 2008 ⁸⁷⁷	Invalid outcomes
Oliveira 2013 ⁸⁷⁶	Invalid comparison
Opal 2014 ⁸⁸⁰	Narrative review
Örtqvist 1995 ⁸⁸⁵	Invalid outcomes
Park 2014 ⁸⁹⁶	Invalid population
Park 2014B ⁸⁹⁴	Invalid diagnostic tests
Pechorsky 2009 ⁹¹⁰	Invalid outcomes
Peduzi 1992 ⁹¹¹	Invalid setting
Peltola 1983 ⁹¹³	Invalid population
Pfitzenmeyer 1995 ⁹¹⁷	Published before 1999
Pinilla 1998 ⁹²²	Invalid population
Povoa 1998 ⁹³³	Published before 1999
Povoa 2002 ⁹³²	Narrative review
Povoa 2005 ⁹³⁴	Invalid analysis
Qu 2015 ⁹⁴³	Invalid country
Ranzani 2013 ⁹⁵¹	Invalid country
Raoofi 2014 ⁹⁵²	Procalcitonin
Rast 2015 ⁹⁵³	Invalid population
Ravishankar 2009 ⁹⁵⁴	Invalid study design
Ravishankaran 2011 ⁹⁵⁵	Invalid study design
Reed 2013 ⁹⁵⁸	Invalid analysis
Resch 2003 ⁹⁶³	
Riche 2003 ⁹⁶⁸	Invalid population Invalid population
Riedel 2011 ⁹⁷⁰	
Riedel 2011 ⁹⁷⁹	Procalcitonin Procalcitonin
Rondina 201 ⁹⁸⁴	
	Invalid diagnostic tests
Rønnestad 1999 ⁹⁸⁵	Invalid analysis
Sakha 2008 ⁹⁹⁵	Invalid country
Samraj 2013 ⁹⁹⁸	Narrative review
Santolaya 2008 ¹⁰⁰⁰	Invalid country
Sauer 2003 ¹⁰⁰³	Invalid intervention
Schreiber 2013 ¹⁰⁰⁹	Invalid outcomes
Schwarz 2000 ¹⁰¹¹	Invalid analysis

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Scott 2012 ¹⁰¹³	Invalid outcomes
Seigel 2012 ¹⁰¹⁴	Invalid outcomes
Shaw 1991 ¹⁰²⁶	Case-control study
Shine 1985 ¹⁰³⁰	Invalid analysis
Shorr 2010 ¹⁰³²	Invalid analysis
Sierra 2007 ¹⁰³⁵	Systematic review
Silveira 1999 ¹⁰³⁹	Invalid population
Simms 1992 ¹⁰⁴⁰	Invalid diagnostic tests
Sivula 2015 ¹⁰⁴⁵	Invalid diagnostic tests
Somech 2000 ¹⁰⁵⁶	Invalid outcomes
Sonawane 2014 ¹⁰⁵⁸	Invalid country
Spasova 2005 ¹⁰⁶³	Invalid outcomes
Steinbach 2007 ¹⁰⁶⁷	Invalid population
Struelens 1988 ¹⁰⁷⁰	Invalid outcomes
Su 2012B ¹⁰⁷⁵	Invalid country
Su 2014 ¹⁰⁷³	Invalid country
Sucilathangam 2012 ¹⁰⁸⁰	Invalid country
Suri 1991 ¹⁰⁸³	Invalid country
Tegtmeyer 1992 ¹⁰⁹²	Invalid outcomes
Toh 2003A ¹⁰⁹⁸	Invalid analysis
Tong 2015 ¹⁰⁹⁹	Invalid diagnostic tests
Tsalik 2012 ¹¹⁰⁷	Inconsistencies regarding units of measurement
Tschaikowsky 2011 ¹¹⁰⁹	Invalid outcomes
Tugrul 2002 ¹¹¹³	Invalid country
Turi 2013 ¹¹¹⁵	Invalid diagnostic tests
Ueda 2014 ¹¹¹⁶	Not relevant to review question
Ulla 2013 ¹¹¹⁸	Invalid diagnostic tests
Van den Bruel 2011 ¹¹²⁴	Invalid study design
Vassiliou 2015A ¹¹³¹	Invalid diagnostic tests
Venkataseshan 2007 ¹¹³⁵	Invalid diagnostic tests
Ventetuolo 2008 ¹¹³⁷	Narrative review
Venugopal 2014 ¹¹³⁸	Narrative review
Verbakel 2014 ¹¹⁴⁰	Study protocol
Viallon 2008 ¹¹⁴¹	Invalid diagnostic tests
Volante 2004 ¹¹⁴⁷	Narrative review
Wacharasint 2012 ¹¹⁵¹	Invalid analysis
Waliullah 2010 ¹¹⁵³	Invalid country
Waliullah 2009 ¹¹⁵⁵	Invalid country
West 2012 ¹¹⁶⁶	Invalid country
Wilkinson 2009 ¹¹⁷⁰	Invalid outcomes
Xie 2013 ¹¹⁸³	Invalid diagnostic tests
Yan 2001 ¹¹⁹⁰	Invalid outcomes
Yentis 1995 ¹¹⁹⁵	Invalid analysis
Yilmaz 2003 ¹¹⁹⁷	Invalid outcomes
Yin 2011 ¹¹⁹⁹	Invalid outcomes
Zant 2014 ¹²⁰⁶	Invalid outcomes Invalid population
Zarkesh 2015 ¹²⁰⁷	
Zarkesn 2015 ¹¹³ Zimmerman 2010 ¹²¹⁸	Invalid country Invalid outcomes
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L.4 Lactate

Table 38: Studies excluded from the clinical review

Study	Reason for exclusion
Aitofella 2012 ⁴¹	AUC data but no sensitivity or specificity data
Berger 2013 ¹⁷⁸	Hyperlactaemia was an outcome not a predictor
Bollaert 2003 ²⁰⁵	No diagnostic accuracy data; relativistic OR/RR data only
Breuling 2015 ²²⁹	No diagnostic accuracy data; relativistic OR/RR data only
Brodska 2013 ²³²	No diagnostic accuracy data; relativistic OR/RR data only
Casagandra 2015 ²⁶²	AUC data but no sensitivity or specificity data
Chen 2014F ³⁰²	Study conducted in non OECD country (China)
Cicarelli 2007 ³¹⁴	Study conducted in a developing country (Brazil)
Contenti 2015 ³²⁶	No protocol outcomes
Gao 2014 ⁴⁸³	Study conducted in a developing country (China)
Giannazzo 2006 ⁴⁹⁹	No diagnostic accuracy data; relativistic OR/RR data only
Giulieri 2015 ⁵⁰²	Target disease was community-acquired meningitis
Gwak 2015 ⁵³⁸	Target disease was community-acquired pneumonia
Hermans 2012 ⁵⁶⁵	AUC data but no sensitivity or specificity data
Hernandez 2012A ⁵⁶⁷	No protocol outcomes
Hisamuddin 2012 ⁵⁷⁴	Study conducted in a developing country (Malaysia)
Howell 2007A ⁵⁸⁹	No diagnostic accuracy data; relativistic OR/RR data only
Jansen 2011 ⁶¹³	Non-systematic review with different inclusion criteria (prognostic value of lactate, non-sepsis specific)
Jones 2010 ⁶²⁸	No relevant to protocol
Kang 2011 ⁶³⁶	Wrong population
Kim 2015B ⁶⁶⁴	Outcomes not relevant to this review
Kobayashi 2001 ⁶⁷⁷	No diagnostic accuracy data; relativistic OR/RR data only
Krishna 2009 ⁶⁸⁹	No protocol outcomes
Kung 2014 ⁶⁹³	No diagnostic accuracy data
Kung 2015 ⁶⁹⁴	AUC data but no sensitivity or specificity data
Lee 2008 ⁷¹⁷	No diagnostic accuracy data; relativistic OR/RR data only
Li 2013A ⁷²⁵	Li 2013A ⁷²⁵
Liu 2015 ⁷³⁷	Target condition was severe pneumonia, and country was non OECD (China)
Linder 2012 ⁷³³	No protocol outcomes
Lorente 2013 ⁷⁴³	No diagnostic accuracy data; relativistic OR/RR data only
Lorente 2014A ⁷⁴⁴	No diagnostic accuracy data; relativistic OR/RR data only
Lorente 2014B ⁷⁴⁵	No protocol outcomes
Lorente 2015A 745	Not protocol biomarker
Lorente 2015 ⁷⁴²	AUC data but no sensitivity or specificity data
Mallat 2014A ⁷⁶³	No diagnostic accuracy data; relativistic OR/RR data only
Manzon 2015 ⁷⁷⁰	AUC data but no sensitivity or specificity data
Mato 2010 ⁷⁸⁴	No protocol outcomes
Matsumura 2014 ⁷⁸⁶	ICU population but did not have sepsis
Mesquida 2015 ⁸⁰²	No diagnostic accuracy data; relativistic OR/RR data only

Study	Reason for exclusion
Miguelbayarri 2012 ⁸⁰⁶	No diagnostic accuracy data; relativistic OR/RR data only
Mikkelsen 2009 ⁸⁰⁷	No diagnostic accuracy data; relativistic OR/RR data only
Muller 2000 ⁸²⁸	Target condition was sepsis – not a worsening of existing sepsis
Musikatavorn 2015 835	No diagnostic accuracy data
Nanda 2009 ⁸³⁹	No protocol outcomes
Nguyen 2010A ⁸⁵⁴	No diagnostic accuracy data; relativistic OR/RR data only
Nguyen 2011 ⁸⁵³	Not relevant to the protocol
Ouillette 2014 ⁸⁸⁸	Case control study
Pandey 2014 ⁸⁹⁰	AUC data but no sensitivity or specificity data
Park 2014 ⁸⁹⁶	Study conducted in a developing country (South Korea)
Puskarich 2012A ⁹⁴²	Insufficient data for analysis
Ryoo 2015 ⁹⁹³	No diagnostic accuracy data; relativistic OR/RR data only
Shapiro 2010 ¹⁰²³	AUC data but no sensitivity or specificity data
Singer 2014 ¹⁰⁴¹	No diagnostic accuracy data
Singh2012A ¹⁰⁴²	Study did not evaluate lactate specifically
Song 2012 ¹⁰⁵⁹	No diagnostic accuracy data; relativistic OR/RR data only
Suarezsantamaria 2010 ¹⁰⁷⁷	AUC data but no sensitivity or specificity data
Tang 2015 ¹⁰⁹⁰	No diagnostic accuracy data; relativistic OR/RR data only
Varpula 2005 ¹¹³⁰	No diagnostic accuracy data; relativistic OR/RR data only
Whittaker 2015 ¹¹⁶⁸	No diagnostic accuracy data; relativistic OR/RR data only
Zanaty 2012 1205	Study conducted in a developing country (Egypt)
Zhang 2014E ¹²⁰⁹	Study conducted in a developing country (China)

L.5 Serum creatinine

Table 39: Studies excluded from the clinical review

Study	Reason for exclusion
Badin 2011 ¹¹⁶	Not protocol biomarker
Bagshaw 2013 ¹¹⁹	Not protocol biomarker
Bagshaw 2010 ¹²⁴	Not protocol biomarker
Bagshaw 2007 ¹²³	No protocol outcomes
Bagshaw 2007 ¹²⁸	Not protocol biomarker
Bagshaw 2006 ¹²²	Not protocol biomarker
Bagshaw 2006 ¹²⁷	Not protocol population
Basu 2011 ¹⁵⁵	No protocol outcomes
Carbonell 2004 ²⁵⁰	Not protocol biomarker
Cartinceba 2012 ²⁶⁰	SR with no protocol outcomes
Chawla 2005 ²⁸⁷	No outcomes of interest
De 2004 ³⁵⁹	Not protocol study type
Desouza 2014 ³⁶⁷	Study conducted in developing country
Dinardo 2013 ³⁸⁸	No protocol outcomes

Drey 2015 ⁴⁰⁰	No protocol outcomes
Elfarghali 2012 ⁴¹³	No protocol outcomes
Glassford 2013 ⁵⁰³	No protocol outcomes
Guo 2011 ⁵³³	Study conducted in developing country
Hamzic-Mehmedbasic 2015 549	Study conducted in non-OECD country
Hoste 2003 ⁵⁸⁶	No protocol outcomes
Iglesias 2003 ⁵⁹⁶	Not protocol population
Kiers 2010 ⁶⁶⁰	No protocol outcomes
Mariano 2008 ⁷⁷²	Not protocol biomarker
Martensson 2010 ⁷⁷⁷	Not protocol biomarker
Martensson 2012 ⁷⁷⁸	No protocol outcomes
Mazulsunko 2004 ⁷⁸⁸	No protocol outcomes
Nejat 2010 ⁸⁴⁵	No protocol outcomes
Nie 2013 ⁸⁵⁷	Not protocol biomarker
Plataki 2011 ⁹²⁴	No protocol outcomes
Poukkanen 2013 ⁹³¹	No protocol outcomes
Soni 2009 ¹⁰⁶⁰	Not protocol population
Su 2011 ¹⁰⁷⁴	Study conducted in developing country
Suh 2013 ¹⁰⁸¹	No protocol outcomes
Terzi 2014 ¹⁰⁹³	No protocol outcomes
Vanmassenhove2013 ¹¹²⁹	Not protocol biomarker
Walshe 2009 ¹¹⁵⁶	No protocol outcomes
Waring 2011 ¹¹⁵⁹	SR with no protocol outcomes
Wheeler 2008 ¹¹⁶⁷	No protocol outcomes
Wong 2015 ¹¹⁷⁸	Not protocol biomarker
Yamashita 2014 ¹¹⁸⁹	Not protocol population
Yegenaga 2004 ¹¹⁹⁴	No protocol outcomes
Zhang 2015 ¹²¹⁰	Not protocol study type
Zhou ¹²¹⁷	Study conducted in non OECD country

L.6 Disseminated intravascular coagulation (DIC)

Table 40: Studies excluded from the clinical review

Study	Reason for exclusion
Angstwurm 2006 ⁸³	Not protocol study design
Brenner 2012 ²²⁵	Not protocol study design
Cauchie 2006 ²⁶⁶	Not protocol population
Dempfle 2004 ³⁷⁷	Not protocol study design
Ersoy 2007 ⁴²⁵	Not protocol risk factor
Gamper 2001 ⁴⁷⁷	Not protocol population
Gando 1999 ⁴⁸⁰	Not protocol study design
Gando 2002 ⁴⁷⁹	Not protocol study design
Gando 2006 ⁴⁷⁸	Not protocol study design
Gando 2009 ⁴⁸¹	Not protocol study design

Study	Reason for exclusion
Gogos 2003 ⁵⁰⁵	Not protocol risk factor
Gomez 2007 ⁵⁰⁹	Not protocol risk factor
Guirgis 2014 ⁵³¹	SR not protocol risk factor
Ha 2015 ⁵³⁹	Not protocol study design
Harbarth 2002 ⁵⁵³	Not protocol study design
Hayakawa 2007 ⁵⁵⁷	Not protocol study design
Hoppensteadt 2014 581	Not protocol study design
lba 2015 ⁵⁹⁵	Not protocol study design
Ishimura 2014 ⁶⁰³	Not protocol study design
Jesmin 2013 ⁶²¹	Not protocol risk factor
Keneka 2012 ⁶⁴⁸	Not protocol study design
Kienast 2006 ⁶⁵⁹	Not protocol study design
Kim 2014 ⁶⁶²	Not protocol risk factor
Kinasewitz 2005 ⁶⁷¹	Not protocol study design
Kinasewitz 2004 ⁶⁷⁰	Not protocol study design
Kobayashi 2001 ⁶⁷⁷	Not protocol study design
Koyama 2014 ⁶⁸⁵	Not protocol risk factor
Kushimoto 2008 ⁶⁹⁵	Not protocol study design
Lavigne-Lissalde 2015 ⁷¹²	Conference abstract
Lin 2006 ⁷³²	Not protocol study design
Lin 2008 ⁷³¹	Not protocol study design
Lissaldelavigne 2008 ⁷³⁴	Not protocol study design
Madoiwa 2006 ⁷⁵⁶	Not protocol risk factor
Massion 2012 ⁷⁸²	Not protocol risk factor
Muller 2014 ⁸²⁹	Not protocol risk factor
Ogura 2014 ⁸⁷²	Not protocol study design
Oh 2010 ⁸⁷³	Not protocol study design
Okabayashi 2004 ⁸⁷⁴	Not protocol population
Ostrowski 2013 ⁸⁸⁶	Not protocol risk factor
Park 1999 ⁸⁹⁷	Not protocol study design
Park 2011 ⁸⁹³	Not protocol study design
Peigne 2013 ⁹¹²	Not protocol study design
Saracco 2011 ¹⁰⁰¹	Not protocol study design
Sawamura 2009 ¹⁰⁰⁵	Not protocol study design
Sawamura 2009 ¹⁰⁰⁴	Not protocol study design
Seki 2013 ¹⁰¹⁶	Not protocol study design
Takahashi 2015 ¹⁰⁸⁷	Not protocol study design
Voves 2006 ¹¹⁴⁹	Not protocol study design
Yamakawa 2013 ¹¹⁸⁷	Not protocol study design

L.7 Antimicrobial treatment

Table 41: Studies excluded from the clinical review

Reference	Reason for exclusion
Bagshaw 2009 ¹²¹	Not relevant outcomes
Band 2011 ¹³⁶	Comparison does not match protocol (patients who presented to the ED by ambulance versus patients who arrived by alternative means)
Barochia 2010 ¹⁴⁷	Setting does not match protocol (review on the use of bundles in patients with septic shock)
Beck 2014A ¹⁶²	Comparison does not match protocol (time to vasopressor initiation in patients with septic shock)
Behrendt 1999 ¹⁶⁵	Comparison does not match protocol (appropriate therapy within 48 hours versus after 48 hours)
Degoricija 2006 ³⁷³	No relevant outcomes, comparison does not match protocol
Erbay 2009 ⁴²³	Comparison does not match protocol (appropriate treatment within 24 hours versus after 24 hours)
Gabram 1993 ⁴⁷³	No relevant outcomes, study population does not match protocol (trauma patients)
Garcia-Saenz 2002 ⁴⁸⁶	Full text not available. Not in English language.
Garnacho-Montero 2003 ⁴⁸⁹	Comparison does not match protocol (adequate versus non-adequate empirical antimicrobial therapy; no time to antibiotics)
Garnacho-Montero 2006 ⁴⁸⁸	Comparison does not match protocol (appropriate treatment within 24 hours versus after 24 hours)
Gordon 2005 ⁵¹²	Comparison does not match protocol (not time to antibiotics)
Hanzelka 2013 ⁵⁵²	Setting does not match protocol (implementation of an EGDT protocol for cancer patients)
Hetem 2011 ⁵⁷⁰	Comparison does not match protocol (under 24 hours versus after 24 hours)
Hortmann 2014 ⁵⁸⁵	Comparison does not match protocol (time to antibiotics not analysed)
Houck 2004 ⁵⁸⁷	Study population does not match protocol (proportion of patients with sepsis not clearly mentioned)
Iscimen 2008 ⁶⁰¹	No relevant outcomes and does not match review protocol
Irwin 2015 ⁶⁰⁰	No relevant outcome
Jacob 2012 ⁶⁰⁶	Wrong population
Kang 2003 ⁶³⁴	Comparison does not match protocol (under 24 hours versus after 24 hours)
Khan 2015 ⁶⁵¹	No relevant intervention (over 24 hours)
Khatib 2006A ⁶⁵⁵	Comparison does not match protocol (not early versus delayed treatment)
Kim 2012C ⁶⁶⁷	Comparison does not match protocol (adequate versus inadequate treatment)
Ko 2015 ⁶⁷⁶	Setting does not match protocol (implementation of a door-to-antibiotics time)
Krediet 2003 ⁶⁸⁷	No relevant outcomes
Lin 2008 ⁷²⁹	Comparison does not match protocol (under 24 hours versus after 24 hours)
Lodise 2007 ⁷⁴¹	Comparison does not match protocol (appropriate treatment up to 52

Reference	Reason for exclusion
	hours)
Lodise 2003 ⁷⁴⁰	Comparison does not match protocol (under 44.75 hours versus after 44.75 hours)
MacArthur 2004 ⁷⁵²	Comparison does not match protocol (adequate versus inadequate treatment)
MacRedmond 2010 ⁷⁵⁵	Setting does not match protocol (implementation of a sepsis management protocol)
Meehan 1997 ⁷⁹⁵	Study population does not match protocol (proportion of patients with sepsis not clearly mentioned)
Natarajan 2014 ⁸⁴¹	No data reported
Nguyen 2006A ⁸⁵⁰	Study design does not match protocol (review with different protocol)
Nguyen 2007B ⁸⁵¹	Setting does not match protocol (implementation of a sepsis bundle)
Nguyen 2010 ⁸⁴⁹	Study design does not match protocol
Nickerson 2009 ⁸⁵⁶	Comparison does not match protocol (median delay is 3 days)
Onder 2008 ⁸⁷⁸	Not relevant outcomes
Parish 2013 ⁸⁹²	Setting does not match protocol (assessing a nurse-led protocol)
Park 2013 ⁸⁹⁵	Comparison does not match protocol (adequate antimicrobial therapy within 3 days)
Paul 2010 ⁹⁰⁵	Study population does not match protocol (12% sepsis)
Paul 2010A ⁹⁰⁷	Comparison does not match protocol (assesses appropriate antibiotics)
Pestana 2010 ⁹¹⁶	No relevant outcomes, study population does not match protocol
Rehmani 2014 ⁹⁶⁰	Setting does not match protocol (assessing an antibiotic protocol)
Rodriguez-Pardo 2015 ⁹⁸¹	No relevant outcomes, study population does not match protocol
Ronnestad 2005 ⁹⁸⁶	Study design does not match protocol (survey), not relevant (no info on antibiotics intervention)
Sainio 1995 ⁹⁹⁴	Not relevant review question
Schweizer 2010 ¹⁰¹²	Comparison does not match protocol (adequate versus inadequate treatment)
Shime 2010 ¹⁰²⁹	Intervention does not match protocol (antibiotics up to 48 hours)
Shorr 2011 ¹⁰³¹	Comparison does not match protocol (appropriate therapy versus inadequate; no time to antibiotics)
Siddiqui 2009 ¹⁰³³	Comparison does not match protocol (no comparison)
Siddiqui 2010 ¹⁰³⁴	Cochrane review does not include RCT evidence
Silber 2003 ¹⁰³⁶	Study population does not match protocol (proportion of patients with sepsis not clearly mentioned)
Sterling 2015 ¹⁰⁶⁸	Unclear methodology
Strang 1992 ¹⁰⁶⁹	Incorrect study design (survey data)
Studnek 2012 ¹⁰⁷²	Setting does not match protocol (EGDT paper)
Sweet 2010 ¹⁰⁸⁴	Setting does not match protocol (study assesses protocol and not timing of antibiotics)
Talmor 2008 ¹⁰⁸⁹	Setting does not match protocol (EGDT paper)
The ProCESS Investigators 2014 ⁹³⁷	Setting does not match protocol (EGDT paper)
Tumbarello 2007 ¹¹¹⁴	Comparison does not match protocol (examines inadequate antibiotics)
Uittenbogaard 2014 ¹¹¹⁷	No relevant outcomes and does not match review protocol

Reference	Reason for exclusion
Vanparidon 2015 ¹¹²⁶	No relevant analysis (effect size per minute)
Venkatesh 2013 ¹¹³⁶	No relevant outcomes
Waterer 2006 ¹¹⁶²	Study population does not match protocol (no sepsis)
Yahav 2013 ¹¹⁸⁵	Review with different inclusion criteria (pneumonia population)
Zahar 2011 ¹²⁰³	Comparison does not match protocol (appropriate treatment within 24 hours versus after 24 hours)

L.8 IV fluid administration

Table 42: Studies excluded from the clinical review

Study	Exclusion reason
Abulebda 2014 ¹⁶	Incorrect interventions
Andre 2010 ⁸¹	Incorrect interventions
Andre 2011 ⁸⁰	Incorrect interventions
Annane 2013 ⁸⁵	Incorrect interventions
Apibunyopas 2014 ⁸⁹	Paper not available
Arnold 2013 ⁹⁷	No relevant outcome
Bagshaw 2013 ¹²⁵	Not guideline condition
Bansal 2013 ¹³⁹	Invalid inclusion criteria
Bayer 2011 ¹⁵⁹	Incorrect interventions
Bayer 2012 ¹⁵⁸	Incorrect interventions
Boldt 1995 ²⁰²	No relevant outcome
Boldt 1996 ²⁰¹	Incorrect interventions
Boldt 1996 ²⁰³	Incorrect interventions
Boldt 1998 ²⁰⁴	Incorrect interventions
Boyd 2011 ²²²	Incorrect interventions
Brunkhorst 2008 ²³⁵	Incorrect interventions
Busund 1993 ²⁴¹	Incorrect interventions
Cardoso 2010 ²⁵²	Incorrect interventions
Carlsen 2011 ²⁵⁴	Incorrect interventions
Casserly 2011 ²⁶³	Incorrect interventions
Castellanos-ortega 2010 ²⁶⁴	Incorrect interventions
Chang 2014 ²⁸⁴	No relevant outcome
Chen 2014 ²⁹⁴	Incorrect interventions
Chong 2014 ³⁰⁸	Incorrect interventions
Chopra 2011 ³⁰⁹	Incorrect interventions
Chuesakoolvanich 2007 ³¹⁰	Not study design
Coen 2014 ³²²	Inappropriate comparison
Crowe 2010 ³³⁹	Inappropriate comparison
Cui 2012 ³⁴¹	Not English
De oliveira 2008 ³⁶⁶	Inappropriate comparison
Delaney 2011 ³⁷⁴	Incorrect interventions
Dubin 2010 ⁴⁰⁷	No relevant outcome
El solh 2008 ⁴¹²	Inappropriate comparison

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Ernest 1999 ⁴²⁴	No relevant outcome
Estrada 2013 ⁴³²	Commentary
Fang 2008 ⁴⁴¹	No relevant outcome
Femling 2014 ⁴⁴³	Incorrect interventions
Ferrer 2009 ⁴⁴⁹	Incorrect interventions
Finfer 2004 ⁴⁵⁴	Incorrect interventions
Ford 2012 ⁴⁶⁵	No relevant outcome
Fuller 2012 ⁴⁷⁰	No relevant outcome
Groeneveld 2011 ⁵²¹	Incorrect interventions
Guidet 2012 ⁵²⁸	Incorrect interventions
Gurnani 2010 ⁵³⁵	Incorrect interventions
Haase 2013 ⁵⁴¹	No relevant outcome
Haase 2013 ⁵⁴⁰	Incorrect interventions
Haase 2014 ⁵⁴²	Incorrect interventions
Holst 2013 ⁵⁸⁰	Study protocol
Jacob 2012 ⁶⁰⁶	Not study population
Jiang 2014 ⁶²³	Incorrect interventions
Jones 2007 ⁶²⁵	Inappropriate comparison
Karam 2011 ⁶³⁷	Incorrect interventions
Lee 2014 ⁷¹⁸	Incorrect interventions
Lefrant 2010 ⁷¹⁹	Incorrect interventions
Lin 2006 ⁷³⁰	Incorrect interventions
Liu 2013 ⁷³⁶	Incorrect interventions
Ma 2015 ⁷⁵¹	Systematic review
Maitland 2011 ⁷⁶⁰	Not guideline condition
Malbrain 2014 ⁷⁶¹	Not guideline condition
Miller 2013 ⁸⁰⁹	Incorrect interventions
Muller 2015 ⁸³⁰	Incorrect interventions
Murphy 2009 ⁸³³	Incorrect interventions
Nunes 2014 ⁸⁶²	No relevant outcome
Nurnberger 1999 ⁸⁶⁴	Incorrect interventions
O'neill 2012 ⁸⁶⁶	Incorrect interventions
Opiyo 2014 ⁸⁸¹	Incorrect interventions
Orbegozo cortes 2014 ⁸⁸²	Not guideline condition
Parsons 2011 ⁹⁰⁰	Incorrect interventions
Patel 2013 ⁹⁰¹	Incorrect interventions
Peake 2014 ⁹⁰⁸	Incorrect interventions
Perner 2012 ⁹¹⁵	Incorrect interventions
Perner 2012 ⁹¹⁴	Incorrect interventions
Purdy 1997 ⁹⁴¹	No relevant outcome
Raghunathan 2014 ⁹⁴⁸	Incorrect interventions
Raza 2015 ⁹⁵⁶	Not review population
Reiter 2013 ⁹⁶²	Incorrect interventions
Rewari 2014 ⁹⁶⁵	Abstract only
Rinaldi 2013 ⁹⁷¹	Incorrect interventions
Rivers 2001 ⁹⁷³	Incorrect interventions
Rochwerg 2014 ⁹⁷⁹	No relevant outcome
Rochwerg 2015 ⁹⁷⁸	Network meta-analysis with different study protocol
Rosland 2014 ⁹⁸⁸	Incorrect interventions

Serpa neto 2014 ¹⁰²⁰	No relevant outcome
Smith 2012 ¹⁰⁵¹	Incorrect interventions
Surat 2014 ¹⁰⁸²	Paper not available
Trof 2010 ¹¹⁰³	Inappropriate comparison
Upadhyay 2005 ¹¹²⁰	No relevant outcome
Vanparidon 2015 ¹¹²⁶	Invalid analysis
Veneman 2004 ¹¹³⁴	No relevant outcome
Wawrzeniak 2015 ¹¹⁶³	Inappropriate comparison
Wiedermann 2008 ¹¹⁶⁹	Incorrect interventions
Wittbrodt 2013 ¹¹⁷⁵	Incorrect interventions
Xu 2014 ¹¹⁸⁴	Incorrect interventions
Yang 2010 ¹¹⁹¹	Not English
Yealy 2014 ¹¹⁹³	Incorrect interventions
Zhang 2015 ¹²⁰⁸	Incorrect interventions
Zhong 2013 ¹²¹³	No relevant outcome

L.9 Escalation of care

Table 43: Studies excluded from the clinical review

Study	Exclusion reason
Alsolamy 2014 ⁶⁹	Invalid intervention
Austin 2014 ¹⁰⁷	Invalid population
Chamberlain 2015 ²⁷⁷	Invalid analysis
Esteban 2007 ⁴³¹	Invalid comparison
Evans 2014 ⁴³⁴	Invalid population
Femling 2014 ⁴⁴³	Invalid comparison
Fendler 2012 ⁴⁴⁴	Invalid intervention
Jaderling 2013 ⁶⁰⁸	Invalid comparison
Junhasavasdikul 2013 ⁶³²	Invalid population
Robert 2000 ⁹⁷⁵	Invalid outcome
Takeyama 2012 ¹⁰⁸⁸	Invalid intervention
Vinson 2014 ¹¹⁴⁶	Invalid intervention

L.10 Inotropic agents and vasopressors

Table 44: Studies excluded from this clinical review

Study	Exclusion reason
Acevedo 2009 ¹⁷	Abstract
Agrawal 2011 ³⁰	No relevant outcome
Agrawal 2012 ³¹	Invalid study design
Albanese 2004 ⁵⁷	No relevant outcome
Albanèse 2005 ⁵⁶	Incorrect interventions

Annatasit 2014 ⁷⁶ Retrospective analysis of VASST trial Anwar 2002 ⁸⁷ Not available Avni 2015 ¹⁶⁸ Systematic review Bahloul 2014 ¹³⁰ Inappropriate comparison Barton 1996 ¹⁵⁰ No relevant outcome Boulain 2009 ²³¹ Invalid study design Cardoso 2010 ²³² Incorrect interventions Cha 2004 ²⁷¹ Not fallish Daley 2013 ³⁴⁸ Invalid study design Dunser 2009 ⁴⁶⁸ No relevant outcome El Solla 2008 ⁴⁷² Incorrect interventions Cha 2004 ²⁷³ Invalid study design Dunser 2009 ⁴⁶⁸ No relevant outcome El Solla 2008 ⁴⁷² Incorrect interventions El Solla 2008 ⁴⁷² Invalid study population Gordon 2010 ²³³ Invalid study population Gordon 2010 ²³⁴ Not relevant outcome Hall 2004 ⁴⁶⁶ Invalid study design Klein 2006 ⁶⁷⁴ Not relevant setting Kumar 2008 ⁶⁷⁹ Inappropriate comparison Lampin 2012 ⁷²⁴ Inappropriate comparison Lampin 2012 ⁷²⁴ Inappropriate comparison Levy 2005 ⁷⁷² Inappropriate comparison Lupei 2009 ⁷⁸⁸ Inappropriate comparison Lupei 2009 ⁷⁸⁸ Inappropriate comparison Marka 2014 ⁷⁷³ Inappropriate comparison Marka 2014 ⁷⁷³ Inappropriate comparison Marka 2014 ⁷⁷³ Inappropriate comparison Marki 2000 ⁷⁸⁶ Inappropriate comparison Marki 2000 ⁷⁸⁷⁶ Inappropriate comparison Marki 2000 ⁷⁸⁷⁸ Inappropriate comparison Marki 2000 ⁷⁸⁸⁸ Inappropriate comparison Morelli 2000 ⁷⁸⁹⁸ Incorrect interventions Morelli 2000 ⁷⁸⁹⁸ Inappropriate comparison Morelli 2000 ⁷⁸⁹⁸ Inappropriate comparison Morelli 2000 ⁷⁸⁹⁸ Inappropriate comparison Morelli 2000 ⁷⁸⁹⁸ Inappropriate comparison Morelli 2000 ⁷⁸⁹⁸ Inappropriate comparison Mulliner 2004 ⁷⁹³¹ Cochrane review (outdated) Oba 2014 ⁶⁸⁹⁷ Systematic review Obritsch 2004 ⁷⁸⁹⁸ Inappropriate comparison Poleial 2012 ⁷⁸⁹⁸ Inappropriate comparison Poleial 2012 ⁷⁸⁹⁸ Inappropriate comparison No relevant outcome Povoa 2009 ⁷⁹⁹² Incorrect interventions Inappropriate comparison Poleiare comparison No relevant outcome Povoa 2009 ⁷⁹⁹³ Incorrect interventions Inappropriate comparison Inappropriate comparison No relevant outcome Inappropriate comparison Inappropriate comparison No relevant	Study	Exclusion reason
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Bahloul 2014 ¹³⁰ Inappropriate comparison Barton 1996 ¹⁵⁰ No relevant outcome Boulain 2009 ²²¹ Invalid study design Cardoso 2010 ²⁵² Incorrect interventions Cardoso 2010 ²⁵² Incorrect interventions Daley 2013 ¹⁴⁸ Invalid study design Dunser 2009 ⁴⁶⁸ No relevant outcome El solh 2008 ⁴¹² Incorrect interventions Elmenesy 2008 ⁴⁴⁷ Not available Gordon 2010 ⁵¹⁰ Invalid study population Gordon 2012 ⁵¹¹ No relevant outcome Hall 2004 ⁵⁴⁶ Invalid study design Klein 2006 ⁵⁷⁴ Not relevant setting Klein 2006 ⁵⁷⁴ Not relevant setting Kumar 2008 ⁶⁰⁰ Inappropriate comparison Lawpin 2012 ⁷⁰⁴ Inappropriate comparison Levy 1999 ⁷²² No relevant outcome Levy 2005 ⁷²⁸ Inappropriate comparison Lupel 2009 ⁴⁴⁷ Inappropriate comparison Mark 2014 ⁷⁷³ Inappropriate comparison Mark 2014 ⁷⁷⁴ Inappropriate comparison Mark 2014 ⁷⁷⁵ Inappropriate comparison Mark 2008 ⁷⁸⁰ Incorrect interventions Matok 2005 ⁷⁸⁵ Incorrect interventions Micek 2007 ⁸⁰⁶ Invalid study design Moon 2010 ⁸²⁰ Not guideline condition Morelli 2008 ⁸²³ Incorrect interventions Morelli 2008 ⁸²³ Incorrect interventions Morimatsu 2004 ⁸²⁵ Inappropriate comparison Mullner 2004 ⁸²¹ Cochrane review (outdated) Oba 2014 ⁸²⁷ Systematic review Obritsch 2004 ⁸²⁸ Inappropriate comparison Portisch 2004 ⁸²⁹ No relevant outcome Povoa 2009 ⁹³⁸ Inappropriate comparison Portisch 2004 ⁸²⁹ No relevant outcome Povoa 2009 ⁹³⁸ Inappropriate comparison Portisch 2004 ⁸²⁹ No relevant outcome Povoa 2009 ⁹³⁸ Inappropriate comparison Portisch 2004 ⁸²⁹ No relevant outcome Povoa 2009 ⁹³⁸ Inappropriate comparison Russell 2013 ⁹³⁸ Inappropriate comparison Russell 2013 ⁹³⁸ Inappropriate comparison Russell 2009 ⁸³⁰ Inappropriate comparison Russell 2009 ⁸³⁰ Inappropriate comparison	Avni 2015 ¹⁰⁸	Systematic review
Barton 1996 ¹⁵⁰ No relevant outcome Boulain 2009 ²²¹ Invalid study design Cardoso 2010 ²⁵² Incorrect interventions Cha 2004 ²⁷⁷ Not English Daley 2013 ³⁵⁴⁸ Invalid study design Dunser 2009 ⁴⁶⁸ No relevant outcome El solh 2008 ⁴¹² Incorrect interventions El solh 2008 ⁴¹⁷ Not available Gordon 2010 ⁵²¹⁰ Invalid study oppulation Gordon 2010 ⁵²¹⁰ Invalid study design Melai 2004 ⁵⁴⁶ Invalid study design Klein 2006 ⁵⁷⁴ Not relevant outcome Hall 2004 ⁵⁴⁶ Invalid study design Klein 2006 ⁵⁷⁴ Not relevant setting Klein 2006 ⁵⁷⁴ Not relevant setting Lampin 2012 ⁷⁰⁴ Inappropriate comparison Lampin 2012 ⁷⁰⁴ Inappropriate comparison Levy 1999 ⁷²² No relevant outcome Levy 2005 ⁷²³ Inappropriate comparison Lupei 2009 ⁷⁴⁸ Inappropriate comparison Lupei 2009 ⁷⁴⁸ Inappropriate comparison Mark 2014 ⁷⁷³ Inappropriate comparison Mark 2014 ⁷⁷³ Inappropriate comparison Mark 2010 ⁷⁷⁹ Incorrect interventions Micke 2007 ⁸⁰⁵ Invalid study design Moon 2010 ²²⁰ Not guideline condition Morelli 2007 ⁸²² Abstract Morelli 2008 ⁸²³ Incorrect interventions Morimatus 2004 ⁸²⁵ Inappropriate comparison Mullner 2004 ⁸²⁵ Inappropriate comparison Mullner 2004 ⁸²⁶ Inappropriate comparison Mullner 2004 ⁸²⁷ Systematic review Obritsch 2004 ⁸⁰⁶ Inappropriate comparison Mullner 2004 ⁸²⁸ Inappropriate comparison Poblisch 2008 ⁸⁰⁸ Inappropriate comparison Mullner 2004 ⁸²⁹ Inappropriate comparison Poblisch 2004 ⁸⁰⁹ Inappropriate comparison Mullner 2004 ⁸²⁹ Inappropriate comparison Poblisch 2004 ⁸⁰⁹ Inappropriate comparison Poblisch 2009 ⁹⁰⁹ Inappropriate comparison Poblisch 2009 ⁹⁰⁹ Inappropriate comparison Russell 2013 ⁹⁰⁹ Inappropriate comparison Russell 2013 ⁹⁰⁹ Inappropriate comparison Russell 2013 ⁹⁰⁹ Inappropriate comparison	Backer 2012 ¹¹⁵	Systematic review
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Levy 1999 ⁷²² No relevant outcome Levy 2005 ⁷²³ Inappropriate comparison Lin 2006 ⁷³⁰ Inappropriate comparison Lupei 2009 ⁷⁴⁸ Inappropriate comparison Mark 2014 ⁷⁷³ Inappropriate comparison Mark 2010 ⁷⁷⁹ Incorrect interventions Matok 2005 ⁷⁸⁵ Incorrect interventions Micek 2007 ⁸⁰⁵ Invalid study design Moon 2010 ⁸²⁰ Not guideline condition Morelli 2007 ⁸²² Abstract Morelli 2008 ⁸²³ Incorrect interventions Morimatsu 2004 ⁸²⁵ Inappropriate comparison Mullner 2004 ⁸³¹ Cochrane review (outdated) Oba 2014 ⁸⁶⁷ Systematic review Obritsch 2004 ⁸⁶⁹ Inappropriate comparison O'neill 2012 ⁸⁶⁶ Inappropriate comparison Patel 2002 ⁹⁰² No relevant outcome Povoa 2009 ⁹³⁵ Inappropriate comparison Prys-picard 2013 ⁹³⁸ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Inappropriate comparison Russell 2003 ⁹⁹² Inappropriate comparison Russell 2003 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Kumar 2008 ⁶⁹⁰	Inappropriate comparison
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Micek 2007 ⁸⁰⁵ Invalid study design Moon 2010 ⁸²⁰ Not guideline condition Morelli 2007 ⁸²² Abstract Morelli 2008 ⁸²³ Incorrect interventions Morimatsu 2004 ⁸²⁵ Inappropriate comparison Mullner 2004 ⁸³¹ Cochrane review (outdated) Oba 2014 ⁸⁶⁷ Systematic review Obritsch 2004 ⁸⁶⁹ Inappropriate comparison O'neill 2012 ⁸⁶⁶ Inappropriate comparison Patel 2002 ⁹⁰² No relevant outcome Povoa 2009 ⁹³⁵ Inappropriate comparison Prys-picard 2013 ⁹³⁸ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Incorrect interventions Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Martin 2000 ⁷⁷⁹	Incorrect interventions
Moon 2010 ⁸²⁰ Not guideline condition Morelli 2007 ⁸²² Abstract Morelli 2008 ⁸²³ Incorrect interventions Morimatsu 2004 ⁸²⁵ Inappropriate comparison Mullner 2004 ⁸³¹ Cochrane review (outdated) Oba 2014 ⁸⁶⁷ Systematic review Obritsch 2004 ⁸⁶⁹ Inappropriate comparison O'neill 2012 ⁸⁶⁶ Inappropriate comparison Patel 2002 ⁹⁰² No relevant outcome Povoa 2009 ⁹³⁵ Inappropriate comparison Prys-picard 2013 ⁹³⁸ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Incorrect interventions Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Matok 2005 ⁷⁸⁵	Incorrect interventions
Morelli 2007 ⁸²² Abstract Morelli 2008 ⁸²³ Incorrect interventions Morimatsu 2004 ⁸²⁵ Inappropriate comparison Mullner 2004 ⁸³¹ Cochrane review (outdated) Oba 2014 ⁸⁶⁷ Systematic review Obritsch 2004 ⁸⁶⁹ Inappropriate comparison O'neill 2012 ⁸⁶⁶ Inappropriate comparison Patel 2002 ⁹⁰² No relevant outcome Povoa 2009 ⁹³⁵ Inappropriate comparison Prys-picard 2013 ⁹³⁸ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Incorrect interventions Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Micek 2007 ⁸⁰⁵	Invalid study design
Morelli 2008 ⁸²³ Incorrect interventions Morimatsu 2004 ⁸²⁵ Inappropriate comparison Mullner 2004 ⁸³¹ Cochrane review (outdated) Oba 2014 ⁸⁶⁷ Systematic review Obritsch 2004 ⁸⁶⁹ Inappropriate comparison O'neill 2012 ⁸⁶⁶ Inappropriate comparison Patel 2002 ⁹⁰² No relevant outcome Povoa 2009 ⁹³⁵ Inappropriate comparison Prys-picard 2013 ⁹³⁸ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Incorrect interventions Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Moon 2010 ⁸²⁰	Not guideline condition
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Oba 2014 ⁸⁶⁷ Systematic review Obritsch 2004 ⁸⁶⁹ Inappropriate comparison O'neill 2012 ⁸⁶⁶ Inappropriate comparison Patel 2002 ⁹⁰² No relevant outcome Povoa 2009 ⁹³⁵ Inappropriate comparison Prys-picard 2013 ⁹³⁸ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Incorrect interventions Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Morimatsu 2004 ⁸²⁵	Inappropriate comparison
Obritsch 2004 ⁸⁶⁹ Inappropriate comparison O'neill 2012 ⁸⁶⁶ Inappropriate comparison Patel 2002 ⁹⁰² No relevant outcome Povoa 2009 ⁹³⁵ Inappropriate comparison Prys-picard 2013 ⁹³⁸ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Incorrect interventions Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Mullner 2004 ⁸³¹	Cochrane review (outdated)
O'neill 2012 ⁸⁶⁶ Inappropriate comparison Patel 2002 ⁹⁰² No relevant outcome Povoa 2009 ⁹³⁵ Inappropriate comparison Prys-picard 2013 ⁹³⁸ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Incorrect interventions Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Oba 2014 ⁸⁶⁷	Systematic review
Patel 2002 ⁹⁰² No relevant outcome Povoa 2009 ⁹³⁵ Inappropriate comparison Prys-picard 2013 ⁹³⁸ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Incorrect interventions Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Obritsch 2004 ⁸⁶⁹	Inappropriate comparison
Povoa 2009 ⁹³⁵ Inappropriate comparison Prys-picard 2013 ⁹³⁸ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Incorrect interventions Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	O'neill 2012 ⁸⁶⁶	Inappropriate comparison
Prys-picard 2013 ⁹³⁸ Inappropriate comparison Rodriguez-nunez 2006 ⁹⁸⁰ Incorrect interventions Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Patel 2002 ⁹⁰²	No relevant outcome
Rodriguez-nunez 2006 ⁹⁸⁰ Incorrect interventions Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Povoa 2009 ⁹³⁵	Inappropriate comparison
Russell 2009 ⁹⁹² Inappropriate comparison Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Prys-picard 2013 ⁹³⁸	Inappropriate comparison
Russell 2013 ⁹⁹¹ Not review population Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Rodriguez-nunez 2006 ⁹⁸⁰	Incorrect interventions
Sakr 2006 ⁹⁹⁶ Inappropriate comparison	Russell 2009 ⁹⁹²	Inappropriate comparison
	Russell 2013 ⁹⁹¹	Not review population
Serpa neto 2012 ¹⁰¹⁹ Incorrect interventions	Sakr 2006 ⁹⁹⁶	Inappropriate comparison
	Serpa neto 2012 ¹⁰¹⁹	Incorrect interventions

Study	Exclusion reason
Shapiro 2006 ¹⁰²⁴	Incorrect interventions
Soong 2011 ¹⁰⁶¹	Inappropriate comparison
Tourneux 2008 ¹¹⁰¹	Inappropriate comparison
Tsapenko 2013 ¹¹⁰⁸	Inappropriate comparison
Tsuneyoshi 2001 ¹¹¹¹	Invalid study design
Vasu 2012 ¹¹³²	Systematic review
Waechter 2014 ¹¹⁵²	Inappropriate comparison
Wilkman 2013 ¹¹⁷¹	Inappropriate comparison
Yildizdas 2008 ¹¹⁹⁶	Incorrect interventions
Zhang 2015 ¹²⁰⁸	Inappropriate comparison
Zhao 2012 ¹²¹²	Not English
Zhou 2013 ¹²¹⁴	Not English
Zhou 2014 ¹²¹⁵	Systematic review
Zhou 2015 ¹²¹⁶	Systematic review

L.11 Supplemental oxygen

Table 45: Studies excluded from the clinical review

Reference	Reason for exclusion
Alia 1999 ⁶³	Inappropriate comparison (therapy with normal targeted value of oxygen delivery versus targeted oxygen delivery index)
Balk 2004 ¹³⁴	Inappropriate study design (narrative paper)
Bellomo 2008 ¹⁶⁸	Inappropriate study design (commentary)
Crone 1994 ³³⁸	Inappropriate study design (letter to the editor)
Duarte 2005 ⁴⁰⁶	Inappropriate study design (narrative review)
Erstad 1994 ⁴²⁶	Review with different protocol
Esen 1992 ⁴³⁰	Inappropriate intervention (artificial ventilation)
Ferrer 200 ⁴⁴⁸	Inappropriate population (acute hypoxemic respiratory failure) and incorrect comparison (non invasive ventilation versus oxygen using high concentration sources)
Freebairn 1997 ⁴⁶⁶	Inappropriate interventions (vecuronium or saline closed-loop infusion)
Ince 1999 ⁵⁹⁸	Review with different protocol
Matuschak 1997 ⁷⁸⁷	Review with different protocol
Rampal 2010 ⁹⁴⁹	Review with different protocol
Russell 1995 ⁹⁹⁰	Inappropriate study design (narrative review)
Textoris 2011 1094	Inappropriate intervention (local hospital protocol)
Vincent 1995 ¹¹⁴²	Inappropriate study design (narrative review)

L.12 Use of bicarbonate

Table 46: Studies excluded from the clinical review

Reference	Reason for exclusion
Kim 2013 ⁶⁶³	Population not relevant to review question (61% of patients had sepsis as cause of lactic acidosis; 67 % of the population received bicarbonate therapy)
Velissaris 2015 ¹¹³³	Literature review

L.13 Early goal-directed therapy (EGDT)

None.

L.14 Monitoring

Table 47: Studies excluded from the clinical review (use of scoring systems)

Reference	Reason for exclusion
Abbott 2015 ¹¹	Intervention does not match protocol (not for monitoring: comparison of NEWS and PARS)
	Population does not match protocol (not sepsis specific: all patients admitted to the acute assessment unit)
Adshead 2009 ²⁸	Incorrect study design (narrative article)
Akre 2010 ⁴⁵	Intervention does not match protocol (not for monitoring: external validation or PEWS and calculation of median time from critical PEWS to rapid response team)
	Population does not match protocol (not sepsis specific: hospitalised paediatric patients, respiratory, infectious disease, cancer, cardiac, digestive)
Alam 2014A ⁵³	Intervention does not match protocol (not for monitoring: systematic review on ability of EWS to identify patients at risk of deterioration)
	Population does not match protocol (not sepsis specific: ED and ward patients)
Alam 2015 ⁵⁴	Intervention does not match protocol (not for monitoring: validation of NEWS to predict adverse outcome)
	Population does not match protocol (not sepsis specific: all ED patients with an emergency severity index of 2 and 3 not triaged to the resuscitation room)
Albert 2011 ⁵⁸	Intervention does not match protocol (not for monitoring: development of a modified EWS)
	Population does not match protocol (not sepsis specific: cardiac, respiratory, neurological, sepsis (1.3%))
Alrawi 2013 ⁶⁸	Intervention does not match protocol (not for monitoring: to assess ability of MEWS to predict mortality)
	Population does not match protocol (not sepsis specific: acutely ill nursing home residents)
Anon 2014B ⁹	Incorrect study design (narrative article)
Armagan 2008 ⁹⁴	Intervention does not match protocol (not for monitoring: validation of MEWS)

Reference	Reason for exclusion
	Population does not match protocol (not sepsis specific: all ED patients)
Ausania 2015 ¹⁰⁶	Intervention does not match protocol (not for monitoring: multivariable analysis of risk factors associated with morbidity and mortality) Population does not match protocol (not sepsis specific: post-operative patients)
Bayer 2015 ¹⁶⁰	Intervention does not match protocol (not for monitoring: development of a new scoring system, not externally validated) Population does not match protocol (not sepsis specific: all patients admitted to ED)
Bradman 2008 ²²⁴	Intervention does not match protocol (not for monitoring: to see if PEWS could determine at triage children who needed admission or who could be discharged at home) Population does not match protocol (not sepsis specific: all children attending the paediatric emergency department)
Badriyah 2014 ¹¹⁷	Intervention does not match protocol (not for monitoring: validation of NEWS) Population does not match protocol (not sepsis specific: all patients admitted to the medical assessment unit)
Breslin 2014 ²²⁶	Intervention does not match protocol (to establish that higher PEWS at time of ED disposition decision is associated with need for higher levels of care at ED disposition, not for monitoring) Population does not match protocol (not sepsis specific: ED patients)
Burch 2008 ²³⁹	Intervention does not match protocol (to evaluate the utility of MEWS as a triage tool, not for monitoring) Population does not match protocol (not sepsis specific: medical patients presenting to the ED)
Chaiyakulsil 2015 ²⁷⁴	Population does not meet protocol (not sepsis)
Cei 2009 ²⁶⁸	Intervention does not match protocol (to identify patients at risk of deterioration, not for monitoring) Population does not match protocol (not sepsis specific: all patients admitted to a medical ward)
Churpek 2013 ³¹²	Intervention does not match protocol (to discuss risk scores for use on the general inpatient wards to predict mortality, ICU transfer and cardiac arrest, not for monitoring) Population does not match protocol (not sepsis specific: patients on general wards)
Cildir 2013 ³¹⁵	Intervention does not match protocol (not for monitoring: to evaluate the ability of MEDS, MEWS and the Charlson comorbidity index (CCI) to predict prognosis in patients who are diagnosed in sepsis)
Corfield 2014 ³²⁸	Intervention does not match protocol (not for monitoring: to determine, in patients with sepsis, whether a single NEWS on ED arrival is a predictor of mortality, or ICU admission)
Correia 2014 ³³⁰	Intervention does not match protocol (not for monitoring: EWS score at -72h, -24h and -12h in patients transferred from the ward to the ER) Population does not match protocol (not sepsis specific: cardiovascular, respiratory, neurological, renal or other clinical reasons)
Dawes 2014 ³⁵⁵	Intervention does not match protocol (not for monitoring: ability of the Worthing PSS score, calculated using VitalPAC, to predict mortality.) Population does not match protocol (not sepsis specific: all patients admitted to the Acute Medical Unit)

Reference	Reason for exclusion
De Meester 2013A ³⁶⁵	Intervention does not match protocol (monitoring for serious adverse events after ICU discharge)
	Population does not match protocol (not sepsis specific: surgical and medical ICU patients)
Ennis 2014 ⁴²²	Intervention does not match protocol (not for monitoring: evaluate the effectiveness of PEWS to early detect clinical deterioration)
	Population does not match protocol (not sepsis specific: acutely ill children in hospital)
Fairclough 2009 ⁴³⁸	Intervention does not match protocol (not for monitoring: use of MEWS to predict mortality in acute medical admission unit)
	Population does not match protocol (not sepsis specific: only 12% of patients had sepsis)
Finlay 2014 ⁴⁵⁵	Intervention does not match protocol (not for monitoring: MEWS to predict mortality)
	Population does not match protocol (not sepsis specific: general medical-surgical patients)
Friedman 2015 ⁴⁶⁷	Incorrect study design (narrative review)
Fuijkschot 2015 ⁴⁶⁹	Intervention does not match protocol (not for monitoring: PEWS to identify patients for PICU admission)
	Population does not match protocol (not sepsis specific: all patients receiving emergency medical interventions at the paediatric wards; all patients admitted to paediatric oncology ward)
Goldhill 2004 ⁵⁰⁷	Intervention does not match protocol (not for monitoring: physiological variables to predict mortality)
	Population does not match protocol (not sepsis specific: all patients in non-obstetric bed area)
Goldhill 2005 ⁵⁰⁸	Intervention does not match protocol (not for monitoring: physiological variables and Patient-At-Risk score to predict mortality)
	Population does not match protocol (not sepsis specific: outreach service database)
Griffiths 2012 ⁵²⁰	Incorrect study design (survey)
Haines 2006 ⁵⁴⁴	Intervention does not match protocol (not for monitoring: to develop and evaluate a clinical and physiologically based for identification of acutely ill children in ward areas) Population does not match protocol (not sepsis specific)
Hammond 2013 ⁵⁴⁸	Intervention does not match protocol (not sepsis specific)
Hammond 2013	change in combination or individual vital signs frequency before and after MEWS implementation)
	Population does not match protocol (not sepsis specific: ICU patients with three diagnostic groups: cardiovascular, respiratory and gastrointestinal)
Henry 2015 ⁵⁶¹	Outcome does not match protocol (diagnostic accuracy data)
Ho 2013 ⁵⁷⁷	Intervention does not match protocol (not for monitoring: MEWS to
	predict mortality and ICU admission) Population does not match protocol (not sepsis specific: critically ill patients who require continuous ECG monitoring)
Holme 2013 ⁵⁷⁹	Intervention does not match protocol (not for monitoring: To design and
Holling 2013	validate an objective clinical scoring system to identify unwell neonates) Population does not match protocol (not sepsis specific: all neonates >35 weeks' gestation admitted to the NICU)
Jarvis 2015A ⁶¹⁶	Intervention does not match protocol (not for monitoring: use of NEWS
Jai VIS ZULJA	intervention does not match protocol (not for monitoring, use of NEWS

Reference	Reason for exclusion
Reference	to calculate risk of death and adverse outcome)
	Population does not match protocol (not sepsis specific: all patients admitted to hospital)
Jo 2013 ⁶²⁴	Intervention does not match protocol (not for monitoring: to examine whether the predictive value of EWS could be improved by including rapid lactate levels, and to compare the modified EWS with the preexisting risk scoring systems)
Kaul 2014 ⁶⁴²	Incorrect study design (survey)
Kellett 2012 ⁶⁴⁵	Intervention does not match protocol (not for monitoring: validation of an abbreviated Vitalpac Early Warning Score) Population does not match protocol (not sepsis specific: includes surgical patients, medical, cardiac, oncology, renal and stroke patients)
Kyriacos 2011 ⁶⁹⁸	Intervention does not match protocol (not for monitoring: review the validity of EWS/MEWS)
	Population does not match protocol (not sepsis specific: population not specified)
Lam 2006 ⁷⁰³	Intervention does not match protocol (not for monitoring: applicability of MEWS for the emergency department observation ward to predict serious outcome)
	Population does not match protocol (not sepsis specific: patients with cardiac or gastrointestinal symptoms, or dizziness)
Liu 2015 ⁷³⁵	Intervention does not match protocol (not for monitoring: validation of National EWS in emergency intensive care unit)
	Population does not match protocol (not sepsis specific: neurological, cardiovascular, respiratory, gastrointestinal and other diseases)
Ludikhuize 2012 ⁷⁴⁷	Intervention does not match protocol (not for monitoring: effectiveness of MEWS to predict cardiopulmonary arrest, ICU admission, death, emergency surgery) Population does not match protocol (not sepsis specific: patients on
	general wards)
Ludikhuize 2014 ⁷⁴⁶	Intervention does not match protocol (not for monitoring: implementation of a RRs protocol) Population does not match protocol (not sepsis specific: hospitalised
	patients)
Mandell 2015 ⁷⁶⁴	Population does not match protocol (not sepsis population)
Moseson 2014 ⁸²⁶	Intervention does not match protocol (not for monitoring: comparison of APACHE II, APACHE III, SAPS II, MEWS, REMS, PEDS to predict mortality)
	Population does not match protocol (not sepsis specific: critically ill patients admitted to the ICU with one of the following diagnosis category: respiratory, cardiovascular, infectious disease, neurology, gastrointestinal, other)
Oldroyd 2011 ⁸⁷⁵	Incorrect study design (narrative article)
Parshuram 2011 ⁸⁹⁸	Intervention does not match protocol (not for monitoring: before-and- after study to evaluate the effect of implementation of the Bedside PEWS)
	Population does not match protocol (not sepsis specific: all paediatric patients)
Parshuram 2011A ⁸⁹⁹	Repeated measures analysis showed that the Bedside PEWS increased over the 24 hours before urgent ICU admission or code blue event from a baseline mean score of 5.3, 20-24h before clinical deterioration, to 8.4 in the last 4 h

Reference	Reason for exclusion
	Population does not match protocol (not sepsis specific: all paediatric patients, case-control study)
Patterson 2011 ⁹⁰³	Incorrect study design (survey)
Pearson 2011 ⁹⁰⁹	Incorrect study design (narrative article)
Prytherch 2010 ⁹³⁹	Intervention does not match protocol (not for monitoring: to develop a validated, paper-based, aggregate weighted track and trigger system (AWTTS) for the detection of patient deterioration)
	Population does not match protocol (not sepsis specific: database of acute medical admissions)
Reini 2012 ⁹⁶¹	Intervention does not match protocol (not for monitoring: to assess ability of MEWS, SAPS III, and SOFSA to predict ICU mortality) Population does not match protocol (not sepsis specific: only 13% of participants had sepsis. ICU setting)
Seiger 2013 ¹⁰¹⁵	Intervention does not match protocol (not for monitoring: review to evaluate ability of PEWS to predict hospitalisation and ICU admission) Population does not match protocol (not sepsis specific: all children presenting to the ED with the following problems: trauma, gastrointestinal, FWS, dyspnea, wounds, neurologic, urinary tract problems, local infection/abscess, rash, ear, nose, throat, other)
Silcock 2015 ¹⁰³⁷	Intervention does not match protocol (not for monitoring: validation of NEWS in identifying patients at risk of death or deterioration in the pre-hospital setting) Population does not match protocol (not sepsis specific: unselected pre-hospital patients)
Skaletzky 2012 ¹⁰⁴⁶	Intervention does not match protocol (not for monitoring: validation of a modified PEWS) Population does not match protocol (not sepsis specific: all patients admitted to medical-surgical wards. Case-control study)
Smith 2013 ¹⁰⁴⁹	Intervention does not match protocol (not for monitoring: evaluate the ability of NEWS to detect mortality and ICU admission) Population does not match protocol (not sepsis specific: patients admitted to the medical assessment unit)
Smith 2014 ¹⁰⁵⁰	Intervention does not match protocol (not for monitoring: review on the validity of EWS) Population does not match protocol (not sepsis specific: medical and surgical inpatients)
So 2015 ¹⁰⁵³	Intervention does not match protocol (to detect weather ED monitoring by MEWS is better than nurse clinical judgement in changing the patient's ED management plan) Population does not match protocol (not sepsis specific: all patients being held in the ED observation area because of access block to the following specialty wards: medical, general surgery, neurosurgery and clinical oncology)
Solevag 2013 ¹⁰⁵⁵	Intervention does not match protocol (not for monitoring: to assess the correlation of modified PEWS results with other indicators of severe illness) Population does not match protocol (not sepsis specific: injury, congenital cardiovascular disease, acquired cardiovascular disease, neurological disease, renal disease including urinary tract infection, gastrointestinal disease, respiratory, other infection, miscellaneous including dehydration and diabetes ketoacidosis)

Reference	Reason for exclusion
Subbe 2001 ¹⁰⁷⁸	Intervention does not match protocol (not for monitoring: validation of a modified EWS)
	Population does not match protocol (not sepsis specific: all medical emergency admissions admitted to the medical admissions unit)
Tafelski 2015 ¹⁰⁸⁶	Intervention does not match protocol (not for monitoring: application of three different PIRO systems)
Tucker 2009 ¹¹¹²	Intervention does not match protocol (not for monitoring: validation of PEWS)
	Population does not match protocol (not sepsis specific: most common diagnosis were asthma exacerbation, bronchiolitis and pneumonia)
Van Rooijen 2013 ¹¹²⁷	Intervention does not match protocol (not for monitoring: evaluation of the threshold value for the EWS on general wards) Population does not match protocol (not sepsis specific: all patients on medical and surgical wards)
Vorwerk 2009 ¹¹⁴⁸	Intervention does not match protocol (not for monitoring: to determine the efficacy of the abbreviated MEDS score (without neutrophil bands), and MEWS in predicting mortality in adult ED patients with sepsis)
Yoo 2015 ¹²⁰⁰	Intervention does not match protocol (not for monitoring: to determine whether use of a combination of MEWS and lactate enhances prediction of ICU transfer and mortality in hospitalized patients with severe sepsis/septic shock)

L.15 Patient education, information and support

Table 48: Studies excluded from the clinical review

Reference	Reason for exclusion
Flynn 2012 ⁴⁶²	SR includes studies in wrong population
Higgins 2008 ⁵⁷¹	Wrong population
Jeon 2012 ⁶¹⁹	Wrong intervention
Obermann 2007 ⁸⁷¹	Wrong intervention
Plowright 2013 ⁹²⁵	Wrong study type
Yamamoto 1997 ¹¹⁸⁸	Wrong intervention

L.16 Education and training

Table 49: Studies excluded from the clinical review

Reference	Reason for exclusion
Adler 2007 ²⁶	Not relevant to review question
Allen 2011 ⁶⁶	Not relevant to review question
Anon 2008 ⁴	Not relevant to review question
Anon 2005A ²	Not relevant to review question
Anon 2007 ³	Comment
Anon 2008F ⁵	Not relevant to review question
Anon 2010 ⁶	Not relevant to review question.

Reference	Reason for exclusion
Anon 2010A ⁷	Comment
Anon 2013D ⁸	Comment
Arabi 2014 ⁹¹	Expert opinion
Assuncao 2010 ¹⁰³	No detail about how training was carried out
Assuncao 2014 ¹⁰⁴	No detail about how training was carried out
Austin 2014 ¹⁰⁷	Not relevant to review question
Bach 1996 ¹¹⁴	Not relevant to review question.
Berger 2010 ¹⁷⁷	Not education/training.
Bond 2013 ²¹¹	No detail about how training was carried out
Bridgewater 2014	Critical care nursing education/degree
Bruce 2011 ⁷⁹³	Protocol. Not on education/training
Buckley 2010 ²³⁷	Implementation of a protocol, not any details of training
Burney 2012 ²⁴⁰	Not relevant to review question
Baez 2013 ¹¹⁸	not relevant to review question/not enough details in paper
Barbieri 2013 ¹⁴⁴	Quality improvement initiatives, do not explain specific training or education
Benczo 2004 ¹⁷⁰	Not related to sepsis
Benson 2014 ¹⁷⁴	Early recognition, not training
Berg 2013 ¹⁸⁰	No details of how implemented/training
Capp 2011 ⁴⁶⁴	No details of training provided
Carlbom 2007 ²⁵³	Survey on barriers which may inform a training intervention but no training intervention
Casserly 2011 ²⁶³	Implementation of a Sepsis Intervention Programme, but no details on training
Chamberlain 2006 ²⁷⁶	Short summary
Carter 2007 ²⁵⁹	Outcomes not adequately measured
Castro2008 ²⁶⁵	Comparison of 2 intervention protocols, but no details on training
Chen 2013 ¹²¹¹	Impact of an education programme on patient outcomes. Details of training and education programme not included
Coba 2011 ³²⁰	Outcomes not adequately measured
Croft 2014 ³³⁷	Not relevant to review question.
Cruz 2012 ³⁴⁰	Not relevant to review question.
Daniels 2010	No details of training provided
Daniels 2011 ³⁵¹	States staff underwent training on sepsis 6 but no details of training provided
De Groot 2012 ³⁶⁰	No details of training provided.
Demmel 2010 ³⁷⁶	Not relevant to review question.
Desmond 2013 ³⁸⁰	Not relevant population.
Deutsch 2014 ³⁸²	Conference abstract
Devita 2007 ³⁸⁴	GDG ref. Comment on review
Fadale 2014 ⁴³⁵	Not relevant to review question. Training about vasopressor titration.
Fitzpatrick 2014 ⁴⁵⁷	Not relevant to review question. Wrong study design.
Fuchs 2015 ⁴⁶⁸	Conference abstract
Funk 2009 ⁴⁷¹	Review proposes and discusses barriers and RRS but does not present actual results of effectiveness of these.

ReferenceReason for exclusionGannon 2011482Not relevant to review question.Gerber 2010493Not relevant to review question.Gerdtz2013495GDG ref. Not relevant to review questionGirardis 2009501Not relevant to review question. Development and implement protocol. No details given on the training and education.	
Gerber 2010 ⁴⁹³ Not relevant to review question. Gerdtz2013 ⁴⁹⁵ GDG ref. Not relevant to review question Girardis 2009 ⁵⁰¹ Not relevant to review question. Development and implement	
Gerdtz2013 ⁴⁹⁵ GDG ref. Not relevant to review question Girardis 2009 ⁵⁰¹ Not relevant to review question. Development and implement	
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protocol. No details given on the training and education.	itation of a
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Granier 1998 ⁵¹⁵ Not relevant to review question.	
Greenspoon 1994 Implementation of a protocol.	
Guerra 2013 ⁵²⁵ No detail about how training was carried out	
Gultepe 2014 ⁵³² Not relevant to review question	
Harrigan 2006 ⁵⁵⁵ GDG ref. Not relevant to review question.	
Herasevich 2011 ⁵⁶³ Not relevant to review question.	
Hitti 2012 ⁵⁷⁵ No details of training provided.	
Huggan 2011 ⁵⁹¹ summary	
Hurtado 2006 ⁵⁹³ summary of bundles in surviving sepsis campaign	
Jeon 2012 ⁶¹⁹ GDG ref. Not relevant to review question. Implementation of No details given on the training and education.	f a protocol.
Jones 1998 ⁶²⁹ Comment on sepsis and SIRS definitions	
Jones 2014 ⁶²⁶ Not relevant to review question.	
Kang 2012 ⁶³⁵ Not relevant to review question.	
Kellie 2014 ⁶⁴⁶ Not relevant to review question.	
Kim 2001 ⁶⁶⁶ Prevention of infection for HCP	
Kim 1999 ⁶⁶¹ Not relevant to review question.	
Kleinpell 2014 ⁶⁷⁵ comment on SSC and bundles, not original research	
Kollef 2010 ⁶⁸³ GDG ref. Not relevant to review question. Implementation of No details given on the training and education.	f a protocol.
Larosa 2012 ⁷⁰⁸ Not relevant to review question. No details given on the train education.	ning and
Launay 2011 No details of training provided.	
Levy 2010 ⁷²⁴ No detail about what was how training/education carried out.	
Levy 2014 No detail about what was how training/education carried out.	
Lobo 2005 ⁷³⁹ GDG ref. Prevention of catheter-related infections, not about awareness of identification/ management of sepsis	raising
Lobo 2010 ⁷³⁸ Prevention of catheter-related infections, not about raising avidentification/ management of sepsis	vareness of
Mackintosh 2012 ⁷⁵⁴ GDG ref. Not relevant to review question. Not about education	n/training
Mahavanakul 2012 ⁷⁵⁸ Not relevant to review question.	
McGaughey 2010 ⁷⁹⁰ GDG ref. Wrong study design (protocol). Not relevant to review Not about education/training	w question.
Mann-Salinas 2014 ⁷⁶⁶ Description of sepsis in theory	
Marshall 2009 ⁷⁷⁵ Conference abstract	
Mckinley 2011 ⁷⁹³ Implemented protocol but no details of how implemented/tra	nining
McNally 2009 ⁷⁹⁴ Not relevant to review question.	
Meyer 2013 ⁸⁰⁴ No training implementation/analysis	
Mok 2014 ⁸¹⁶ Not relevant to review question	
Monette 2007 ⁸¹⁸ Not relevant to review question.	

Reference	Reason for exclusion	
Moore 2009 ⁸²¹	Sensitivity and specificity of sepsis screening protocol	
Nassau 2003 ⁸⁴⁰	Summary/comment, not original research	
Nelson 2011 ⁸⁴⁶	Not relevant to review question.	
Nguyen 2014 ⁸⁵⁵	Not relevant to review question.	
Nguyen 2009 ⁸⁵²	Not relevant to review question.	
Noritomi 2014 ⁸⁶¹	Protocol implementation. No detail about what was included/how training/education carried out	
Orji 2007 ⁸⁸⁴	Not relevant to review question	
Ottestad 2007 ⁸⁸⁷	Scores performance in identifying sepsis but not implementing any training	
Patocka 2014	No details of training provided.	
Phua 2012 ⁹¹⁸	Not relevant to review question.	
Phua 2013 ⁹¹⁹	Not relevant to review question.	
Plambech 2012 ⁹²³	Protocol implementation. No detail about what was included in the training.	
Potter 2011 ⁹³⁰	Editorial article	
Prasas 2010	Not relevant to review question.	
Puntis 1991 ⁹⁴⁰	Prevention of catheter-related infections, not about raising awareness of identification/management of sepsis	
Reuben 2006 ⁹⁶⁴	Not relevant to review question.	
Rincon 2011 ⁹⁷²	No details of how implemented/training	
Robson 2008 ⁹⁷⁷	Not relevant to review question.	
Robson 2007 ⁹⁷⁶	Not relevant to review question.	
Salluh 2008 ⁹⁹⁷	Not relevant to review question.	
Santana 2008 ⁹⁹⁹	Prevention of catheter-related infections, not about raising awareness of identification/management of sepsis	
Sarani 2008 ¹⁰⁰²	Not relevant to review question	
Sawyer 2011 ¹⁰⁰⁶	Not training/education.	
Scheer 2015 ¹⁰⁰⁷	Conference abstract	
Schramm 2011 ¹⁰⁰⁸	Implementation of a protocol, not any details of training	
Semelsberger 2009 ¹⁰¹⁷	Prevention of catheter-related infections, not about raising awareness of identification/management of sepsis	
Seoane 2013 ¹⁰¹⁸	Implementation of a protocol, not any details of training	
Shearer 2012 ¹⁰²⁷	Not relevant to review question.	
Sherertz 2000 ¹⁰²⁸	GDG ref. Prevention of infection, not about raising awareness of identification/ management of sepsis	
Smith 2012 ¹⁰⁵¹	Implementation of a protocol, not any details of training	
Tromp 2009 ¹¹⁰⁴	No details of training.	
Tromp 2010 ¹¹⁰⁵	Implementation of a protocol, not any details of training	
Tromp 2011	Implementation of a protocol, not any details of training	
Tafelski 2010 ¹⁰⁸⁵	Not relevant to review question.	
van Zanten 2014	No detail about what was how training/education carried out.	
van Dijck 2009 ¹¹²⁸	Not relevant to review question.	
Wallgren 2014 ¹¹⁵⁴	Implementation of two sepsis screening tools, not any details of training	
Warren 2003 ¹¹⁶⁰	Specific sepsis prevention programme	

Reference	Reason for exclusion	
Weaver 2003 ¹¹⁶⁴	States what can be done but does not show results of it being done	
Weinert 2008 ¹¹⁶⁵	General ICU not Sepsis	
Wolbrink 2014 ¹¹⁷⁷	describes platform but no results of effect in practice	
Winters 2013 ¹¹⁷⁴	GDG ref. Comment. No study undertaken	
Winterbottom 2011 ¹¹⁷³	Implementation of bundle of care for managing patients with severe sepsis/septic shock. No details of education programme	
Yilmaz 2007 ¹¹⁹⁸	Prevention of catheter-related infections, not about raising awareness of identification/management of sepsis	
Yurkova 2011	No details of training provided.	
Zaffar 2009	Not relevant to review question.	
Zuhlke 2013 ¹²¹⁹	Public education, not health professionals.	

Appendix M: Excluded health economic studies

M.1 Scoring systems

None.

M.2 Signs and symptoms

None.

M.3 Blood tests

None.

M.4 Lactate

None.

M.5 Serum creatinine

None.

M.6 Disseminated intravascular coagulation (DIC)

None.

M.7 Antimicrobial treatment

None.

M.8 IV fluid administration

Table 50: Studies excluded from the economic review

Reference	Reason for exclusion
GUIDET 2007 ⁵²⁷	This study was selectively excluded due to a combination of applicability and methodological limitations.
	Health outcomes were not expressed as QALYs. Time horizon may not be sufficient to capture all benefits and costs if benefits persist beyond 5 years. The associated RCT (SAFE Study) is just 1 of 7 included studies in the clinical review, which also has limitations because the treatment effect used in the cost effectiveness paper is a post hoc analysis and the treatment effect in the severe sepsis group was not found to be significant.

M.9 Escalation of care

None.

M.10 Inotropic agents and vasopressors

None.

M.11 Supplemental oxygen

None.

M.12 Use of bicarbonate

None.

M.13 Early goal-directed therapy

None.

M.14 Monitoring

None.

M.15 Patient education, information and support

None.

M.16 Education and training

None.

Appendix N: Research recommendations

N.1 Epidemiological study on presentation and management of sepsis in England

Research question:

What is the incidence, presentation and management of sepsis in the United Kingdom?

Why this is important:

The lack of robust UK based epidemiological studies on the incidence and outcomes from sepsis have been clear throughout the guideline development process. A large epidemiological study to collect information about where sepsis is being treated, patient interventions and patient outcomes would provide population based statistics on epidemiology of sepsis which are necessary to support evaluation of interventions, planning of services and service redesign. The mortality and morbidity and service complexity associated with severe infection and sepsis, and the need to use broad spectrum antimicrobials to treat sepsis, justifies the cost required to set up such a study.

Criteria for selecting high	i-priority research recommendations.
PICO question	The questions that a registry could help answer are: What is the epidemiology of life threatening sepsis in the UK?
	How and where is life threatening sepsis treated? What important safety monitors need to be in place to capture unintended consequences?
	Would co-ordinated service evaluation linked to a Sepsis Registry lead to better patient care?
Importance to patients or the population	The interventions recognised in this guideline as a standard of care for sepsis require timely, coordinated, and robust healthcare services. Process and patient outcome improvement can only occur if based on standardised data systems that inform us of epidemiological, clinical and outcome trends. There is a lack of evidence to support any particular service improvement methodology in sepsis but coordinated efforts to provide longitudinal data on process and outcome would help with this.
Relevance to NICE guidance	Provide baseline data on impact of sepsis in UK population and help inform future guidance on effective service improvement methodologies
Relevance to the NHS	Will provide assurance of guideline implementation which (along with mechanisms such as CQUIN) will drive service improvement. Will provide measures of local and population based epidemiology to inform service design and resourcing.
National priorities	National Sepsis CQUIN
	NHS Ombudsman Report into Sepsis Sign up to Safety NHSE priorities
Current evidence base	There is a lack of current national sepsis statistics with poor coding of episodes of sepsis and limited knowledge of UK sepsis epidemiology
Equality	None Relevant
Study design	Service evaluation and audit Epidemiological primary research
Feasibility	Information governance and Caldicott issues will need to be addressed.

	Centralised registry will need to be funded in line with other similar databases.
Other comments	A variety of known local service evaluation and audit methods could be adapted for national use.
Importance	 High: the research is essential to inform future updates of key recommendations in the guideline.

N.2 A complex service evaluation of implementation of NICE Sepsis guideline

Research question:

What effect will the NICE Sepsis guideline have on patient care processes and outcome in the UK over the next 5 years?

Why this is important:

Implementation of the NICE Sepsis guideline will be a challenge to the NHS. A robust evaluation of how NHS service providers adhere to the recommended care processes and the effect of implementation needs to be carried out.

A complex evaluation is required to understand the effect of guidelines on services and on patient outcomes. Evaluation should include assessment of costs and cost effectiveness, the use of a universal audit tool for sepsis patient care that includes evaluation of pre-hospital and secondary care and monitoring of broad spectrum antibiotic use, development of multi-resistant organisms and incidence of antibiotic related infection such as C. Difficile.

PICO question	What effect will the NICE Sepsis guideline have on patient care processes and outcome in the UK over the next 5 years?
Importance to patients or the population	The interventions recognised in this guideline as a standard of care for sepsis require timely, coordinated, and robust healthcare services. This is a complex intervention that needs assessment as such to allow changes to care to be monitored and evaluated to ensure improvement in care for people with sepsis.
Relevance to NICE guidance	Inform NICE of clinical effectiveness of guideline implementation and inform guideline updates.
Relevance to the NHS	Will provide information on guideline implementation which (along with mechanisms such as CQUIN) will drive service improvement.
National priorities	National Sepsis CQUIN NHS Ombudsman Report into Sepsis Sign up to Safety NHSE priorities
Current evidence base	Not applicable
Equality	None Relevant
Study design	Complex evaluation using the principles of process evaluation
Feasibility	Information governance and Caldicott issues will need to be addressed. The evaluation is feasible.
Other comments	
Importance	 High: the research is essential to inform NICE and local commissioners in gaps or difficulties in implementation of the guideline

N.3 Use of biomarkers to diagnose and initiate treatment

Research question:

What is the clinical and cost effectiveness of procalcitonin (PCT) point-of-care tests at initial triage compared for diagnosis of serious infection and the initiation of appropriate antibiotic therapy?

Why this is important:

There is an urgent clinical need for accurate biomarkers of serious bacterial infection (SBI) which provide early diagnosis of SBI, and prompt clinical interventions to improve outcomes. The current tests used in the NHS (white cell count and C-reactive protein) are non-specific and not sensitive enough. Biomarker-guided initiation and termination of antibiotic therapy might be an effective strategy to reduce unnecessary antibiotic use and help prevent further multidrug resistance. The recent NICE Diagnostic Guidance (DG18) on Procalcitonin for diagnosing and monitoring sepsis has shown there is not enough evidence in this area.

	F - 7
PICO question	Population: Adults and children with suspected sepsis at triage in the UK Index test: PCT Comparison: CRP Outcomes: time to diagnosis of sepsis, antibiotic exposure (initiation of appropriate antibiotic therapy), duration of hospital stay, duration of ICU stay, adverse clinical outcomes (for example mortality, antibiotic-related adverse events)
Importance to patients or the population	The rapid and accurate determination of the presence or absence of systematic infection is important for patients' clinical outcomes and also to reduce unnecessary exposure to antibiotics.
Relevance to NICE guidance	Further research on PCT would provide a stronger evidence base in order for NICE to issue clear guidance for diagnosis of children, young people and adults with suspected sepsis
Relevance to the NHS	Antimicrobial stewardship is important for the NHS and accurate identification of the need for antibiotics would allow more targeted use of antibiotics. Better stratification of disease severity will reduce morbidity and mortality, and reduce NHS costs.
National priorities	National Sepsis CQUIN NHS Ombudsman Report into Sepsis Sign up to Safety NHSE priorities UK Five Year Antimicrobial Resistance Strategy 2013 to 2018
Current evidence base	The current evidence for PCT is is limited. The current evidence for CRP is considered in Chapter 8 of the full guideline.
Equality	There are no equality issues.
Study design	PCT and CRP would be evaluated by standard methods including specificities, sensitivities, receiver operator curves (ROCs) or area under the curves (AUC) for diagnosis of sepsis. Assessment of initiation of appropriate antibiotic therapy would be evaluated by hazard ratios, odds ratios and/or relative risk for duration of hospital stay, duration of ICU stay, and adverse clinical outcomes.
Feasibility	The study is feasible as currently CRP is routinely tested in people with suspected sepsis.
Other comments	The study may attract commercial funders in the diagnostics arena including companies developing novel PCT assays.

Importance

• High: the research is essential to inform future updates of key recommendations in the guideline.

N.4 Validation of clinical early warning scores in pre-hospital and emergency care settings

Research question:

Can early warning scores for example NEWS (national early warning scores for adults) and PEWS (paediatric early warning score) be used to improve the detection of sepsis and facilitate prompt and appropriate clinical response in pre-hospital settings and in emergency departments?

Why this is important:

Delay in detecting and treating sepsis increases mortality. Early detection and appropriate management will reduce morbidity and mortality and will reduce NHS costs by reducing critical care admissions, inappropriate antimicrobial use and length of hospital stay. No high quality data exist on the validation or use of early warning scores in pre-hospital settings or in the emergency department settings. The use of scores might improve communication between pre-hospital settings and hospital settings and allow recognition of people who need more urgent assessment.

	• •
PICO question	Population: non-hospital based patients (both those totally managed in primary care and those who are transferred to secondary care), and patients managed in the emergency room with suspected sepsis in the UK. Intervention: (1) NEWS and (2) PEWS scores to direct care Comparison: No use of score to direct care Outcomes: referral rates, adverse clinical outcomes (for example mortality)
Importance to patients or the population	Timely diagnosis of sepsis and detection of worsening symptoms will improve patient outcomes.
Relevance to NICE guidance	Research would provide evidence to enable NICE to make recommendations on the use of NEWS and PEWS in the pre-hospital setting, emergency room or secondary care setting.
Relevance to the NHS	Prompt and early recognition of people with sepsis is critical to reducing morbidity and mortality and reducing NHS costs.
National priorities	NICE CG 50 Acutely ill patient in hospital: research recommendation re the sensitivity and specificity of track and trigger systems in various clinical settings NCEPOD Think Sepsis: recommends a standardised approach to vital signs monitoring in primary care, such as NEWS to help in the prioritisation of emergency care Ombudsman report 'Time to Kill': recommends the development of clinical tools highly predictive of sepsis to be used in primary care
Current evidence base	The development of the NICE guideline on sepsis found no evidence for use of validated tools in the pre-hospital or emergency room settings, and limited evidence in the emergency room and secondary care setting (chapter 6 of the guideline)
Equality	There are no equality issues.
Study design	Cluster randomised trial, or, if not feasible due to widespread NHS implementation following NCEPOD recommendation, observational score validation to establish: whether scores taken in primary/community care or the emergency room can
	differentiate patients requiring immediate escalation of care from those who can

	be managed less aggressively
	whether scores taken solely in the community can add to GPs or other health professional add to their assessments and clinical experience
	whether scores help communication between primary and secondary care and ambulances
	Whether scores in emergency room stings reduce the volume of empirical antimicrobial prescription, reduce critical care admissions, reduce length of stay or mortality
Feasibility	Baseline physiological measurements are already routinely taken in primary care but it is not usual practice to measure all the parameters and calculate a NEWS or PEWS score. It would require education and training of clinicians.
	In emergency room is feasible as baseline physiological measurements are routinely taken.
Other comments	
Importance	 High: The research is essential to inform future updates of key recommendations in the guideline.

N.5 Derivation of clinical decision rules in suspected sepsis

Research question:

Is it possible to derive and validate a set of clinical decision rules or a predictive tool to rule out sepsis which can be applied to patients presenting to hospital with suspected sepsis.

Why this is important:

In primary care and emergency departments people with suspected sepsis are often seen by relatively inexperienced doctors. Many of these people will be in low and medium risk groups but evidence is lacking as to who can be sent home safely and who needs intravenous or oral antibiotics. The consequences of getting the decision making wrong can be catastrophic and therefore many patients are potentially over-investigated and admitted inappropriately. Current guidance is dependent on use of individual variables informed by low quality evidence.

Criteria for selecting high	i-priority research recommendations.
PICO question	Population: Adults and children presenting to hospital with suspected sepsis in UK. Intervention: Derivation of history and physiological variables as well the application of diagnostic testing to be applied to patients fulfilling the inclusion criteria. Comparison: Normal practice/ guidelines. Outcome: diagnosis of sepsis, length of hospital stay, adverse clinical events (for example mortality)
Importance to patients or the population	Errors are still made with clinical decisions making in patients with suspected sepsis. Delays in initiating treatments can unfortunately lead to life-threatening consequences. Evidence based clinical decision rules would support safer decision making and improve patient safety
Relevance to NICE guidance	Would help to influence future guidelines in the moderate to low risk group.
Relevance to the NHS	Safer patient care. Cost reductions to allow early discharge of appropriate patients.
National priorities	Sepsis is high on the national agenda. Mortality rates are high and life- threatening treatments are occasionally omitted or delayed due to poor clinical decision making.
Current evidence base	The development of the NICE guideline on sepsis suggested that the current available evidence in this area is of poor quality and not fit for purpose.
Equality	None Relevant
Study design	Prognostic observational cohort study to identify risk factors for developing sepsis, and then validation of derived prediction tool in separate cohorts.
Feasibility	The research is feasible as comparable research has been achieved for other presentations, for example chest pains, DVTsGI Bleeds, headache, and head injuries
Other comments	The difficulty of diagnosing sepsis is the lack of an acceptable, recognised gold standard from which to work. Gold standard for a study may need to be developed by a Delphi method.
Importance	 High: the research is essential to inform future updates of key recommendations in the guideline.

Appendix O: NICE technical team

Name	Role
Sharon Summers-Ma	Guideline Lead
Martin Allaby	Clinical Advisor
Judith Thornton	Technical Lead (until November 2015)
Bhash Naidoo	Technical Lead (HE)
Caroline Keir	Guideline Commissioning Manager
Helen Dickinson	Guideline Coordinator
Gareth Haman	Editor
Rachel O'Mahony	Technical Lead (December 2015-present)
Laura Sadler	PIP Lead
Andrew Gyton	Project Manager

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