Effect of non-sucrose sweet tasting solution on neonatal heel prick responses

Luca A Ramenghi, Gillian C Griffith, Christopher M Wood, Malcolm I Levene

Abstract
A substance commercially described as 'sugar free,' used as a sweetener for paracetamol suspension, was evaluated on measures of neonatal pain. Sixty infants were randomly allocated to receive one of four solutions before heel stab blood sampling: sterile water (placebo); 25 or 50% sucrose (weight/volume); and the commercial sweet-tasting solution. There was a significant reduction in crying time and pain score 3 minutes after the painful stimulus in all groups compared with the controls.

It is concluded that this sweet-tasting solution has analgesic effects as potent as those of concentrated sucroce solutions. (Arch Dis Child 1996; 74: F129–F131)

Keywords: analgesia, sweet-tasting solution, sucrose, behavioural response, heel prick.

All healthy newborn infants born in Britain are exposed to iatrogenic painful procedures. We estimate that 15% of term newborns undergo an additional two to five heel prick sampling procedures in the first week of life. These painful procedures may be considered minor, but recent research has suggested that babies' early pain experience may affect adversely their pain response in later infancy.

We have recently reported that concentrated sucrose solution significantly reduces the duration of cry in newborn babies when given immediately before a heel prick blood sampling for serum bilirubin and this may be mediated by release of endogenous opiates. The use of concentrated sucrose may be criticised in light of its high toxicity and its cariogenic effect on erupted teeth. We therefore investigated a sweet-tasting solution commercially described as 'sugar free' and used as a vehicle for paracetamol (Calpol) for its effect on reducing pain response following heel prick blood sampling.

Method
Healthy term newborn infants who required

heel prick blood sampling for serum bilirubin estimations were recruited. Criteria for inclusion were: birthweight above 2500 g; gestation 37–42 weeks; Apgar score of 7–10 at 5 minutes and no previous exposure to naloxone.

Infants were fully clothed apart from the foot which was used for sampling. Before skin preparation a pulse oximeter was applied to the baby's hand or contralateral foot to monitor heart rate throughout the study period. Test solution (2 ml) was syringed into the baby's mouth for 1 minute. The solution was applied mainly to the anterior part of the tongue to best promote taste perception. Two minutes after beginning to administer the solution the heel prick was performed by licking and gently squeezing the heel which had been cleaned with a sterile swab. Because the method of performing blood samples may affect the nociceptive response, all heel pricks were inflicted by one experienced nurse standardising the procedure and recording the time spent squeezing. Changes in four facial expressions (brow bulge, eye squeeze, nasolabial furrow and open mouth) and the presence of crying related to the heel prick were recorded by the same observer at −2, −1, 0, 1, 2, 3, and 5 minutes on a 0–5 scale, giving a score of 1 if present and 0 if absent to each item for each criterion. The baby's behavioural state was scored before heel lancing.

Crying during sampling and in the 3 minutes after heel prick were recorded on to a tape recorder. The first cry was defined as the duration of audible distressed vocalisations with a continuous pattern before a quiet interval of 5 seconds from the time of the painful stimulus.

Babies were allocated at random to receive one of four solutions: sterile water (control); 25 or 50% sucrose (weight/volume); and the sweet-tasting commercial Calpol solution without paracetamol (Burroughs Wellcome). This contains Lycasin (hydrogenated glucose syrup) as the sweetening agent in a 40% w/v solution. Fifteen babies were required in each group to achieve the 80% power necessary to show a 50% reduction in crying time (P=0.05).

Investigators were blind to the nature of the

Table 1 Details of neonates studied and changes in pain score in response to heel lance

<table>
<thead>
<tr>
<th>Solution</th>
<th>Median gestational age (weeks) (range)</th>
<th>Median postnatal age (days) (range)</th>
<th>Median arousal state score (0–5) (range)</th>
<th>Median time from last feed (minutes) (range)</th>
<th>Median time spent squashing heel (seconds) (range)</th>
<th>Pain score 1 minute before heel prick (range)</th>
<th>Pain score at time of heel prick (range)</th>
<th>Pain score 3 minutes after heel prick (range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>50% Sucrose</td>
<td>15 (38–40)</td>
<td>3 (2–4)</td>
<td>2 (1–3)</td>
<td>45 (30–80)</td>
<td>80 (70–115)</td>
<td>0 (0–0)</td>
<td>3 (3–5)</td>
<td>0 (0–3)</td>
</tr>
<tr>
<td>Water (control)</td>
<td>15 (38–40)</td>
<td>3 (2–4)</td>
<td>1 (1–3)</td>
<td>90 (20–150)</td>
<td>90 (50–148)</td>
<td>2 (1–3)</td>
<td>4 (3–5)</td>
<td>2 (1–5)</td>
</tr>
<tr>
<td>25% Sucrose</td>
<td>15 (38–40)</td>
<td>4 (3–5)</td>
<td>1 (1–3)</td>
<td>90 (70–198)</td>
<td>85 (60–98)</td>
<td>0 (0–0)</td>
<td>4 (3–5)</td>
<td>0 (0–2)</td>
</tr>
<tr>
<td>Calpol solution</td>
<td>15 (39–40)</td>
<td>3 (2–3)</td>
<td>3 (0–3)</td>
<td>60 (45–120)</td>
<td>90 (68–108)</td>
<td>0 (0–0)</td>
<td>4 (3–5)</td>
<td>0 (0–1)</td>
</tr>
</tbody>
</table>
control and sucrose solutions, but the Calpol vehicle could not be disguised because of its pink colour. The recorded cry of each baby was later analysed without knowledge of the solution group.

Parents gave informed and signed consent and were present if they wished, but avoided interaction with their infant during the study.

The study was approved by the research ethics board of Leeds General Infirmary.

Results were analysed using the Mann-Whitney U test (Astute Program).

**Results**

There were no significant differences between the groups in sex, gestational age, postnatal age, mode of delivery, behavioural state at rest and time spent squeezing the heel (table 1). Changes in the babies' pain scores in each group are shown in table 1. The median crying time and the median duration of first cry are illustrated in fig 1. Figure 2 shows the duration of crying in the first 3 minutes after heel prick in each treatment group. Heart rate changes are illustrated in fig 3.

There was a significant reduction in both the duration of first cry and the percentage crying time during the first 3 minutes in all groups (25 and 50% sucrose; Calpol vehicle solution) compared with controls (P=0.02).

The pain score was significantly higher in the placebo group compared with the other groups 3 minutes after (P=0.05) and 1 minute before lancing (P=0.04).

There was a significant increase in heart rate between 3 minutes after the painful procedure in the control group compared with the 50% sucrose and Calpol vehicle solution group (P=0.009).

**Discussion**

Cry is still considered the most sensitive behavioural response to a noxious stimulus in neonates.9 We found that neonates who had been given oral placebo (sterile water) before heel pricks had significantly longer crying time than babies given a small volume of concentrated sucrose and a sweet-tasting, non-sucrose solution. The significant reduction in heart rate and pain score 3 minutes after heel prick in the babies receiving the Calpol vehicle solution as well as concentrated sucrose solution tends to confirm the analgesic effects of these forms of treatment.

The major finding of this study is to extend to a non-sucrose sweet-tasting solution our recent observation concerning the apparent analgesic effects of concentrated sucrose solutions.1 We had subjectively observed that young infants given Calpol solution for perceived pain or fever rapidly settled within seconds of its administration. This action is too rapid to be attributable to the analgesic effects of the paracetamol. We were intrigued to discover whether the sweet, non-sucrose vehicle of Calpol solution may have independent effects on pain.

The sweet taste of Calpol solution is achieved by hydrogenated glucose. The manufacturers of Calpol describe their solution as 'sugar free' because it does not contain sucrose. The speed of analgesic response of the Calpol solution is consistent with a preabsorptive mechanism which stimulates endogenous opiate release, as has been proposed as an explanation for the analgesic effects of sucrose.3 10-12

The reduction in pain perception in the neonate may be achieved by a variety of effects with possibly cumulative action. Previous studies have shown that non-nutritive sucking on a pacifier attenuates behavioural distress during heel stick procedures.13

Newborns' behaviour may often be influenced by maternal contact. We have chosen to investigate the effect of a sweet solution in the

---

**Figure 1** Duration of first crying after heel prick in 60 babies given sterile water (controls), 25% or 50% sucrose, and non-sucrose sweet-tasting Calpol solution. Points are individual values. Horizontal lines and boxes represent median values and interquartile ranges.

**Figure 2** Percentage time spent crying in the first 3 minutes after heel prick in the 60 babies grouped as in fig 1.
absence of close maternal contact in an attempt to tease out these factors. If mothers wished to cuddle their baby during or immediately after the procedure then the baby was not enrolled in the study. Very few mothers expressed a wish to do this. Sweet solutions stimulate sucking and we are undertaking further studies to investigate whether there is an underlying reinforcement between non-nutritive sucking and the effect of sweet substances on behavioural response to the heel prick stimulus.

LR was supported by the Special Trustees of the Leeds General Infirmary. We are grateful to Burroughs Wellcome for supplying the Calpol solution.

Effect of non-sucrose sweet tasting solution on neonatal heel prick responses.

L. A. Ramenghi, G. C. Griffith, C. M. Wood and M. I. Levene

Arch Dis Child Fetal Neonatal Ed 1996 74: F129-F131
doi: 10.1136/fn.74.2.F129

Updated information and services can be found at:
http://fn.bmj.com/content/74/2/F129

These include:

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/