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## Glucocorticosteroids for sepsis: systematic review with meta-analysis and trial sequential analysis

Received: 23 December 2014  
Accepted: 27 May 2015  
Published online: 23 June 2015  
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**Take-home message:** There is no evidence for a beneficial effect on mortality of glucocorticosteroids in patients with sepsis.

**Electronic supplementary material**  
The online version of this article (doi:10.1007/s00134-015-3899-6) contains supplementary material, which is available to authorized users.

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**Abstract** *Introduction:* Glucocorticosteroids (steroids) are widely used for sepsis patients. However, the potential benefits and harms of both high and low dose steroids remain unclear. A systematic review of randomised clinical trials with meta-analysis and trial sequential analysis (TSA) might shed light on this clinically important question.

*Methods:* A systematic review was conducted according to a published protocol and The Cochrane Handbook methodology including meta-analyses, TSA of randomised clinical trials, and external validity estimation (GRADE). Randomised clinical trials evaluating steroids were included for sepsis patients (systemic inflammatory response syndrome, sepsis, severe sepsis or septic shock) aged >18 years. Cochrane Central Register of Controlled Trials (CENTRAL), PubMed/Medline, Embase, Web of Science and Cinahl were searched until 18 February 2015. No language restrictions were applied. Primary outcomes were mortality at longest follow-up and serious adverse events.

*Results:* A total of 35 trials randomising 4682 patients were assessed and reviewed in full text. All trials but two had high risk of bias. No statistically significant effect was found for any dose of steroids versus placebo or

no intervention on mortality at maximal follow-up [relative risk (RR) 0.89; TSA adjusted confidence interval (CI) 0.74–1.08]. Two trials with low risk of bias also showed no statistically significant difference (random-effects model RR 0.38, 95 % CI 0.06–2.42). Similar results were obtained in subgroups of trials stratified according to high (>500 mg) or low (≤500 mg) dose hydrocortisone (or equivalent) (RR 0.87; TSA-adjusted CI 0.38–1.99; and RR 0.90; TSA-adjusted CI 0.49–1.67, respectively). There were also no statistically significant effects on serious adverse events other than mortality (RR 1.02; TSA-adjusted CI 0.7–1.48). The effects did not vary according to the degree of sepsis. TSA showed that many more randomised patients are needed before definitive conclusions may be drawn. *Conclusion:* Evidence to support or negate the use of steroids in any dose in sepsis patients is lacking. The results of ongoing and future well-designed, large randomised clinical trials are needed.

**Keywords** Steroids · Sepsis · Meta-analysis · Trial sequential analysis

## Introduction

In the intensive care unit (ICU), up to 25 % of admissions are related to sepsis and an additional 12 % of patients develop sepsis during their stay [1]. Sepsis and, if deteriorating, septic shock have a high morbidity and mortality. Depending on the definition of sepsis, mortality varies from 27 to 54 % [1]. To decrease this high rate of morbidity and mortality, several interventions have been suggested. Bundled interventions, like those of the Surviving Sepsis Campaign [2], aim to improve outcome of patients by, among others, early antibiotics, glucocorticosteroids (steroids) and supportive care.

Based on supposed pathophysiological mechanisms, two rationales for steroids in sepsis have been put forward. The first rationale is that high dose steroids may suppress the excess in inflammatory response in sepsis. In the 1970s and the early 1980s high dose steroids (30 mg/kg methylprednisolone or equivalent dose) were used in sepsis [3]. In the late 1980s the use declined on the basis of negative results of randomised clinical trials [3]; however, in some centres, high dose steroids are still used in clinical practice today. The second rationale, introduced in the 1990s, is that low dose steroids may recover a relative adrenal insufficiency [4]. Many trials have been conducted, but the pathophysiological basis of the second rationale is still questioned [5]. Possibly both high and low dose steroids can have beneficial or harmful effects in sepsis.

Despite the lack of evidence for the underlying mechanism, the use of low doses of 200–300 mg hydrocortisone is recommended in patients with septic shock not responding to fluid and vasopressor therapy [2]. This recommendation is likely based on results of one systematic review, which found a statistically significant 16 % relative risk reduction (RRR) of mortality (relative risk 0.84; unadjusted 95 % confidence interval 0.72–0.97) in favour of prolonged low dose steroids [6]. The beneficial effect found might be a subgroup effect in more severely ill patients, a spurious finding due to a type I random error as a consequence of repetitive testing as the information size required for showing a 16 % RRR was far from being reached, or an overestimation of a treatment effect due to bias and suboptimal trial methodology [7–12].

A sound methodology in a systematic review is as important as in any type of study to avoid critical errors in the analyses and conclusions [13]. We therefore decided to conduct a new systematic review evaluating the effects of steroids for sepsis in patients with systemic inflammatory response syndrome (SIRS), sepsis, severe sepsis or septic shock as previous meta-analyses fall short on several aspects of rigorous methodology.

## Objective

The objective was to perform a systematic review according to a published protocol following guidelines from PRISMA [14] and *The Cochrane Handbook for Systematic Reviews of Interventions* [15]. We also planned to execute meta-analyses and trial sequential analyses (TSA) of randomised clinical trials that compared the benefits and harms of high and/or low dose steroids for patients with SIRS, sepsis, severe sepsis or septic shock. Our primary outcome was mortality at longest follow-up and serious adverse events.

Available evidence was to be evaluated in the perspective of the three dimensions of possible risks of errors: systematic errors (bias), design errors (also leading to systematic errors due to outcomes, comparators, etc.) and random errors ('the play of chance') [16].

## Methods

The systematic review was conducted following the recommendations of *The Cochrane Handbook for Systematic Reviews of Interventions* [15] and reported according to the PRISMA statement ([www.prisma-statement.org](http://www.prisma-statement.org)). The protocol was published on PROSPERO (<http://www.crd.york.ac.uk/PROSPERO>, ID: CRD42013005617).

In addition to an overall analysis including all doses, two separate analyses were conducted of high and low dose steroids. The cut-off value of high and low dose steroids was chosen arbitrarily. High doses were defined as daily doses of more than 500 mg hydrocortisone; low doses were defined as a daily intake of equal or less than 500 mg. Other steroids were recalculated into the equivalent hydrocortisone dose [17]. When doses were expressed in milligram per kilogram body weight, doses were calculated assuming a body weight of 75 kg.

## Eligibility criteria

Randomised clinical trials that included adult patients (age >18 years) with SIRS, sepsis, severe sepsis or septic shock, or any combinations thereof (Table S1). Trials with patients with SIRS were excluded when SIRS criteria were not explicitly described in their methods section. Trials were also excluded when evaluating steroids for the prevention of the occurrence of SIRS. No limitations were made regarding underlying cause of illness. There were no restrictions on duration of treatment (days) or whether administration occurred continuously or intermittently. All types of steroids were included.

Trials were included independently of the type of control intervention: placebo, no intervention, or any other control intervention. Co-interventions were allowed provided that similar administration occurred in the intervention groups. Trials were included irrespective of chosen outcomes. No limitations were made on the basis of language or publication status. The following study types were excluded: quasi-randomised studies, observational studies, cross-over studies, and studies comparing different doses or different types of steroids in both trial intervention groups.

### Search strategy

We searched the Cochrane Central Register of Controlled Trials (CENTRAL) in The Cochrane Library, PubMed/Medline, Embase, Web of Science and Cinahl. We also hand-searched the reference lists of included trials and systematic reviews for further trials. Ongoing trials were sought through trial registries ([www.clinicaltrials.gov](http://www.clinicaltrials.gov), [www.controlled-trials.com](http://www.controlled-trials.com), [www.centerwatch.com](http://www.centerwatch.com)). No time restrictions were applied. The electronic literature search strategies are listed in Table S2.

### Study selection and data extraction

Two authors independently reviewed all identified titles and abstracts and excluded clearly irrelevant hits. The remaining hits were evaluated in full text. Disagreements were resolved through discussion and the hits excluded on the basis of full text were all listed.

Characteristics of patients and trials and data for analyses were extracted by two authors independently from the included reports. A summary of the recorded patient data is presented in Table 1. A thorough recording of patient data is listed in Table S5.

We contacted corresponding authors for unreported data.

### Outcomes

Primary outcomes were mortality at longest follow-up and serious adverse events. Serious adverse events were a composite outcome, summarizing all serious events excluding mortality, necessitating an intervention, operation or prolonged hospital stay.

Secondary outcomes were persistent dependence on haemodialysis and duration of mechanical ventilation. Additionally, time-specific analyses of mortality at 30 and 90 days were conducted as secondary outcomes according to availability of data. Data of trials reporting 28-day mortality were included in 30-day mortality analyses. All outcomes were classified according to the patients'

perspective according to GRADE Working Group (Table S3) [18]. Although used in previous trials and meta-analyses, shock reversal was not considered, as it is a surrogate outcome, which is not important according to the patients' perspective [19].

### Risk of bias assessment

We assessed the risk of bias according to *The Cochrane Handbook for Systematic Reviews of Interventions* [15], including all eight domains: random sequence generation, allocation concealment, blinding of participants and personnel, blinding of outcome assessment, incomplete outcome data, selective reporting, and other bias (academic or funding bias). If one or more of the domains were judged as having high or unclear risk of bias, the trial was classified as having a high risk of bias. Only two trials had low risk of bias (in all domains) with a pooled information size of 591 patients (nearly all data came from one trial [20]). Therefore, we formulated a group of trials with lower risk of bias. These trials had at least low risk of bias in sequence generation, allocation concealment, blinding of participants and personnel, and blinding of outcome assessment. This does not, however, exclude risk of bias from other domains.

### Statistical analysis

Review Manager 5.1.6 was used for statistical analyses. We used the TSA program version 0.9 beta ([www.ctu.dk/tsa](http://www.ctu.dk/tsa); [21]) to control random errors and assess imprecision. For each included trial we calculated the relative risk (RR) with 95 % confidence intervals (CI) for dichotomous outcomes. We report risk differences if statistically significant different from relative risk. We calculated the numbers needed to treat or numbers needed to harm with 95 % CI based on a statistically significant RR.

Heterogeneity among trials was explored by the Chi-squared test with significance set at  $P$  value of 0.10, and quantified with inconsistency factor ( $I^2$ ) statistics. We reported the results from the random-effects model anticipating abundant clinical heterogeneity (in populations, interventions and settings). We reported the results from a fixed-effect model if one or two trials dominated the available evidence [22].

The following subgroup analyses were planned: (1) the stratification of bias risk of trials (lower risk of bias compared to high risk of bias); (2) the duration of steroid treatment [long-term ( $\geq 4$  days) compared to short-term ( $< 4$  days) use]; (3) patients with SIRS or sepsis compared to patients with severe sepsis or septic shock; (4) stratification based on the aetiology of sepsis; (5) trials using dexamethasone compared with trials using any other

**Table 1** Characteristics of included trials

Trial	Number of patients	Inclusion criteria	Treatment	Mortality data
High dose steroids (>500 mg hydrocortisone or equivalent dose) in experimental intervention group				
Bernard [35]	99	Sepsis	MP 30 mg/kg every 6 h for a total of 4 doses	45 days
Bone [41]	382	Severe sepsis	MP bolus 30 mg/kg repeated every 6 h up to a total of 4 doses	14 days and ICU
De Gans [36]	301	Sepsis	D 10 mg every 6 h for 4 days	8 weeks
Hoffman [37]	38	Sepsis	D 3 mg/kg, followed by 1 mg/kg every 6 h for 8 times	16 months
Hughes [28]	57	Septic shock	Group A ( <i>n</i> = 14): bolus N 10 µg/kg and a single dose of 30 mg/kg MP and 30 min later: a second dose of 100 µg/kg N	Not reported
			Group B: ( <i>n</i> = 15) single bolus of 30 mg/kg MP	
			Group I ( <i>n</i> = 7): 30 µg/kg bolus of N, followed by 30 µg/kg/h for 1 h plus a single dose of 30 mg/kg MP	
			Group II ( <i>n</i> = 7), bolus 30 µg/kg N, followed by 30 µg/kg/h for 1 h without MP	
			Group III ( <i>n</i> = 7): MP, 30 mg/kg	
Klastersky [29]	85	Sepsis	B 0.5 mg/kg every 12 h, during 3 consecutive days	30 days
Luce [42]	87	Septic shock	MP 30 mg/kg per dose every 6 h, four times MP diluted in 50 ml of 5 % dextrose in water	Hospital
Marik [38]	30	Sepsis	Once HC 10 mg/kg iv 30 min prior to starting antibiotic therapy	Not specified
Scarborough [30]	465	Sepsis	D 16 mg in 4 ml of sterile water intravenously for 4 days	10 days, 40 days, 6 months
Schumer [43]	172	Septic shock	Single bolus infusion of either 3 mg/kg D in 100 ml IS or 30 mg/kg of MP in 100 ml IS. If 'necessary' repeated once after 4 h	Not specified
Sprung [44]	59	Septic shock	Group 1 ( <i>n</i> = 21): MP 30 mg/kg Group 2 ( <i>n</i> = 22): D 6 mg/kg	7 days, 14 days and 20 days
			Dose was repeated once in both groups after 4 h if shock persisted	
Thwaites [20]	545	Sepsis	D week 1, 0.4 mg/kg/day; week 2, 0.3 mg/kg/day; week 3, 0.2 mg/kg/day; week 4, 0.1 mg/kg/day; week 5, fixed dose 4 mg/day, decreased by 1 mg each week in following weeks <sup>a</sup>	9 months
VASSC [45]	223	Sepsis	MP 30 mg/kg bolus followed by 5 mg/kg/h for 9 h Maximal dose 10 g	14 days
Wan [31]	81	SIRS	D iv 1 mg/kg 3 times/day for 3 days	30 days
Low dose steroids (≤500 mg hydrocortisone or equivalent dose) in experimental intervention group				
Annane [46]	300	Septic shock	HC 50 mg every 6 h and FC 50 µg every 24 h Treatment duration 6 days	1 year
Arabi [47]	75	Septic shock and cirrhosis	50 mg HC every 6 h. After shock resolution, defined as a MAP > 65 mm Hg without vasopressors for 24 h: dose reduction by 10 mg every 2 days until discontinuation	28 days, ICU and hospital
Bollaert [48]	41	Septic shock	HC 75 mg every 6 h for 5 or more days. In case of shock reversal after 5 days: for 3 days 150 mg HC, then 75 mg for the next 3 days. Treatment was stopped after 5 days in the absence of shock reversal	28 days

Table 1 continued

Trial	Number of patients	Inclusion criteria	Treatment	Mortality data
Briegel [49]	40	Septic shock	HC bolus of 100 mg, followed by a continuous infusion of 0.18 mg/kg/h. After shock reversal dose reduction to 0.08 mg/kg/h. Dose was kept constant for 6 days. HC was tapered in steps of 24 mg/day after successful treatment of infection or serum [sodium] >155 mmol/l	30 days, 90 days, 1 year and ICU
Chawla [50]	44	Septic shock	HC 100 mg every 8 h for 72 h, followed by tapering over 4 days	Not reported
Cicarelli [32]	29	SIRS with or without sepsis	D 0.2 mg/kg (in a single dose)	7 days
Cicarelli [51]	29	Septic shock	D 0.2 mg/kg given three times at intervals of 36 h	7 and 28 days
Confalonieri [33]	46	Sepsis	HC bolus 200 mg, followed by an infusion rate of 10 mg/h for 7 days	7, 28, 60 days and hospital
Gordon [52]	61	Septic shock	HC 50 mg every 6 h for the first 5 days, 50 mg every 12 h for the next 3 days, 50 mg every 24 h for the last 3 days. HC was weaned quicker if shock had resolved	28 days, ICU and hospital
Hu [53]	77	Septic shock	HC 50 mg every 6 h for the first 7 days, 50 mg every 8 h for the next 3 days, 50 mg every 12 h for the next 2 days every 24 h for the next 2 days	14 days
Kaufmann [54]	30	Septic shock	HC bolus of 100 mg followed by a continuous infusion of 10 mg/h for 24 h	Not reported
Mirea [55]	171	Septic shock	Group A: bolus 200 mg HC every 24 h during 4 days Group B: continuous HC 200 mg/day during 4 days	Not reported
Meduri [56]	80	Severe sepsis	HC infusion (10 mg/h) for 7 days	ICU and hospital
Nafae [39]	80	Sepsis	HC 200 mg loading bolus dose followed by a continuous infusion of 240 mg/24 h for 7 days	7 days
Oppert [57]	48	Septic shock	HC (500 mg/50 ml) bolus 50 mg, followed by continuous infusion of 0.18 mg/kg/h. 1 h after cessation of vasopressor support: 0.06 mg/kg/h for 24 h, followed by a reduction of 0.02 mg/kg/h every day	28 days
Rinaldi [58]	40	Severe sepsis	Standard therapy and HC 300 mg/day diluted in 50 ml of IS for 6 days via continuous intravenous infusion. After the study period HC therapy was tapered	Hospital
Ruolan [59]	60	Sepsis	Chinese herb group: 100 ml Sini decoction <sup>b</sup> every 24 h for 7 days	28 days
Snijders [40]	213	Sepsis	HC group (GOJ): 200 mg HC every 24 h for 7 days	30 days
			P 40 mg once daily for a total of 7 days	
Sprung [60]	499	Septic shock	HC 50 mg iv bolus every 6 h for 5 days, then tapered to 50 mg iv every 12 h for days 6–8, 50 mg every 24 h for days 9–11, and then stopped	Mortality rate at 28 days in patients who did and did not have RC
			P iv at 06.00 (5 mg) and 18.00 (2.5 mg) for 10 days P iv 3 times a day at 06.00 (10 mg), 14.00 (5 mg), and 22.00 (5 mg) for 10 days	
Yildiz [61]	40	Sepsis		Overall 28 days, ICU, hospital, 1 year
Yildiz [62]	55	Sepsis		28 days 28 days

B betamethasone, HC hydrocortisone, D dexamethasone, FC fludrocortisone, GOJ group of interest, MP methylprednisolone, iv intravenous, IS isotonic saline, MAP mean arterial pressure, N naloxone, ND not defined, P prednisolone, RC response to corticotrophin

<sup>a</sup> 1/3 of patients received a lower dose of dexamethasone, but still a high dose

<sup>b</sup> Sini decoction: decoction of monkshood 15 g, dried ginger 15 g, honey-fried licorice 10 g



steroid possessing mineralocorticoid properties (unlike dexamethasone).

### Trial sequential analysis

We conducted trial sequential analysis (TSA). Conventional meta-analysis runs the risks of random errors due to sparse data and repetitive testing [15, 18]. TSA adjusts the confidence intervals, if data are sparse or repeatedly analysed as a result of multiple updates, to allow firm conclusions. TSA is similar to interim analysis in a single trial where monitoring boundaries are used to decide whether the trial should be terminated early or whether the confidence interval and the adjacent  $P$  value are sufficiently narrow or small respectively to show the anticipated effect [23]. In the same manner, trial sequential monitoring boundaries can be applied to meta-analyses [9–12, 24].

TSA depends on the quantification of the required information size. We calculated a diversity-adjusted ( $D^2$ ) required information size, since the heterogeneity adjustment with  $I^2$  underestimates the required information size [25]. TSA was conducted with the intention to maintain an overall 5 % risk of a type I error and a power of 90 %. For the calculation of the required information size, we anticipated an intervention effect of a 10 % RRR using the control event proportion calculated from the actual meta-analyses. We provided the TSA-adjusted CI for sparse data and repetitive testing, which we described as the trial sequential analysis adjusted CI. We also performed sensitivity analyses with a power of 80 % and assuming a 20 % RRR.

### Grade assessment of outcomes

Data on the outcomes of all trials were assessed for the risk of bias measured by the level of evidence, the risk of random error measured by standard error, and the design error measured by grading the outcomes according to GRADE [16, 26]. Data were presented in a three-dimensional Manhattan error matrix that facilitates the overview of available evidence at a glance and may identify possible lacunae [16]. A GRADE assessment of all outcomes considering risk of bias, inconsistency, imprecision and risk of publication bias was conducted as well [22].

## Results

The search retrieved 5366 hits (Fig. S1). A total of 48 articles were included describing 35 distinct randomised clinical trials. One ongoing trial was identified [27]. Two

papers were translated from Chinese by a native Chinese medical doctor. The excluded trials and reasons for exclusion are listed (Table S4).

### Characteristics of trials

Thirty-five randomised trials were included. Fourteen trials evaluated high dose (>500 mg hydrocortisone or equivalent) steroids and 21 trials used low dose ( $\leq$ 500 mg hydrocortisone or equivalent) steroids (Table 1 and Table S5). Fifteen trials included patients with septic shock and two trials included patients with SIRS. Duration of steroid treatment varied between one single dose and 8 weeks. The daily dose of steroids varied between 30 mg and 600 mg/kg (total 45 g) hydrocortisone (or equivalent) (Fig. 1). In the high dose trials the daily doses varied between 10 mg/kg (total 750 mg) and 600 mg/kg (total 45 g) hydrocortisone (or equivalent). In the low dose trials daily doses varied between 30 mg and 440 mg hydrocortisone (or equivalent).

There was insufficient data to evaluate the outcomes persistent dependence on haemodialysis and duration of mechanical ventilation.

### Bias risk assessment

Bias risk of trials was assessed according to *The Cochrane Handbook for Systematic Reviews of Interventions* (Fig. S2) [15]. Only two trials scored low risk of bias in all domains. Therefore, we used the prespecified group of trials with lower risk of bias in four domains. A total of 16 studies were assessed to have lower risk of bias (Fig. S2).

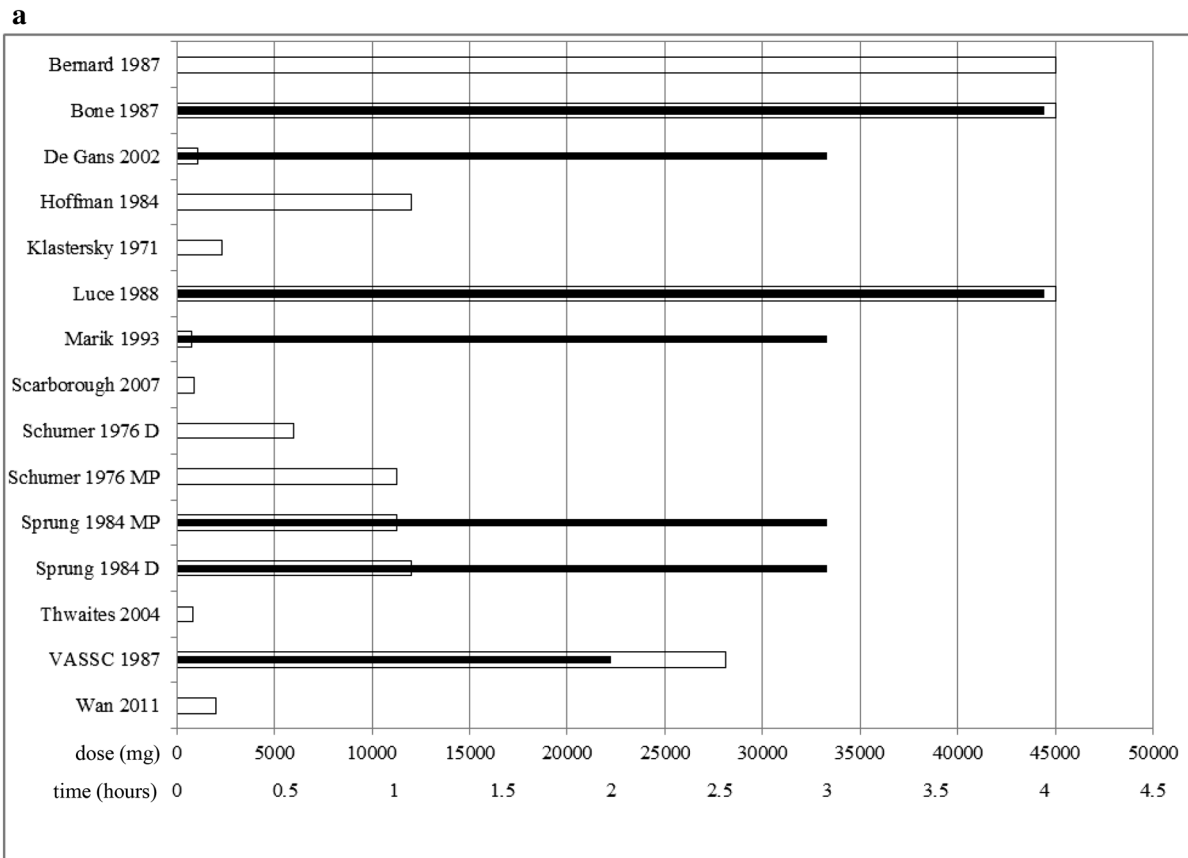
### Effects of interventions

All pooled intervention effects with their 95 % CI of all trials along with subgroup effects and all TSA are listed in Table 2.

#### *Comparison 1: steroids versus placebo or no intervention*

Thirty-five trials randomised 4682 patients and evaluated any dose of steroids in patients with sepsis. Two trials had low risk of bias. We considered 16 trials to have lower risk of bias.

*All-cause mortality within longest follow-up.* Thirty-one trials (including 4290 patients) provided mortality data. Mortality within longest follow-up was 37.6 % in the steroids group and 41.0 % in the control group. Substantial heterogeneity was found ( $I^2 = 54$  %). There was



**Fig. 1** Randomised clinical trials with mortality data showing for each trial the hydrocortisone dose (on the first day) (white bars) and the time interval from sepsis/septic shock onset until randomisation/start treatment (black bars); **a** high dose steroids (>500 mg hydrocortisone or equivalent) and **b** low dose ( $\leq$ 500 mg hydrocortisone or equivalent). When other steroids were used, the equivalent hydrocortisone dose was calculated using the table in the *Oxford Handbook of Critical Care* [17]. When doses were

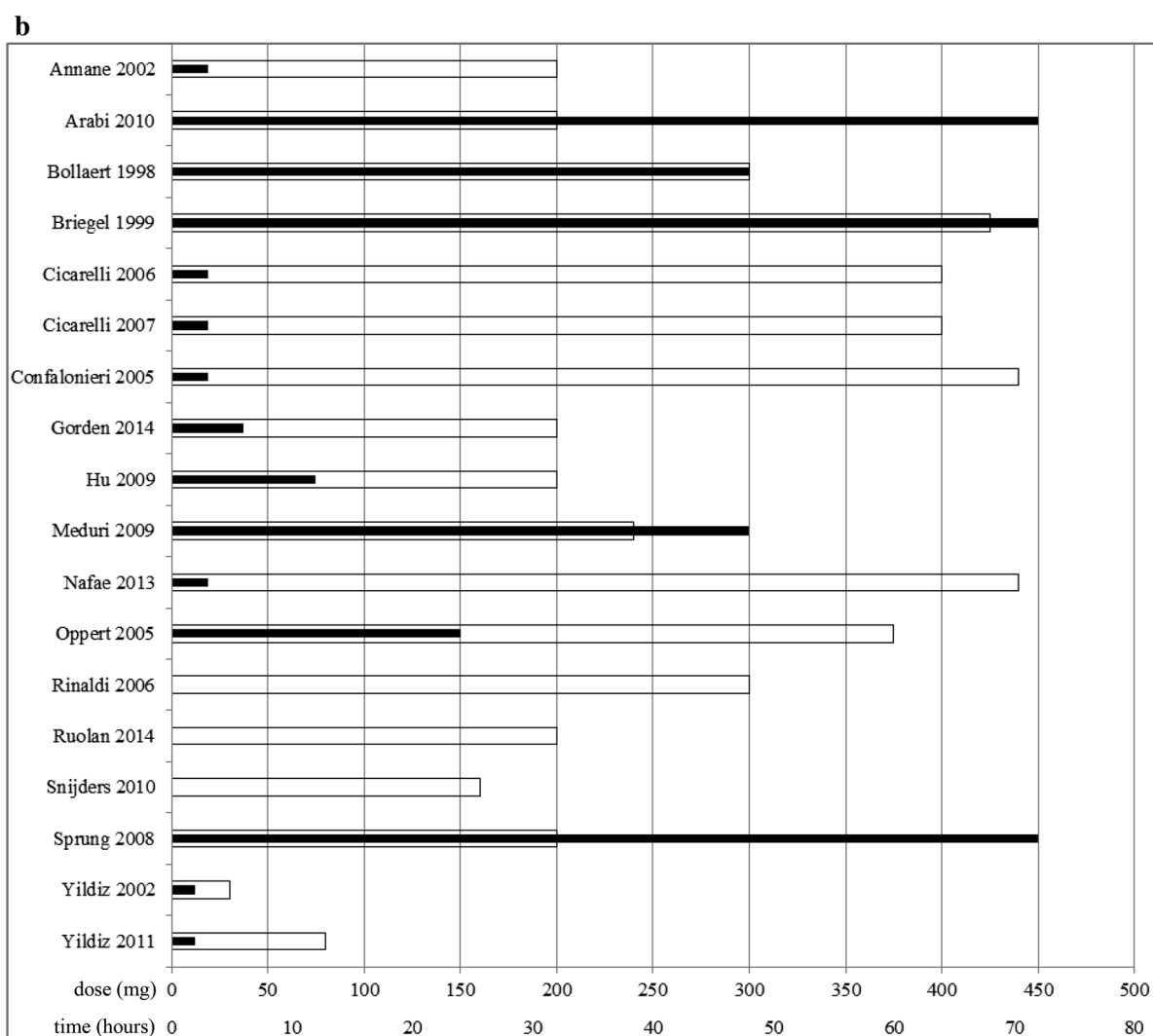
expressed in milligrams per kilogram body weight, daily doses were calculated assuming a body weight of 75 kg. The trials by Hoffman [37], Klastersky [29], Scarborough [30], Schumer [43], Snijders [40], Rinaldi [58], Ruolan [59] and Wan [31] did not provide information on the time interval. The trials by Schumer [43] and Sprung [44] included two intervention groups using different doses of steroids: *D* dexamethasone, *MP* methylprednisolone

no statistically significant difference (random-effects model RR 0.89, 95 % CI 0.79–1.01; TSA-adjusted CI 0.74–1.08; Figs. 2, S3). TSA on the two low risk of bias trials appeared impossible because of insufficient data; the conventional model showed no statistically significant effect (Table 2). Subgroup analysis based on the pre-specified lower risk of bias trials revealed no differential effect. TSA of all trials (RRR 10 %; power 90 %) showed that the accrued and the required information size were far apart (Fig. S3) and that more than 17,000 additional patients may need to be randomised before firm conclusions can be drawn regarding the effect on mortality. However, futility has been reached in our sensitivity trial sequential analysis (RRR 20 %; power 80 %) (Fig. S3) refuting a 20 % RRR to be shown with 80 % power.

Analyses stratified by risk of bias (Fig. 2), treatment duration (long- compared to short-course steroids; Fig. S4), severity of illness (SIRS and sepsis compared to

severe sepsis and septic shock; Fig. S5) and type of steroids (excluding trials using dexamethasone) all showed no statistically significant effects (Table 2) and were in line with the other analyses suggesting that many more patients need to be randomised before firm conclusions may be drawn.

*Serious adverse events excluding mortality.* No statistical difference was found in serious adverse events including all trials that evaluated steroids for sepsis (random-effects model RR 1.02, 95 % CI 0.92–1.15; TSA-adjusted CI 0.7–1.48; Fig. 3, S6). TSA showed that nearly 50,000 additional patients may need to be randomised before firm conclusions can be drawn on the effect on serious adverse events (Fig. S6). The types of serious adverse events are listed in Table S6. The incidence of serious adverse events did not vary according to the degree of sepsis (Table 2).



**Fig. 1** continued

*Other outcomes.* Time-specific analyses of mortality were conducted for 30-day (16 trials) and 90-day (two trials) follow-up. We found no statistically significant treatment effect when evaluating 30-day mortality (random-effects model RR 0.96, 95 % CI 0.85–1.08; TSA RR 0.98, TSA-adjusted CI 0.83–1.17; point estimates were different as a result of different handling of zero event trials) (Fig. S7). Data for 90-day mortality were too sparse to perform TSA-adjusted analysis; in a conventional analysis no statistically significant effect was found (random-effects model RR 0.36, 95 % CI 0.04–2.90) (Fig. S8).

GRADE assessment considering risk of bias, inconsistency, imprecision and risk of publication bias showed very low quality of evidence (Table S7).

*Comparison 2: high dose steroids versus placebo or no intervention*

Fourteen trials (2624 patients) evaluated high doses (>500 mg hydrocortisone or equivalent) of steroids for sepsis. Only one trial had low risk of bias in all domains [20]. Six trials were considered to have lower risk of bias (lower risk of bias in four domains). One trial did not report mortality (Table 1) [28].

*All-cause mortality within longest follow-up.* Six trials with lower risk of bias and seven trials with high risk of bias evaluated mortality in 2537 patients at different lengths of follow-up. No statistically significant beneficial effect from steroid treatment was found (random-effects



**Table 2** Conventional and trial sequential analysis (TSA)-adjusted relative risks (RR) with 95 % confidence intervals for the primary and secondary outcome of mortality and serious adverse events

		All trials		High dose trials (>500 mg hydrocortisone or equivalent)		Low dose trials (≤500 mg hydrocortisone or equivalent)	
		Conventional meta-analysis RR (95 % CI)	TSA-adjusted meta-analysis RR (CI) Model a or b	Conventional meta-analysis RR (95 % CI)	TSA-adjusted meta-analysis RR (CI) Model a or b	Conventional meta-analysis RR (95 % CI)	TSA-adjusted meta-analysis RR (CI) Model a or b
<i>Primary outcomes</i>							
Mortality at longest follow-up	All trials	0.89 (0.79–1.01) 0.95 (0.84–1.08) <sup>b</sup> 0.97 (0.86–1.09) <sup>h</sup>	a: 0.89 (0.74–1.08) b: 0.89 (0.78–1.02)	0.87 (0.70–1.07) 1.08 (0.92–1.28) <sup>h</sup>	a: 0.87 (0.38–1.99) b: 0.87 (0.62–1.22)	0.90 (0.77–1.05) 0.95 (0.84–1.06) <sup>f</sup> 0.91 (0.78–1.06) <sup>h</sup>	a: 0.9 (0.49–1.67) b: 0.9 (0.73–1.1)
Subgroup: risk of bias	Lower risk	0.92 (0.82–1.03) 0.98 (0.87–1.09) <sup>b</sup> 0.38 (0.06–2.42) <sup>c</sup>	a: 0.92 (0.69–1.24) b: 0.92 (0.81–1.05)	0.94 (0.74–1.19)	a: 0.94 (0.36–2.44) b: 0.94 (0.61–1.46)	0.92 (0.80–1.05) 0.94 (0.84–1.05) <sup>g</sup>	a: 0.92 (0.53–1.58) b: 0.92 (0.78–1.08)
Subgroup: treatment duration	High risk	0.79 (0.58–1.08)	–	0.70 (0.45–1.09)	–	0.88 (0.53–1.48)	–
	Short course	0.89 (0.67–1.17)	a: ID	0.86 (0.65–1.15)	–	–	–
	Long course	0.88 (0.77–1.01)	b: 0.89 (0.48–1.64) a: 0.88 (0.51–1.51)	0.82 (0.57–1.19)	–	0.89 (0.76–1.04)	a: 0.89 (0.48–1.67) b: 0.89 (0.72–1.11)
Subgroup: disease severity	SIRS and sepsis	0.86 (0.70–1.04) 0.85 (0.69–1.04) <sup>a</sup> 0.91 (0.77–1.07)	a: 0.86 (0.39–1.89) b: 0.86 (0.63–1.17) a: 0.91 (0.47–1.76) b: 0.91 (0.72–1.15)	0.91 (0.74–1.12)	a: 0.91 (0.4–2.11) b: 0.91 (0.65–1.29) a: ID b: ID	0.68 (0.40–1.15) 0.60 (0.34–1.08) <sup>a</sup> 0.95 (0.83–1.08)	a: ID b: 0.68 (0.08–5.54) a: 0.95 (0.56–1.61) b: 0.95 (0.81–1.10)
Subgroup: disease severity	Severe sepsis and shock	1.01 (0.88–1.16) 1.24 (0.90–1.70)	a: ID a: ID	1.07 (0.89–1.27) 1.72 (0.72–4.13)	a: 1.06 (0.53–2.13) a: ID	0.73 (0.41–1.29) 1.20 (0.74–1.92)	a: ID a: ID
<i>Secondary outcomes</i>							
30-day mortality	All	0.96 (0.85–1.08) 0.94 (0.83–1.08) <sup>e</sup>	a: 0.98 (0.83–1.17) <sup>d</sup> b: 0.98 (0.86–1.12) <sup>d</sup>	1.03 (0.90–1.17) 1.00 (0.87–1.15)	a: 1.02 (0.76–1.37) a: 0.99 (0.66–1.51)	0.96 (0.73–1.27) 0.92 (0.59–1.44)	a: 0.94 (0.55–1.62) <sup>d</sup> b: 0.94 (0.81–1.09) <sup>d</sup>
Subgroup: risk of bias	Lower risk	0.91 (0.75–1.10)	–	1.31 (0.67–2.6)	–	0.42 (0.01–21.21)	–
Subgroup: risk of bias	High risk	1.02 (0.87–1.18)	–	1.04 (0.89–1.22)	–	0.79 (0.48–1.30)	–
90-day mortality	All	0.36 (0.04–2.90)	a: ID b: ID	–	–	0.36 (0.04–2.90)	a: ID b: ID
Subgroup: risk of bias	Lower risk	0.36 (0.04–2.90)	–	–	–	0.36 (0.04–2.90)	–
Subgroup: risk of bias	High risk	–	–	–	–	–	–

All estimates are reported with relative risks and calculated using a random-effects model

SIRS systemic inflammatory response syndrome, DIS diversity-adjusted information size, ID insufficient data (<5 % DIS) for calculation of an information size and TSA-adjusted confidence interval, – no data available

Model a = TSA with the predefined  $\alpha = 0.05$  (two sided),  $\beta = 0.10$  (power 90 %), and an anticipated relative risk reduction of 10 %

Model b = TSA with the predefined  $\alpha = 0.05$  (two sided),  $\beta = 0.20$  (power 80 %), and an anticipated relative risk reduction of 20 %

<sup>a</sup> Results with sepsis trials only

<sup>b</sup> Results after exclusion of the following trials: Bernard [35], Confalonieri [33], de Gans [36], Hoffman [37], Marik [38], Nafae [39], Scarborough [30], Snijders [40], Thwaites [20] and Wan [31] (inclusion can be disputed)

<sup>c</sup> Low risk of bias trials only: Confalonieri, Thwaites

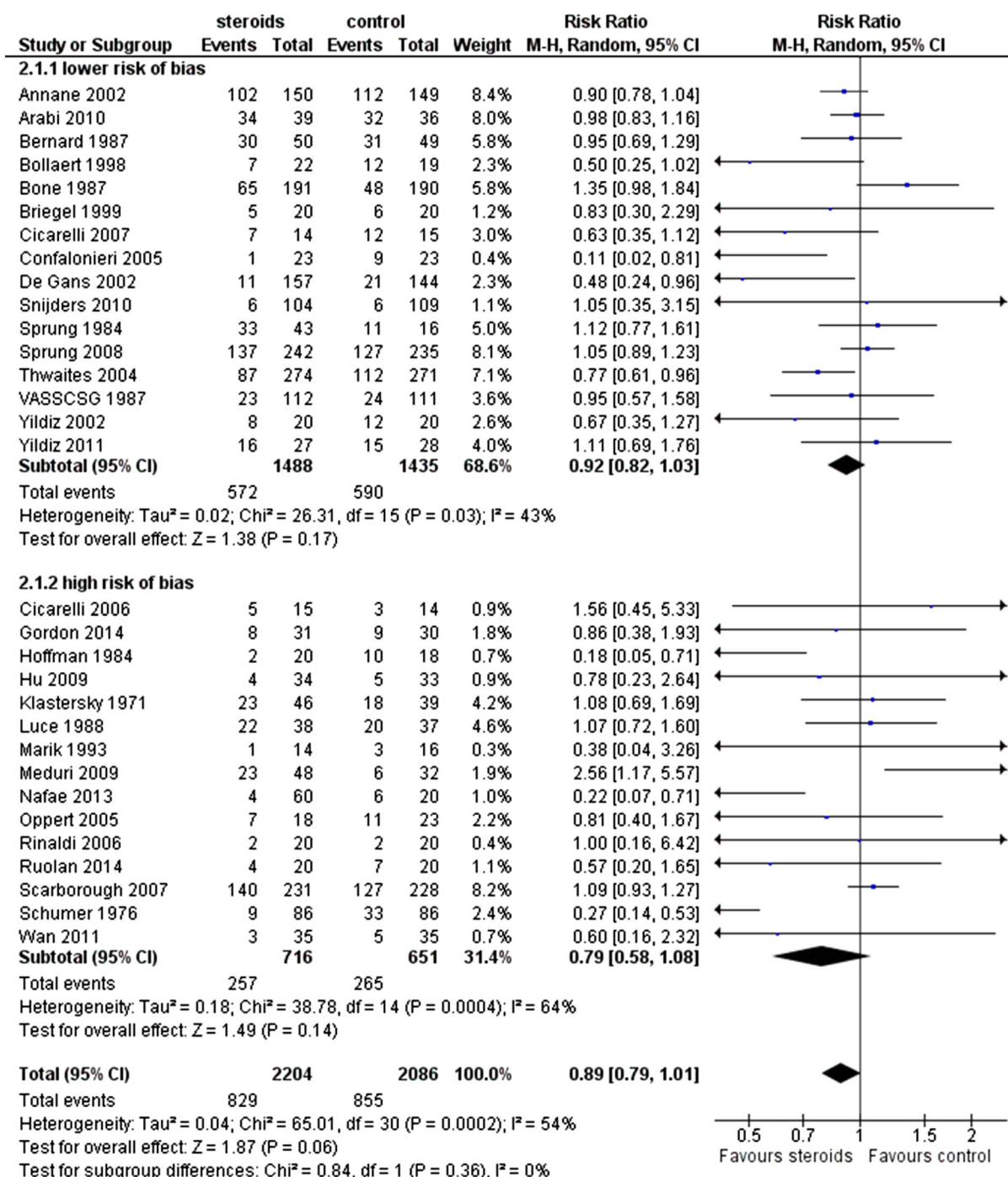
<sup>d</sup> Point estimate varies to some extent since Review Manager software (used for conventional analysis) uses a constant continuity correction for zero events and TSA software uses an empirical continuity correction for zero events

<sup>e</sup> Results after exclusion of the following trials: Confalonieri, Scarborough and Snijders (inclusion can be disputed)

<sup>f</sup> Results after exclusion of the following trials: Confalonieri, Snijders, Nafae (inclusion can be disputed)

<sup>g</sup> Results after exclusion of Confalonieri and Snijders (inclusion can be disputed)

<sup>h</sup> Results after exclusion of trials using dexamethasone (dexamethasone has no mineralocorticoid activity)



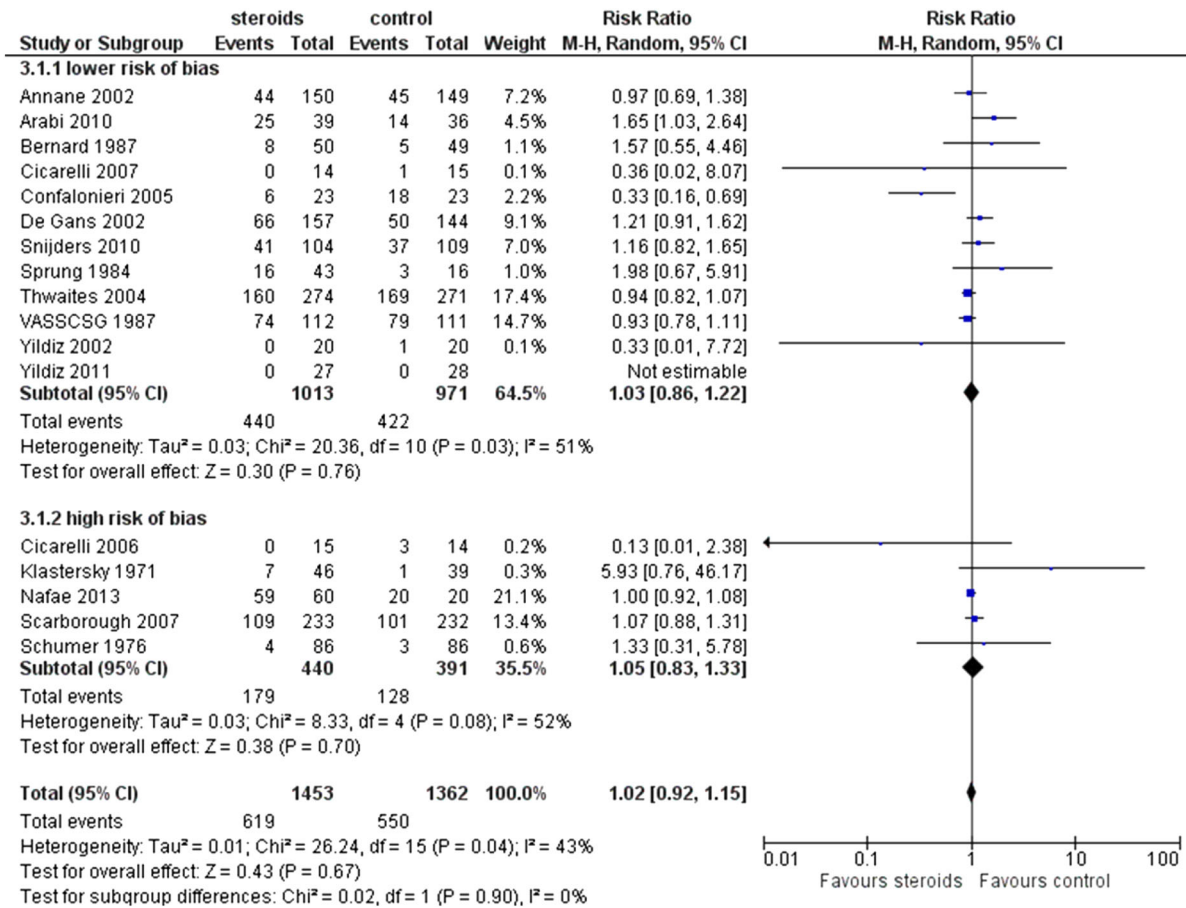
**Fig. 2** Forest plot of mortality at longest follow-up of all trials evaluating steroids for sepsis with subgroups according to risk of bias (random-effects model)

model RR 0.87, 95 % CI 0.70–1.07; TSA-adjusted CI 0.38–1.99; Fig. S9).

Analyses stratified by risk of bias (Fig. S9), severity of illness (SIRS and sepsis compared to severe sepsis and septic shock; Fig. S10) and type of steroids (excluding

trials using dexamethasone) all showed no statistically significant effects (Table 2).

*Serious adverse events excluding mortality.* No statistical significant difference was found in the overall



**Fig. 3** Forest plot of serious adverse events of all trials evaluating steroids for sepsis with subgroups according to risk of bias (random-effects model)

proportion of serious adverse events (random-effects model RR 1.03, 95 % CI 0.90–1.17; TSA-adjusted RR 1.02, CI 0.70–1.48). No statistically significant differences were found in subgroups according to bias risk and disease severity (Fig. S11, Table 2).

**Other outcomes.** Three trials with high risk of bias [29–31] reported mortality at 30 days; no 90-day follow-up data was reported (Table 2). There was no statistically significant effect.

Error matrix plots were constructed for overview of all available evidence for high dose steroids at a glance (Fig. S17). GRADE assessment considering risk of bias, inconsistency, imprecision and risk of publication bias showed very low quality of evidence for all outcomes (Table S8).

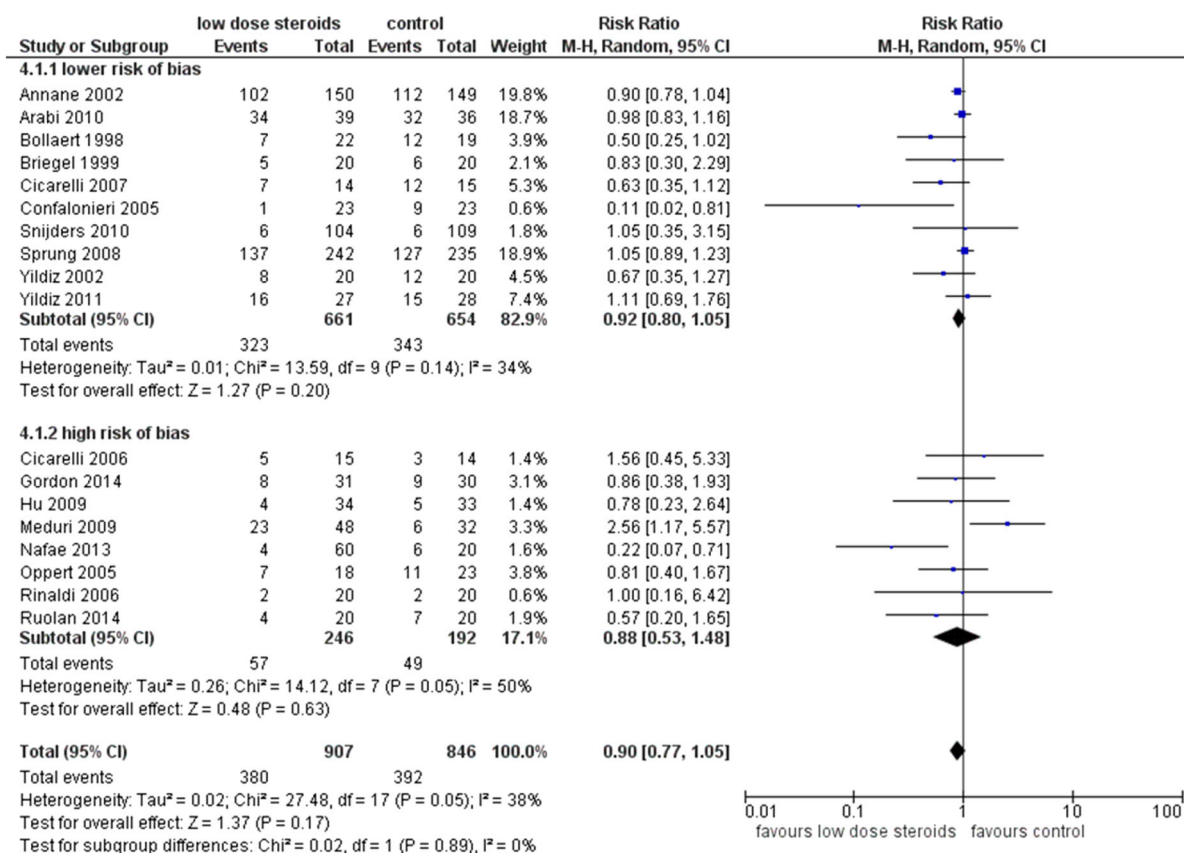
*Comparison 3: low dose steroids versus placebo or no intervention*

Twenty-one trials randomised 2058 patients for low dose ( $\leq 500$  mg hydrocortisone or equivalent) steroids for

sepsis. There were large differences between the trials in the time interval between sepsis onset and initiation of steroids treatment: 2–72 h (Fig. 1). Only one trial applied short-course low dose steroids [32].

**All-cause mortality within longest follow-up.** Only one trial had low risk of bias in all domains [33]. Ten trials with lower risk of bias (1315 patients) and eight trials with high risk of bias (438 patients) evaluated mortality at different lengths of follow-up. The overall pooled estimate showed no statistical difference (random-effects model RR 0.90, 95 % CI 0.77–1.05; TSA-adjusted CI 0.49–1.67) (Figs. 4, S12).

Analyses stratified by risk of bias (Figs. 4, S12), treatment duration (long- compared to short-course steroids; Fig. S13), severity of illness (SIRS and sepsis compared to severe sepsis and septic shock; Fig. S14) and type of steroids (excluding trials using dexamethasone) all showed no statistically significant effects (Table 2). Sensitivity analyses (TSA, RRR 20 %; power 80 %) based on lower risk of bias trials and separately all trials evaluating severe sepsis and septic shock showed futility for treatment with low dose steroids (Figs. S12, S14).



**Fig. 4** Forest plot of mortality at longest follow-up of low dose steroids ( $\leq 500$  mg hydrocortisone or equivalent) use according to risk of bias subgroups (random-effects model)

*Serious adverse events excluding mortality.* Nine trials (with 866 patients) evaluated serious adverse events. One trial had low risk of bias [33]. Seven trials had lower risk of bias. Data were too sparse to perform a TSA. A conventional analysis found no statistically significant intervention effect (random-effects model RR 0.96, 95 % CI 0.73–1.27; Fig. S15). The incidence of serious adverse events did not vary according to the degree of sepsis (Table 2).

*Mortality at 30 days.* Thirteen trials (1479 patients) evaluated mortality at 30-day follow-up. One trial had low risk of bias [33]. Ten trials had lower risk of bias. No significant benefit was found from steroids treatment (random-effects model RR 0.91, 95 % CI 0.77–1.07; TSA RR 0.94, TSA-adjusted CI 0.55–1.62; point estimates were different as a result of different handling of zero event trials; Fig. S16). Subgroup analysis based on bias risk showed no differential effect. TSA estimated that many more randomized patients are needed before firm conclusions can be drawn. A sensitivity analysis (TSA, RRR 20 %; power 80 %) showed futility for low dose steroids treatment (Fig. S16).

*Mortality at 90 days.* The results of 90-day mortality in low dose steroids are equal to the overall 90-day mortality, since only trials that evaluated low dose steroids provided data on 90-day follow-up.

*Other outcomes.* Error matrix plots were constructed for overview of all available evidence for low dose steroids at a glance (Fig. S18). Data suggest that there are considerable risks of systematic errors (bias) and random errors (large standard errors on average). GRADE assessment considering risk of bias, inconsistency, imprecision and risk of publication bias showed very low quality of evidence for all outcomes (Table S9).

## Discussion

We did not find evidence for a beneficial effect of an intervention with steroids in patients with SIRS, sepsis, severe sepsis or septic shock in this systematic review with meta-analyses and TSA, including 35 randomised trials with 4682 patients. High ( $>500$  mg hydrocortisone



or equivalent) and low dose ( $\leq 500$  mg hydrocortisone or equivalent) steroids were evaluated in separate comparisons and no evidence of a beneficial effect was found. Moreover, TSA suggested that more than 17,000 patients need to be randomised before firm conclusions can be drawn on any present or absent intervention effect with a 10 % RRR. Evidence has been reached to refute a 20 % RRR with a power of 80 %.

Our conclusion contrasts with previous publications suggesting beneficial effects associated with use of a long course of low dose steroids [6]. Differences might be explained by another search strategy and different analyses with improved accounting for risks of systematic, design and random errors. There is accumulating evidence that random error plays an important role in premature conclusions of spurious significant findings [11]. In simulation studies up to 30 % of premature declarations of significant effects are in fact overestimations of intervention effects, once sufficient evidence has been reached [34].

A substantial clinical heterogeneity existed between the included trials, even after separation into high and low doses of steroids. Within both high and low dose steroid groups the administered daily dose of steroids differed importantly (Fig. 1). Moreover, there were substantial differences regarding the time interval between sepsis onset and administration of the first steroid dose, e.g. in antibiotic therapy timing appears to be an essential feature for any beneficial effect. Therefore, lack of both optimised dosing and optimised timing might obscure potential beneficial effects of steroids. As dexamethasone has no mineralocorticoid activity, analyses were repeated excluding trials using dexamethasone and results appeared to be similar (Table 2).

Retrospectively, our study protocol could have been more specific on exclusion criteria for trials evaluating patients with SIRS and sepsis. We excluded trials that evaluated steroids for prevention of diseases and for autoimmune diseases. We also excluded trials that evaluated steroids for treatment of localized oedema. Furthermore, we only included trials that evaluated patients with SIRS if the criteria for SIRS were explicitly stated in the methods section of the report; however, SIRS criteria may not predict patient important outcomes and may be too inclusive. Although we reached consensus through discussions, the inclusion of some trials can still be discussed [20, 30, 31, 33, 35–40]. To test the

robustness of our conclusions, we therefore conducted several sensitivity analyses by excluding these trials and they all resulted in similar findings (Table 2).

Trial sequential analyses, an error matrix to evaluate risks of errors and GRADE assessment add to the strength of the conclusions. However, our systematic review mirrors the lack of quality and quantity of the included randomised clinical trials. The included trials fall short on bias protection, included numbers of patients and chosen outcomes. Therefore, the evidence of steroids for sepsis (both high and low dose) is characterized by high risks of both systematic errors and random errors. We used wide inclusion criteria to evaluate the effect of steroids in the broad spectrum of disease. Although none of the analyses showed a statistically significant effect, we cannot exclude a small beneficial intervention effect of steroids in a specific subgroup.

## Conclusion

In this systematic review with meta-analyses and TSA we did not find any statistically significant beneficial effect that could support propagating the use of steroids for sepsis. Further, our TSA suggests that many thousands of randomised patients are needed in order to change this perspective. Therefore, steroids should no longer be recommended for patients with sepsis outside ongoing [27] or future well-designed randomised clinical trials with low risks of both systematic error and random error.

**Acknowledgments** We thank Y. J. Gu for translation of the Chinese papers and W. Bruins for language suggestions. We also thank Dr. Confalonieri for providing additional unpublished follow-up data.

**Conflicts of interest** None of the authors has a conflict of interest to declare.

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