Very long apnea events in preterm infants

Mary A. Mohr,1 Brooke D. Vergales,2 Hoshik Lee,1,3 Matthew T. Clark,4 Douglas E. Lake,4,5 Anne C. Mennen,1 John Kattwinkel,2 Robert A. Sinkin,2 J. Randall Moorman,4,6,7 Karen D. Fairchild,2 and John B. Delos1

1Department of Physics, College of William and Mary, Williamsburg, Virginia; 2Department of Pediatrics (Neonatology), University of Virginia, Charlottesville, Virginia; 3Samsung Advanced Institute of Technology, Suwon, South Korea; 4Division of Cardiovascular Medicine, University of Virginia, Charlottesville, Virginia; 5Department of Statistics, University of Virginia, Charlottesville, Virginia; 6Department of Biomedical Engineering, University of Virginia, Charlottesville, Virginia; and 7Department of Molecular Physiology, University of Virginia, Charlottesville, Virginia

Submitted 13 February 2014; accepted in final form 22 December 2014

Very long apnea events in preterm infants. J Appl Physiol 118: 558–568, 2015. First published December 30, 2014; doi:10.1152/japplphysiol.00144.2014.—Apnea is nearly universal among very low birth weight (VLBW) infants, and the associated bradycardia and desaturation may have detrimental consequences. We describe here very long (>60 s) central apnea events (VLAs) with bradycardia and desaturation, discovered using a computerized detection system applied to our database of over 100 infant years of electronic signals. Eighty-six VLAs occurred in 29 out of 335 VLBW infants. Eighteen of the 29 infants had a clinical event or condition possibly related to the VLA. Most VLAs occurred while infants were on nasal continuous positive airway pressure, supplemental oxygen, and caffeine. Apnea alarms on the bedside monitor activated in 66% of events, on average 28 s after cessation of breathing. Bradycardia alarms activated late, on average 64 s after cessation of breathing. Before VLAs oxygen saturation was unusually high, and during VLAs oxygen saturation and heart rate fell unusually slowly. We give measures of the relative severity of VLAs and theoretical calculations that describe the rate of decrease of oxygen saturation. A clinical conclusion is that very long apnea (VLA) events with bradycardia and desaturation are not rare. Apnea alarms failed to activate for about one-third of VLAs. It appears that neonatal intensive care unit (NICU) personnel respond quickly to bradycardia alarms but not consistently to apnea alarms. We speculate that more reliable apnea detection systems would improve patient safety in the NICU. A physiological conclusion is that the slow decrease of oxygen saturation is consistent with a physiological model based on assumed high values of initial oxygen saturation.

apnea; neonatal; hypoxia; bradycardia

MORE THAN TWO DECADES AGO, Southall et al. (60) reported prolonged apnea and bradycardia events in preterm infants cared for in neonatal intensive care units (NICUs). In their study of 14 babies observed for 24-h periods, the authors were disturbed to report cessations of breathing as long as 213 s and to report that extended apnea events were often not detected or reported. Present practices (13, 18, 36) hold that any cessation of breathing longer than 10 s, if accompanied by bradycardia and oxygen desaturation, is an event that should be considered clinically significant (as should any apnea longer than 20 s regardless of bradycardia or desaturation). Most events last less than half a minute, either self-resolving or terminating when a caregiver stimulates the infant in response to a monitor alarm. Very long apnea (VLA) events may reflect abnormal physiology, impending illness, alarm failure, and/or failure of bedside clinicians to rescue infants from prolonged apneic spells.

Nearly all very low birthweight infants (VLBW; <1,500 g) experience apnea due to inherent brainstem and peripheral chemoreceptor immaturity, and pathologic processes such as sepsis, respiratory failure, intracranial hemorrhage, and seizures may increase the number or severity of apnea events. Apnea events may lead to tissue hypoxemia and, especially if prolonged, severe, or frequent, might contribute to adverse short- and long-term outcomes, including neurodevelopmental impairment. Preventing or minimizing bradycardia and desaturation related to apnea should therefore be a priority in NICU care. Steps toward this goal include understanding physiologic associations with severe apnea and optimizing monitor alarm systems and caregiver responses.

Apnea research was previously hindered by inaccurate and inefficient recording of events. Our group recently developed an automated apnea detection system that analyzes chest impedance (CI) waveform signals from all NICU bedside monitors, filters out cardiac and motion artifacts, and accurately identifies central apnea events accompanied by bradycardia and oxygen desaturation (ABD) (34). Using this novel system, together with a database consisting of over 100 infant years of electronic signals, we sought to quantitate and characterize VLA events in VLBW infants, defined as cessation of breathing >60 s accompanied by bradycardia (heart rate <100 beats/min) and oxygen desaturation (SpO2 <80%).

In this article we report descriptive clinical aspects and quantitative physiological aspects of very long central apnea events. We seek answers to the following questions. 1) Do infants really stop breathing for more than 60 s? How frequent are such events in our population? 2) Are the babies who have VLAs in some way different from those who do not, or is it the events that are different? 3) How significant are these events for the health of the infants? 4) Why do these events still occur? 5) We observe in these events very slow decline of heart rate and oxygen saturation. Can we give a physiological explanation for either of them? This article is organized such that the primary clinical results are in the main text, and everything mathematical is in APPENDICES B–D.

Neonatal apnea has previously been addressed from a variety of perspectives: physiology of respiration (8, 25), apnea of prematurity (6, 7, 13, 18, 27, 31, 35, 37, 41, 44, 46, 63), long apnea events (1, 10, 30, 45, 61), failure of monitoring (9, 22, 39, 60, 62, 66), the effect of saturation on apnea (23, 24), and...
the effect of apnea on SpO₂, and bradycardia (16, 20, 28, 42, 43, 57). However, no previous workers have had access to such a large dataset (~100 infant years, 335 VLBW infants, ~20,000 apnea events), and none have connected clinical observations to the quantitative physiological theory that we give in the appendices.

METHODS

Patients

We studied all 335 VLBW (<1,500 g) infants admitted from January 2009 through March 2011 to the 45 bed University of Virginia (UVa) NICU, a quaternary care facility that has more than 500 admissions per year, about one quarter of them VLBW. The study was approved by the UVa Institutional Review Board with waiver of consent and subsequently by the William and Mary Protection of Human Subjects Committee.

Bedside Monitor Data Collection

We collected and stored all waveform, vital sign, and alarm data from all NICU bedside monitors (GE Medical, models Solar 8000M and I and Dash 3000) via a BedMaster central network server (Excel Medical, Jupiter, FL). CI waveforms were collected at 60 Hz, three channels of electrocardiogram waveforms were collected at 240 Hz, and the photoplethysmographic waveform was collected at 120 Hz. In addition, vital sign data (heart rate, CI-derived respiratory rate, and pulse oximetry-derived O₂ saturation recorded every 2 s) were collected. Also, all bedside monitor alarms were collected. Bradycardia alarms sound when the heart rate goes below a set limit (typically 90 beats/min) for at least eight heartbeats, oxygen alarms sound when SpO₂ is lower than a set limit (typically 85%) for at least 15 s, and apnea alarms sound after 15 s of no monitor-detected breaths in the CI signal.

Apnea Analysis

We previously developed and validated a new algorithm for detecting central apnea (34). The algorithm filters the cardiac artifact and low-frequency fluctuations from the CI signal, analyzes the residual fluctuations, and produces, at quarter-second intervals, the probability that the infant is having a central apnea event at that moment. Careful validation indicated that the algorithm agrees with the expert opinion of clinicians in ~90% of central apnea events. For the current analysis, we focused on central apnea events associated with bradycardia and desaturation. The algorithm’s threshold for bradycardia is a heart rate <100 beats/min and for oxygen desaturation SpO₂ <80%, for any duration. The events are categorized as ABdN, with “n” signifying the minimum length of cessation of breathing in seconds (e.g., ABD60 means cessation of breathing was 60 s or more, and ABd30–40 means cessation of breathing was between 30 and 40 s). For all infants in our database, we have recorded the date, time, and duration of every ABD10 event (10 s or more). Those ABD60 events that survived the validation described below are called VLAs.

We also collected several random samples of 10- to 20-s and 30- to 40-s apnea events to compare their properties with those of VLAs.

Identification of VLAs

In VLBW infants the computer algorithm identified 461 very long central apnea events (~60 s) accompanied by bradycardia and desaturation in which all waveforms were available in VLBW infants. Each such event was examined by at least two clinicians who are expert on interpretation of the signals. Two groups of two clinicians independently reviewed half of the candidate VLA events, followed by consensus review in equivocal cases. It was found that some algorithmically detected events were artifacts caused by weak signals. In many other cases, the CI waveform signal was not completely flat but had some small or brief fluctuation that might represent one or several shallow breaths. Such events appear to be clinically relevant, and we discuss them in APPENDIX A. Here, however, we wish to focus on the less common indisputable events. Candidate events were accepted as VLAs only if there was an acceptable CI tracing before, during, and after the event, the event was accompanied by any duration of bradycardia (<100 beats/min) and desaturation (<80%), the cessation of breathing lasted at least 60 s, and the filtered CI had no artifacts, no indication of shallow breaths, and no twitches in CI having an associated increase in heart rate. With this stringent process, 86 VLA events were identified in 29 infants (Fig. 1A and APPENDIX A). Study Group (I) and Comparison Group (N)

The result of this validation was that algorithmically-detected ABD60 events were divided into three categories: 1) VLAs, which we regard as indisputable cessations of breathing of 60 s or more; 2) “questionable” 60-s events involving, for example, long apnea-hypopnea combinations, which we regard as clinically significant but not indisputable VLAs (discussed in APPENDIX A); and 3) excluded events. The same process identified the study group (I), consisting of those 29 infants who had “Indisputable” events. We chose a comparison group (N), consisting of those infants whose electronic signals “Never” showed a 60-s event of any type, validated, questionable, or excluded. In RESULTS we compare apnea rates and other characteristics of these
Table 1. Infant characteristics

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>VLA (n = 29)</th>
<th>No VLA (n = 306)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gestational age, wk</td>
<td>26.5 ± 1</td>
<td>27.5 ± 3.1</td>
<td>0.09</td>
</tr>
<tr>
<td>Birth weight, g</td>
<td>952 ± 249</td>
<td>908 ± 299</td>
<td>0.53</td>
</tr>
<tr>
<td>ELBW</td>
<td>19 (66%)</td>
<td>150 (49%)</td>
<td>0.12</td>
</tr>
<tr>
<td>Male</td>
<td>14 (48%)</td>
<td>154 (50%)</td>
<td>0.85</td>
</tr>
<tr>
<td>Mechanical ventilation at any time</td>
<td>23 (79%)</td>
<td>214 (70%)</td>
<td>0.39</td>
</tr>
<tr>
<td>Death before NICU discharge</td>
<td>1 (3%)</td>
<td>31 (10%)</td>
<td>0.34</td>
</tr>
<tr>
<td>Length of NICU stay, days</td>
<td>78 ± 27</td>
<td>63 ± 42</td>
<td>0.02</td>
</tr>
</tbody>
</table>

VLA, very long apnea; ELBW, extremely low birth weight; NICU, neonatal intensive care unit.

groups. (We also report the number of infants who never had a computer-detected ABD10.)

**Medical Record Review**

Demographic data collected from the medical record included gestational age at birth, postmenstrual age (PMA) at the time of events, birthweight, and gender. Respiratory data collected from the medical record included the type of respiratory support, supplemental oxygen, administration of caffeine (if any), and the time on and off respiratory support. Times on mechanical ventilation were excluded. Clinical conditions or events potentially associated with VLA events were recorded, including severe intraventricular hemorrhage, severe gastroesophageal reflux, suspected sepsis (cultures obtained and antibiotics initiated within 24 h of VLA), respiratory failure (intubation and mechanical ventilation within 12 h after the VLA), and death. Survival to NICU discharge and length of NICU stay were recorded.

**RESULTS**

**Patient Characteristics and Clinical Associations**

Data from 335 VLBW infants were analyzed, and 86 VLA events in 29 infants were validated by the process described in METHODS and in APPENDIX A. The estimated frequency of these VLA events is 1 per 70 infant days. (If questionable events were included, the estimate of the frequency would be about four times larger; see APPENDIX A.) Characteristics of the infants are given in Table 1. The number of events per infant ranged from 1 to 19 (Fig. 2). In most respects, infants with VLA were not significantly different from other VLBW infants (Table 1). Infants with VLA events had significantly longer NICU stays than those without (P = 0.02), and their average gestational age was slightly lower (P = 0.09; Table 1). Over half of the VLA infants fell in the gestational age (GA) range of 25–27 wk (Fig. 3A) and 84% of VLAs occurred <4 wk from birth (Fig. 3B). Seventy-nine percent of the VLAs occurred while infants were on continuous positive airway pressure (CPAP), usually with supplemental oxygen (Fig. 4). All 29 infants were on caffeine therapy at some time during their NICU stay, one

**Fig. 3.** A: histogram of number of VLA events and number of VLA infants vs. gestational age (GA) at birth. For comparison the number of infants vs. GA for the entire population of very low birth weight (VLBW) infants in our database is also shown. For example, among 26-wk GA infants, 8 of the 37 VLBW infants had at least 1 event, and there were 15 total events. B: histogram of number of VLA events vs. chronological age (CA) at time of event for those infants having 1 or more VLAs (left scale). For comparison the number of VLBW infants who were in the University of Virginia neonatal intensive care unit (NICU) at that CA is also shown (right scale). For example, at 1 wk of age, there were 25 VLAs among 266 VLBW infants.

**Fig. 4.** Histogram showing respiratory support of VLA infants at the times of their events. From left to right: room air, nasal cannula (NC) without supplemental oxygen (sup O2), nasal cannula with supplemental oxygen, continuous positive airway pressure (CPAP) without supplemental oxygen, and CPAP with supplemental oxygen. Most VLA events occurred while infants were on CPAP with supplemental oxygen.
Table 2. Clinical conditions

<table>
<thead>
<tr>
<th>#Infants</th>
<th>Clinical Association</th>
<th>GA</th>
<th>#VLAs</th>
<th>PMA</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Died 48 h after last VLA</td>
<td>27</td>
<td>19</td>
<td>28–32</td>
</tr>
<tr>
<td>1</td>
<td>Posthemorrhagic hydrocephalus, ventricular tap performed before VLA cluster</td>
<td>26</td>
<td>3</td>
<td>31</td>
</tr>
<tr>
<td>1</td>
<td>Grade IV intraventricular hemorrhage</td>
<td>26</td>
<td>2</td>
<td>30</td>
</tr>
<tr>
<td>1</td>
<td>On prostaglandins for congenital heart disease</td>
<td>29</td>
<td>3</td>
<td>31</td>
</tr>
<tr>
<td>2</td>
<td>On medication for severe GE reflux</td>
<td>25</td>
<td>3–12</td>
<td>27–29</td>
</tr>
<tr>
<td>4</td>
<td>Antibiotics started for suspected sepsis within 24 h of VLA</td>
<td>23–29</td>
<td>1–19</td>
<td>26–31</td>
</tr>
<tr>
<td>10</td>
<td>Receiving antibiotics at time of event</td>
<td>25–30</td>
<td>1–19</td>
<td>25–32</td>
</tr>
<tr>
<td>7</td>
<td>Intubated within 12 h of VLA</td>
<td>23–29</td>
<td>1–4</td>
<td>24–31</td>
</tr>
<tr>
<td>11</td>
<td>None identified, other than prematurity</td>
<td>24–31</td>
<td>1–5</td>
<td>27–34</td>
</tr>
</tbody>
</table>

GA, gestational age; PMA, postmenstrual age; GE, gastroesophageal.

VLA occurred 4 days after caffeine was discontinued. Two infants had VLA on the day of birth, before caffeine being initiated. No VLA events occurred within 10 days of NICU discharge.

For 18 of the 29 infants with VLAs, medical record review identified clinical conditions (separate from prematurity, which was present in all subjects), which are possibly associated with apnea (Table 2). Ten infants were receiving antibiotics at the time of the VLA, four were started on antibiotics within 24 h of the VLA for suspected sepsis (although none had proven sepsicemia), seven were intubated within 12 h of the VLA, and six had other conditions. The infant with the greatest number of events (19) died suddenly and unexpectedly for unclear reasons. He was stable on full feeds, CPAP, and caffeine at the time of death, and autopsy was declined. The other infant with a large number of events (12 VLAs at 2–3 wk of age) was on erythromycin for severe gastroesophageal reflux. No VLAs occurred beyond 18 days of age, and he was discharged home at 78 days of age. We could see no systematic differences in the characteristics of the VLAs associated with these different clinical situations.

For the other 11 infants (38% of the VLA infants), we could not identify a clinical condition that appeared to be a cause or effect or to have an association with the VLA. These 11 differ only slightly from the 18 with clinical associations. The infants without clinical associations had slightly higher GAs, but the difference was not significant ($P = 0.16$). Not surprisingly, these 11 infants also had shorter stays in the NICU than the other 18 infants ($P = 0.02$). There were no differences in the rate of fall of heart rate and $SpO_2$ between VLA infants with clinical associations and those without.

Table 3. Study group characteristics

<table>
<thead>
<tr>
<th>Groups</th>
<th>Number of ABD10 Tags</th>
<th>Number of Infants</th>
<th>%All VLBW Babies</th>
<th>Tags per Baby-Day of Data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group I: infants with indisputable 60-s VLAs</td>
<td>6,617</td>
<td>29</td>
<td>9</td>
<td>4.29</td>
</tr>
<tr>
<td>Group N: infants who Never had a ABD60</td>
<td>6,300</td>
<td>199</td>
<td>59</td>
<td>1.52</td>
</tr>
<tr>
<td>Infants who never had a computer-detected ABD10 (included in group N)</td>
<td>0</td>
<td>62</td>
<td>19</td>
<td>0</td>
</tr>
<tr>
<td>All VLBW infants</td>
<td>30,793</td>
<td>335</td>
<td>100</td>
<td>2.94</td>
</tr>
</tbody>
</table>

ABD, apnea events accompanied by bradycardia and oxygen desaturation.

Apnea Rates in Different Groups of Infants

ABD10 tags in our database identify every computer-detected cessation of breathing lasting at least 10 s, when also accompanied by bradycardia and desaturation. The majority of VLBW infants in our population never had any computer-detected 60-s ABD event (group N). This group of infants had a low rate of short apnea events, and 31% of them never had even an ABD10 event. On the other hand, infants who had indisputable VLAs (group I) also had many short apnea events. The total number of ABD10 tags for the study group (I) and the comparison group (N) of VLBW infants are in Table 3.

Included in the 199 infants in group N are 62 infants who never had a computer-detected apnea lasting at least 10 s with bradycardia and desaturation (ABD10). Thus in our population, 81% of VLBWs had at least one detected ABD10, while 19% did not.

Bedside Monitor Alarms in VLA Events

Apnea alarms are supposed to activate 15 s from the beginning of cessation of breathing; in fact they activated in 66% of these VLAs but failed to activate in 34% of them. Since bradycardia alarms activated in 57% of VLAs, and low oxygen alarms activated in 81%, the end result is that at least one alarm activated in 97% of VLAs. The monitors and the algorithm use slightly different criteria for bradycardia and desaturation, so it is not surprising that these alarms did not go off during all VLA events. (For additional comparisons among algorithm detection, monitor alarms, and nursing records, see Ref. 66.)

For the cases in which the bedside monitor gave an apnea or bradycardia alarm, Fig. 5 shows histograms of lengths of time from the start of the computer-detected apnea to the alarm and of times from the alarm to the end of the detected apnea. If an apnea alarm activated, it did so rather promptly, usually <25 s from the beginning of the computer-detected apnea. If a bradycardia alarm sounded, it usually occurred long after the cessation of breathing (Fig. 5), consistent with the slow fall of heart rate in these events. Shortly after the bradycardia alarm sounded (but frequently quite long after the apnea alarm), the apnea ended. We do not know which events were self-resolved and which were ended by staff response, but this observation is consistent with many others that suggest that NICU personnel respond quickly to bradycardia alarms and not necessarily to apnea alarms (9, 22, 39, 60, 62, 66).

Characteristics of Short Apnea Events in Infants with and without VLA

To determine whether short apnea events in infants who have VLAs are different from short apnea events in other infants we collected a random sample of 100 ABD30–40
compared with the decline in heart rate and SpO2 during shorter events. In VLA events, the heart rate falls slowly, so the bradycardia alarm is much delayed, typically sounding 60–70 s after the apnea begins. Apneas usually end a short time after the bradycardia alarm.

Comparison of Short and Long Events in Infants Having VLAs

A striking characteristic of the long apnea events is that the heart rate and oxygen saturation fall later and more slowly compared with the decline in heart rate and SpO2 during shorter ABD events. This is shown in Fig. 7; 100 ABD10–20 and 100 ABD30–40 events were randomly selected only from infants who had indisputable VLAs (group I). The events were chosen such that the infant’s PMA was <31 wk at the time of the event. The average of multiple events is plotted, and the standard error of the means is marked. The heart rate fell below 100 beats/min and the oxygen saturation fell below 80% in every individual event. However, these thresholds were crossed at different times in different events, so the averages do not necessarily fall below the thresholds.

Just before the VLA begins, the oxygen saturation is significantly higher than the saturation before shorter ABDs (other somewhat related studies include Refs. 16, 20, 23, 24, 28, 42, 43, 57). The median oxygen saturations in the 60 s before the apnea begins are 92% for ABD10–20 events; 93% for ABD30–40 events; 96% for VLAs where apnea lasts from 60 to 80 s; and 97% for VLAs where apnea lasts longer than 80 s. Oxygen saturations before VLAs was significantly greater than saturation before the shorter apnea events (Wilcoxon rank-sum, P << 0.001).

Additional details about the rate of fall of SpO2 are given in APPENDIXES B–D. 1) Because of the slow fall, it is not immediately clear whether the severity of apnea events increases in proportion to their duration; measures of severity of apnea events of various durations are discussed in APPENDIX B, and we conclude that the duration is indeed a fair measure of the deficit of oxygen transported in the blood. 2) In APPENDIX C, we show that before a VLA, the infants are, on average, breathing slightly faster than they are before a short apnea, and we show that the median SpO2 is gradually rising in the 5 min before the VLA. 3) A quantitative theory for the rate of fall of SpO2 is given in APPENDIX D; the results of this theoretical calculation are shown in Fig. 7.

DISCUSSION

While apnea is a normal developmental phenomenon in preterm infants, exaggerated apnea is an important problem that may reflect or lead to adverse events or outcomes. These events are interesting from a physiologic perspective and important from a patient safety perspective; thus we turn to the questions posed in the Introduction.

1) How frequent are the events? VLA events are not rare. We used very stringent criteria to define a VLA. We found that infants show no evidence of breathing for durations of up to 132 s. These events occur with a frequency of 1 per 70 infant days in our population.

2) Are the infants with VLA different? VLA infants differ from non-VLA infants in that they have a much higher rate of short apnea events and presumably have less developed respiratory control. We found short apnea events in VLA infants to be indistinguishable from short apnea events in non-VLA infants.

3) How clinically significant are the events? It is difficult to establish the effect of VLAs on the short-term or long-term health of infants. In APPENDIX B we show that, despite the slow drop of SpO2 in VLAs, the available measure of deficit in oxygen transport is approximately proportional to the duration of the apneas.
of the cessation of breathing, i.e., 90-s apnea events result in approximately three times the oxygen transport deficit of 30-s apnea events (which are already considered by some to be “extreme”; Ref. 30). Our information on clinical consequences leads to ambiguous results. On one end of the VLA spectrum, an apparently healthy preterm infant had 19 events over a 3-wk period and eventually died following a respiratory arrest from which he could not be resuscitated. However, other VLA infants had normal courses in the NICU. Lacking postdischarge objective neurodevelopmental follow-up testing on infants, we can only say that, with the possible exception of the infant who died, no clinical consequences could be proved to be caused by VLAs. Nevertheless, VLAs cannot be good for infants, and their frequency is disturbing.

4) Why do VLAs occur? Part of the explanation of why VLAs occur is that apnea alarms fail to activate in about one-third of these events. In addition, we previously showed that about two-thirds of all apnea alarms are false alarms (34), so alarm fatigue or alarm distrust may also be a factor (66). Clinical experience and our data suggest that bedside caregivers respond quickly to bradycardia alarms, which are “crisis” alarms on the monitors in the UVa NICU (in contrast to apnea “warning” alarms), but as a result of the slow fall of heart rate these alarms are delayed. We believe that more reliable apnea alarms, with fewer false negatives and false positives, would be helpful to NICU personnel and might have a significant beneficial effect on the health of infants by reducing hypoxemia and bradycardia.

5) What is the physiological mechanism? We show that the respiratory rate is somewhat elevated and $SpO_2$ steadily rises in the 5 min before a VLA. Accordingly, $SpO_2$ is unusually high at the commencement of the event (see Comparison of Short and Long Events in Infants Having VLAs and APPENDIX C). Combining this observation with the hypothesis that venous oxygen is also somewhat elevated leads to a quantitative description of the average rate of fall of $SpO_2$ with time (presented in APPENDIX D).

CONCLUSION

Prolonged apnea events should be preventable. Since these events could have detrimental consequences and are often not noticed and/or not recorded in the medical record, improvements in bedside monitor apnea detection, alarm systems, and caregiver responses are key to providing a safer environment for hospitalized preterm infants.

APPENDIX A: QUESTIONABLE AND EXCLUDED EVENTS

Clinicians reviewed 461 ABD60 events in VLBW infants for which all waveforms were available. Of these, 20 had poor and uninterpretable signals, and 38 were recognized as apnea events of less than a 60-s duration. These 58 events were excluded. The remaining 317 events failed our strict VLA criteria for the following reasons: there were fluctuations in the CI with a frequency lower than typical breathing, the origin of these fluctuations is unknown, but they certainly do not represent normal breathing; there were small shallow breaths present (hypopnea); the event was actually two separate apnea events; or there was a small wave in the CI which was followed by a rise in the heart rate. In Fig. 1B the two small fluctuations in the CI are followed by an increase in heart rate, so they are possibly small breaths and this event was excluded.

We consider these 317 “questionable” cases to be examples of failure of respiratory regulation lasting at least 60 s, but we do not include them in our category of indisputable VLAs. If these 317 events have clinical significance comparable to that of the indisputable VLAs, then the total number of events is more than four times larger than stated in Patient Characteristics and Clinical Associations, i.e., 1 per 16 infant days in our population.

APPENDIX B: MEASURES OF SEVERITY OF A VLA

A balanced perspective on the significance of VLAs is needed. One difficulty in assessing their impact arises from the slow fall in oxygen saturation and heart rate during these events; consequently, it is not clear that the duration of the apnea is a good measure of the severity of a VLA. Indeed, Alvarro and Rigatto (2) suggested that “The length of the respiratory pause is not a very useful indicator of [its] severity. . . . For this reason, many centers . . . have decided to rely on heart rate as the primary indicator of severity.” Furthermore, the optimal oxygen saturation for infants is a matter of current debate (4, 5, 11, 15, 40, 53, 58, 59, 65, 70).

To compare VLAs with shorter apnea events, we defined and computed measures of the oxygen deficit and the heartbeat deficit. These measures are areas on graphs of $SpO_2$ or heart rate vs. time.

Method

Clinicians note the depth and the duration of desaturation events. However, automated data analysis systems could provide superior measures, for example, the area enclosed by the $SpO_2$ curve below a chosen threshold. This area is related to the deficit in the quantity of oxygen carried by the blood. If normal oxygen transport in the blood is M ml/min, and the measured area under the baseline is A % min, then $MA/100$ is the oxygen deficit resulting from the apnea. For example, if normal transport were 5 ml O2/min (Ref. 14, p. 970), then an area of 22% min means that over the duration of the apnea event, the blood transported 1.1 ml of O2 less than it would have in the absence of the apnea.
Results

We examined long and short apnea events in “Infant 1,” the infant who had 19 VLAs. Random sets of ABD10–20 and ABD30–40 events from this infant were collected. With the use of the method described above, the “oxygen deficit” and the “heartbeat deficit” were calculated in these shorter events and in the 19 VLAs. The results are shown in Fig. B1B. The areas increase approximately linearly with the duration of the apnea. The sample correlation coefficients are 0.81 for heart rate ($P < 0.001$) and 0.88 for oxygen saturation ($P < 0.001$). This indicates that the area measure is providing some information independent of the duration, but area and duration are well correlated.

Lower baselines give smaller areas. A baseline of 80% for oxygen saturation gives a slope that is about half of what our baseline gave (0.18%-min$^{-1}$), but the correlation coefficient is comparable (0.79, $P < 0.001$).

Discussion

The graphs in Fig. B1B show measures of the relative significance of VLAs vs. short apnea events. The roughly linear trend suggests that a 90-s apnea results in about three times the oxygen deficit and three times the heartbeat deficit of a 30-s apnea. (This fact is not obvious a priori, because heart rate and SpO$_2$ decrease slowly in long apnea events.) Such deficits could be reduced by earlier and more accurate identification of apnea events.

APPENDIX C: OXYGENATION BEFORE VLAS

Here we give evidence that infants are raising the oxygen level in their blood before a VLA. 1) We show that those infants who have VLAs breathe more rapidly before a VLA than before a short apnea. 2) We show that the median SpO$_2$ rises gradually in the minutes before a VLA.

Method

The respiration rate reported by monitors is notoriously unreliable because of artifacts in the CI signal. Using our filtered CI (34), from which most artifacts have been removed, we found zero-crossings of the filtered CI and maxima and minima between each zero-crossing. We define the breath length to be the time between two successive minima. To prevent residual fluctuations around zero to be mistakenly interpreted as breaths, we applied two additional criteria. 1) The probability of apnea must be less than 0.5 at the times of the minima. 2) Consecutive maxima and minima must be at least 0.1 times the standard deviation of the signal.

We examined apnea events only in those infants who have VLAs. We obtained the respiration rate for 100 breaths preceding each of the 86 VLAs and also for 100 randomly selected apnea events of duration 10–20 s in the same infants.
The best way to make the comparison is with the cumulative histograms of breath lengths (Fig. C1A). This shows for each duration $T$ the fraction of breaths having duration less than $T$. That curve was fit to a rational function of $T$, and the derivative of that rational function was computed, giving what we call the continuous-fit histogram.

For $\text{SpO}_2$, we calculated the median in the 5 min leading up to the 86 VLAs and in the 5 min leading up to the same randomly selected apnea events of 10- to 20-s duration (Fig. C1B).

**Result**

The resulting graphs are shown in Fig. C1. From the cumulative histograms it is clear that the breath durations leading up to a VLA are generally shorter than those leading up to a 10- to 20-s apnea. Comparing the two distributions using the Kolmogorov-Smirnov test gives a $P$ value of 0.0024, confirming that the two distributions are different. Barely visible behind the cumulative distributions in Fig. C1A is a smooth curve representing the rational-function fit. The derivatives of these, the continuous-fit histograms, are also shown. Breaths before VLAs are on average 3% shorter, in median 3.3% shorter, and in mode 9% shorter than breaths before 10- to 20-s apnea events.

Also we see that median $\text{SpO}_2$ rises gradually in the 5 min leading up to VLAs.

**APPENDIX D: A MATHEMATICAL MODEL FOR THE RATE OF DECLINE OF $\text{SpO}_2$**

Here we present the theory represented by the theoretical curves in Fig. 7 showing $\text{SpO}_2$ as a function of time. The most striking fact about the VLAs is that oxygen saturation falls more slowly in these events than in shorter apnea events (Fig. 7). Why does this happen? We propose that the explanation of the slow fall of $\text{SpO}_2$ is that both arterial and venous oxygenation are unusually high at the start of the event. The high value of venous saturation ($\text{Sv}$) causes the partial pressure of oxygen in the alveoli to drop rather slowly. The high initial values of $\text{SpO}_2$ correspond to values at which the oxygen-hemoglobin dissociation curve $\text{SpO}_2(\text{PO}_2)$ is flat; as oxygen is absorbed from alveoli into blood, and the partial pressure of oxygen in the alveoli falls, the associated drop in arterial oxygen saturation is small. Here we give a quantitative model that shows this effect.

**Method**

An excellent study of the theory of arterial oxygen desaturation during apnea in preterm infants was carried out by Sands et al. (50), following a long tradition of closely related work (17, 25, 32, 38, 54, 55, 64, 68, 69) (see also Refs. 48, 49, 51, 52, 67). We use this as the basis for the following theory which we compare with our observations. The assumptions and approximations used are 1) during an apnea, the only loss of oxygen in the alveoli arises from diffusion into the blood. 2) In each heartbeat, gaseous oxygen in the alveoli equilibrates with oxygen in pulmonary capillaries. 3) Venous blood is homogeneous, with uniform $\text{SpO}_2$. 4) The oxygen saturation in the pulmonary vein leaving the lungs is equal to the oxygen saturation measured by the pulse oximeter a few seconds later, in the finger or toe. 5) All $\text{O}_2$ in the blood is carried by hemoglobin. 6) During the apnea, the cardiac output does not change significantly. 7) $\text{SpO}_2$ is related to partial pressure of oxygen in the alveoli by an equation first given by Hill (29) and quoted (with a slight change) by Delivoria-Papadopoulos and McGowan (14)

$$\text{SpO}_2 = 100kP_{O_2}^n/(1 + kP_{O_2}^n) \quad (1)$$

In this equation, we took $n = 2.9$ (14) and calculated $k$ from reasonable values of $P_{50}$, the value of pressure at which $\text{SpO}_2 = 50\%$.

These assumptions lead to two differential equations with a time delay:

$$dS(t)/dt = C D(S)[S(t) - Sv(t)] \quad (2)$$

$$dSv(t)/dt = \begin{cases} 0 & t \leq T \\ \frac{d}{dt}[S(t - T) - Sv(t)] - e^{-T} & t \geq T \end{cases} \quad (3)$$

Fig. C1. A: continuous histograms and cumulative probability distributions with rational-function fit of 100 breaths right before an apnea. Distributions are shown for VLAs and ABD10–20 events in the same infants. B: median $\text{SpO}_2$ in the 5 min leading up to a VLA for the 46 ABD60–80 events and the 40 ABD80 events. Apnea starts at $t = 0$. Vertical lines indicate ±1SE.
Equation 2 is equivalent to Eq. 10 of Sands et al. (50), and Eq. 3 can be derived from Eq. 6 in Sands et al. Here S(t) is the oxygen saturation leaving the lungs, which by assumption is equal to the measured $S_{O2}$, a few seconds later; $S_{v}(t) = Sv_{O2}$, the venous oxygen saturation, C is a constant, proportional to hemoglobin concentration in the blood and to the cardiac output and inversely proportional to lung volume, and

$$D(S) = dS_{O2}/dP_{O2}$$

which can be obtained from Eq. 1, and which must be expressed as a function of $S_{O2}$ rather than $P_{O2}$. $T$ is arterial transit time, $d$ is equal to cardiac output divided by venous blood volume, and $e$ is the rate at which oxygen is utilized by the tissues. We treat $C$, $T$, $d$, and $e$ as constants, independent of time, and independent of the duration of the apnea event. From their values, and from the initial conditions $S(t = 0)$ and $S_v(t = 0)$, the above equations determine the evolution of $S(t)$.

These four constants and the initial conditions were determined by first obtaining preliminary estimates and then refining by fitting to the observations. The preliminary estimates were obtained by the following process. Sands et al. (50) showed that oxygen saturation during an apnea frequently follows a two-stage pattern. In the first stage, $t \leq T$, the arterial oxygen saturation $S(t)$ drops rapidly, while in the second stage, it drops more slowly and at a constant rate (Fig. D1). In cases where the two stages can be easily identified on a graph of $S(t)$, the arterial transit time $T$ can be determined from the graph. Also, in the first stage $S_v(t)$ is constant.

In this stage, $S_v(t)$ contains two unknowns: $S_v$ and $C$. These can be determined by choosing two points in time early in the apnea, $t_j = (t_1, t_2)$, extracting $S(t_j)$ and $dS/dt$ from the observations, and then solving the two equations for $C$ and $S_v$.

To obtain estimates of $d$ and $e$ we set $t = T$ in Eq. 3 and find

$$d[S(0) - S_v(T)] - e = d[S(0) - S_v(0)] - e = 0$$

In the second stage, the arterial oxygen saturation falls linearly with time at a rate approximately equal to $(50)$

$$dS(t)/dt = -e/(1 + DT)$$

From the observed rate of decline $dS(t)/dt$ in the second stage, and with $S_v(0)$ determined as above from stage 1, we have two equations for $d$ and $e$ with two unknowns.

Two issues remain. 1) There is a time delay between oxygen leaving the lungs and oxygen detected in the finger or toe. Part of this time delay is related to circulation time of blood and part of it is related to the averaging time of the oxygen monitor. A time delay of $20 \, \text{s}$ was chosen for all of the average curves in Fig. 7 and gave satisfactory results. 2) The average saturation curves do not show any clear division between stage 1 and stage 2. Therefore, individual apnea events that show this distinction were used to obtain estimates of $T$, $d$, and $e$. There are several such cases, one of which is shown in Fig. D1.

Finally, the values of these parameters were adjusted to fit the observations.

Results

It is very important to note that we used the same constant values of $C$, $T$, $d$, and $e$ for short and long apnea events. We adjusted only the initial conditions $S(0)$ and $S_v(0)$ to fit the individual curves.

Table D1 lists the constants (except $e$, which can be found using Eq. 5) and the initial conditions. The resulting theoretical curves are shown in Fig. 7. The agreement with the average $S_{O2}$ curves is remarkable: over the whole of each of the ranges where $S_{O2}$ is falling, the theoretical curves are only barely discernible from the observed ones and differ from them by less than the standard error of the means of the observations.

In parentheses in Table D1 we report the uncertainty in the values of the parameters. These were obtained by finding the standard deviation of the mean of each set of observations and adjusting parameters one-at-a-time to find values where the theoretical curves would disagree with observations by more than one standard error of the mean.

Discussion

We have shown that the theory given by Sands et al. (50) leads to remarkable agreement with observation. Thereby, it provides a credible explanation of why $S_{O2}(t)$ falls slowly in long apnea events (whereas other attempted explanations fail). Two factors are at work. At high oxygen saturations, the curve relating oxygen saturation to partial pressure of oxygen $S_{O2}(P_{O2})$ is flat, and the function $D(S)$ is near zero. In the alveoli and in the arteries, a large change in partial pressure of oxygen gives only a small change in $S_{O2}$. As a result, $S_{O2}$ falls very slowly near the beginning of an apnea when oxygen saturation is high. In this regard the slow fall of $S_{O2}$ from high levels is a natural consequence of well-known physiology. The second factor is the inferred somewhat elevated oxygen saturation in the veins. We have no direct measurements of venous oxygen saturation; it is obtained only by fitting to the observations. However, this is consistent with the result shown in APPENDIX C showing the steady rise of $S_{O2}$ over at least 5 min before VLAs.

Assumptions and approximations. To understand the limits of the theory (50) as we apply it, let us reexamine some of the approximations. During an apnea, the CO$_2$ concentration and pH of the blood may change, and these shift the hemoglobin saturation curve, $S_{O2}(P_{O2})$. This shift can be characterized by a changed value of $P_{50}$, the pressure at which hemoglobin is 50% saturated. This change, however, turns out to have only a small effect on the time dependence $S_{O2}(t)$. One can include the effect approximately by changing the value of $k$ in Eq. 1. However, when calculating $S_{O2}(t)$ as described above using Eq. 1, the value of $k$ exactly cancels (for fixed $C$ and $S_v$), so at this level of approximation, $P_{50}$ has no effect. [In a more accurate treatment, again holding $C$ and $S_v$ fixed, $P_{50}$ would have a nonzero but small effect; Sands et al. (50), who used the Severinghaus formula (56) instead of Eq. 1, found that a change of $P_{50}$ by 6–12

<table>
<thead>
<tr>
<th>Parameter</th>
<th>ABD10–20</th>
<th>ABD30–40</th>
<th>ABD60–80 (VLAs)</th>
<th>ABD80 (VLAs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$S$</td>
<td>0.870 (0.005)</td>
<td>0.915 (0.007)</td>
<td>0.950 (0.005)</td>
<td>0.970 (0.003)</td>
</tr>
<tr>
<td>$S_v$</td>
<td>0.70 (0.01)</td>
<td>0.74 (0.01)</td>
<td>0.800 (0.005)</td>
<td>0.825 (0.005)</td>
</tr>
<tr>
<td>$C$</td>
<td>13.3 (0.6)</td>
<td>0.2 (0.6)</td>
<td>12 (1)</td>
<td></td>
</tr>
</tbody>
</table>

S, saturation; $S_v$, venous saturation; $C$, constant proportional to hemoglobin concentration in the blood and to the cardiac output and inversely proportional to lung volume; $d$, cardiac output divided by venous blood volume; $T$, arterial transit time.
mmHg gave significant changes to \(\text{SpO}_2(t)\), but part of the change they obtained resulted from changed initial values of \(\text{SpO}_2\) and \(\text{SaO}_2\).

Another effect we neglected is extrapulmonary right-to-left shunt. Since we find that the median oxygen saturation before a VLA is 96–98%, it appears that shunted blood is a small fraction of arterial blood and would have little effect on the calculation of \(\text{SpO}_2\).\(^{(50)}\)\ldots

Further study of assumption (3), that venous blood is well-mixed and homogeneously oxygenated would be beneficial. This should be accurate in steady state, but it is not clear that it would hold during the rapid changes of oxygenation that occur during apnea.

**Alternative hypotheses.** We arrived at this theory only after elimination of a number of other hypotheses, which we examine here. 1) Infants who have VLAs are different in some way from other infants of comparable age. For example, they may have unusually high amounts of fetal hemoglobin, which holds oxygen more strongly than adult hemoglobin, and therefore oxygen saturation may drop more slowly when they have apnea. However, this hypothesis can be excluded, because these same infants (in the same epochs) have short apnea events that are indistinguishable from short apnea events in other infants. 2) There is some mechanism acting in some instances that, analogous to the “diving seal” reflex, reduces oxygen consumption in these infants. This hypothesis is not credible because these events look nothing like the dive reflex. This reflex acts within seconds; one observation of the dive reflex in a human infant showed the heart rate dropping from >150 to <100 beats/min in ~1 s (21). Furthermore, reduced oxygen consumption would affect venous oxygen saturation, which would slow the rate of fall of arterial oxygen only after a time delay, but we see arterial oxygen saturation falling slowly from the beginning. 3) Noting that most of the infants who have VLAs are on CPAP, one might hypothesize that the infant is somehow getting oxygen from the CPAP even though there is no detectable chest movement. CPAP is expected to maintain functional residual capacity; in this regard, this hypothesis is consistent with the model presented above. In addition, one might hypothesize that CPAP is providing a flow of fresh air to the alveoli that is not visible in the CI signal (12). “Apneic oxygenation” is flow of fresh air into alveoli caused by removal of oxygen into the blood stream (19, 26). Rough estimates suggest that this process would have a small effect. We have no decisive evidence for or against this hypothesis, but it does not explain why the same infant in the same epochs has short apnea events in which oxygenation falls quickly and long ones in which it falls slowly. Also, a quantitative theory of this process is not presently available, it would be complex, and it would depend on a number of unknown parameters. In contrast, the theory given in Sands et al. (50) and applied above is fully developed, and we can extract all parameters by fitting to observations.

**Conclusion**

This theoretical method might turn out to provide useful physiological information. \(C\) and \(d\) are both proportional to cardiac output, \(e\) is proportional to metabolic rate, and \(S_v\) is the venous oxygen saturation, none of which are continuously available in NICU monitoring. By fitting the time course of \(\text{SpO}_2\) in individual apnea events to the theory, it may be possible to get estimates of these parameters during each apnea. This will be the topic of a future study.

**ACKNOWLEDGMENTS**

Present address of M. T. Clark: Adult Medical Predictive Devices, Diagnostics, and Displays, LLC, Charlottesville, VA 22903.

**REFERENCES**


