Interface leakage during neonatal CPAP treatment: a randomised, cross-over trial

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ABSTRACT
Objective To determine leakage for two neonatal continuous positive airway pressure (CPAP) interfaces and evaluate leak-corrective manoeuvres.
Design The ToNIL (Trial of NCPAP Interface Leakage) study was a randomised, clinical, cross-over trial with data collection between August 2018 and October 2019. The primary outcome was blinded to the treating staff.
Setting One secondary, 8-bed neonatal intensive care unit (NICU) and three larger (>15 beds), academic NICU referral centres.
Patients Newborns with CPAP were screened (n=73), and those with stable spontaneous breathing, low oxygen requirement, postmenstrual age (PMA) over 28 weeks and no comorbidities were eligible. In total, 50 infants were included (median PMA 33 completed weeks).
Interventions Leakage was measured for both prongs and nasal mask, before and after leak-corrective manoeuvres. Interface application was performed in a randomised order by a nurse, blinded to the measured leakage.
Main outcome measures 30 s average leakage, measured in litres per minute (LPM).
Results Analyses showed a significantly lower leakage (mean difference 0.86 LPM, 95% CI 0.07 to 1.65) with prongs (median 2.01 LPM, IQR 1.00–2.80) than nasal mask (median 2.45 LPM, IQR 0.99–5.11). Leak-corrective manoeuvres reduced leakage significantly for both prongs (median 1.22 LPM, IQR 0.54–1.87) and nasal mask (median 2.35 LPM, IQR 0.76–4.75).
Conclusions Large leakages were common for both interfaces, less with prongs. Simple care manoeuvres reduced leakage for both interfaces. This is the first report of absolute leakage for nasal interfaces and should encourage further studies on leakage during CPAP treatment.

INTRODUCTION
The beneficial effects of continuous positive airway pressure (CPAP) for newborns in respiratory distress have been thoroughly confirmed since its clinical introduction. CPAP with short nasal prongs or nasal mask is currently the recommended first-line treatment for neonates in respiratory distress.
Several randomised trials have compared mask and prong interfaces using CPAP failure and nasal injuries as primary outcomes. Three recent systematic reviews favour nasal masks for both of these outcomes. The heterogeneity of the included studies, and the different CPAP interfaces used, was recognised in all reviews. Nasal trauma has been described with both interfaces, most commonly at the philtrum or glabella with nasal mask and at the septum with prongs. The rotational use of prongs or nasal mask to minimise trauma has not shown to reduce injuries.
The reason for CPAP failure reduction with nasal masks is not fully understood. Suggested explanations were that masks have less resistance and provide more stable CPAP. An increased resistance, together with a leakage, leads to a reduction in pharyngeal pressure, which was another suggested explanation by Kieran et al. To achieve an efficient delivery of CPAP, a tight fit without leakage has been assumed to be of importance. This might increase the risk of nasal trauma, a conflict well recognised. The link between leakage and outcome type, design and other factors has not been well studied.
The suggested method for reporting leakage by the European Respiratory Society/American Thoracic Society task force is a ratio between expiratory and inspiratory volumes. Previous CPAP studies reported that high levels of leakage were incompatible with determining tidal volumes and.
breathing patterns and resulted in high exclusion rates. In 2009, Schmalisch et al. suggested that a solution would be to report leakage flow rather than tidal volume ratios. Our Trial of NCPAP Interface Leakage (ToNIL) used improved methods allowing measurements of high leakage levels and was a randomised, clinical, two-period cross-over study on leakage during CPAP with nasal prongs and nasal mask. Our primary aim was to compare absolute leakage for prongs and nasal mask. Our secondary aim was to evaluate if leakage could be reduced by common care manoeuvres.

METHODS

Study design

The cross-over design was chosen to use each patient as its own control thereby reducing the number of infants needed for the study. The risk for carry-over effects was considered negligible.

Study population

Patients were included between August 2018 and October 2019 at four Swedish sites: an 8-bed neonatal intensive care unit at the Östersund hospital and three larger (>15 beds) tertiary referral centres at the Karolinska University Hospital, Stockholm.

Whenever investigators were available, newborn infants with CPAP were screened for eligibility. Infants older than 28 weeks of postmenstrual age (PMA) with stable spontaneous breathing were included after informed parental consent. Exclusion criteria were cardiac or respiratory malformations, facial defects or injuries, FiO2 ≥0.5, circulatory instability, recent surgery or recently extubated.

The starting interface was randomised using sequentially numbered, sealed envelopes, opened when the system was calibrated for measuring. The envelopes were prepared by a researcher with no connection to the study, and the 1:1 sequence was generated by sealedenvelope.com, stratified on age (<34 weeks or ≥34 weeks) and presence of nasogastric tube using permuted blocks of variable sizes.

The study was ethically approved, conducted according to good clinical practice (GCP) standards and registered at the US National Institutes of Health (ClinicalTrials.gov #NCT03586856). No changes to the study design, eligibility criteria or trial outcomes were made after commencement.

Study interventions

A schematic equipment setup figure is provided as online supplemental figure 1. The ordinary flow generator was replaced with an oxygen mixer and a rotameter to avoid problems with gas supply and potential zeroing discrepancies. Flow and pressure was recorded for 30 s, labelled, stored and processed using customised software (LabVIEW 2015; National Instruments). Recordings were labelled, stored and processed using customised software (LabVIEW 2015; National Instruments).

At 500 Hz using single-board microcontrollers (Arduino Uno, Arduino, Italy) and a data acquisition device (NI-6451; National Instruments, Austin, TX, USA). Recordings were labelled, stored and processed using customised software (LabVIEW 2015; National Instruments).

For each study, data were collected with the infant in an incubator, cot or during skin-to-skin care and at least 30 min after feeding (if not continuous). Parents were invited to be present and comfort the infant. A nurse applied the randomised first interface with a size of her or his choice, blinded to leakage measurements at all times.

Flow and pressure was recorded for 30 s after securing the interface and the infant was breathing quietly. If the measured leakage exceeded 0.2 LPM, the nurse was asked to leave before corrective manoeuvres were attempted by the investigators, guided by the measured leakage. Not all manoeuvres were tested on each infant and selection was based on previous attempts and experience. Each manoeuvre was evaluated with a 30 s recording until leakage was below 0.2 LPM or no further reduction was possible. The same procedure was then repeated with the second interface. No wash-out period was included as no carry-over effects were expected.

Sample size and statistical analyses

No sample size calculations were possible as absolute leakage during neonatal CPAP treatment has not been previously measured and published. A study sample size of 50 infants was chosen, based on the sizes of previous, experimental neonatal CPAP cross-over trials.

Data were imported to SPSS V.26 and analyses were both parametric (independent samples and paired samples t-test) and non-parametric (Wilcoxon signed-rank test and Mann-Whitney U).

In the regression analyses, PMA was removed due to co-linearity with weight and the stratification on nasogastric tube was removed due to a highly asymmetrical distribution. Univariable linear regression analyses were performed on leakage for each interface against the remaining strata, weight and CPAP level. A p value below 0.05 was considered statistically significant.

RESULTS

Seventy-three infants were screened for eligibility and 52 were enrolled after consent (figure 1). Due to equipment problems, two infants were never measured. One was never randomised, and one was randomised but not measured. All analyses included the remaining 50 study participants, equally randomised to start with either patient interface (table 1).

Measurements were well tolerated with no adverse events or need to increase the oxygen inspiratory fraction during the study. For each infant, the median time connected to the measuring equipment was 25 min (IQR 21–30). Analyses showed no significant carry-over or period effect (not reported).

There was a significantly lower leakage (mean difference 0.86 LPM, 95% CI 0.07 to 1.65, p = 0.034) with nasal prongs (median 2.01 LPM, IQR 1.00–2.80) than nasal mask (median 2.45 LPM, IQR 0.99–5.11). Leaks ranged from none to 9.9 litres per minute (LPM). A leakage above 0.2 LPM was present with at least one interface in all patients (figure 2). Regression analyses showed that leakage was associated to the set CPAP level for both prongs (β 0.78, 95% CI 0.15 to 1.41, p = 0.02) and nasal mask (β 1.17, 95% CI 0.13 to 2.20, p = 0.03), but was not significantly associated to the PMA stratification or weight.
With an initial leakage above 0.2 LPM, attempted non-blinded leak-corrective manoeuvres were successful in 96% of the cases with prongs (45/47) and 98% with nasal mask (48/49). The median reduction in leakage was 1.22 LPM (IQR 0.54–1.87) for prongs and 2.35 LPM (IQR 0.76–4.75) for nasal mask, both statistically significant (p<0.001) (figure 3).

DISCUSSION
As far as we know, this is the first study that reports absolute CPAP interface leakage measured in litres per minute. We found that after applying a CPAP interface (prongs or mask) in a randomised order, followed by a cross-over, less leakage was seen with prongs compared with nasal mask. The amount of leakage could be reduced by simple care manoeuvres for both interfaces when guided by the measured leakage. For each patient, leakage was rarely the same for both interfaces (dashed line figure 2) which indicates that leakage was related to the type of interface.

Our results confirm that interface leakages are common, often high and consistent with the high exclusion rates due to leakages reported in previous studies: Hückstädt et al excluded 49 of 69 patients with an excessive leakage (1.3 to 2.0 L/min) and Fischer et al reported that 31 of 32 infants displayed leakages exceeding what was possible to measure (>1.4 L/min). Our median leakage was higher than their equipment limits and could explain their high exclusion rates.

Leakage is a suggested factor that may affect CPAP quality and clinical outcome. Other potentially important factors are CPAP level, imposed work of breathing and pressure oscillations present in bubble CPAP systems. Little is known about how these factors interact and their role in optimising CPAP treatment.

Recent meta-analyses on clinical outcome measures, such as CPAP failure, have indicated that nasal mask is more favourable compared with nasal prongs, but the reasons have not yet been fully explained. Our study measured leakage over a short period of time in stable infants with a minimal risk of failure. Relating our findings to results in previous clinical trials with CPAP failure as an outcome is difficult. If there is a link between leakage and clinical outcome, it could be indirect through reductions in delivered pressure, pressure stability, imposed work of breathing or related to leakage compensation when using more complex CPAP drivers. This is speculative since leakage flow has, as far as we know, not been measured in any trials with clinical outcomes.

Leak-corrective manoeuvres
Leakage location cannot be determined from our measurements but can occur at the patient interface, through the mouth or to the stomach. Reducing leakage with simple manoeuvres was possible in most infants (figure 4). Even if it was easy to reduce leakage, we considered it not possible to go through a full list of standardised adjustments and compare their relative effectiveness.

The most successful leak-corrective manoeuvre was adjusting the seal of the interface and was achieved by adjusting the angle, straps, bonnet or manually changing the shape of the interface.
Manual mouth closure during measurements and adding or removing pacifiers was tested in some, but not all, infants. Mouth closure was the third most successful manoeuvre. Early studies stated that neonates are obligate nose breathers, but later results show that neonates are able to breathe orally but prefer to breathe through the nose. The effect of oral breathing on CPAP treatment and leakage is not known. In a calm infant with closed mouth, inserting a pacifier was not feasible. No definitive conclusions can be drawn from the data on pacifier addition or removal.

In our setting, care was based on the advice that prongs should fill the nares comfortably and a mask with a snug fit was preferred. In the majority of patients, the most appropriate interface size was already selected which explains the low number of attempts with changing interface sizes. When attempted, leakage reduction by interface size change was more successful with prongs than mask.

Evaluating CPAP leakage in clinical practice includes listening to the sound the device makes, looking at the compensatory increase in driver flow and evaluating the clinical effect over time. These techniques are subjective, time consuming and seem outdated in modern intensive care of newborn infants. New tools for measuring and controlling leakage could be of great value, possibly contributing to less CPAP failure, minimising the need for mechanical ventilation and subsequently lead to less chronic lung disease.

Limitations

All infants were treated with the original Infant Flow device design. The results would likely be different if other types of CPAP devices or interfaces were used.

The time on each interface was short and leakage over time or during feeding and care was not investigated. Conclusions on long-term leakages cannot be made. The current method did not allow for identification of leakages split on exhalation and inhalation. Theoretically, a pressure-unstable system carries an increased risk for leakages during exhalation.

The second part of the study, exploring ways to reduce leakage, was not randomised and not all infants underwent the same interventions. This limits the possibility to draw any conclusions about the relative effect of each intervention but shows that a reduction was possible in most infants. The second part of the study should be considered hypothesis generating.

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Future directions

Our study method could be useful for researchers and possibly also for clinical use. We suggest that leakage should be considered a variable affecting treatment outcome and, together with interface type, resistance to breathing and CPAP level, could add information in both animal and human research trials.

Leakage monitoring during non-invasive support might also allow optimisation of nebulised drugs, including surfactant. As high levels of leakage will likely lead to the dilution of aerosols. Our findings suggest that a better fit, especially together with a nasogastric tube, might reduce leakage. If a reduced leakage improves the clinical outcome remains an open question.

Future studies should address the two major limitations of our trial: short registration time and measurements on stable infants only. To be clinically meaningful, leakage needs to be studied over longer periods, in unstable infants and linked to delivered CPAP and clinical outcome.

CONCLUSION

This is the first report on absolute leakage for CPAP interfaces. In our randomised, short-duration, cross-over trial, nasal prongs had less leakage than nasal mask. High levels of leakage were common for both interfaces. When guided by measured leakage, simple manoeuvres could reduce leakage. The findings encourage studies on interface design and clinical effects of optimising leakage.

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Deidentified participant data and full trial protocol available from corresponding author to researchers who provide a methodologically sound proposal. Reuse of data permitted to achieve aims in the approved proposal.

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