Pulse oximetry, severe retinopathy, and outcome at one year in babies of less than 28 weeks gestation

W Tin, D W A Milligan, P Pennefather, E Hey

Abstract

Aim—To determine whether differing policies with regard to the control of oxygen saturation have any impact on the number of babies who develop retinopathy of prematurity and the number surviving with or without signs of cerebral palsy at one year.


Results—Babies given enough supplemental oxygen to maintain an oxygen saturation of 88–98%, as measured by pulse oximetry, for at least the first 8 weeks of life developed retinopathy of prematurity severe enough to be treated with cryotherapy four times as often as babies only given enough oxygen to maintain an oxygen saturation of 70–90% (27.2% v 6.2%). Surviving babies were also ventilated longer (31.4 v 13.9 days), more likely to be in oxygen at a postmenstrual age of 36 weeks (46% v 18 %), and more likely to have a weight below the third centile at discharge (45% v 17%). There was no difference in the proportion who survived infancy (53% v 52%) or who later developed cerebral palsy (17% v 15%). The lowest incidence of retinopathy in the study was associated with a policy that made little use of arterial lines.

Conclusions—Attempts to keep oxygen saturation at a normal “physiological” level may do more harm than good in babies of less than 28 weeks gestation.

Keywords: prematurity; retinopathy of prematurity; cerebral palsy; oxygen saturation; pulse oximetry

Since the first large collaborative controlled trial showed that excess oxygen could cause permanent cicatricial retinopathy of prematurity in 1956, many papers have attempted to define what constitutes a safe level of arterial oxygenation. Indwelling arterial lines and sensors have been widely used to monitor arterial oxygen tension, but no controlled trial has defined what constitutes a safe level of arterial oxygen. Most had also been intermittently monitored with a transcutaneous oxygen tension monitor in the period immediately after birth, but transcutaneous monitoring was often discontinued after a few days, especially in the most immature babies with delicate, easily damaged skin. Saturation settings (which were always documented at least once an hour) have been used throughout in the analysis that follows because the transcutaneous monitors were less consistently used, especially after 1991, and often had their alarm limits muted, or adjusted so that the saturation alarm was triggered before the tension alarm.

Policy with regard to oxygen saturation was ascertained retrospectively by questioning senior medical and nursing staff, and confirmed by reference to written ward protocols and by sampling the case notes. One unit changed its policy once during the study period, and outcome in this unit has been assessed by reference to the policy in operation at the time the baby was under care. Staff always aimed to maintain saturation in the top half of the target range (particularly when the lower limit of this range was less than 85%). No formal attempt was made to document how often saturation fell outside the recommended management.
Oxygen saturation and retinopathy of prematurity

Table 1 Outcome at one year in babies of 23–27 weeks gestation at birth and its relation to target levels of fractional oxygen saturation

<table>
<thead>
<tr>
<th>Target O2 saturation</th>
<th>No of babies admitted</th>
<th>One year survivors with cerebral palsy</th>
<th>One year survivors with threshold retinopathy</th>
</tr>
</thead>
<tbody>
<tr>
<td>88–98%*</td>
<td>123</td>
<td>65 (52.8)</td>
<td>11 (16.9)</td>
</tr>
<tr>
<td>85–95%</td>
<td>235</td>
<td>128 (54.5)</td>
<td>20 (15.6)</td>
</tr>
<tr>
<td>84–94%</td>
<td>84</td>
<td>37 (44.0)</td>
<td>6 (16.2)</td>
</tr>
<tr>
<td>70–90%</td>
<td>126</td>
<td>65 (51.6)</td>
<td>10 (15.4)</td>
</tr>
</tbody>
</table>

Values in parentheses are percentages.

*Measurements made using a Nellcor pulse oximeter (functional saturation).

Ventilatory support was defined as the length of time the baby spent with an airway tube below the vocal cords, and gestational age calculated on the basis of the mother’s menstrual history as confirmed by an ultrasound assessment of fetal size at 15–19 weeks. Retinopathy was assessed by one of a small group of collaborating ophthalmologists and “threshold” retinopathy defined as stage 3 plus acute retinopathy extensive enough to merit cryotherapy using the entry criteria for the CRYOROP trial, or permanent (cicatricial) retinal scarring in a few of the preterm babies born in 1990–1991 in whom early serial screening was suboptimal. All retinal treatment was undertaken in one of two units.

Results

Babies of less than 28 weeks gestation were the focus of this study because review of a larger group showed that these are at greatest risk of threshold retinopathy (fig 1). All had been monitored with a pulse oximeter continuously throughout the time they were in receipt of respiratory support or receiving supplemental oxygen in hospital. Monitoring for hyperoxia did not usually begin until the baby was about 2–3 hours old, and babies with bronchopulmonary dysplasia who were at least 8 weeks old and whose retinal vasculature was mature received liberal oxygen supplementation. With these two exceptions, policy and practice with regard to oximeter monitoring had conformed at all times to one of four clear patterns (table 1).

Differences in policy with regard to oxygen saturation had no effect on survival or on the proportion of survivors who developed cerebral palsy, but did have a major impact on the proportion who developed threshold retinopathy. Babies nursed with a monitor set to alarm if oxygen saturation fell outside the range 70–90% had a 6.2% (95% confidence interval (CI) 1.7 to 15.0%) chance of developing retinopathy as severe as this; those nursed with a monitor set to maintain a saturation of 88–98% had a 27.7% (17.3 to 42.2%) chance of developing this degree of retinopathy (fig 2). Standardisation for minor differences in the distribution of gestational age in the four study groups did not abolish the significance of these findings (p < 0.01).

Surviving babies required ventilation and continuous supplemental oxygen for longer when the target saturation was high. Babies given enough oxygen to maintain an oxygen saturation of 88–98% were ventilated for an average of 31.4 days, and given supplemental oxygen for an average of 96 days; those maintained at saturations of 70–90% were ventilated for 13.9 days and in oxygen for 40 days (fig 3). Examination of the four management policies showed a positive correlation between target saturation and mean time to extubation.

All the 65 long term survivors nursed in enough oxygen to maintain an oximeter reading of 88–98% in the neonatal period had an arterial line inserted, and regular samples taken for an average of 12 days with a view to keeping the arterial PO2 below 12 kPa; only six of the 65 survivors managed at saturations of 70–90% ever had an arterial line inserted, or any arterial blood gas measurements taken, and only two had a line in place for more than a week. Four of the 65 long term survivors nursed in enough oxygen to maintain a saturation of 88–98% were registered as blind due to retinopathy at follow up, but none of the 65 cared for at saturations of 70–90% were so registered. The weight centiles of the babies nursed in enough oxygen to maintain a saturation of 88–98% for at least the first 8 weeks of life fell more between birth and discharge.

Figure 1 Incidence of threshold (grade 3+) retinopathy and its relation to gestation at birth in the 566 one year survivors from a population based study of 781 babies of 23–31 weeks gestation born in 1990–1991.

Figure 2 Relation between the target range for oxygen saturation and the proportion of survivors so nursed who later developed severe (grade 3+) retinopathy of prematurity. A comparison of four policies. Staff aimed to keep saturation in the upper half of the target range.

Figure 2 Limits within which oxygen saturation was allowed to vary.
(p < 0.01) than those of the babies cared for at saturations of 70–90% (fig 4). Twenty nine (45%) of the former and 11 (17%) of the latter had a weight below the third centile at discharge. Table 2 outlines other differences in management and outcome.

**Discussion**

There is, as yet, no controlled trial evidence that any of the invasive or non-invasive techniques used to monitor oxygenation in the neonatal period have any measurable impact on clinical outcome.1 We do not even know what level of arterial oxygen tension we should be aiming for. Because mortality went undocumented in the first of the large trials of oxygen administration, we do not even know if there is a price to be paid for controlling administration strictly enough to minimise the risk of severe retinopathy.2

Once equipment for monitoring transcutaneous oxygen tension became widely available, it became standard practice to aim for an oxygen tension of 6–10 kPa,11 (limits that were slightly lower than those generally used during the years when regular arterial sampling had been the main means of minimising the risk of hyperoxia12 and common to set the alarm limits at 5 and 11 kPa. However, when pulse oximeters first became available, policy became more diverse. Saturation monitors are relatively poor at detecting hyperoxia,13–25 and not all commercial oximeters have similar performance characteristics. Monitors designed to record functional saturation, such as the Nellcor oximeter, consistently read about 1.5% higher than monitors that record fractional saturation, such as the Ohmeda and Radiometer oximeters.26 Many units in the United Kingdom started setting the upper alarm limit at 95%, even though this is sometimes associated with an arterial oxygen tension of more than 90 mm Hg (12 kPa).17 It also became common practice to set the lower limit at 85%, even though a tension of 5 kPa equates more nearly to a saturation of 80%, a level still above that at which overt clinical cyanosis usually starts to become detectable. Similar divergence of practice has been documented in the United States, where surveys suggest that over a third of units currently set alarm limits that allow saturation to exceed 95%,27,28

Only a randomised control trial would establish whether some other unrecognised factor was responsible for the difference in the incidence of retinopathy found in this study. The present observational study does, however, strengthen the case for such a trial. The only trial designed to compare two strategies of oxygen administration from birth undertaken to date has been interpreted as suggesting that a restrictive policy may reduce pulmonary oxygen toxicity without increasing mortality.21–24 The recent STOP-ROP trial, in which babies were nursed in enough oxygen to maintain a saturation of 96–99% rather than 89–94% once prethreshold retinopathy had been detected (in the hope that this would limit the progression of retinal disease), also uncovered evidence that liberal oxygen use may cause pulmonary damage.25
Oxygen delivery depends on cardiac output and on tissue perfusion as much as on arterial saturation, so it is arguable that, although arterial tension (or some reliable surrogate of this) is probably the best way of minimising the retinal hazard associated with hypoxaemia in the very preterm baby, measurement of central venous saturation provides a better measure of the adequacy of tissue delivery. The babies in this study who were managed at saturations of 70–90% seldom had any arterial lines inserted or arterial blood gas measurements taken. Ventilator support was managed by taking, every six to eight hours, capillary or venous blood gas samples supplemented on occasion by transcutaneous PCO₂ measurements, while Doppler ultrasound was used to monitor systolic blood pressure non-invasively. Because relatively little blood was taken for laboratory purposes, only 13 of the 65 long term survivors needed more than two blood transfusions before discharge (table 2). The result certainly calls into question the unsubstantiated belief that arterial sampling is essential if the risk of retinopathy is to be minimised. Oximeters and indwelling sensors are actually of comparable accuracy when saturation is in the range 70–90%, Preterm babies requiring many blood transfusions have an increased incidence of retinopathy, and it has been suggested that this association could be causal, although a recent small controlled trial found no support for such a view. Were such a link to be established, it could even be argued that regular arterial sampling, by increasing the number of transfusions required, actually increases the risk of retinopathy.

Whatever the explanation for the variation in the amount of threshold retinopathy seen in this observational study, there can be little doubt that policy differences with regard to oxygenation had an impact on the amount of ventilatory support provided and therefore on the total cost of care before discharge (fig 3). In this respect the range of saturation deemed acceptable by the most permissive of the four policies may have been as influential as the degree of desaturation allowed because the mean saturation achieved with this policy, in the first two days after extubation, exceeded 82% in all but eight of the 65 long term survivors. It has been suggested that fluctuation in oxygenation, rather than the absolute level experienced, does most to render the preterm retina vulnerable to severe retinopathy. While such a suggestion may seem hard to reconcile with the finding that threshold retinopathy was least often seen with the policy that allowed the widest range of saturation, it needs to be remembered that, because of the shape of the dissociation curve, a small change in saturation masks a wider change in partial pressure when saturation exceeds 90% than when saturation is lower than this.

It has been reported that babies with subclinical cyanosis grow less well, but that was not true of the babies in this observational study (fig 4). The present finding does not invalidate this general proposition, but does imply that a sustained period of ventilatory support can seriously compromise growth in the period immediately after birth, even if the baby is better oxygenated during this period and also offered parenteral nutrition. The wide range of clinical practice shown by this study has other implications. It has recently become common practice to adjust for differences in initial disease severity when assessing the care provided by different neonatal units, or groups of units. The simplest and most widely used system (the Clinical Risk Index for Babies (CRIB)) uses information on the amount of oxygen given to the baby in the first 12 hours of life, but any baby cared for in a unit that aims for an oxygen saturation of 70–90% will inevitably generate a CRIB score that is lower than would have been assigned to the same baby in a unit that aims for an oxygen saturation of 88–98%. As information on oxygen requirement accounts for up to 39% of the current score, failure to allow for this difference undermines the generalisability of the score.

Current guidelines suggest that retinopathy should be looked for in all babies of less than 32 weeks gestation, implying that these are the babies most in need of monitoring for hyperoxia. However, no case of acute threshold retinopathy has been seen in a baby of more than 30 completed weeks gestation in the north

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**Table 2** Early nursing management and cranial ultrasound findings in the one year survivors nursed in enough supplemental oxygen to maintain an oximeter saturation of 88–98% or 70–90% throughout the first 8 weeks of life.

<table>
<thead>
<tr>
<th>88–98%</th>
<th>70–90%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of one year survivors</td>
<td>65</td>
</tr>
<tr>
<td>Gestation (weeks)</td>
<td>26.4 (25.8–27.3)</td>
</tr>
<tr>
<td>Birth weight (g)</td>
<td>910 (810–1018)</td>
</tr>
<tr>
<td>Male sex (%)</td>
<td>55</td>
</tr>
<tr>
<td>Number given 20 µg/kg IM vitamin E at birth</td>
<td>65</td>
</tr>
<tr>
<td>Number given IV amino acids ± lipid</td>
<td>62</td>
</tr>
<tr>
<td>Mean time given (in those so managed) (days)</td>
<td>19.2</td>
</tr>
<tr>
<td>Number with an arterial line in place ≥7 days</td>
<td>49</td>
</tr>
<tr>
<td>Blood transfusions given</td>
<td>5 (3–8)</td>
</tr>
<tr>
<td>Number given &gt;2 transfusions</td>
<td>55</td>
</tr>
<tr>
<td>First fully enterally fed (days)</td>
<td>25 (18–34)</td>
</tr>
<tr>
<td>Surgery for necrotising enterocolitis</td>
<td>0</td>
</tr>
<tr>
<td>Cerebral ultrasound findings</td>
<td></td>
</tr>
<tr>
<td>Intraventricular haemorrhage</td>
<td>4</td>
</tr>
<tr>
<td>Haemorrhagic ventriculomegaly</td>
<td>5</td>
</tr>
<tr>
<td>Periventricular leucomalacia</td>
<td>5</td>
</tr>
</tbody>
</table>

Where applicable, values are median (interquartile range). IM, Intramuscular; IV, intravenous.
of England in the last eight years, and only two cases at 30 weeks gestation (one in every 300 births at this gestation). A six year study in Birmingham, also found much the same thing, and did studies from Tennessee in America and New South Wales in Australia. As there is no evidence that light for dates babies are any more prone to severe retinopathy than normal weight babies, there is no logic in using birth weight, as well as gestational age, when defining which babies merit monitoring for hyperoxia (or ophthalmic screening) as long as gestational age has been reliably determined.

Although healthy preterm babies maintain a saturation in excess of 95% most of the time, we conclude that artificial attempts to keep arterial oxygenation at a “physiological” level may do more harm than good in the first few weeks of life. The issue has been raised previously and certainly deserves more study, particularly in babies of less than 30 weeks gestation. Such levels are certainly higher than the retina, or the brain, would normally experience during fetal life.

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