Analgesic efficacy and safety of morphine in the Procedural Pain in Premature Infants (Poppi) study: randomised placebo-controlled trial

Caroline Hartley*, Fiona Moultrie*, Amy Hoskin, Gabrielle Green, Vaneesha Monk, Jennifer L Bell, Andrew R King, Miranda Buckle, Marianne van der Vaart, Deniz Gursul, Sezgi Goksan, Edmund Juszczak, Jane E Norman, Richard Rogers, Chetan Patel, Elier Adams, Rebecca Slater

Summary

Background Infant pain has immediate and long-term effects but is undertreated because of a paucity of evidence-based analgesics. Although morphine is often used to sedate ventilated infants, its analgesic efficacy is unclear. We aimed to establish whether oral morphine could provide effective and safe analgesia in non-ventilated premature infants for acute procedural pain.

Methods In this single-centre masked trial, 31 infants at the John Radcliffe Hospital, Oxford, UK, were randomly allocated using a web-based facility with a minimisation algorithm to either 100 μg/kg oral morphine sulphate or placebo 1 h before a clinically required heel lance and retinopathy of prematurity screening examination, on the same occasion. Eligible infants were born prematurely at less than 32 weeks' gestation or with a birthweight lower than 1501 g and had a gestational age of 34–42 weeks at the time of the study. The co-primary outcome measures were the Premature Infant Pain Profile–Revised (PIPP-R) score after retinopathy of prematurity screening and the magnitude of noxious-evoked brain activity after heel lancing. Secondary outcome measures assessed physiological stability and safety. This trial is registered with the European Clinical Trials Database (number 2014-003237-25).

Findings Between Oct 30, 2016, and Nov 17, 2017, 15 infants were randomly allocated to morphine and 16 to placebo; one infant assigned placebo was withdrawn from the study before monitoring began. The predefined stopping boundary was crossed, and trial recruitment stopped because of profound respiratory adverse effects of morphine without suggestion of analgesic efficacy. None of the co-primary outcome measures differed significantly between groups. PIPP-R score after retinopathy of prematurity screening was mean 11·1 (SD 3·2) with morphine and 10·5 (3·4) with placebo (mean difference 0·6, 95% CI –0·16 to 1·4; p=0·30). Noxious-evoked brain activity after heel lancing was median 0·99 (IQR 0·40–1·56) with morphine and 0·75 (0·33–1·22) with placebo (median difference 0·25, 95% CI –0·16 to 0·80; p=0·25).

Interpretation Administration of oral morphine (100 μg/kg) to non-ventilated premature infants has the potential for harm without analgesic efficacy. We do not recommend oral morphine for retinopathy of prematurity screening and strongly advise caution if considering its use for other acute painful procedures in non-ventilated premature infants.

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We studied infants who required both a routine blood test and retinopathy of prematurity screening on the same morning. We aimed to test whether one dose of oral morphine sulphate (100 μg/kg) administered to non-ventilated premature infants before heel lancing and retinopathy of prematurity screening would provide analgesia, reduce physiological instability, and be safe.15 We chose an oral dose of 100 μg/kg (with an estimated peak effect at 1 h) based on extrapolation from guidance in the British National Formulary for children, local practice guidelines for neonatal eye surgery, and findings of a previous incomplete trial.21 We assessed multimodal pain measures: noxious-evoked brain activity, reflex activity, physiology, and behaviour. To provide insights into how both morphine and retinopathy of prematurity screening alter infant physiology, we monitored infants’ heart rate, respiratory rate, blood pressure, and oxygen saturation for 24 h before and after the clinical procedure. We evaluated drug safety by assessing the incidence of hypotension requiring ionotropes and of apnoeic episodes requiring resuscitative non-invasive positive pressure ventilation (NIPPV), both potential adverse effects of morphine.

Methods

Study design and participants

We undertook a single-centre, masked, randomised, placebo-controlled trial at the Newborn Care Unit at the John Radcliffe Hospital (Oxford University Hospitals NHS Trust, Oxford, UK). This trial was supported by the National Perinatal Epidemiology Unit Clinical Trials Unit (NPEU CTU).

Each infant was studied on a single test occasion. Infants were eligible for inclusion if they were born prematurely at less than 32 weeks’ gestation or with a birthweight less than 1501 g (fulfilling UK retinopathy of prematurity guidelines),17 were both inpatients at the time of the study and aged 34–42 weeks’ gestation, and required a heel lance and retinopathy of prematurity screening on the same test occasion (referred to hereafter as the clinical procedure). Exclusion criteria are provided in the appendix. All infants were assessed for eligibility by a senior clinician. We reassessed eligibility at randomisation, study commencement, and before administration of morphine or placebo.

We obtained written informed parental consent for all infants. Approval was obtained from the Medicines and Healthcare products Regulatory Agency (MHRA) and Northampton Research Ethics Committee (15/EM/0310). This trial conformed to the standards set by the Declaration of Helsinki.

Randomisation and masking

We randomised infants to receive either morphine or placebo, using a web-based facility hosted by the NPEU CTU, with a minimisation algorithm to ensure
approximate balance of key demographics between the
groups (gestational age at birth, gestational age at time of
randomisation, intraterine growth restriction, time on
ventilation, time since morphine last given, presence of a
gastric tube, and history of surgery). Morphine sulphate
(at a concentration of 200 µg/mL) and placebo solutions
were indistinguishable by colour, odour, and flow, and
were dispensed in 10 mL glass amber bottles with tamper-
evident caps and a pack identification label (appendix).
Researchers, clinicians, outcome assessors, and parents
were masked to treatment allocation. In the event of an
emergency, treatment allocation could be unmasked by
a member of the clinical team using a single-use
access code on the randomisation website. In the event of
an infant becoming ineligible after randomisation, the
study was postponed and recommenced when the infant
became eligible, without rerandomisation.

Procedures
A study timeline is provided in the appendix. Continuous
electronic data capture of heart rate, respiratory rate, and
oxygen saturation began approximately 24 h before the
clinical procedure to establish a baseline of clinical
stability for every infant. We recorded all changes in
oxygen requirement, and measured blood pressure
every 6 h.

Approximately 60 min before the clinical procedure,
we gave infants one dose of either morphine sulphate
(100 µg/kg) or placebo (of equivalent volume). We
calculated the volume of the dose using the infant’s
working weight (the most recent weight in the
infant’s medical notes and used on their current drug
prescription chart). We administered the dose orally
(via syringe) or via a nasogastric tube (flushed with
aspirate). Mydriatic eye drops (tropicamide 1% and
phenylephrine 2·5%) were administered at 60 min and
again at 45 min before the clinical procedure.
Electroencephalography (EEG) and electromyography
(EMG) electrodes were then sited, as described in the
appendix.

Shortly before the clinical procedure, we swaddled the
infant (to provide non-pharmacological pain relief), began
video monitoring, and did a control heel lance (lancet was
rotated and held against the foot, with the blade released
into the air). This procedure was followed by the clinically
required heel lance, approximately 60 min post admini-
stration of morphine or placebo. No non-essential or
additional blood tests were done. After the heel lance and
blood collection, we ensured that infants were fully settled
and did not exhibit behavioural or physiological signs of
distress before the retinopathy of prematurity examination.
A senior ophthalmologist performed all retinopathy of
prematurity examinations. Topical local anaesthetic
(proxymetacaine 0·5%) eye drops were instilled before
insertion of an eyelid speculum, and binocular indirect
ophthalmoscopic examination was completed using a
Flynn style indenter.

After the clinical procedure, EEG and EMG leads were
removed and physiological recordings and documentation
of oxygen requirements continued for 24 h. A skilled
neonatal nurse or paediatric doctor from the research
team remained present for a minimum of 6 h after drug
administration.

Outcomes
The co-primary outcome measures were a behavioural
pain score calculated after retinopathy of prematurity
screening using the Premature Infant Pain Profile–
Revised (PIPP-R), and the magnitude of noxious-evoked
brain activity in response to heel lancing, measured using

Figure 1: Trial profile
EEG=electroencephalography. EMG=electromyography. PIPP-R=Premature Infant Pain Profile–Revised.
ROP=retinopathy of prematurity. *One excluded because of artifact.
We calculated the PIPP-R score in the 30 s period after retinopathy of prematurity screening (after removal of the speculum following examination of the second eye). A PIPP-R score of 6 or lower indicates little or no pain and a score greater than 12 indicates moderate-to-severe pain.19 Secondary outcome measures were reflex withdrawal and the PIPP-R score after heel lancing (appendix). To assess the nociceptive specificity of outcome measures, we also assessed the PIPP-R score, magnitude of noxious-evoked brain activity, and reflex withdrawal for the control heel lance. Background brain activity and reflex withdrawal activity were also assessed in a baseline period during which the infant’s foot was gently held but no stimuli were applied.

We assessed the clinical stability of infants by considering episodes of oxygen desaturation, bradycardia, tachycardia, and apnoea, and requirements for an increase in respiratory support, during the 6 h and 24 h periods after the clinical procedure, which were also secondary outcomes of the study. Episodes of oxygen desaturation were identified from the peripheral oxygen saturation signal as periods during which oxygen saturation fell below 80% for at least 10 s. Episodes of bradycardia were identified as periods during which the heart rate fell below 100 beats per min (bpm) for at least 15 s. Episodes of tachycardia were defined as periods during which the heart rate was greater than 200 bpm for at least 15 s (appendix). Apnoeic episodes were identified from clinical records or by retrospective review of the impedance pneumograph for breathing pauses longer than 20 s during bradycardic episodes (appendix). Increases in respiratory support were defined as a significant increase in oxygen requirement or an increase in respiratory support modality (appendix).

We assessed drug safety by calculating the incidence of apnoea requiring NIPPV and the incidence of hypotension requiring inotropes in the 24 h period post administration of morphine or placebo. Clinicians on the research team recorded a description of adverse events that occurred in the 24 h period post administration of morphine or placebo, including action taken, severity, and causality of the event, identified by consultation of the clinical team and review of clinical records (appendix). All serious adverse events were reviewed by the Data Monitoring Committee.

### Statistical analysis

We calculated that a sample size of 132 infants would allow detection of a clinically meaningful reduction in pain scores (a clinically significant reduction in PIPP-R scores was defined as 2 points),17 from a conservative post-retinopathy of prematurity screening mean PIPP-R score of 8·3 (SD 3·5) from a previous study,20 with power of 90% (p<0·05; two-tailed). We considered a 40% reduction in noxious-evoked brain activity clinically meaningful, since a similar reduction in adults corresponds to significantly lower verbally reported pain scores.19 A sample size of 132 infants was also required for this co-primary outcome measure, with power of 90% (p<0·05; two-tailed). We inflated the sample size to 156 infants (78 per arm) to account for

<table>
<thead>
<tr>
<th>Characteristics at birth</th>
<th>Morphine (n=15)</th>
<th>Placebo (n=15)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gestational age (weeks)**†</td>
<td>28·1 (26·3–30·1)</td>
<td>28·6 (27·9–29·7)</td>
</tr>
<tr>
<td>Birthweight (g)</td>
<td>1107 (329)</td>
<td>1173 (350)</td>
</tr>
<tr>
<td>Birthweight Z-score</td>
<td>-0·4 (0·9)</td>
<td>-0·2 (1·0)</td>
</tr>
<tr>
<td>Intrauterine growth restriction*</td>
<td>2 (13%)</td>
<td>3 (20%)</td>
</tr>
<tr>
<td>Apgar score at 10 min</td>
<td>10·0 (9·0–10·0)</td>
<td>10·0 (8·0–10·0)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Characteristics at time of randomisation</th>
<th>Morphine (n=15)</th>
<th>Placebo (n=15)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gestational age (weeks)**</td>
<td>34·7 (34·1–35·1)</td>
<td>34·7 (34·1–35·1)</td>
</tr>
<tr>
<td>Time on ventilation (days)*§</td>
<td>8·0 (1·0–20·0)</td>
<td>3·5 (2·0–19·5)</td>
</tr>
<tr>
<td>Time since morphine last given (days)*¶</td>
<td>46·5 (33·5–49·0)</td>
<td>19·0 (15·0–39·0)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Characteristics at time of clinical procedure</th>
<th>Morphine (n=15)</th>
<th>Placebo (n=15)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gestational age (weeks)†</td>
<td>35·0 (34·3–35·4)</td>
<td>34·9 (34·3–36·3)</td>
</tr>
<tr>
<td>Postnatal age (days)</td>
<td>50 (28–58)</td>
<td>49 (43–59)</td>
</tr>
<tr>
<td>Weight (g)</td>
<td>2049 (428)</td>
<td>2127 (331)</td>
</tr>
<tr>
<td>Duration of ROP screening (s)</td>
<td>97 (82–108)</td>
<td>91 (83–110)</td>
</tr>
<tr>
<td>Diagnosis of ROP</td>
<td>2 (13%)</td>
<td>2 (13%)</td>
</tr>
<tr>
<td>Level of care</td>
<td>2 (13%)</td>
<td>2 (13%)</td>
</tr>
<tr>
<td>Intensive care unit</td>
<td>1 (7%)</td>
<td>1 (7%)</td>
</tr>
<tr>
<td>High-dependency unit</td>
<td>5 (33%)</td>
<td>9 (60%)</td>
</tr>
<tr>
<td>Low-dependency unit</td>
<td>9 (60%)</td>
<td>5 (33%)</td>
</tr>
<tr>
<td>Respiratory support modality</td>
<td>1 (7%)</td>
<td>1 (7%)</td>
</tr>
<tr>
<td>High-flow oxygen therapy</td>
<td>9 (60%)</td>
<td>8 (53%)</td>
</tr>
<tr>
<td>Time between IMP administration and heel lance (min)</td>
<td>61 (57–66)</td>
<td>63 (58–70)</td>
</tr>
</tbody>
</table>

Data are median (IQR), mean (SD), or number (%). IMP = investigational medicinal product. ROP = retinopathy of prematurity. *Criteria used in minimisation algorithm for randomisation. †Postmenstrual age is often used in neonatal practice. In our unit, the infants’ gestational age is recorded each day in the medical and nursing notes; therefore, we have used this nomenclature. ‡Data are for six infants in the morphine group and 12 in the placebo group. §Data are for four infants in the morphine group who previously received morphine and seven in the placebo group. ¶Data are for four infants in the morphine group who previously received morphine and seven in the placebo group.

Table 1: Infant demographics

a validated EEG template (appendix).18 We calculated the PIPP-R score in the 30 s period after retinopathy of prematurity screening (after removal of the speculum following examination of the second eye). A PIPP-R score of 6 or lower indicates little or no pain and a score greater than 12 indicates moderate-to-severe pain.19

Secondary outcome measures were reflex withdrawal and the PIPP-R score after heel lancing (appendix). To assess the nociceptive specificity of outcome measures, we also assessed the PIPP-R score, magnitude of noxious-evoked brain activity, and reflex withdrawal for the control heel lance. Background brain activity and reflex withdrawal activity were also assessed in a baseline period during which the infant’s foot was gently held but no stimuli were applied.

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multiple births (assuming a 25% rate and an intra-cluster correlation coefficient of 0.5) and 10% loss to follow-up.

Trial outcomes were analysed and reported according to the trial protocol (version 6.0) and statistical analysis plan (version 2.0). Changes to the protocol (because of early trial cessation) are listed in the statistical analysis plan and the appendix. Analysis was per protocol, and a p value of 0.05 (two-sided 5% significance level) was deemed significant for all outcome measures.

We reported mean (SD) or median (IQR) values according to whether data were normally distributed or skewed. We calculated mean or median differences with 95% CIs and p values. We compared PIPP-R scores between groups using t tests. The magnitude of noxious-evoked brain activity and reflex withdrawal activity were compared between groups using a Wilcoxon rank-sum test, with the Hodges-Lehmann estimator used to calculate median differences with 95% CIs. Intra-rater reliability was 0.98 (95% CI 0.97–0.99) for heel lance PIPP-R scores and 0.97 (0.94–0.99) for retinopathy of prematurity screening. Inter-rater reliability was 0.98 (0.95–0.99) for heel lance and 0.89 (0.79–0.95) for retinopathy of prematurity screening.

For episodes of bradycardia, tachycardia, and oxygen desaturation, we standardised the difference in counts in the 6 h and 24 h periods before and after the clinical procedure, to adjust for the number of preprocedure episodes for every infant. We defined the standardised difference in number of episodes in the periods before and after the clinical procedure as the difference in number of episodes, as a proportion of the total number of episodes, for every infant, symmetrically in both the 6 h and 24 h periods (eg, in the 24 h period post procedure relative to the 24 h period preprocedure). To avoid issues caused by zero counts, we added a negligible constant term (0.01) to each count before standardisation. We compared standardised differences between treatment groups using a Wilcoxon rank-sum test, with the Hodges-Lehmann estimator used to calculate median
average time courses of physiological variables (heart rate, respiratory rate, and oxygen saturation) over the 48 h trial period and compared them between groups using non-parametric cluster analysis (appendix). We analysed data with Stata SE (version 13.1) and MATLAB (R2017a), and East (version 6.4) was used for the stopping boundary and safety analysis.

A trial stopping boundary was predefined based on the event rate of apnoeic episodes requiring resuscitative NIPPV [bag valve mask [visionary single patient use manual resuscitator; Marshall Airway Products, Radstock, UK] or Neopuff [Fisher & Paykel Healthcare, Auckland, New Zealand]]. The boundary was chosen by the Data Monitoring Committee before any analyses, after review of hypothetical trial scenarios and defined using a group sequential method with a one-sided gamma spending function (γ=4·5, type I error rate=0·2, estimated power=0·79). The selected boundary was based on a control group event rate of 7% and a difference between the group event rates of 12%. After 25 infants were randomised and studied, the Data Monitoring Committee convened for a safety review as planned and were provided with the stopping boundary graph, clinical stability and safety data summarised by arm, and detailed summaries of safety and adverse events, to consider evidence for benefit and harms.

This trial is registered with the European Clinical Trials Database (number 2014-003237-25).

Role of the funding source
The funder had no role in study design, data collection, data analysis, data interpretation, or writing of the report. The corresponding author had full access to all data in the study and had final responsibility for the decision to submit for publication.

Results
Between Oct 30, 2016, and Nov 17, 2017, 31 infants aged 34–39 weeks’ gestation (at the time of the study), who required retinopathy of prematurity screening and heel lancing on the same occasion, were enrolled and randomly assigned either 100 μg/kg of oral morphine (n=15) or placebo (n=16); one infant assigned placebo was withdrawn from the study before monitoring began (figure 1).

The Data Monitoring Committee reviewed safety data after recruitment of 25 infants (on Dec 4, 2017), and the predefined stopping boundary had been crossed. The Data Monitoring Committee requested data for all 31 infants who had been randomised. They concluded that the trial should not continue in its present form and recommended that the investigators review the data to guide

differences with 95% CIs. Infants with new-onset apnoea or an increased number of apnoeic episodes after the clinical procedure were compared using risk ratios (RRs). The number of infants requiring increased respiratory support after the clinical procedure was compared using risk differences (although we planned to use RR analysis according to the statistical analysis plan, this could not be done because no infants in the placebo group needed increased respiratory support). In a post-hoc analysis, we calculated the
Figure 4: Assessments of physiological stability

(A) Median (SE) of the standardised difference in number of episodes of desaturation in the 6 h period after the clinical procedure compared with the 6 h period before. (B) Median (SE) of the standardised difference in number of episodes of desaturation in the 24 h period after the clinical procedure compared with the 24 h period before. (C) Median (SE) of the standardised difference in number of episodes of bradycardia in the 6 h period after the clinical procedure compared with the 6 h period before. (D) Median (SE) of the standardised difference in number of episodes of bradycardia in the 24 h period after the clinical procedure compared with the 24 h period before. (E) Mean (SE) heart rate during the 48 h monitoring period. (F) Mean (SE) respiratory rate during the 48 h monitoring period. (G) Mean (SE) oxygen saturation during the 48 h monitoring period. (E–G) Individual infant traces are baseline-corrected to the average baseline across all infants. Time zero is the point of the clinical procedure. Grey vertical dashed line indicates the time of administration of morphine or placebo. Grey boxes indicate periods during which the treatment groups differed significantly.

Further investigation. The trial was stopped by the Central Monitoring Team on March 15, 2018 (appendix).

There were no deviations from exclusion or inclusion criteria. Infant demographics and clinical characteristics at birth and at the time of intervention are reported in table 1 according to group allocation. Most baseline characteristics were well balanced between treatment groups; however, a few variables were less well balanced (eg, the number of infants who required ventilatory support), which is expected in view of the small sample size. Figure 2 shows example data for one infant for each recording modality.

The co-primary outcome measures used to assess morphine analgesic efficacy were PIPP-R score after the heel lance and the magnitude of reflex withdrawal activity evoked by the heel lance did not differ between the two groups. Mean PIPP-R score after heel lance was 7·9 (SD 3·4) with morphine and 8·5 (3·9) with placebo (mean difference –0·6, 95% CI –3·3 to 2·1; p=0·66).

The magnitude of reflex withdrawal was median 22·4 (IQR 19·7–44·8) with morphine and 12·4 (6·1–46·3) with placebo (median difference 8·9, 95% CI –12·0 to 22·4; p=0·48; figure 3). The magnitude of each pain-related outcome measure increased significantly after the clinical procedure compared with control stimuli and non-noxious background activity (appendix), showing that the measures were discriminative and appropriate for assessing analgesic efficacy in this population.

The clinical stability of each infant was assessed over 48 h (24 h before and after the clinical procedure). Infants assigned morphine had significantly more episodes of oxygen desaturation in the 6 h and 24 h periods after the clinical procedure, and significantly more episodes of bradycardia in the 24 h period after the clinical procedure, compared with those allocated placebo (table 2; figure 4). There were no differences in the number of episodes of tachycardia.

Eight (53%) of 15 infants who received morphine developed new-onset apnoea or an increase in the number of apnoeic episodes in the 24 h period after the clinical procedure compared with control stimuli and non-noxious background activity (appendix), showing that the measures were discriminative and appropriate for assessing analgesic efficacy in this population.
Table 3: Adverse events

<table>
<thead>
<tr>
<th>Onset of event post drug (h, min)</th>
<th>Treatment</th>
<th>Grade</th>
<th>Attribution</th>
<th>Allocation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nasal congestion</td>
<td>11 h, 56 min</td>
<td>Saline drops</td>
<td>Mild</td>
<td>Not related</td>
</tr>
<tr>
<td>Rash</td>
<td>4 h, 4 min</td>
<td>Cream</td>
<td>Mild</td>
<td>Not related</td>
</tr>
<tr>
<td>Profound desaturation episodes</td>
<td>17 h, 59 min</td>
<td>Facial oxygen</td>
<td>Mild</td>
<td>Not related</td>
</tr>
<tr>
<td>Recurrent desaturation episodes</td>
<td>8 h, 9 min</td>
<td>Stimulation</td>
<td>Mild</td>
<td>Possibly</td>
</tr>
<tr>
<td>Recurrent desaturation episodes</td>
<td>1 h, 58 min</td>
<td>Facial oxygen</td>
<td>Mild</td>
<td>Possibly</td>
</tr>
<tr>
<td>Apnoea</td>
<td>2 h, 13 min</td>
<td>NIPPV; increase high-flow oxygen</td>
<td>Moderate</td>
<td>Possibly</td>
</tr>
<tr>
<td>Recurrent apnoeic episodes</td>
<td>2 h, 39 min</td>
<td>Stimulation; increase low-flow oxygen</td>
<td>Moderate</td>
<td>Possibly</td>
</tr>
<tr>
<td>Recurrent apnoeic episodes</td>
<td>1 h, 28 min</td>
<td>Stimulation (× 3); NIPPV (× 3)</td>
<td>Moderate</td>
<td>Possibly</td>
</tr>
<tr>
<td>Recurrent desaturation, bradycardia, and apnoeic episodes</td>
<td>2 h, 3 min</td>
<td>Commenced high-flow oxygen; feed volume reduction</td>
<td>Moderate</td>
<td>Possibly</td>
</tr>
</tbody>
</table>

Serious adverse events

| Persistent hypoventilation and desaturation | 6 h, 0 min | Moved to high-dependency unit; commenced high-flow oxygen | Moderate | Possibly | Morphine |
| Recurrent apnoeic episodes | 6 h, 24 min | Unmasked by clinical team; moved to high-dependency unit; commenced high-flow oxygen; naloxone (× 2) | Moderate | Probably | Morphine |

Adverse events are shown that occurred during the 24 h period post administration of morphine or placebo. NIPPV=non-invasive positive pressure ventilation.

Discussion

This trial shows that administration of 100 μg/kg oral morphine before acutely painful clinical procedures in infants born prematurely (and aged 34–39 weeks’ gestation at study) has a profound negative effect on respiratory stability, without any suggestion of analgesic
efficacy (video). A multimodal approach was used to assess analgesic efficacy and safety, providing detailed evidence of the effects of morphine on infant nociceptive and physiological activity. Behavioural pain scores, noxious-evoked brain activity, and reflex withdrawal activity did not differ between morphine and placebo groups. The study was underpowered for the co-primary outcome measures, because of early trial cessation, and we therefore cannot conclude whether morphine provided effective analgesia at this dose. However, a trend was noted across modalities that infants who received morphine had greater noxious-evoked activity, suggesting that even if the trial had continued to completion, we would have been unlikely to observe an analgesic effect of morphine.

This trial suggests that oral morphine at a dose of 100 μg/kg in non-ventilated infants has the potential for harm. The profound respiratory effects observed justified trial cessation and lead us to recommend that oral morphine (at this dose) should not be given to non-ventilated premature infants for acute pain relief during retinopathy of prematurity screening. The age range in this study was restricted to infants requiring retinopathy of prematurity screening at 34–42 weeks' gestation, so we cannot ascertain the effects of oral morphine in younger or older infants. However, international paediatric formularies—eg, the British National Formulary for children—recommend an oral dose of 50–100 μg/kg every 4 h in infants aged 1–2 months, and 50 μg/kg of intravenous morphine every 6 h to treat pain in neonates, which is roughly equivalent to the 100 μg/kg oral dose administered in this study, assuming an oral bioavailability of approximately 50%.22 Our data suggest that in non-ventilated premature infants of 1–2 months' postnatal age, these doses could cause substantial respiratory adverse effects, requiring resuscitative respiratory support or a change in respiratory support modality. This effect might be due to immature glucuronidation and reduced clearance of morphine metabolites, because these processes are not mature in the first 2 months of life.21

Although an intravenous morphine dose of 10–30 μg/kg provided effective analgesia in infants receiving continuous positive airway pressure in a previous study,26 severe apnoeic episodes requiring substantial intervention were reported in 9% of participants (who were very premature), and consistent with the findings of our study, a reduction in heart rate and respiratory rate was suggested. Similarly, a dose of 100 μg/kg of intravenous morphine in ventilated premature infants provided effective pain relief for central line placement in a previous study,23 but with significantly increased ventilation requirements compared with infants who received tetracaine. Findings of another retrospective study24 showed that five of 43 infants who received approximately 50–100 μg/kg of intravenous morphine for central line placement had respiratory depression requiring intervention or increased respiratory support, compared with none in a control group.

It is possible that if we had lowered the dose of oral morphine that the adverse respiratory outcomes could have been reduced. However, it seems unlikely that a lower dose would have provided effective analgesia. Although our national drug formulary recommends 50–100 μg/kg of morphine orally for pain, neonatal drug guidelines from other countries (eg, Australia) recommend higher oral doses of 100–200 μg/kg for pain in neonates.25 Our local practice guidelines also recommend that an oral dose of 100 μg/kg is given to infants requiring laser eye surgery. In a pilot trial that was started but not completed,26 six non-ventilated infants were administered a much larger oral dose of morphine (200 μg/kg) for retinopathy of prematurity screening. These researchers did not report any adverse effects. We therefore determined that, on balance, 100 μg/kg was a justifiable dose.

The acceptable balance of benefit and harm for any treatment is contextually dependent. All infants in our trial were clinically stable before the study started, and most infants who received morphine were cared for in a low-dependency setting and self-ventilating in air. Although ventilatory support can be routinely and expertly provided in neonatal care, escalation of oxygen therapy or level of care can be costly, result in prolongation of admission and distress to parents, and be viewed as a considerable setback. Ideally, morphine should be titrated to provide patients with optimum analgesic benefit and minimum adverse effects. The likely requirement for increased respiratory support would need to be expected, manageable, and justified by analgesic benefits. In our study, we noted considerable adverse effects before any suggestion of benefit, suggesting a non-existent therapeutic window in this context. Doses of morphine that are routinely administered intravenously to ventilated infants27 might provide effective analgesia in premature infants, albeit limited analgesic efficacy has been reported.3 Because respiratory adverse effects can be managed well in ventilated infants, the benefit of morphine administration for sedation might outweigh the risks, and results from this study cannot be interpreted to suggest that ventilated infants should not be given morphine.

Controversy remains over the analgesic efficacy of intravenous morphine for procedural pain in ventilated infants,1 and comprehensive assessment of pain-related brain activity could help settle this debate. The single bolus dose of morphine administered in our trial produced clinically significant cardiorespiratory effects; the heart rate and respiratory rate were significantly lower in infants assigned morphine for 6–8 h after the clinical procedure. This finding corresponds with the half-life of morphine in premature infants.28 A comparable degree of cardiorespiratory depression has been seen in ventilated infants receiving intravenous morphine infusions.29 This finding highlights the importance of comprehensively evaluating the side-effects of pain-relieving drugs through detailed physiological recordings and clinical observations.
Due to early trial cessation, our study was underpowered to detect significant effects in the co-primary outcome measures. Although mydriatic eye drops were given at approximately the same time as the drug, which might delay gastric emptying, this does not seem to have prevented morphine absorption. A limitation of our study design was that timing of peak analgesic efficacy of oral morphine is unknown; however, the time courses of cardiac and respiratory effects suggest a drug effect at the time of the clinical procedure. It is unlikely that an absence of analgesic efficacy is related to a lack of absorption or poor timing of drug administration.

In view of the challenges in measuring analgesic efficacy in non-verbal infants, the recorded increases in the measures of noxious-evoked brain activity, reflex withdrawal, and PIPP-R scores in response to the clinical procedure confirm the suitability of these approaches to assess analgesic efficacy. The multimodal approach used in our trial to assess both analgesic efficacy and drug safety can provide detailed mechanistic insight into the precise time course of physiological effects of potential analgesics. Although this multimodal methodology cannot be implemented easily into standard clinical practice, this work highlights the importance and feasibility of using this approach in future clinical trials of analgesics in infants.

Our trial provides further evidence that retinopathy of prematurity screening is a painful and destabilising procedure. In a meta-analysis of pain relief for retinopathy of prematurity screening, topical anaesthetic the median PIPP score was 15, reducing to 11 with the addition of sweet taste. In our placebo group, the mean PIPP median PIPP score was 15, reducing to 11 with the addition of sweet taste. This finding corresponds with previous work.

In conclusion, oral morphine at a dose of 100 μg/kg has the potential for harm with no suggestion that it provides analgesic efficacy for acute clinical procedures in non-ventilated infants born prematurely (and aged 34–39 weeks’ gestation at study). However, because of early trial cessation, we cannot draw conclusions about the analgesic efficacy of oral morphine; obtaining a large enough sample size to test this objective would have required exposing infants to an unacceptable risk of respiratory adverse events. We do not recommend this dose of morphine for use as pain relief during binocular indirect ophthalmoscopy retinopathy of prematurity screening. Using multimodal outcome measures, along with detailed physiological recordings, provides a rigorous approach to assess analgesic efficacy and adverse effects, leading to a greater mechanistic understanding of the effects of a drug, and is desirable in future clinical trials of analgesics in infants.

Contributors
RS conceived the idea for the study. EA was the chief investigator and RS was the principal investigator. Funding proposals were developed by RS, EA, EJ, JEN, RR, and CH. The study was designed by RS, CH, FM, AH, GG, EJ, JEN, RR, and EA. VM was the trial manager. JLB was the trial statistician. ARK created the data collection forms and programmed the trial database. Participant recruitment was undertaken by RS, AH, FM, GG, and CH. Data collection was done by RS, CH, FM, GG, AH, MvdV, DG, SG, and CP. Retinopathy of prematurity screening was done by CP, EA, AH, FM, and GG provided clinical care for the participants during the trial. EA, FM, and GG reported adverse events. Data entry was completed by RS, CH, FM, JLB, DG, and AH. RS, CH, and FM wrote the first draft of the manuscript. The manuscript was critically reviewed by all authors.

Declaration of interests
We declare no competing interests.

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