Nocturnal oximetry in infants with cystic fibrosis

M P Villa, J Pagani, V Lucidi, S Palamides, R Ronchetti

Abstract

Aim—To investigate whether children with cystic fibrosis under 3 years of age have disordered breathing and episodes of oxygen desaturation during sleep.

Methods—We studied 19 infants (9 boys and 10 girls) with cystic fibrosis, mean age 13.1 months (range 3–36 months) and 20 age and sex matched healthy subjects. Patients and controls underwent an overnight polysomnographic study and respiratory function testing on the following morning.

Results—Seven patients with ongoing respiratory tract inflammation had disordered breathing and episodes of oxygen desaturation during sleep. Pulse oximetry showed a significantly lower mean oxygen saturation (SaO₂) and a higher percentage of total sleep time spent with SaO₂ less than 93% in symptomatic children than in controls.

Conclusion—Results suggest that infants and young children with cystic fibrosis and mild airways inflammation (rhinitis, cough, red throat) have episodes of oxygen desaturation during sleep.

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Keywords: cystic fibrosis; sleep; oxygen desaturation

There have been several reports of severe episodes of arterial oxygen desaturation during sleep in adult and adolescent patients with cystic fibrosis (CF). Hypoxaemia may be an arousing stimulus that disturbs these patients’ normal sleep pattern and their quality of life. Some evidence suggests that nocturnal hypoxaemia plays a role in the pathogenesis of lung damage and cor pulmonale in patients with CF.

A number of investigators have suggested that even brief desaturation episodes can increase pulmonary artery pressure.

The largest falls in saturation occur during rapid eye movement (REM) sleep and are associated with decreases in tonic electromyographic activity of intercostal muscles and with an irregular respiratory pattern. Ballard et al showed that desaturation episodes during sleep in patients with CF are associated with hypventilation caused by a reduction in tidal volume and minute ventilation.

Reduced ventilation during REM sleep in adults with CF has also been described by Tepper et al, who propose that this could account for most of the sleep hypoxaemia in CF. A decrease in the contribution of the rib cage to tidal volume breathing leads to a decrease in lung volume and to episodes of desaturation. Because desaturation episodes coincided not with obstructive or central apnoeas but with a reduced end expiratory volume during REM sleep they were attributable to airway closure and to the development of underventilated lung regions. To explain the mechanism responsible for sleep induced oxygen desaturation in patients with CF, Muller et al suggest that a cyclic reduction in functional residual capacity (FRC) may determine a ventilation–perfusion mismatch.

The sleep induced changes in oxygenation in patients with CF resemble those reported in adults with chronic bronchitis and chronic obstructive pulmonary disease. In patients with chronic bronchitis and chronic obstructive pulmonary disease the quality of sleep is particularly disturbed in the REM sleep stage. As most studies to date that have examined ventilation and oxygenation changes during sleep in patients with CF have concentrated on adults and adolescents, we sought to examine these changes in CF patients under 3 years, an age when lung function remains largely unaffected by the disease. We designed this study to investigate whether CF patients less than 3 years of age have disordered breathing and oxygen desaturation during sleep and whether acute respiratory tract inflammation affects ventilation during sleep.

Subjects and methods

We studied 19 infants and young children (9 boys and 10 girls), mean age 13.1 months (range 3–36 months), referred to our respiratory and sleep laboratory from September 1998 to May 1999 whose CF had been diagnosed before 6 months of age. Patients seen during routine follow up visits at the CF centre of Bambino Gesù and considered clinically well, without acute exacerbation of respiratory symptoms, were scheduled for lung function testing and a sleep study. The interval between clinical assessment and lung function testing ranged from seven to 14 days. In 12 infants CF had been identified between 1 and 2 months following newborn screening and confirmed by three abnormal sweat tests; in seven infants the diagnosis of CF was first suggested by symptoms including meconium ileus (n = 1), meconium peritonitis (n = 1), metabolic acidosis (n = 1), diarrhoea and vomiting with poor growth (n = 2), and bronchial asthma (n = 1) and confirmed by three abnormal sweat tests. As a control group we studied 20 age and sex matched healthy subjects.

Patients with CF and controls underwent clinical assessment, a polysomnographic study, and lung function testing on the following morning. When studied, seven children had symptoms compatible with mild respiratory tract inflammation—that is, rhinorrhea, cough, and red throat. At the time of the study
Table 1 Anthropometric variables in children with cystic fibrosis

<table>
<thead>
<tr>
<th>Subject no. (sex)</th>
<th>Age (mth)</th>
<th>Weight (centile)</th>
<th>Height (centile)</th>
<th>Diagnosis (type)</th>
<th>Inflammatory index†</th>
</tr>
</thead>
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<tr>
<td>Symptomatic* (n = 7)</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>1 (F)</td>
<td>18</td>
<td>77</td>
<td>97</td>
<td>Screening</td>
<td>1.99</td>
</tr>
<tr>
<td>2 (F)</td>
<td>10</td>
<td>80</td>
<td>90</td>
<td>Screening</td>
<td>4.88</td>
</tr>
<tr>
<td>3 (M)</td>
<td>7</td>
<td>25</td>
<td>90</td>
<td>Screening</td>
<td>6.8</td>
</tr>
<tr>
<td>4 (M)</td>
<td>7</td>
<td>60</td>
<td>25</td>
<td>Symptoms</td>
<td>8.3</td>
</tr>
<tr>
<td>5 (F)</td>
<td>3</td>
<td>3</td>
<td>3</td>
<td>Symptoms</td>
<td>7.2</td>
</tr>
<tr>
<td>6 (F)</td>
<td>13</td>
<td>80</td>
<td>25</td>
<td>Symptoms</td>
<td>0.92</td>
</tr>
<tr>
<td>7 (F)</td>
<td>7</td>
<td>25</td>
<td>50</td>
<td>Screening</td>
<td>5.14</td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>9.6 (4.5)</td>
<td>50 (31.8)</td>
<td>54 (38.6)</td>
<td></td>
<td>5.1 (2.8)‡</td>
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<tr>
<td>Asymptomatic* (n = 12)</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>8 (M)</td>
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<td>25</td>
<td>25</td>
<td>Screening</td>
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</tr>
<tr>
<td>9 (M)</td>
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<td>80</td>
<td>90</td>
<td>Screening</td>
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<tr>
<td>10 (F)</td>
<td>36</td>
<td>97</td>
<td>80</td>
<td>Symptoms</td>
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</tr>
<tr>
<td>11 (M)</td>
<td>25</td>
<td>27</td>
<td>50</td>
<td>Screening</td>
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</tr>
<tr>
<td>12 (F)</td>
<td>9</td>
<td>30</td>
<td>50</td>
<td>Screening</td>
<td>6.6</td>
</tr>
<tr>
<td>13 (M)</td>
<td>30</td>
<td>12</td>
<td>4</td>
<td>Screening</td>
<td>1.2</td>
</tr>
<tr>
<td>14 (F)</td>
<td>5</td>
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<td>75</td>
<td>Symptoms</td>
<td>7.2</td>
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<tr>
<td>15 (M)</td>
<td>32</td>
<td>3</td>
<td>3</td>
<td>Symptoms</td>
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<tr>
<td>16 (M)</td>
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<td>90</td>
<td>Symptoms</td>
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<td>6</td>
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<td>50</td>
<td>Screening</td>
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<tr>
<td>18 (M)</td>
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<td>3</td>
<td>3</td>
<td>Screening</td>
<td>1.9</td>
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<tr>
<td>19 (F)</td>
<td>9</td>
<td>50</td>
<td>10</td>
<td>Screening</td>
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</tr>
<tr>
<td>Mean (SD)</td>
<td>15.2 (12.1)</td>
<td>39.7 (29.6)</td>
<td>47.7 (33.7)</td>
<td></td>
<td>1.8 (2.5)</td>
</tr>
<tr>
<td>Total subjects</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>13.1 (10.2)</td>
<td>43.5 (29.6)</td>
<td>50.4 (34.4)</td>
<td></td>
<td>3.1 (3.0)</td>
</tr>
</tbody>
</table>

Statistical analyses used two tailed t test.
*Clinical condition on testing with or without symptoms of respiratory inflammation.
†Number of exacerbations per month × 12.
‡p = 0.02 versus asymptomatic.

No abnormal findings were shown on chest x ray in any patient.

We reviewed each child’s respiratory history to assess the number of respiratory tract infections during life, normalised for age (inflammatory index: number of pulmonary exacerbations per month of life × 12).14

Parents gave informed consent for their children to take part in the study; none of the procedures were invasive or traumatic.

POLYSOMNOGRAPHIC STUDY

Polysomnograms were recorded with a Medilog SAC 800–Oxford apparatus, in 16 infants during the night (between 2100 and 0600), and in three during the afternoon (between 1400 and 1800). No infant had sleep pharmacologically induced or was sleep deprived. Polysomnographic recordings included the following channels: electroencephalogram (C3–A2; C4–A1) recorded through surface electrodes attached to the scalp; electrooculogram; and electromyography (submental electrodes attached to the scalp; electrooculogram; and electromyography (submental and exterior tibial). All variables were monitored continuously and were used to determine sleep stages according to the criteria of Rechtschaffen and Kales.21 Chest and abdominal respiratory movements were determined by impedance plethysmography; nasal and mouth air flows were recorded by thermocouples placed near the nostrils and the mouth. SaO2 was monitored by finger pulse oximetry; we defined oxygen desaturation as a 4% fall from baseline oxygen values. We calculated total sleep time (TST), and the sleep time spent with oxygen saturation lower than 93%, 90%, and 85%. The mean low SaO2 and the lowest SaO2, and the duration of desaturation episodes were also measured. The electrocardiogram was recorded through precordial electrodes. Polysomnographic studies took place in the sleep laboratory, a comfortable temperature controlled (21–22°C) room equipped to allow children and parents to sleep together.

We defined sleep disordered breathing as follows: apnoea index >1 event/h, apnoea + hypopnea index >5 events/h, mean low SaO2 <90%, and lowest SaO2 <85%.

Respiratory function was studied (Sensor Medics 2600 equipment) during quiet sleep induced by chloral hydrate (50 mg/kg) in patients and in 14 age and sex matched healthy subjects. Partial flow–volume curves were obtained by the chest compression technique. In brief, an inflatable plastic bag was wrapped around the infant’s abdomen. The bag pressure was increased in increments of 5–10 cm water from a minimum of 20 to a maximum of 70 cm water. On each occasion, the inflation of the bag was begun at end tidal inspiration and maintained until a spontaneous inspiratory effort was evident or no additional airflow could be obtained. The maximal pressure applied was that which produced no further increase in flow at the end tidal expiration point (functional residual capacity, FRC). This manoeuvre was performed three to five times in each infant, and the greatest flow obtained at the end tidal expiration point (Vmax/FRC) was used in the analyses. We also calculated the expiratory time constant (TC). Compliance and resistance of the respiratory system was determined with the method of single occlusion, evoking the Hering–Breuer reflex to bring on a passive expiration. FRC was measured by the nitrogen washout method (FRCneo) by the N2 dilution technique using 100% oxygen, a mixing chamber, and a fast N2 analyser (>50 ms) (Giesler tube).

STIATISTICAL ANALYSIS

Data are expressed as mean (SD). Student’s t tests for independent data (univariate analysis) were used to compare continuous variables in patients and control subjects. The Pearson correlation was used to evaluate the association between SaO2 and respiratory function. Multiple regression analysis was used to determine the importance of these variables in determining desaturation during sleep. Probability values equal to or less than 0.05 were considered statistically significant.

Results

Sixteen of those studied had grown normally and three infants were on the 3rd percentile for weight and height (table 1).

When tested, seven infants (two boys and seven girls) had mild respiratory symptoms (runny nose, cough, and red throat), and 12 (seven boys and five girls) were asymptomatic. Symptomatic infants had a significantly higher inflammatory index than asymptomatic infants (5.1 (2.8) v 1.8 (2.5); p < 0.01). Five symptomatic (5/7; 71%) and two asymptomatic subjects (2/12; 17%) had an inflammatory index greater than 4. In 4/7 symptomatic (57%) and 8/12 asymptomatic infants (66%) CF was diagnosed at newborn screening.

No difference was found between the sleep architecture of patients with CF or control subjects (REM = 27.0 (19.1)% v 28.0 (9.6)%).
Symptomatic subjects spent a smaller percentage of the total sleep time in REM sleep than controls (16.9 (8.7)% vs 28.0 (9.6)%; p < 0.001). Symptomatic and asymptomatic patients had normal apnoea indexes (number of apnoeas per hour of sleep) and normal apnoea + hypopnoea indexes (table 2).

In asymptomatic patients and controls, pulse oximetry showed no significant differences in mean oxygen saturation, mean lowest oxygen saturation, or percentage of total sleep time spent with SaO2 less than 93%. Symptomatic patients had significantly lower mean oxygen saturation than controls (p < 0.0001). In symptomatic patients, the mean oxygen saturation in REM sleep was lower than in non-REM (NREM) sleep. The percentage of total sleep time spent with SaO2 less than 93% was higher in symptomatic children than in controls (p < 0.0001). The percentage of total sleep time spent with SaO2 less than 90% and 85% was higher in symptomatic than in asymptomatic patients. The lowest SaO2 values were lower in symptomatic patients than in controls (p < 0.001). Sleep disordered breathing was shown in some symptomatic patients but in no asymptomatic patients or control subjects.

No differences were found in polysomnographic variables in patients diagnosed by screening or by symptoms. Lung function testing (table 3) showed a higher respiratory rate in symptomatic than in control subjects (p < 0.01). Values for compliance and resistance of the respiratory system came within normal ranges and no significant differences emerged between the symptomatic and asymptomatic groups and controls. FRC<sub>2</sub> values did not differ significantly in patients or in controls. Plethysmography yielded a significantly lower mean V<sub>max</sub>FRC in the symptomatic than in the control group (p < 0.001).

Pearson’s correlation between the respiratory function variables and polysomnographic variables and clinical findings showed that respiratory frequency correlated with the mean low SaO2 during sleep (r = −0.592; p < 0.05), and the FRC<sub>2</sub> values correlated with the number of respiratory exacerbations (inflammatory index) (r = −0.593; p < 0.05).

Multiple regression among the dependent variables (mean SaO2, clinical findings) and the clinical and respiratory function variables (respiratory rate, compliance and resistance of the respiratory system, V<sub>max</sub>FRC, FRC<sub>2</sub> and TC) identified the variables most likely to be associated with desaturation during sleep as the presence of respiratory symptoms during testing (p < 0.001) and higher TC values (p < 0.01).

Discussion

Of the 19 CF infants and children less 3 years of age, only the seven with ongoing respiratory tract inflammation had sleep disordered breathing and oxygen desaturation during sleep. Even in normal individuals, sleep induces changes in the respiratory pattern, consisting of a reduction in lung volume and minute ventilation.22–25 Whereas these changes have no metabolic consequences in children with healthy lungs, in children with chronic lung disease they can provoke ventilatory anomalies and oxygen desaturation episodes.26

In infants and young children such as those studied, ventilatory mechanics differ from those in children and adolescents. The shape of a young infant’s rib cage, for example, is unlike that of an adult. Owing to the horizontal insertion of the ribs and the position of the diaphragm in infancy, the rib cage is more compliant and the lung is mechanically less efficient. Increased rib cage compliance and higher respiratory resistance are partly responsible for respiratory events during sleep. In addition, the respiratory muscle hypotonia that characterises the REM sleep stage favours and aggravates the nocturnal hypoventilation typical of obstructive lung diseases.

The normal sleep induced fall in FRC, which may drop even to 40% of the baseline value in some sleep stages,27 reduces O2 reserve. These effects become even more apparent if the fall in FRC coincides with the paradoxical breathing characteristic of newborn babies.
and infants with chronic lung diseases. These conditions predispose subjects with compromised lung mechanics to a sleep induced fall in O₂.

When respiratory tract inflammation intervenes, respiratory system resistance and hypventilation may increase, thus inducing desaturation during sleep. This is presumably what happened in our symptomatic patients. Polysomnograms showed that the symptomatic subjects spent more sleep time with SaO₂ less than 93% than controls (p < 0.0001). In addition, they spent more sleep time with desaturation (SaO₂ less than 90%). They also had lower mean SaO₂ during sleep (p < 0.001). Our observations refer exclusively to non-complicated respiratory tract infections. None of the infants we studied had severe pulmonary exacerbations or fever, neither did they have clinical evidence of infection or abnormal chest x ray findings.

In a study conducted in 1983 in adolescent CF patients, Tepper et al showed that episodes of arterial oxygen desaturation occurred chiefly during REM sleep. REM sleep worsened sleep induced respiratory changes, including hypventilation, and increased the variability of the sleep pattern. The desaturation episodes coincided strictly with reduced minute ventilation. Our data agree with the observation of Tepper et al of lower SaO₂ values in REM than in NREM sleep, but only in subjects with respiratory symptoms.

In a study comparing nocturnal hypoxia with lung function values, Stokes et al observed smaller SaO₂ changes in patients with moderate to severe illnesses (FEV₁ = 31–63% of the predicted value) than in patients with severe illnesses (FEV₁ = 17% of the predicted value). In adults, as Pond and Conway have shown, low FeO₂ values may be useful in identifying patients at risk for nocturnal hypoxaemia, even though no predictive cut off value has been defined for FEV₁.

Except for the significantly lower V_a/FRC in children who were symptomatic when tested, our 19 patients had normal lung function values. In a study of CF infants without respiratory symptoms, Beardsmore et al concluded that flow–volume curves do not disclose lung dysfunction at this young age. Our lung function findings confirm this conclusion. Apart from the correlation between respiratory frequency and mean SaO₂ during sleep, we found no correlation between oxygen saturation and lung function variables in our patients. One reason could be that CF first reduces alveolar ventilation and only later affects lung function variables, as Coates et al have proposed. More subtle anomalies such a change in the ventilation/perfusion ratio would be better revealed by more sensitive methods, including lung scintigraphy, than by lung function tests. Sleep could be a challenge that triggers these anomalies. An interesting finding in our study is that even in CF patients younger than 3 years, pulmonary exacerbations can alter lung function, reducing V_a/FRC.

It is known that infants with CF already have airways inflammation, even in the absence of acute infections. Tepper et al have proposed that this inflammation helps to aggravate the lower respiratory tract infections that develop early in life and may be responsible for airways obstruction. Our data are in total agreement, showing mild airway inflammation and signs of obstruction in early months of life, as shown by the low V_a/FRC values and high TC values on the flow–volume curves.

In the multiple regression including all the polysomnographic, clinical, and respiratory variables studied we identified two variables as risk factors for nocturnal desaturation: respiratory symptoms and higher TC values. In addition, we found lower FRC values in patients who had more pulmonary exacerbations. Another predisposing factor for nocturnal desaturation was a high respiratory rate, an expression of possible lung impairment. Accordingly, subjects who had higher respiratory rates also had lower SaO₂ values during sleep. Infants who had respiratory symptoms when tested were mainly (5/7) also those who had more pulmonary exacerbations. Hence CF infants with more severe clinical disease may be those who have nocturnal desaturation.

Repeated episodes of nocturnal hypoxia are a stimulus for the development of pulmonary hypertension and later cor pulmonale. Nevertheless, definitive data are lacking on the natural history of sleep induced hypoxia and its role in the pathogenesis of cor pulmonale. If hypoxic stimuli begin at such an early age, their consequences could be more severe. Hence the importance of assessing the severity and duration of hypoxia even at this age.

We conclude that infants with CF in whom mild airways inflammation develops during the first months of life have episodes of oxygen desaturation during sleep. Hence these children may benefit from nocturnal SaO₂ monitoring. Controlled studies are needed to determine the usefulness of O₂ supplementation for sleep induced respiratory symptoms.

References

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