Safety and efficacy of nitric oxide in chronic lung disease

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Background: Therapies for neonatal chronic lung disease (CLD) of prematurity have had limited success. Aims: To determine whether inhaled nitric oxide (INO) administered to very low birthweight infants with developing CLD might improve oxygenation without adverse effects. Methods: Subjects were 10–30 days of age, birth weight <1250 g, with developing or established CLD, and requiring mechanical ventilation with mean airway pressure $\geq 7$ cm H$_2$O and FIO$_2$ $\geq 0.40$. We monitored changes in oxygenation and FIO$_2$ requirement during treatment with INO (initial dose 20 ppm). Tracheal aspirate samples obtained before, during, and after treatment were analysed for interleukin 1$\beta$ (IL-1$\beta$), IL-8, 8-epi-prostaglandin F$_{\alpha}$ (8-epi-PGF$_{\alpha}$), laminin, and endothelin 1 (ET-1) to assess any potential effects of INO on markers of inflammation/oxidation, basement membrane injury, or vasoactivity. Results: Thirty three patients met entry criteria. Mean gestational age was 25 (SD 2) weeks; birth weight was 736 (190) g; age of study infants was 19 (6) days (range 9–29). Mean FIO$_2$ decreased from baseline (0.75) to 0.58 at 72 hours. Duration of therapy was seven days. Tracheal aspirate concentrations of IL-1$\beta$, IL-8, 8-epi-PGF$_{\alpha}$, ET-1, and laminin were unchanged between baseline and 48 hours of INO, and 48 hours after discontinuation of INO. No new cases of, nor extension of, intraventricular haemorrhage occurred. Four infants died. Conclusion: INO (≤20 ppm) improved oxygenation in most infants with early CLD, without inducing changes in markers of inflammatory or oxidative injury.

Neonatal chronic lung disease (CLD), defined as the continuing need in preterm infants for supplemental inspired oxygen at 36 weeks postconceptional age, affects an estimated 12 000 infants per year in the USA and many more infants worldwide. Limitations of current treatments for CLD have prompted a search for other interventions. One potentially useful therapy, on the basis of its known effectiveness to improve oxygenation in near term neonates with hypoxic respiratory failure, is inhaled nitric oxide (INO). No reports, to our knowledge, describe INO use in extremely preterm infants beginning at 10 days of age, when signs of CLD are becoming apparent.

The impact of INO, administered for a clinically meaningful time, on an already inflamed, immaturely developed lung, is largely unknown. Inflammation contributes to CLD, and nitric oxide can have both proinflammatory and anti-inflammatory effects. In the presence of high FIO$_2$, NO is converted to NO$_2$, peroxynitrite, and other oxides of nitrogen, which may initiate or exacerbate pulmonary inflammation. However, nitric oxide may modulate the pulmonary inflammatory response by downregulating the production of inflammatory cytokines, and by decreasing lung neutrophil accumulation.

Because we considered it premature to conduct a prospective randomised trial of INO in the very low birthweight (VLBW) infant population with early CLD, based on the limited data available regarding dosage, efficacy, and especially safety, we conducted an open trial of INO in CLD, emphasising safety issues and short term efficacy in a severely ill group of patients.

METHODS
The Pediatric Institutional Review Board of the University of Missouri–Kansas City and the Children's Mercy Hospital approved the study protocol.

Subjects
All patients admitted to the Intensive Care Nursery from June 1997 to June 1999 were reviewed for eligibility. Parental consent was obtained for all eligible candidates identified, with no parental refusals. Inclusion criteria included birth weight <1250 g; age <30 days, but ≥10 days; need for assisted ventilation; need for FIO$_2$ ≥0.4 without fluctuations of >0.25 in the preceding 24 hours; and clinical course and radiographic findings compatible with CLD.

Exclusion criteria included: initiation of systemic corticosteroid or inhaled $\beta$ agonist therapy within the preceding 48 hours; new diagnosis of sepsis (two blood cultures yielding growth of a single pathogenic organism) within the preceding 48 hours; thrombocytopenia (<100 000/mm$^3$); progressive intraventricular haemorrhage; obviously lethal congenital anomaly; and complex congenital heart disease. The clinical course and chest radiographs of potential subjects were reviewed prior to enrolment by at least two investigators not serving as the patient’s attending physician.

Intervention
Inhaled nitric oxide was administered initially at 20 ppm. Arterial oxygen saturation was monitored by pulse oximetry, and transcutaneous levels of CO$_2$ and O$_2$ were measured over the next three hours to ascertain response. FIO$_2$ was adjusted to keep SpO$_2$ at 89–96%. No change in mean airway pressure occurred during this period. At the end of the initial three hour treatment period, a decision was made by the investigators regarding continuation of INO. If there was evidence of efficacy (FIO$_2$ decreased by ≥0.10, SpO$_2$ increased by ≥10%, or TcPO$_2$ improved by 1.3 kPa), INO was continued at 20 ppm.

Abbreviations: CLD, chronic lung disease; CMV, conventional mechanical ventilation; ELISA, enzyme linked immunoassay; HFOV, high frequency oscillatory ventilation; INO, inhaled nitric oxide; IVH, intraventricular haemorrhages; PCA, postconceptional age; PVR, pulmonary vascular resistance; sSC-IgA, soluble secretory component of IgA; VLBW, very low birthweight.
After 36 hours, the dose of INO was reduced to 15 ppm. The dose was further reduced every 12 hours by 2–3 ppm to a dose of 2 ppm as long as there was no worsening of oxygenation. A trial of cessation of INO was then undertaken. A reduction of INO followed by an absolute decrease of SpO₂ of >10% or need for increase in FIO₂ of ≥0.2 to sustain the previous SpO₂ reading within ≤30 minutes of changing the INO dose was considered an indication of lack of tolerance to the change, and the prior dose was reinstituted. Treatment with INO was discontinued by seven days if no evidence of dependency was detected. If INO dependency had developed, INO was continued with repeated trials of discontinuation every 24–48 hours.

All patients were treated with assisted ventilation with time cycled, pressure limited, and/or patient triggered ventilation (Dräger Babylog 8000), or high frequency oscillatory ventilation (Sensormedics 3100A). Changes from one mode of ventilation to another were allowed, but were made in only one case between 3 and 72 hours. Tracheal aspirate fluid samples were collected by standard techniques for analysis of pulmonary inflammatory cytokines (interleukin 1β (IL-1β), IL-8); a marker of peroxidative injury (8-epi-prostaglandin F₉ (8-epi-PGF₉α)); a proliferative and vasoconstrictor substance (endothelin 1 (ET-1)); and a marker of basement membrane injury (laminin). Samples were immediately frozen at −70°C until batch analysis was carried out.

Blood methaemoglobin and inspired nitrogen dioxide were continually measured. Surveillance was conducted for haemorrhage in any organ, infection, thrombocytopenia, necrosis of enterococci, and retinopathy of prematurity. Bloodstream infections were defined as clinical deterioration plus blood culture yielding a pathogenic organism. Pneumonia was defined as deterioration of respiratory status, concomitant with tracheal aspirate material yielding growth of a pathogenic organism. Cranial ultrasound examinations were performed before initiation of INO, during treatment if clinically indicated, and after discontinuation of INO therapy.

Echocardiographic analysis
Echocardiography was performed within the 48 hours preceding INO whenever possible. Elevation in pulmonary vascular resistance was considered to be present if there was a detectable tricuspid valve regurgitation jet.

Respiratory monitoring
The FIO₂, ventilator pressures (peak inflating pressure and positive end expiratory pressure for patients on conventional ventilation, and mean airway pressure and amplitude for those on high frequency oscillatory ventilation (HFOV)), tidal volume, minute ventilation, and rates were recorded hourly. The SpO₂ and the most recent blood gas tensions and pH were recorded. All variables were averaged from hourly readings for the six hours before for baseline measurements, and six hours before and six hours after the 24 and 72 hour time points.

Tracheal aspirate analysis
Tracheal aspirate samples were assayed for IL-1β, IL-8, ET-1, and 8-epi-PGF₂α utilising available enzyme linked immunoassay (ELISA) kits (Quantikine, R&D Systems, Minneapolis, Minnesota). Laminin concentrations were assayed by competitive inhibition ELISA (Chemicon International, Temecula, California). The soluble secretory component of IgA (sSC-IgA) was assayed by methods established in our laboratory. All samples were assayed at least in duplicate and the results averaged. Coefficients of variation for each of these assays is <10% in our laboratory.

Statistical analysis
Statistical testing for continuous variables was performed by paired t test with Bonferroni correction for multiple comparisons. For non-parametric data comparison, Wilcoxon rank sign test was performed. Correlation coefficients were tested for significance by Spearman and/or Kendall tau beta tests and linear regression where appropriate. SPSS 8.0 statistical software was used for the statistical computations (SPSS Inc., Chicago, Illinois).

RESULTS
Patients
Trials were performed in 33 consecutive infants who met the criteria for entry. Twenty were male; 21 were black, 11 white, and one Hispanic. Mean birth weight was 736 g (range 509–1250 g); mean gestational age was 25.3 weeks (range 23–29); mean age at enrolment was 19 days (range 9–29). Twenty one infants were treated with antenatal steroids, 18 with antenatal magnesium sulphate; 14 were delivered by caesarean section and 29 received surfactant treatment.

Pulmonary gas exchange
All infants tolerated the initial three hour trial with evidence of improvement. However, two infants did not continue to receive INO beyond 48 hours as there was no evidence then of a decrease of FIO₂ needs. For the remaining group of 31 infants, there was a significant reduction in FIO₂ at 3 hours and at 72 hours of therapy (at which time the median dose of INO was 10 ppm) compared with baseline values (p < 0.05 for each comparison, paired t test and Wilcoxon rank sign test). Mean FIO₂ decreased from 0.75 (0.23) to 0.63 (0.22) at 3 hours and to 0.58 (0.18) at 72 hours of INO therapy. The calculations include data on all 33 patients, including the two infants not receiving INO at 72 hours. The mean FIO₂ at 72 hours was also lower than that at 3 hours (p < 0.05, Wilcoxon rank sign test). The reduction in FIO₂ occurred even as mean SpO₂ showed an upward trend. The SpO₂ was 90 (6)% at baseline, 91 (4)% at 3 hours, and 93 (5)% at 72 hours. In spite of the reduced dose of INO by 72 hours, 14 of the 33 trials (45%) resulted in an absolute reduction in FIO₂ of >0.2. Only five infants (including the two whose trial stopped before 72 hours) showed no change or an increase in FIO₂ by 72 hours. Magnitude of reduction in FIO₂ correlated inversely with the baseline FIO₂ (p < 0.05; r = −0.72). The decrease in FIO₂ showed no significant correlation with postnatal or gestational age, prior or concurrent usage of corticosteroids, antenatal corticosteroid use, gender, race, or presence of pulmonary air leak. The mean reduction in FIO₂ was similar for infants treated with conventional mechanical ventilation (CMV) (n = 15) or HFOV (n = 18) at baseline. For the subset of 15 infants with baseline FIO₂ ≥0.9, the mean FIO₂ at 72 hours had decreased to 0.68 (range 0.97–0.35).

Measurements of tcPCO₂, during INO administration were unchanged. Tidal volume, minute ventilation, and mean airway pressure did not change significantly between initiating INO and 72 hours of administration, as the emphasis was to reduce FIO₂. Mean airway pressure at baseline was 7.3 (1.1) cm H₂O for the CMV treated infants and 12.2 (3.1) cm H₂O for the HFOV treated infants. Only one patient underwent a change in type of assisted ventilation (from synchronised intermittent mandatory ventilation to HFOV) during the first 72 hours of INO. FIO₂ needs did not change after initiation of HFOV.

Eight patients failed to tolerate the first attempt at discontinuation of INO at the end of their trials and required repeated attempts at discontinuation of the inhaled NO. The magnitude of initial response was not different in this group compared to that of the other 25 infants. The longest duration of treatment was 13 days.

Twenty one of 33 infants were treated with dexamethasone at the time of initiation of the INO trial. All had been receiving dexamethasone for more than 48 hours but still met entry criteria. There was no difference in mean decrease in FIO₂ between the steroid pretreated group and those not using steroids.
Echocardiographic analysis

Echocardiography was performed on 27 infants within 48 hours preceding initiation of treatment with INO. Eleven showed indirect evidence of increased pulmonary vascular resistance (PVR), in the form of a measurable tricuspid regurgitation jet. There was no correlation between the presence or absence of echocardiographic evidence of raised PVR and the magnitude of response to INO.

Serial tracheal aspirate sample analysis

No difference was found in concentrations of IL-1β, IL-8, 8-epi-PGF$_{2a}$, ET-1, and laminin when values of each substance were compared between baseline and during INO or after INO therapy (figs 1, 2, and 3). There was wide variation in interpatient concentrations, but not in intrapatient concentrations, over time. When comparing concentrations across time in patients with a history of air leak (pulmonary interstitial emphysema or pneumothorax, n = 5), there again was no difference.

Complications

Five subjects showed intraventricular haemorrhages (IVH) of grade 3 (n = 3) or grade 4 (n = 2) at baseline. All other infants had grade 2 or less IVH prior to enrolment. No cases of new intracranial haemorrhage nor extension of old haemorrhage occurred. No subject experienced a clinically evident pulmonary haemorrhage. There were no sudden or unexplained decreases in haemoglobin or haematocrit, nor were there any unexplained episodes of thrombocytopenia. No concentration of methaemoglobin above 2.8% was detected at any time during INO use. Bloodstream infection occurred in five patients (coagulase negative staphylococci in three, Candida sp. in two) during or following therapy with INO. Five patients developed pneumonia (ureaplasma/mycoplasma species in three, Candida sp. in one, and coagulase negative staphylococci in one) during or following INO.

Outcome

Four of the 33 infants died before discharge (see table 1). No additional infants died during mean period of 22 months (range 15–36). At 36 weeks postconceptional age (PCA), one child continued to be treated with assisted ventilation, two were treated with continuous positive airway pressure, and 22 were treated with nasal cannula supplemental oxygen. Three infants tolerated breathing room air. By 44 weeks PCA, only three infants remained hospitalised, two of whom were treated with nasal cannula oxygen; 10 infants were breathing room air, and 19 were utilising nasal cannula oxygen. At six months PCA, 25 infants had accessible records available for evaluation. Ten infants continued treatment with supplemental oxygen by nasal cannula; 15 were breathing room air.

Discussion

Inhaled NO acutely improved pulmonary oxygen uptake in most infants with early CLD. This is the first report of administration of INO in premature infants without evidence of increasing airway inflammation, peroxidation, or basement membrane breakdown. We studied a specific but important population of infants. These extremely low birthweight infants had very significant but early CLD, probably characterised by substantial pulmonary inflammation. Use of INO has not been reported in this specific patient population, yet the pulmonary morbidity and mortality for 750 g infants still requiring assisted ventilation at 19 days of age with a mean $F_{iO_2}$ of 0.75 would be predicted to be considerable.
Neonatal CLD appears to represent the effects of mechanical distortion of the lung, oxidative injury, and resultant inflammatory response in immature lungs. Evidence for early inflammatory changes in CLD has been documented. Increased concentrations of IL-1β and IL-6 have been found in bronchoalveolar lavage fluid from patients with CLD, and IL-1β has been suggested as a participant in the development of CLD. The neutrophil chemotactic cytokine, IL-8, may participate in CLD, and both its production and that of IL-1β may be regulated directly or indirectly by NO and by peroxynitrites. Our findings of no effect of INO on tracheal aspirate concentrations of IL-1β and IL-8 suggest that proinflammatory lung cytokine production was not worsened by INO administration. Inhaled NO could exert anti-inflammatory effects by means of its effect on cytokines and neutrophil migration. Inhaled NO has also been shown to reduce neutrophil accumulation in the lung.

There is evidence that 8-isoprostanes such as 8-epi-PGF_2α, which are generated by oxidation of membrane phospholipids, serve as markers for pulmonary oxidative injury, and may themselves contribute to pulmonary hypertension. Concentrations of the pro-fibrotic and constrictor substance ET-1 have been found to be increased in tracheal aspirate fluid of infants developing CLD. Tracheal aspirate concentrations of 8-epi-PGF_2α and ET-1 were unchanged with NO treatment. Concentrations of laminin, an indicator of basement membrane disruption, were also unchanged with use of INO in our patients. Thus, with relatively short term use of INO we found no evidence of additional peroxidative or inflammatory damage or of matrix disruption.

Microvascular obliteration and disorganised vasculogenesis in CLD may lead to pulmonary hypertension and contribute to poor matching of ventilation and perfusion, leading to hypoxaemia, hypercarbia, and exacerbation of pulmonary hypertension. Indirect evidence suggests that PVR is increased in patients who develop CLD. The pulmonary vasculature of premature infants can respond to nitric oxide with vasodilatation. Our results imply that in early CLD, INO may improve ventilation–perfusion matching by mechanisms including, but not limited to, reduction in PVR, as the response to INO did not depend on presence of raised PVR as measured by echocardiography. Some of the improvement in oxygenation may occur because NO may also function as a bronchodilator.

Inhaled NO has been reported as a “rescue” therapy based on short term trials in preterm infants with severe RDS and hypoxaemic respiratory failure. Van Meurs and colleagues and Kinsella and colleagues tested the efficacy of NO in preterm infants beginning at 1–2 days of age with total treatment duration of seven days in almost all infants. Subedar and colleagues treated 10 infants with RDS from day 4 to day 7 with INO. As CLD may arise following mild or no previous RDS, these previously reported results do not predict the effect of INO on early CLD. Our study design also differs from that of Longist and colleagues, who evaluated INO in nine infants with severe CLD by treating them for 40 minutes total, and from those of Banks and colleagues, who treated 16 infants with “end stage” CLD beginning at a median age of 3 months. Longist and colleagues did not address safety issues, while Banks and colleagues did not measure directly markers of lung inflammation. No new intraventricular haemorrhage or extension of previous haemorrhage occurred in our study, in contrast to the results of Cheung and colleagues, in a study of the “rescue” use of INO in very premature infants treated at less than 24 hours of life. Cheung and colleagues found new haemorrhages or significant increases in extent of hemorrhage in 15 of 23 patients receiving INO who survived to undergo repeat ultrasound examination. During the time period encompassed by this study, we treated an additional six infants who were 2–14 months of age with INO for exacerbations of their underlying CLD. Our results were similar to those already well characterised by Banks and colleagues.

Long term efficacy cannot be inferred from our study. We terminated NO usage, by design, at 7 days when possible, and had decreased the dose to 10 ppm by 3 days. It is possible that a more substantial reduction in FIO2 could have been achieved with continuation of the starting dose of 20 ppm for 72 hours or more, or the same reduction achieved by starting at 10 ppm without lowering the dose.

Chronic lung disease is now recognised as an independent contributor to poor neurodevelopmental outcome. In addition, diminished long term pulmonary function remains a major long term sequela of CLD. Our findings support the performance of a definitive trial of INO in early or evolving CLD, but do not offer sufficient support to warrant widespread use of INO in infants remaining on assisted ventilation for early CLD.

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REFERENCES


Table 1 Details of the four infants who died

<table>
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<tr>
<th>Case</th>
<th>BW (g)</th>
<th>DOL at start to CMH</th>
<th>DOL at start of INO</th>
<th>Baseline FIO2/SpO2</th>
<th>INO response at 72 h</th>
<th>INO stopped, DOL</th>
<th>Age at death (days)</th>
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<td>17</td>
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<td>24</td>
<td></td>
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</tbody>
</table>

BW, birth weight; DOL, day of life; CMH, Children’s Mercy Hospital; INO, inhaled nitric oxide.
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