Hypoxic ventilatory defence in very preterm infants: attenuation after long term oxygen treatment

Miriam Katz-Salamon, Hugo Lagercrantz

Abstract
The activity of peripheral chemoreceptors was studied in 19 preterm very low birthweight infants at the postconceptional age of 36 and 40 weeks using the hyperoxic test. The infants were in a healthy condition and did not receive any extra oxygen or medication when tested. The inhalation of pure oxygen caused a decrease in mean (SE) ventilation by 16-1 (2·6)% and 15·1 (2·1)% at the 36th and 40th gestational week respectively. At the 36th gestational week the ventilatory response was significantly lower than at 40 weeks (10·9 (6) and 7·3 (3) sec). Six infants who had been on supplemental oxygen for more than 21 days (from 21 to 56 days) responded with significantly lower response to hyperoxia at the 36th gestational week (7·9 (3·6)% of those receiving oxygen treatment for a shorter period of time, 0 to 16 days (19·9 (3·2)%). The 'low responding' group included three infants who had suffered from chronic lung disease. Those infants showed the lowest hyperoxic response (4·3 (3·9)%). There was no difference in the response among healthy preterm infants (eight infants) and infants with respiratory distress syndrome. At the 40th gestational week the differences, even though showing the same characteristics, were not statistically significant. No statistically significant relationship was found between the strength of the ventilatory response to oxygen versus gestational, postnatal age, nor the time interval between the termination of supplemental oxygen treatment and the test. No relationship was found between the number of apnoeic/bradycardiac spells and the strength of the ventilatory depression caused by hyperoxia. In conclusion we found that the very preterm infants, with the exception of those who received long periods of oxygen treatment, have stronger peripheral chemoreceptor responses than those reported for 2-4 day old full term infants. However, infants who had suffered from chronic lung disease show a depressed hyperoxic response.

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The peripheral chemoreceptors play a crucial part in the defence against hypoxia and in the control of breathing. They have been found to be adapted to the low intrauterine oxygen tension (PO₂) levels and do not react until PO₂ drops below 5 kPa, as shown in the sheep fetus. It has been demonstrated in newborn sheep, rats, and man, during the first days after birth, that the receptors reset to the higher PO₂ of the extraterine environment. Older preterm infants have been reported to react to hypoxia in the same manner as adult man. When the peripheral chemoreflex was tested by using a single breath of 100% oxygen a similar response was observed at 1, 2, and 3 months of age in full term infants. The postnatal changes of the chemoreflexes in the newborn rat are attributed to a decrease in dopamine turnover. However, the postnatal adjustment of the chemoreflex can be delayed by chronic hypoxia possibly due to a sustained high dopamine turnover.

The aim of this study was to investigate to what extent preterm extrauterine life affects the maturation of the peripheral chemoreceptors in infants. We studied the function of peripheral chemoreceptors in very preterm low birthweight infants in relation to: (a) gestational and postnatal age, (b) the duration and character of ventilatory and oxygen treatment, and (c) the maturity of the respiratory control mechanisms reflected by the incidence of recurrent apnoea.

As the inhalation of hyperoxic gas causes an almost immediate fall in minute ventilation, due to suppression of a chemoreceptor oxygen drive, the inhalation of pure oxygen has been used to gauge the response to hypoxia.

Subjects and methods

SUBJECTS
Nineteen infants from the neonatal unit at the Karolinska Hospital born at 24–34 gestational weeks (mean (SD) 28·8 (2·7) weeks), weighing 711–1482 g at birth (1098·6 (251·6) g), were studied at the 36th and 40th week of gestation (see table). All but two infants needed ventilatory support, that is treatment on ventilator or continuous positive air way pressure. Seven infants had respiratory distress syndrome, five had transient tachypnoea, and eight recurrent apnoea that required treatment with theophylline. Three infants had chronic lung disease defined as the requirement for supplemental oxygen for more than 28 days and by specific radiographic findings. One infant with transient tachypnoea required oxygen treatment for 56 days because of several instances of septicaemia. At the time of the examination, however, no infant was receiving oxygen or theophylline treatment. The postnatal age at the first test was 50·7 (4·5) days. Informed consent from the
Clinical data on infants studied

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AR=apnoea; *CILD=chronic lung disease; RDS=respiratory distress syndrome; TTN=transient tachypnoea.

parents was obtained. The examination was performed in the presence of at least one parent. The study was approved by the local ethical committee at the Karolinska Hospital.

RESPIRATORY RECORDINGS
Oxygenation was measured by a pulse oximeter probe (Respitrace, Studley Data System, Oxford). Ventilatory measurements were made with a Greer Electrospirometer CSS5 (Mercury Electronics, Scotland). A pneumotachograph was connected to a facemask. The system was calibrated with 5, 10, and 20 ml syringes of warmed, humidified air and oxygen. The flow head was connected via a T tube to a stream of 4 litres/min humidified, warmed air which could be rapidly switched to the 4 litres/min 100% humidified oxygen. The dead space of the mask, flow head, and tubing was 2–3 ml and the resistance 5 cm H2O/l/min. The carbon dioxide build up in the spirometric system and the inspired and expired carbon dioxide were analysed at the end of the hyperoxic test with an infrared carbon dioxide analyser (AMETEK CD-3A) sampling gas from the mask at 500 ml/min. Movements of the chest and the abdomen were recorded by inductance plethysmography (Respitrace). The flow signal, its integral tidal volume, movements of the chest and abdomen, inspired and expired carbon dioxide were recorded on an eight channel inkjet recorder.

EXPERIMENTAL PROCEDURE
The pulse oximeter probe was taped to the infant’s foot while the infant was wrapped in blankets in its cot. All examinations were performed in quiet sleep characterised by the absence of eye and body movements and by regular respiration. The facemask was gently placed on each infant’s face. The experimental trial consisted of a period of 3–5 minutes of stable breathing with warmed, humidified room air. Thereafter the inspiratory line was rapidly switched and the infant breathed 100% humidified oxygen for 30 seconds. Thereafter the oxygen was substituted by air. After a period of 5–10 minutes the test was repeated.

ANALYSIS OF THE RESULTS
The recordings were only considered satisfactory for analysis if the baby remained in the same position, was not startled, and the sleep state did not change.

Response to hyperoxia
Tidal volume, breath duration, and minute ventilation (tidal volume×respiratory rate) were analysed on a breath by breath basis for each individual. The ventilatory parameters were calculated during 30 seconds of air breathing before the hyperoxic test and during 30 seconds of oxygen administration. The mean value from two trials was calculated and used for the analysis. Percentage changes in tidal volume, breath duration, and minute ventilation from normoxic to hyperoxic conditions were determined for each infant and tested for significance by Student’s t test.

Profile of the hyperoxic response
The response time to the administration of 100% oxygen was estimated as follows: breath-by-breath data were split into groups of two breaths. The response time was defined as the time elapsed between the start of hyperoxic challenge and the first statistically significant change in ventilation (analysis of variance). The results from 36th and 40th gestational week infants were compared.

Relationships between the hyperoxic response and the neonatal status of the infant
The hyperoxic changes in ventilation, tidal volume, and breath duration were related to: (a) the postnatal age and the amount of apnoeic/bradycardic episodes, (b) the length of ventilatory treatment, (c) the duration (1, 2, 3, and 4 weeks) and the intensity of supplemental oxygen treatment (median concentrations), (d) the time elapsed from the termination of oxygen treatment, and (e) results from blood samples (PO2) multiple regression analysis and/or Spearman rank correlation.

Results
Oxygen saturation during air breathing and 100% oxygen varied between 92%–96% and 98%–99% respectively. The accumulation of carbon dioxide at the end of the hyperoxic test ranged from 0.2% to 0.4%.

RESPONSE TO HYPEROXIA AT 36TH AND 40TH GESTATIONAL WEEKS
The spontaneous variation in resting ventilation preceding the hyperoxic challenge was 11 (3%). The mean ventilation at normoxia did not increase significantly (p=0.37, t test) at the end of the control period in spite of a slight
increase in carbon dioxide (0.2%-0.4%) in the spirometric system. Figure 1 shows group data on changes in minute ventilation, tidal volume, and breath duration in response to hyperoxic test. The results are expressed as percentage change from normoxic breathing. At the 36th gestational week minute ventilation decreased by mean (SE) 16.1 (2.6)% in 47% of all tests this was caused by a decrease in tidal volume alone, whereas a concomitant change in both breath duration and in tidal volume was observed in the other 47% of the tests. Only in 6% of the hyperoxic expositions were the changes in ventilation significantly correlated with the increase in breath duration. At the 40th gestational week the ventilation decreased by 15.1 (2.1)%.

Changes in ventilation were due to changes in tidal volume or to variations in both duration and amplitude of breath. Thus, the strength of the response expressed in percentage changes in ventilation, tidal volume, and breath duration did not alter between the 36th and 40th gestational weeks.

PROFILE OF THE HYPEROXIC RESPONSE

Changes in breathing patterns caused by hyperoxia were characterised by wide individual variability. In most of the infants, however, the reaction pattern could be described generally either as a depression in ventilation followed by a return to control levels (see fig 2, patients 3 and 10). In two infants the initial decrease in ventilation was followed by a successive increase and then a decrease (fig 2, patients 4 and 13). In three infants at the gestational age of 36 weeks the hyperoxic inhalation gave rise to a prolonged apnoea of 5–30 seconds. Only one child responded with an apnoea (5 seconds) at the gestational age of 40 weeks. This infant had the most pronounced apnoea (30 seconds) at the 36th gestational week. The response time expressed in seconds and breath passed between the start of oxygen administration and the ventilatory depression was significantly longer (p=0.01) at 36th

![Figure 1](image-url)

*Figure 1* Changes in minute ventilation, tidal volume, and breath duration in response to hyperoxic test expressed as percentage of normoxic values.

![Figure 2](image-url)

*Figure 2* Breath by breath analysis of the ventilatory changes during 30 seconds of normoxic breathing and 30 seconds inhalation of 100% oxygen at (A) the 36th and (B) the 40th gestational week. The arrows indicate the application of 100% oxygen.
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Ventilatory Response

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gestational week (10-9 (6) seconds, 10-2 (5)

breaths) than at 40th week of gestation (7-3 (3)

seconds, 6-3 (3-3) breaths). In two infants

there was no significant decrease in ventilation

so the response time could not be estimated

(see fig 3, patient 5).

POSTNATAL AGE AND RESPIRATORY RESPONSE

TO HYPOXIA

The postnatal age at gestational week 36th

varied from 14 to 86 days (mean 51 (20)

days). There was no statistically significant

correlation between changes in ventilation,
tidal volume, and cycle duration in response to

hypoxia and postnatal age.

VENTILATORY RESPONSE TO HYPOXIA AND

OXYGEN CONCENTRATIONS DURING

SUPPLEMENTAL OXYGEN TREATMENT

The median values of supplemental oxygen

varied from 24% to 35%. Neither at the 36th

nor the 40th gestational week was there any

correlation between oxygen concentrations

and the hypoxic response (p=0.9 and

p=0.27 respectively).

HYPOXIC RESPONSE AND THE TIME AFTER

END OF OXYGEN TREATMENT

The mean number of days elapsed after the

end of oxygen treatment and the hypoxic test

was 26 (13) (from 1–58 days). There was no

correlation between the strength of the response

and the number of days after withdrawal from oxygen treatment.

HYPOXIC RESPONSE AND DURATION OF

OXYGEN TREATMENT

The mean length of supplemental oxygen

treatment was 18 (19) days (minimum 0,

maximum 56 days). When all infants were

tested as a group the length of supplemental

oxygen treatment did not affect the ventilatory

hypoxic response (Spearman rank correlation

coefficient, p=0.18). However, when the infant

group was split into those who received

supplemental oxygen for less than 1, 2, 3, and

4 weeks respectively there was a marked

difference in the ventilatory response between

infants receiving supplemental oxygen for

more than 21 days (six infants) and those

treated for a shorter period of time (14

infants). At the 36th gestational week the mean

decrease in ventilation in infants treated with

oxygen for more than 21 days was significantly

lower (−7.8 (12)%), than the ventilatory fall in

the other group (−19.9 (8.4)%). At the 40th

gestational week the infants dependent on

supplemental oxygen for more than 21 days

still had a lower response to hyperoxia (−10.4

(11)%) than infants treated with oxygen for a

shorter period of time (−17.2 (7)%) but the

difference was not statistically significant.

HYPOXIC RESPONSE AND THE RESPIRATORY

MORBIDITY

At the 36th week of gestation both healthy

preterm infants (eight infants) and those with

respiratory distress syndrome (eight infants)

had significantly higher response than infants

who had suffered from chronic lung disease

(−20.6% −16.1%, and −4.3% respectively; see

fig 4).

HYPOXIC RESPONSE AND THE STABILITY

(MATURITY) OF VENTILATORY CONTROL

The stability of respiration measured by the

number of apnoeic/bradycardic spells (33 (49)

...
Discussion

The general principle of ‘physiological denervation’ of peripheral chemoreceptors by the inhalation of 100% oxygen, used in this study, was outlined by Dripps and Comroe in 1947\textsuperscript{12} and used for the first time by Girard and Dejours in newborn infants in 1969\textsuperscript{14} and later on by a number of other groups.\textsuperscript{15,16} The hypoxic ‘denervation’ of peripheral chemoreceptors can be accomplished by breath-by-breath alternating exposure to 100% oxygen or longer lasting inhalation of 100% oxygen (between 30 seconds and 5 minutes). In the present study we decided to use 30 seconds of exposure to 100% oxygen for several reasons. As described elsewhere,\textsuperscript{17,18} even normal preterm infants show an uneveness in the ratio between alveolar ventilation and vascular perfusion (VA/Q), whereas in infants with respiratory distress syndrome and chronic lung disease the value of VA/Q is even more disturbed due to the maldistribution and reduction of the perfusion area. This may be of some concern with respect to the possible effect of the facemask on the breathing pattern, which may in turn influence the ventilatory response to hyperoxia. As shown in our previous paper,\textsuperscript{19} however, the application of a facemask causes only temporary changes in the breathing pattern, which disappear within 40–50 breaths. As the ventilatory response to hyperoxia was measured after 3–5 minutes of breathing through a facemask, the possible effect of the spirometric device on the respiratory response could be excluded. The small retention of carbon dioxide (0.2–0.4%) in the spirometric system did not influence the response as the ventilation at the beginning and end of air breathing did not change.

The group of preterm infants investigated in this study represented a clinically heterogeneous group, with seven infants who had had respiratory distress syndrome, five with transient tachypnoea, three with chronic lung disease, and eight with recurrent apnoea. By using 30 seconds of continuous stimulation with oxygen instead of the alternate breath method we excluded the risk of insufficient stimulation of the peripheral chemoreceptors due to a delayed response time caused by structural differences in the infants’ lungs. This duration is also assumed to be sufficient to avoid the effect of hyperoxia on central respiratory control mechanisms.\textsuperscript{20}

Almost all preterm infants showed a decrease in ventilation when subjected to 100% oxygen in accordance with previous studies. The decrease in ventilation for the whole group as well as for the healthy preterm infants was almost the same at the 36th and 40th gestational week. The healthy preterm infants showed a much higher hyperoxic response (–20–3%) than healthy newborn term infants (–9–8%) tested by the same method at the age of 2–4 days. These findings indicate that the tonic peripheral chemoreceptor drive during normoxia is higher in preterm than in term infants. This is in agreement with earlier studies.\textsuperscript{15,21} The high hyperoxic sensitivity found in preterm infants has been suggested to be due to a low PO\textsubscript{2}.\textsuperscript{5} It seems to be less likely that the increased chemoreceptor drive is due to a lower resting PO\textsubscript{2}. This would instead lead to a lower drive as a tonic hyperoxia lowers the set point.\textsuperscript{8,9}

The decrease in ventilation in infants in this study was achieved mainly by a highly significant decrease in tidal volume only or by simultaneous changes in breath duration and amplitude. A similar outcome was reported by Aizad et al.\textsuperscript{15} In three infants, however, the hyperoxic inhalation resulted in apnoeas of 4 to 30 sec in duration. Neither the strength of the response nor the breathing pattern was correlated with the infants’ postnatal age and the duration of ventilatory treatment. This is in contrast with the results obtained by Alvaro et al in preterm infants at gestational week 29; they described a profound decrease in respiratory rate associated with hyperoxic inhalation.\textsuperscript{22}

As shown in our study there was no correlation between the maturity in ventilatory control, expressed by the frequency of apnoeic spells and the strength of the hyperoxic response. These results are in agreement with findings by Glotzbach et al, who were unable to find any significant relations between periodic breathing and chemoreceptor activity.\textsuperscript{23} We must be aware, however, of the fact that not all hypoxic spells are recognised in the neonatal ward and that the number of apnoeic episodes is likely to be underestimated and thus not very reliable as a measurement of the maturity of ventilatory control mechanisms.\textsuperscript{24}

We did not find any correlation between the hyperoxic response and the postnatal age, which at the 36th gestational week varied between 14 and 86 days (mean 50.7 (4) days). Our results support the proposal that the peripheral chemoreceptors mature within three weeks after birth.\textsuperscript{5,6}

The profile of the ventilatory response to hypoxia was characterised by individual variability. Most infants reacted with a decrease in ventilation and a recovery to normoxic ventilation. Only three infants showed a shallow hypoventilation followed by an increase in ventilation when subjected to hyperoxic breathing, as described by Alvaro et al.\textsuperscript{22}

The most striking finding in this study is the significantly lower hyperoxic response in infants tested at gestational week 36 who were dependent on supplemental oxygen for more than 21 days (including three infants who had chronic lung disease and one infant who was receiving supplemental oxygen for 56 days because of several instances of septicaemia) than in infants receiving supplemental oxygen for less than 21 days. When tested at
gestational week 40 the response was still lower, but the difference was not statistically significant. The indication for oxygen treatment was desaturation (oxygen saturation <85%) when breathing room air.

One possible explanation for these findings is that the oxygen breathing has attenuated the hypoxic reflex. We believe, however, that these infants sustained hypoxic episodes and, as a consequence, the peripheral chemoreceptors were set at a lower sensitivity. The significantly higher variations in blood gases in infants requiring longer oxygen treatment also indicates an instability in ventilatory control. Our findings are corroborated by the observation that newborn rats exposed to postnatal hypoxia were found to have a weak chemoreflex.8

The latency of the ventilatory response to hypoxia is mainly due to the circulation time from the alveoli to arterial chemoreceptors. This study has shown a significantly longer latency in the response to hypoxia at the 36th gestational week compared with the 40th week of gestation. It may be suggested that the differences in latency could be due to dissimilarities in circulation time. The circulation time is very short (1 ml/sec), however, and therefore seems unlikely to influence the reaction time of several seconds in duration. An alternative explanation seems possible. The adequate adaptation of respiration to changes in metabolic demands is based on an intricate interaction between the activity of the peripheral chemoreceptors and central respiratory control mechanisms. Arterial chemoreceptors afferents are known to excite respiratory neurones via specific pathways, whereas counteracting descending pathways inhibit stimulatory effects of peripheral chemoreceptors.24 Brainstem and the higher central nervous system levels play a crucial part in the control of breathing. The fact that synaptic conductivity is not yet mature26 may explain the longer reaction time to hypoxic challenge at the 36th gestational week.

This study has shown that the three infants who had chronic lung disease belonged to the group of 'low responders'. It would therefore be of great clinical interest to study further the hypoxic response in this group as the deficient hypoxic response might contribute to the high prevalence of sudden infant death syndrome among infants who have had chronic lung disease during the neonatal period.28

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