A systematic review and trial sequential analysis of intravenous vs. oral peri-operative paracetamol

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Summary
Postoperative pain might be different after intravenous vs. oral paracetamol. We systematically reviewed randomised controlled trials in patients >15 years that compared intravenous with oral paracetamol for postoperative pain. We identified 14 trials with 1695 participants. There was inconclusive evidence for an effect of route of paracetamol administration on postoperative pain at 0–2 h (734 participants), 2–6 h (766 participants), 6–24 h (1115 participants) and >24 h (248 participants), with differences in standardised mean (95%CI) pain scores for intravenous vs. oral of −0.17 (−0.45 to 0.10), −0.09 (−0.24 to 0.06), 0.06 (−0.12 to 0.23) and 0.03 (−0.22 to 0.28), respectively. Trial sequential analyses suggested that a total of 3948 participants would be needed to demonstrate a meaningful difference in pain or its absence at 0–2 h. There were no differences in secondary outcomes. Intravenous paracetamol is more expensive than oral paracetamol. Substitution of oral paracetamol in half the patients given intravenous paracetamol in our hospital would save around £38,711 (£43,960 or US$47,498) per annum.

Introduction
Postoperative pain management aims to facilitate optimal recovery and patient satisfaction [1]. Good postoperative analgesia is associated with shorter hospital stay, fewer re-admissions after same-day surgery and fewer postoperative complications, including chronic pain conditions and myocardial ischaemia [2–4].

Paracetamol is a synthetic, non-opioid, centrally-acting analgesic [5]. It is one of the most used and safest drugs available [6, 7], and may avoid the use and undesirable side-effects of opioids [5, 8–10]. Adverse events caused by paracetamol are usually mild, transient and comparable in frequency with placebo [11]. The onset and duration of the analgesic action of paracetamol varies with the route of administration. However, despite several differences in peak plasma concentration after intravenous and oral administration, pain relief is usually similar after 45 min and subsequently could be superior after oral administration [12]. Intravenous paracetamol is more expensive than a bioequivalent dose of oral paracetamol [13, 14].

We aimed to systematically review whether there is a difference in the efficacy, safety and costs of intravenous vs. oral peri-operative paracetamol in adults.

Methods
We searched MEDLINE, Epub, Embase, CENTRAL, Web of Science, LILACs and Google Scholar to February 2020 for trials that compared intravenous with oral peri-operative paracetamol, published in any language (online Supporting Information, Appendix S1). We scanned clinical trial
registries (http://www.who.int/ictrp and clinicaltrials.gov) and the reference lists and citations of included trials and relevant systematic reviews for further trials. When necessary, we contacted trial authors for additional information. Two authors (MM and AV) independently determined trials eligible for inclusion and extracted data, with disagreements resolved by consensus with the help of a third author (JC).

We included parallel group, randomised controlled trials of intravenous vs. oral paracetamol for any operation. Paracetamol could be given before, during or after surgery, as one or multiple doses. We excluded trials with <10 participants or with any participants <15 years old. We chose postoperative pain as the primary outcome, measured by a validated pain scale. We categorised pain measured: during 0–2 postoperative hours; 2–6 postoperative hours; 6–24 postoperative hours; and after 24 postoperative hours. We used the latest available estimate in each time period. Secondary outcomes were: opioid consumption during the first 24 h; time to first analgesic request or rescue dosage; participant satisfaction; time to discharge from the recovery unit and the hospital; nausea or vomiting; pruritus; sedation; and plasma paracetamol concentration (online Supporting Information, Table S1).

Two authors (MM and AV) independently assessed risk of bias in each methodological domain as low, unclear or high [15, 16]. We downgraded the strength of evidence for pooled data from trials at risk of bias and with heterogeneous results, particularly when imprecise. We derived the standard deviations for mean values from standard errors, confidence intervals and p values where necessary and we transformed median (range or interquartile range) to mean (SD) [17–23]. We converted postoperative opioid consumption to intravenous

Figure 1 Flow chart of the literature search.
morpheine milligram equivalents (http://opioidcalculator.practicalpainmanagement.com).

We pooled outcomes reported by two or more trials with a random effects model. We calculated the $I^2$ statistic to assess trial heterogeneity. We considered a $p$ value $< 0.05$ statistically significant. We used trial sequential analysis and a funnel plot for the primary outcome (TSA; version 0.9.5.10 Beta, Copenhagen Trial Unit, Copenhagen, Denmark) [24]. We calculated the required information size allowing for a type-1 error of 0.05 and type-2 error of 0.20.

### Table 1 Details of 14 randomised controlled trials of intravenous vs. oral peri-operative paracetamol.

| Trial                    | Number of patients | Surgery                        | Paracetamol dose and timing | Primary outcome | Peri-operative analgesia               |
|-------------------------|--------------------|--------------------------------|----------------------------|----------------|----------------------------------------|
| Brett et al. [18]       | 10                 | Knee arthroscopy               | 1 g just before surgery    | Plasma concentration | Intra-operative fentanyl               |
| Polito et al. [19]      | 63                 | Hip and knee arthroplasty      | 1 g before surgery and 6-hourly for 24 h | Opioid dose | Pre-operative celecoxib and oxycodone. Intra-operative bupivacaine. Postoperative hydromorphone, oxycodone, oxycotin and celecoxib |
| Plunkett et al. [20]    | 32                 | Cholecystectomy                | 1 g 1 h before surgery and 4 h later | Pain scores differences from baseline first 24 h (NRS) | Intra-operative fentanyl and hydromorphone and subsequent narcotic doses |
| Fenlon et al. [21]      | 63                 | Third molar                    | 1 g 45 min before surgery  | Pain (10 cm VAS) at 1 h after surgery | Intra-operative fentanyl. Postoperative rescue diclofenac |
| Westrich et al. [22]    | 77                 | Total hip arthroplasty         | 1 g 30 min after admission to the PACU | Pain scores (NRS) with activity POD 1 | Intra-operative ketorolac. Postoperative ketorolac, meloxicam and patient-controlled epidural analgesia with bupivacaine and clonidine |
| Bhoga et al. [23]       | 50                 | Endoscopic sinus surgery       | 1 g 1 h before surgery end | Pain scores (10 cm VAS) 1 h postoperative | Pre-operative celecoxib |
| Pettersson et al. [25]  | 40                 | Coronary artery bypass graft   | 1 g 6-hourly after induction until 0900 next morning | Opioid dose | Pre-operative morphine or ketobemidone. Intra-operative fentanyl. Postoperative ketobemidone and aspirin |
| Plunkett et al. [26]    | 47                 | Elective caesarean section     | 1 g postoperative and 8-hourly × 2 | Opioid dose to 24 h | Intra-operative spinal bupivacaine with fentanyl and morphine. Postoperative ketorolac, oxycodone and morphine |
| Hickman et al. [27]     | 245                | Knee or hip arthroplasty       | 1 g intra-operative        | Opioid dose to 24 h postoperative | Pre-operative celecoxib, pregabalin paracetamol (1 g). Postoperative paracetamol (1 g), methocarbamol, tramadol, oxycodone and hydromorphone |
| Van der Westhuizen et al. [28] | 54         | Ear, nose and throat or orthopaedic | 1 g on induction of anaesthesia | Plasma concentration every 30 min for 240 min | Not specified |
| Mahajan et al. [29]     | 50                 | Elective caesarean section     | 10-15 mg kg$^{-1}$ 20 min before surgery end | Analgesia duration Pain (10 cm VAS) 2-hourly to 24 h postoperative | Spinal bupivacaine. Rescue diclofenac |
| O’Neal et al. [30]      | 57                 | Knee arthroplasty              | 1 g at the end of surgery | Pain scores (NRS 11 point) every 15 min for up to 4 h | Pre-operative celecoxib and oxycodone. Intra-operative percutaneous ropivacaine, ketorolac, clonidine |
| Pettersson et al. [31]  | 7                  | Varicose vein, hernia, knee arthroplasty | 2 g propacetamol postoperative | Plasma concentration at 80 min | Lornoxicam |
| Patel et al. [32]       | 44                 | Laparoscopic unilateral hernia repair surgery | 1 g after induction of anaesthesia | Pain scores (NRS 0-10) at rest and 1 h on PACU, and 6 h postoperative Opioid use intra-operatively in the PACU | Intra-operative opioids and bupivacaine for infiltration prior and on closure of the incision site. Postoperative oxycodone and fentanyl; in some cases, used hydromorphone |

i.v., intravenous; p.o., oral; VAS, visual analogue scale; NRS, numerical rating scale; PACU, post-anaesthesia care unit; POD, postoperative days.
Results
We included 14 trials (Fig. 1; Table 1) [18–23, 25–32]. We asked authors of three trials to supply additional information [20, 23, 26]. We derived standard deviations for three trials [18, 19, 26] and mean (SD) for two trials [25, 27]. Most methodological domains were poorly reported by most trials, whereas the provided information revealed high risks of bias for three trials (Fig. 2).

Route of paracetamol administration did not affect postoperative pain (Figs. 3 and 4). There were insufficient trials to interrogate small studies effects (online Supporting Information, Figure S1). Route of paracetamol administration did not affect any of the secondary outcomes (online Supporting Information, Figure S2). We graded the quality of evidence as ‘low’ for an effect of route of paracetamol administration on postoperative pain.

We did not pool plasma paracetamol concentrations, which were reported at different times and in different units by three trials [18, 28, 31]. Intravenous paracetamol administration may increase plasma concentration more than oral administration 20–240 min later, although one trial reported higher plasma concentration 80 min after oral administration [28].

Intravenous paracetamol is approximately 10 times more expensive than an equivalent oral dose, for instance, £1.95 (£2.21, US$2.39) vs. £0.19 (£0.22, US$0.23). We estimate that we spend £85,910 (£97,558, US$120,420) on intravenous paracetamol per annum in our hospital. We would save around £38,711 (£43,960, US$47,498) per annum if we used oral instead of intravenous paracetamol for half of these patients (online Supporting Information, Table S2).

Discussion
We found that the peri-operative route of paracetamol administration, intravenous vs. oral, did not affect pain or any other postoperative outcome. There was insufficient evidence to exclude important clinical effects and overall, the quality of evidence was poor.

Two systematic reviews similarly reported no effect of paracetamol administration route on clinical and pharmacokinetic outcomes [33, 34]. The conclusions of both reviews were limited by the poor methodological or reporting quality of the included trials. This is consistent with large observational studies [35]. Important differences between administration routes could not be excluded.

Most of the included trials gave paracetamol prophylactically and some did not clearly describe whether paracetamol was given as prophylaxis or treatment [22, 25, 26, 30, 31]. We excluded one trial that gave paracetamol as treatment [36].

Previous trials have shown bioequivalence of paracetamol 1 g and propacetamol 2 g [6, 37]. Head-to-head trials of intravenous paracetamol vs. intravenous propacetamol have shown no differences in the proportion of participants achieving at least 50% pain relief during 4 postoperative hours [6, 36, 38, 39]. Only one included trial gave propacetamol compared with five doses of oral paracetamol [31]. This study did not report any clinical outcomes that we could analyse.

All systematic reviews are limited by the trials they observe. Most trials incompletely reported their methods and outcomes were often different. We had to transform some results that were reported as median (range or interquartile range) or as mean without variance. We had to standardise pain scores, limiting their direct clinical interpretation. Our cost analysis is specific to our hospital in the Netherlands, but we believe it is generally applicable [40]. We did not compare the rectal route with others, nor pharmacokinetic profiles of included routes, which remain uncertain. We decided not to extend the pooling of results...
with network meta-analyses, given the heterogeneity of trials and the variability of plasma concentrations [41].

Our review summarises the lack of evidence to justify the expense of peri-operative intravenous paracetamol. It remains possible that there might be an important clinical difference for the route of paracetamol administration. We believe that intravenous paracetamol should only be used in clinical trials or when the oral route is contra-indicated.

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Figure 4  Trial sequence analysis (TSA) for intravenous vs. oral peri-operative paracetamol for postoperative pain: (a) 0–2 h (734 participants); (b) 2–6 h (766 participants); (c) 6–24 h (1115 participants); (d) >24 h (248 participants). The point of interest is whether the cumulative evidence for an effect (Z-curve, blue line) breaches the TSA boundaries (red line) in favour of intravenous paracetamol (above the top red line) or in favour of oral paracetamol (below the bottom red line). The cumulative evidence favours neither route. Additional evidence might breach a boundary for effect, or it might breach the boundaries for clinical futility, set at a Z-score < 1.96 (wedged red lines to the right). At this limit definitive answers could be expected after studying a total of 3948 participants (0–2 h), 14,336 participants (2–6 h), an undetermined number of participants (6–24 h), and 4514 participants (>24 h), assuming alpha 0.05 and beta 0.20.

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Supporting Information
Additional supporting information may be found online via the journal website.

Figure S1. Funnel plot for small studies effects on postoperative pain: (a) 0–2 h; (b) 2–6 h; (c) 6–24 h; (d) >24 h.

Figure S2. Forest plots for secondary postoperative outcomes: (a) opioid consumption; (b) time to first analgesic request or rescue; (c) length of stay in the recovery area (min); (d) length of hospital stay (hours); (e) satisfaction; (f) nausea or vomiting.

Table S1. Definitions for outcomes extracted from included randomised controlled trials.

Table S2. Economic analysis for intravenous vs. oral paracetamol.

Appendix S1. Search strategy for randomised controlled trials of peri-operative intravenous vs. oral paracetamol.