A multicentre, randomised controlled, non-inferiority trial, comparing high flow therapy with nasal continuous positive airway pressure as primary support for preterm infants with respiratory distress (the HIPSTER trial): study protocol

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ABSTRACT

Introduction: High flow (HF) therapy is an increasingly popular mode of non-invasive respiratory support for preterm infants. While there is now evidence to support the use of HF to reduce extubation failure, there have been no appropriately designed and powered studies to assess the use of HF as primary respiratory support soon after birth. Our hypothesis is that HF is non-inferior to the standard treatment—nasal continuous positive airway pressure (NCPAP)—as primary respiratory support for preterm infants.

Methods and analysis: The HIPSTER trial is an unblinded, international, multicentre, randomised, non-inferiority trial. Eligible infants are preterm infants of 28–36+6 weeks gestational age (GA) who require primary non-invasive respiratory support for respiratory distress in the first 24 h of life. Infants are randomised to treatment with either HF or NCPAP. The primary outcome is treatment failure within 72 h after randomisation, as determined by objective oxygenation, blood gas, and apnoea criteria, or the need for urgent intubation and mechanical ventilation. Secondary outcomes include the incidence of intubation, pneumothorax, bronchopulmonary dysplasia, nasal trauma, costs associated with hospital care and parental stress. With a specified non-inferiority margin of 10%, using a two-sided 95% CI and 90% power, the study requires 375 infants per group (total 750 infants).

Ethics and dissemination: Ethical approval has been granted by the relevant human research ethics committees at The Royal Women’s Hospital (13/12), The Royal Children’s Hospital (33144A), The Mercy Hospital for Women (R13/34), and the South-Eastern Norway Regional Health Authority (2013/1657). The trial is currently recruiting at 9 centres in Australia and Norway. The trial results will be published in peer-reviewed international journals, and presented at national and international conferences.

Strengths and limitations of this study

- This is the first study that is appropriately designed and powered to assess the efficacy of high flow therapy as primary respiratory support for preterm infants.
- The use of a non-inferiority design is appropriate given the advantages of high flow over nasal continuous positive airway pressure. A narrow non-inferiority margin (10%) has been chosen to ensure the study results will be convincing to clinicians.
- Blinding of the allocated respiratory support modes is not possible, but subjective criteria are specified for the primary outcome of treatment failure.
- Some infants in the high flow group will have initially received a brief period of nasal continuous positive airway pressure prior to randomisation.

INTRODUCTION

Background

Preterm birth is the leading cause of newborn death worldwide. Every year, 15 million infants are born preterm and >1 million die from complications.1 Respiratory distress syndrome (RDS) is one such complication occurring in 44% of very low birthweight infants (<1500 g);2 therefore, identifying the optimal method for providing
breech delivery shortens the delivery process. A smaller size, decreased forceps usage, and a lower cesarean section rate have been observed in breech deliveries. However, issues like difficulty in breech extraction, birth asphyxia, and the risk of neonatal injury must be considered.

A Cochrane review has compared the outcomes of vaginal breech delivery compared to cesarean section. The review, which included 24 studies, found no significant difference in the risk of adverse perinatal outcomes, such as death or serious injury, between the two groups. However, the review notes that cesarean section was associated with lower rates of vaginal delivery and higher rates of emergency cesarean section, as well as a higher risk of infection and blood loss. Therefore, it is important to establish a comprehensive plan to reduce the risk of complications during breech delivery, ensuring that the patient and the baby receive appropriate care.
have much higher pneumothorax rates of up to 9%,^6,8^ the risk of pneumothorax during primary HF support is unknown.

Three RCTs have convincingly demonstrated that HF results in less nasal trauma than NCPAP.^18–20^ Further studies have shown HF is preferred by parents^27^ and by nursing staff.^28^ Other perceived advantages of HF, such as greater infant comfort and better establishment of feeding, remain unproven.

Rationale and aim

Neonatal HF use, including use as primary support, is rapidly increasing around the world. It is crucial that HF therapy is applied without causing harm by making appropriate assessment of its use before it becomes widely accepted in the neonatal practice. If HF does provide comparable support to NCPAP for preterm infants with early respiratory distress, then it is likely that it will be widely adopted in preference to NCPAP in NICUs as it is easier to use, more comfortable for infants,^29^ reduces nasal trauma, and is preferred by clinicians and parents.^14,17,27^ The aim of this study is to assess whether HF is non-inferior to NCPAP in preventing treatment failure, when used as primary respiratory support for preterm infants.

METHODS

Study design

HIPSTER is an international, multicentre, randomised, non-inferiority trial, conducted in preterm infants ≥28 weeks’ gestational age (GA) requiring primary non-invasive respiratory support for respiratory distress in the first 24 h of life.

Blinding

The intervention in this study cannot be blinded. To limit bias, predefined, objective criteria for the primary outcome of treatment failure are specified so as to provide clear directions to clinicians for the decision to escalate respiratory support.

Primary outcome

The primary outcome is treatment failure within 72 h after randomisation. Treatment failure is reached once an infant is receiving maximal therapy for their allocated treatment (NCPAP 8 cm H₂O or HF 8 L/min), plus at least one of:

1. Sustained increase in oxygen requirement above ≥40% to maintain oxygen saturation in the target range for that centre;
2. Frequent apnoea: six or more apnoeas requiring intervention in a 6-h period, or two or more apnoeas requiring facemask positive pressure ventilation in a 24-h period;
3. Respiratory acidosis: blood pH ≤7.20 and carbon dioxide >60 mm Hg of mercury on capillary/arterial blood, taken at least an hour after commencing the assigned treatment.

Treatment failure will also be adjudged to have occurred in any infant requiring urgent intubation and mechanical ventilation, as determined by the treating clinician.

Secondary outcomes

1. Reason(s) for ‘treatment failure’
2. Intubation rate in first 72 h, and at any time
3. Incidence of radiologically confirmed pneumothorax or other air leak
4. Incidence of significant nasal trauma (as measured using a validated nasal trauma scoring chart)
5. Incidence of bronchopulmonary dysplasia (supplemental oxygen requirement and/or need for respiratory support at 36 weeks’ postmenstrual age)
6. Use of postnatal steroids for the treatment of lung disease
7. Discharged home with supplemental oxygen
8. Duration of admission, days of each respiratory support mode, death before discharge
9. Incidence of important neonatal morbidities including: late-onset sepsis, patent ductus arteriosus, necrotising enterocolitis, intestinal perforation, severe intraventricular haemorrhage, and treated retinopathy of prematurity
10. Days to reach full enteral feeds and full suck feeds, method of feeding at discharge, and weight gain until discharge
11. Economic analyses ( overseen by a trial health economist)
12. Parental stress and perception of infant’s treatment, as measured by a validated questionnaire: (‘Parental Stress Scale: Neonatal Intensive Care Unit’, PSS: NICU^30^).

Setting

The trial will be conducted in nine tertiary level NICUs (4 centres in Australia and 5 centres in Norway). All centres routinely care for preterm infants with respiratory distress, and use NCPAP as their standard mode of primary respiratory support.

Eligibility criteria

Inclusion: Infants will be included if
1. They are born at 28–36^6^ weeks GA, AND
2. They are admitted to a participating NICU (inborn or outborn) when <24 h old, AND
3. The decision has been made by the attending clinician, to commence or continue (from stabilisation at birth) non-invasive respiratory support (this does not include the provision of supplemental oxygen alone), AND
4. They have not previously been intubated or received surfactant, AND
5. At randomisation, the infant has received <4 h of NCPAP support (respiratory support may need to
start prior to consent being obtained and if so, this will be with NCPAP).

**Exclusion:** Infants will be excluded if
1. They immediately require intubation and ventilation (determined by attending clinician), OR
2. They already satisfy ‘treatment failure’ criteria, OR
3. They have a known major congenital anomaly or air leak (pneumothorax).

**Randomisation**
Prerandomisation stratification is by GA (<32 and ≥32 weeks\(^2\)) and by study centre. Multiple births will be randomised individually. The randomisation sequence is computer generated with variable block sizes; assigned randomly.

**Clinical management**
Eligible infants will be randomised to treatment with either HF or NCPAP. Infants with birth weight ≤1250 g will receive caffeine for apnoea prevention\(^3\) at enrolment if not already given, to be continued at least during the primary outcome period. Apnoeic infants >1250 g may receive caffeine as per the clinician’s discretion.

**Standard care: control group (NCPAP)**
1. NCPAP will be delivered using any NCPAP delivery device and short binaural prongs; pressure will start at 6–8 cm H\(_2\)O (clinician discretion). Pressure changes will be made with 1 cm H\(_2\)O increments/decrements in the range 5–8 cm H\(_2\)O. Weaning will be reviewed at least daily with cessation considered once the infant is stable on NCPAP 5 cm H\(_2\)O, in <30% oxygen, for >24 h. Subsequently, unconditioned ‘low flow’ oxygen may be given to maintain oxygen saturation.
2. Infants in the NCPAP group will not receive HF if the infant is already a widely accepted mode of respiratory support in many tertiary and non-tertiary neonatal units; for example, blood tests, X-rays, antibiotics, intravenous fluid/nutrition and enteral feeds.

**Intervention group (HF)**
1. HF will be given using either Optiflow Junior (Fisher & Paykel Healthcare, New Zealand) or Vapotherm (Vapotherm, Exeter, USA). Gas flow will start at 6–8 L/min (clinician discretion), and flow changes will be made with 1 L/min increments/decrements in the range 4–8 L/min. Weaning will be reviewed at least daily with cessation considered once the infant is stable on 4 L/min, in <30% oxygen, for >24 h. Subsequently, unconditioned ‘low flow’ oxygen may be given to maintain oxygen saturation.
2. Infants who reach treatment failure criteria while receiving maximal HF (8 L/min) within the primary outcome period (72 h) will receive NCPAP at 7–8 cm H\(_2\)O (clinician discretion).
3. Infants who again reach treatment failure criteria while receiving maximal NCPAP (8 cm H\(_2\)O), while still within the 72-h primary outcome period, will be intubated and ventilated.
4. If further non-invasive respiratory support is required later during admission (eg, for clinical deterioration), infants should receive HF. However, if they have previously reached treatment failure criteria during HF, they may be treated with NCPAP at clinician discretion.

**Sample size calculation**
A review of preterm infants >28 weeks’ GA receiving NCPAP as their initial mode of respiratory support at the participating Australian centres (unpublished data) showed that 17% of such infants were subsequently intubated and ventilated, within 72 h of starting treatment. We, therefore, chose an expected NCPAP ‘treatment failure’ rate of 17%.

We have set the margin of non-inferiority for the trial at 10%. That is, HF will be considered non-inferior to NCPAP if the risk difference for treatment failure and upper limit of its two-sided 95% CI is <10%\(^3\) (eg, if the NCPAP treatment failure rate is 17%, both the risk difference and upper limit of its two-sided 95% CI must be <27%). To demonstrate this with 90% power, we require a sample size of 375 infants per group, 750 infants in total. We chose this margin of non-inferiority with consideration to the following factors:
- HF is already a widely accepted mode of respiratory support in many tertiary and non-tertiary neonatal units;
- Infants in whom HF treatment fails will receive NCPAP, and we hypothesise that this will ‘rescue’ some of these infants from intubation;
- The primary outcome of this study is treatment failure, as opposed to an outcome like death or severe disability, when a lower margin of non-inferiority would be necessary;
- This non-inferiority margin was thought to be appropriate, and was agreed on by all neonatologists in all the participating centres, and by parent representatives consulted during the trial design phase.

**Statistical analysis**
The incidence of the primary outcome will be compared using risk difference and two-sided 95% CI. Planned subgroup analyses by GA strata will be performed for the primary outcome. Secondary outcomes will be compared using risk difference (95% CI) and \(\chi^2\) tests, or the
appropriate parametric (t test) or non-parametric (Mann-Whitney U) tests. Statistical analyses will be by intention to treat, conforming to the Consort reporting guidelines.

Cost-effectiveness analysis will incorporate the costs of the device and of hospital care; a decision analysis will be constructed based on the primary outcome and associated hospital costs. Univariate and probabilistic sensitivity analyses will be conducted as a cost per additional treatment failure avoided for HF versus NCPAP.

ETHICS AND DISSEMINATION

Research ethics approval

The HIPSTER trial has received multisite ethical approval from the relevant governing bodies for all participating centres.

Recruitment and consent

Written parental consent is required for all infants participating in the trial. Consent will be sought in the antenatal period when possible, at all sites. When antenatal consent is in place, infants will be randomised as soon as possible after meeting eligibility criteria.

When antenatal consent has not been obtained, infants judged to require non-invasive respiratory support will receive standard treatment (NCPAP) until consent has been given. Families of infants meeting eligibility criteria will be approached at the earliest opportunity after birth, and before their infant has received 4 h of NCPAP treatment.

Additionally, at the lead centre (The Royal Women’s Hospital, Melbourne), the Human Research Ethics Committee (HREC) has approved a retrospective consent process. Eligible infants who have not been consented antenatally can be randomised as soon as they meet eligibility criteria. Their parents will then be approached for consent in the first few days after trial entry, at which point the parents may choose to give consent for their infant to remain in the trial, or remove them and opt for standard treatment.

The consent process, whether antenatal or postnatal, will include both a full verbal explanation of the trial and the use of the written patient information and consent form.

Data collection and storage

Outcome data, birth details and parental demographics will be collected from the infant’s records and mother’s medical records, and by parental interview. Data will be de-identified and entered into a paper case report form. Data will subsequently be entered into REDCap (Research Electronic Data Capture)33—a secure, password-protected electronic database.

Monitoring and safety

A data safety monitoring committee (DSMC) comprising of two independent neonatologists and an independent statistician has been appointed. Set DSMC review points on the progress and safety of the trial are after the primary outcome is known for 250 and 500 infants. While no formal stopping rule will be used, the DSMC may recommend ceasing the trial if there is a statistically significant difference (p<0.001) in primary outcome between the treatment groups overall or within prespecified GA subgroups,34 or in case of serious adverse events (identified as pneumothorax or other air leak from the lung while receiving the assigned treatment, and death before discharge). Cessation of the trial may also be recommended if there is equipment failure or recall, or if other evidence becomes available that would make continuing the trial unethical. All serious adverse events are to be reported to the lead centre’s Human Research and Ethics Committee, and will be reviewed by the DSMC at the prespecified monitoring points. The first review point was reached in October 2014 and the DSMC recommended that the trial continue without modification.

Dissemination of results

Trial results will be published in peer-reviewed international journals, and presented at relevant national and international conferences. A plain language summary of the results will be sent to the parents of participants.

Current status and study duration

The trial began single-site recruitment in May 2013; it became multicenter in January 2014 and was extended to international sites in September 2014, and is currently recruiting in all nine participating centres. It is expected that recruitment will be completed in 2016.

Trial registration

The HIPSTER trial is registered with the Australian New Zealand Clinical Trials Registry (ID: ACTRN12613000303741).

DISCUSSION

HF therapy has been widely adopted in neonatal practice due to its desirable qualities such as ease of use, reduced nasal trauma, and parental and nursing preference.12 13 However, it is of concern that HF is being used as primary respiratory support for preterm infants in the absence of good quality evidence of its efficacy in this setting. The HIPSTER trial is the first appropriately powered and designed trial to assess HF as primary support for preterm infants.

Non-inferiority trials are relatively uncommon in neonatal practice, but appropriate in this case due to the advantages associated with HF, which would make it preferable to NCPAP provided that it is non-inferior in efficacy. The choice of non-inferiority margin is important in such a trial, and our margin of 10% was chosen in view of the fact that the primary outcome was...
treatment failure and not a more critical outcome, such as death, and that infants who have treatment failure on HF will be offered NCPAP, which may ‘rescue’ them from intubation and ventilation. This non-inferiority margin is half the size of that used in a previous post-extubation trial of HF published by our group\(^6\) given that the expected rate of treatment failure in the population of The HIPSTER trial, a study of primary respiratory support, is lower, and therefore the criteria for non-inferiority should be stricter.

A potential limitation to this trial is that blinding is not possible. We have attempted to minimise this by setting objective treatment failure criteria, which were agreed on by all participating centres. Some infants randomised to HF will have received a brief period of NCPAP before randomisation, which conceivably could affect interpretation of the results. However, we have aimed to restrict the impact of this by making any infant who has received four or more hours of NCPAP ineligible for the trial and by the use of antenatal consent when possible, and a retrospective consent process at the lead centre. Acceptance of such a process requires the approval of both the HREC and the treating clinical team, and this may vary from site to site. We feel retrospective consent is appropriate in this trial given that HF has already been adopted into standard practice as a mode of primary respiratory support by some neonatologists\(^12\) and that along with the inclusion of ‘rescue’ NCPAP, the HREC adjudged that infants in the HF group were not exposed to additional risk in comparison to those treated with NCPAP.

The use of HF in neonatal practice is now well established, but good quality evidence is required to determine in which clinical settings this is appropriate. If this trial demonstrates that HF is non-inferior to NCPAP as primary support, then this practice is likely to be widely adopted around the world. However, if HF is inferior to NCPAP, then this study will ensure that preterm infants, who require non-invasive respiratory support, receive the optimal treatment.

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Contributors CTR conceived and designed the trial protocol, wrote the first draft and revised the manuscript for intellectual content. LSO, BJM and PGD conceived and designed the trial protocol and revised the manuscript for important intellectual content. SMD designed the protocol statistical analysis and revised the manuscript for important intellectual content. All the authors have read and approved the final manuscript, and are accountable for its accuracy.

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Competing interests None declared.

Ethics approval The trial is approved by: the Royal Women’s Hospital Human Research Ethics Committee (Reference: 13/12), the Royal Children’s Hospital Human Research Ethics Committee (Reference: 33144A), the Mercy Hospital for Women Human Research Ethics Committee (Reference: R13/34), and South-Eastern Norway Regional Health Authority Committee for Medical and Health Research (Reference: 2013/1657).

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Data sharing statement Further information on the study protocol may be requested from the corresponding author.

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