Sleep-disordered breathing and comorbidities: role of the upper airway and craniofacial skeleton

Abstract: Obstructive sleep-disordered breathing (SDB), which includes primary snoring through to obstructive sleep apnea syndrome (OSAS), may cause compromise of respiratory gas exchange during sleep, related to transient upper airway narrowing disrupting ventilation, and causing oxyhemoglobin desaturation and poor sleep quality. SDB is common in chronic disorders and has significant implications for health. With prevalence rates globally increasing, this condition is causing a substantial burden on health care costs. Certain populations, including people with sickle cell disease (SCD), exhibit a greater prevalence of OSAS. A review of the literature provides the available normal polysomnography and oximetry data for reference and documents the structural upper airway differences between those with and without OSAS, as well as between ethnicities and disease states. There may be differences in craniofacial development due to atypical growth trajectories or extramedullary hematopoiesis in anemias such as SCD. Studies involving MRI of the upper airway illustrated that OSAS populations tend to have a greater amount of lymphoid tissue, smaller airways, and smaller lower facial skeletons from measurements of the mandible and linear mental spine to clivus. Understanding the potential relationship between these anatomical landmarks and OSAS could help to stratify treatments, guiding choice towards those which most effectively resolve the obstruction. OSAS is relatively common in SCD populations, with hypoxia as a key manifestation, and sequelae including increased risk of stroke. Combatting any structural defects with appropriate interventions could reduce hypoxic exposure and consequently reduce the risk of comorbidities in those with SDB, warranting early treatment interventions. 

Keywords: obstructive sleep apnea, sickle cell, polysomnography, desaturation, MRI, airway, adenoids

Introduction

This review aims to explore the relationship between sleep-disordered breathing (SDB) and health in the general population, and to focus on anemias, including sickle cell disease (SCD). We address three key areas. Firstly, we review reports including polysomnography (PSG) and limited channel sleep studies undertaken in the general population of adults and normally developing children. We subsequently compare these sleep studies against those in children and young adults with anemia, specifically SCD. Secondly, we review MRI head and neck studies to compare structural anatomical variations in the upper airway in populations exhibiting a high prevalence of SDB. We identify and summarise potential targets for treatment for SDB in patients with anemia such as SCD. Finally, we discuss the implications of SDB for patients with SCD to understand the importance of managing this population.
Normal respiratory physiology in sleep

In typically developing children, as well as in adults, small changes in arterial oxygen saturations (SaO₂; ≤2% decrease), typically measured by pulse oximetry (SpO₂), and partial pressure of carbon dioxide (pCO₂; 4–6 mmHg increase), from end-tidal capnography, are common during sleep. This is related to the reduced metabolic demands of sleep, resulting in lower respiratory and pulse rates, and changes in breathing patterns and muscle tone, which differ between wakefulness and sleep, and during the different stages of sleep. To limit physical activity in response to the vivid dreams of rapid eye movement sleep, inhibition of tonic muscle tone in respiratory intercostal muscles occurs in this stage of sleep, reducing lung capacity to its lowest levels and making the upper airway more susceptible to resistance and collapse. This is usually not a problem in children with adequate upper airway space and normal sleep. However, even in typically developing children, average oxygen saturations are lower during sleep than when awake.

Clinical algorithms have relatively low sensitivity and specificity for diagnosing SDB. PSG is the gold standard for monitoring sleep, and includes measurements of electroencephalography (EEG), electrooculography (EOG), and electromyography (EMG), to determine sleep stages and limb movements. Respiration in sleep is also assessed, with measurements of electrocardiography (ECG), oronasal airflow, thoracic and abdominal respiratory effort, pulse oximetry, and snoring. End-tidal pCO₂ from capnography and video recordings are also often made. Many centers now use cardiorespiratory or limited polygraphy studies as screening or diagnostic tools, as PSGs are expensive. The limited polygraphy studies exclude EEG, EOG, and EMG, so cannot specifically define sleep stage or sleep quality, but give a reasonable estimation of SDB severity and response to treatment. Oxygen saturation studies (pulse oximetry), sometimes recorded with a measure of pCO₂ (oxicapnography), cannot be used to diagnose SDB, but are used for screening and risk assessment in some circumstances.

The limited population-based normative data, lack of consensus regarding respiratory event scoring parameters between full PSG and limited channel studies, and changes in standards over time have been significant barriers to description of respiration in normal sleep in infants and adults. However, the American Academy of Sleep Medicine (AASM) now have consensus definitions for scoring polysomnographic studies, and overnight oximetry, particularly for mean or median across the whole night. These can be compared across studies using oximeters that reduce motion artifact, although summary statistics may not fully describe hypoxic exposure and thus graphical solutions are under development. We will present here literature using pre-AASM and post-AASM scoring.

There are few daytime or overnight oxygen saturation data from the general population living at sea level. In a population-based study of adults in Tromsø, Norway, 6.3% of 6317 adults aged 38–87 years had a daytime saturation of ≤95% and mortality over the subsequent 10 years was higher in this group than in those with SpO₂>96%. In 350 normal adults, minimum overnight saturation was 90.4±3.1% with a median of 96.5%±1.5%. Overnight saturation does appear to decrease with age (Table 1).

In children, average daytime oxygen saturations ≥95% are considered normal, and values below this are considered acutely associated with signs suggesting illness. Overnight medians or means are typically ≥97% in infants and children, rarely dipping to less than 90% in sleep in typically developing children, although lower minimum values are seen in normal infants. Nocturnal oxygen desaturation (NOD) is not the norm in infants, children, or adults under the age of 60 years (Table 1).

Sleep-disordered breathing

Defining normal respiratory patterns in sleep paves the way for the definition of abnormal respiratory patterns in sleep. PSG features of pediatric OSAS differ from adults. Children may have obstructive hypoventilation (persistent partial upper airway closure, sometimes with paradoxical breathing or increased work of breathing, with elevated pCO₂) rather than frank obstructive apneas or hypopneas. In addition, they are less likely to arouse from apnea, hypopnea, or oxygen desaturation. In both adults and children, severity of OSAS is classified according to the number of obstructive apneas and hypopneas recorded per hour of sleep and reported as an Apnea/Hypopnea Index (AHI) or Obstructive Apnea/Hypopnea Index (OAHI).

Obstructive SDB/OSAS is a condition caused by narrowing or obstruction of the upper airways that disrupts ventilation, causing oxyhemoglobin desaturation and often reduced sleep quality. There is a spectrum of obstructive SDB, ranging from primary snoring through upper airways resistance syndrome, to the more severe obstructive sleep apnea syndrome (OSAS). Identifying obstructive
Table 1 Normal data for mean and minimum overnight oxygen saturation and polysomnography variables in infants, children, and adults

<table>
<thead>
<tr>
<th>Author</th>
<th>Age (years) – mean (SD) except*</th>
<th>Mean overnight oxygen saturation % mean (SD) except‡**</th>
<th>Nadir overnight oxygen saturation % mean (SD) except‡**</th>
<th>Apnea Hypopnea Index (events/hr)</th>
<th>Obstructive Apnea Index (events/hr)</th>
<th>Central Apnea Index (events/hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Terrill*</td>
<td>0.04</td>
<td>98.0 (95–100)*</td>
<td>80.4 (78.8–82.0)</td>
<td>0.03 (0.1)</td>
<td>0.62 (0.73)</td>
<td>0.95 (0.82)</td>
</tr>
<tr>
<td>Terrill*</td>
<td>0.25</td>
<td>99.0 (97–100)*</td>
<td>84.7 (83.3 to 86.1)</td>
<td>0.05 (0.11)</td>
<td>0.45 (0.49)</td>
<td>0.90 (0.82)</td>
</tr>
<tr>
<td>Terrill*</td>
<td>0.5</td>
<td>98.0 (95–100)*</td>
<td>92 (3)</td>
<td>0.01 (0.03)</td>
<td>0.08 (0.14)</td>
<td>0.25 (0.17)</td>
</tr>
<tr>
<td>Terrill*</td>
<td>1</td>
<td>99.0 (96–100)*</td>
<td>91.1 (4.2)</td>
<td>0.2 (0.3)</td>
<td>0.34 (0.16)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Terrill*</td>
<td>2</td>
<td>99.0 (97–100)*</td>
<td>94.6 (2.2)</td>
<td>0.1 (0.2)</td>
<td>1.0 (0.99)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Evans3</td>
<td>0.08*</td>
<td>97.1 (96.6–97.5)*</td>
<td>1.5 (0.8)</td>
<td>1.0 (0.99)</td>
<td>0.9 (0.8)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Evans3</td>
<td>0.25–0.33*</td>
<td>97.7 (97.2–98.1)*</td>
<td>1.1 (0.5)</td>
<td>0.8 (0.7)</td>
<td>0.7 (0.6)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Montgomery-Downs*</td>
<td>3–5*</td>
<td>&gt;95</td>
<td>0.8 (0.32)</td>
<td>0.0 (0.0)</td>
<td>0.1 (0.08)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Montgomery-Downs*</td>
<td>6–7*</td>
<td>&gt;95</td>
<td>1.1 (0.5)</td>
<td>0.0 (0.0)</td>
<td>0.1 (0.08)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Traeger23</td>
<td>6.6 (1.9)</td>
<td>97 (1)</td>
<td>92 (3)</td>
<td>0.0 (0.0)</td>
<td>0.1 (0.08)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Straus40</td>
<td>6.6 (3.4)</td>
<td>97.1 (0.9)</td>
<td>91.1 (4.2)</td>
<td>0.2 (0.3)</td>
<td>0.33 (0.16)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Uliel22</td>
<td>7.9 (4.4)</td>
<td>97 (0.8)</td>
<td>94.6 (2.2)</td>
<td>0.1 (0.2)</td>
<td>1.5 (1.1)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Uliel recommended*</td>
<td>8 (1–15)*</td>
<td>92</td>
<td>93 (2.2)</td>
<td>0.1 (0.5)</td>
<td>0.06 (0.16)</td>
<td>0.85 (0.9)</td>
</tr>
<tr>
<td>Uliel recommended*</td>
<td>9.3 (0.6)</td>
<td>97 (1.6)</td>
<td>92.2 (2.7)</td>
<td>0.0 (0.0)</td>
<td>0.1 (0.08)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Urschitz21</td>
<td>9.6 (0.7)</td>
<td>96 (90–99)*</td>
<td>91.6 (73–95)*</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Moss15</td>
<td>9.6 (0.7)</td>
<td>97 (0.6)</td>
<td>91.8 (2.7)</td>
<td>2.8 (0.8–3.7)</td>
<td>0.0 (0.0)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Rosen37</td>
<td>11.7 (2.6)</td>
<td>98 (96–98.5)*</td>
<td>92 (87.6–95)</td>
<td>0.0 (0.0)</td>
<td>1.5 (0.7–6.9)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Verhulst44</td>
<td>1.4 (1.1–1.9)*</td>
<td>99 (95.4–99.5)*</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>2.8 (0.8–3.7)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Scholle42</td>
<td>3.0 (2.2–3.5)*</td>
<td>98 (96–98)*</td>
<td>91.8 (2.7)</td>
<td>0.0 (0.0)</td>
<td>1.5 (0.7–6.9)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Scholle42</td>
<td>5.0 (4.3–5.8)*</td>
<td>98 (97–99)*</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>1.1 (0.5–3.2)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Scholle42</td>
<td>8.0 (6.3–10.1)†T1</td>
<td>98 (96–98)*</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>0.9 (0.3–2.7)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Scholle42</td>
<td>11.5 (9.7–13.2)†T2</td>
<td>98 (96–98)*</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>0.4 (0.1–1.8)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Scholle42</td>
<td>12.5 (11.4–15.2)†T3</td>
<td>97 (96–98)*</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>0.5 (0.1–1.8)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Scholle42</td>
<td>15.2 (13.4–17.2)†T4</td>
<td>97 (96–98)*</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>0.1 (0.0–0.8)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Scholle42</td>
<td>16.9 (15.8–17.9)†T5</td>
<td>97 (96–98)*</td>
<td>0.0 (0.0)</td>
<td>0.0 (0.0)</td>
<td>0.1 (0.0–1.3)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Mitterling Women 2012†13</td>
<td>43 (19–77)*</td>
<td>94.4 (90.0–97.0)*</td>
<td>89.0 (81.0–98.0)*</td>
<td>0.7 (0.0–21.5)</td>
<td>2.1 (0.1–25.0)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Mitterling Men 2007†13</td>
<td>31–40</td>
<td>95.5 (92.5–98.0)*</td>
<td>91.0 (86.0–95.0)*</td>
<td>0.7 (0.0–11.7)</td>
<td>1.0 (0.0–22.5)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Mitterling Men 2007†13</td>
<td>&lt;30</td>
<td>95.2 (93.0–98.0)*</td>
<td>90.0 (84.0–95.0)*</td>
<td>0.1 (0.0–3.9)</td>
<td>0.5 (0.0–8.8)</td>
<td>0.17 (0.11)</td>
</tr>
<tr>
<td>Mitterling 2007†13</td>
<td>31–40</td>
<td>95.5 (92.5–98.0)*</td>
<td>91.0 (86.0–95.0)*</td>
<td>0.7 (0.0–11.7)</td>
<td>1.0 (0.0–22.5)</td>
<td>0.17 (0.11)</td>
</tr>
</tbody>
</table>

(Continued)
### Table 1 (Continued)

<table>
<thead>
<tr>
<th>Author</th>
<th>Age (years) – mean (SD) except€</th>
<th>Mean overnight oxygen saturation % mean (SD) except±*</th>
<th>Nadir overnight oxygen saturation % mean (SD) except±*</th>
<th>Apnea Hypopnea Index (events/ hr)</th>
<th>Obstructive Apnea Index (events/hr)</th>
<th>Central Apnea Index (events/hr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mitterling 2007‡</td>
<td>41–50</td>
<td>95.0 (90.0–98.0)*</td>
<td>89.5 (81.0–94.0)*</td>
<td>0.4 (0.0–15.9)*</td>
<td>1.9 (0.0–18.4)*</td>
<td></td>
</tr>
<tr>
<td>Mitterling 2012‡</td>
<td>41–50</td>
<td>93.8 (91.0–96.3)*</td>
<td>88.0 (78.0–93.0)*</td>
<td>1.2 (0.0–19.1)*</td>
<td>2.6 (0.0–22.6)*</td>
<td></td>
</tr>
<tr>
<td>Hertenstein‡</td>
<td>19–30 Female</td>
<td>93.5 (89.3–96.6)*</td>
<td>87.0 (80.0–98.0)*</td>
<td>0.9 (1.0)</td>
<td>1.7 (2.9)</td>
<td></td>
</tr>
<tr>
<td>Hertenstein‡</td>
<td>19–30 Male</td>
<td></td>
<td></td>
<td>1.0 (0.8)</td>
<td>1.0 (0.8)</td>
<td></td>
</tr>
<tr>
<td>Hertenstein‡</td>
<td>31–40 Female</td>
<td></td>
<td></td>
<td>2.8 (2.7)</td>
<td>2.8 (2.7)</td>
<td></td>
</tr>
<tr>
<td>Hertenstein‡</td>
<td>31–40 Male</td>
<td></td>
<td></td>
<td>2.1 (2.1)</td>
<td>2.1 (2.1)</td>
<td></td>
</tr>
<tr>
<td>Hertenstein‡</td>
<td>41–50 Female</td>
<td></td>
<td></td>
<td>2.2 (2.7)</td>
<td>2.2 (2.7)</td>
<td></td>
</tr>
<tr>
<td>Hertenstein‡</td>
<td>41–50 Male</td>
<td></td>
<td></td>
<td>3.5 (3.5)</td>
<td>3.5 (3.5)</td>
<td></td>
</tr>
<tr>
<td>Hertenstein‡</td>
<td>51–60 Female</td>
<td></td>
<td></td>
<td>5.5 (7.8)</td>
<td>5.5 (7.8)</td>
<td></td>
</tr>
<tr>
<td>Hertenstein‡</td>
<td>51–60 Male</td>
<td></td>
<td></td>
<td>4.9 (5.4)</td>
<td>4.9 (5.4)</td>
<td></td>
</tr>
<tr>
<td>Hertenstein‡</td>
<td>61–73 Female</td>
<td></td>
<td></td>
<td>2.1 (2.5)</td>
<td>2.1 (2.5)</td>
<td></td>
</tr>
<tr>
<td>Hertenstein‡</td>
<td>61–73 Male</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gries and Brooks§</td>
<td>≤1</td>
<td>96.4 (1.2)</td>
<td>90.7 (2.6)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gries and Brooks§</td>
<td>1–10</td>
<td>96.8 (1.4)</td>
<td>90.1 (3.6)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gries and Brooks§</td>
<td>10–20</td>
<td>96.5 (1.6)</td>
<td>90.4 (2.7)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gries and Brooks§</td>
<td>20–30</td>
<td>96.3 (1.0)</td>
<td>92.0 (3.4)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gries and Brooks§</td>
<td>30–40</td>
<td>96.3 (1.1)</td>
<td>91.5 (2.2)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gries and Brooks§</td>
<td>40–50</td>
<td>96.0 (1.3)</td>
<td>91.1 (2.1)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gries and Brooks§</td>
<td>50–60</td>
<td>95.8 (1.7)</td>
<td>90.4 (1.9)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gries and Brooks§</td>
<td>≥60</td>
<td>95.1 (2.0)</td>
<td>89.3 (2.8)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Notes:** €Range ± median (95% confidence intervals); *median (range); ¥median (SD) where median is SAT50; ‡Tanner stage; †American Academy of Sleep Medicine (AASM) 2007 criteria for Apnea Hypopnea Index, Obstructive Apnea Hypopnea Index, Central Apnoea Index.
SDB is important as there are significant impacts on health related to the condition, and thus a significant global burden on health care services. In adults, OSAS correlates with increased incidence of cardiovascular and autonomic dysfunction, cerebrovascular complications, metabolic derangements, insulin resistance, malignancy, and overall reduced quality of life, while the effect on cognition and behavior is of concern in children.

The prevalence of SDB in the general adult population ranges between 9% and 38% and may be as high as 49%. Risk factors for development of the condition include obesity, male gender, increasing age, menopause, genetics, ethnicity, craniofacial abnormalities, as well as lifestyle factors such as alcohol consumption and smoking. OSAS is a worldwide condition with remarkably similar prevalence profiles across continents, including North America, Europe, Australia, and Asia. There seem to be distinct associations, however, between geographical regions related to the condition – population demographics may play a role.

SDB is less common in typically developing children than in adults, with prevalence of obstructive SDB in the general pediatric population ranging from 1% to 5% and depending in part on the diagnostic method and threshold, both of which continue to be debated.

Archetypally most studies, regardless of methodology and populations studied, have found that central apneas (pauses in breathing, typically not related to airway obstruction) are more common than obstructive apneas and hypopneas in children, with indices (number of events/hours of sleep) of close to 1/hr and <0.5/hr, respectively. This is highlighted in Table 1, which shows the normal ranges for PSG variables in children from a selection of studies using pre-AASM scoring. AHIs >1 are probably abnormal, as are pCO2 values of >50mmHg.

In addition to the tabulated pre-AASM studies, the Scholle studies were particularly pertinent in attempting to clearly describe sleep using AASM criteria in typically developing children according to Tanner developmental stages across different age groups. In this prospective study of PSG in 209 healthy German children of White ethnicity, aged 1–18 years, obstructive apneas and hypopneas were very rare, with mean OAI of 0.0 across all age groups. Central apneas were more common but decreased with increasing age.

Obesity is the primary risk factor associated with OSAS in adults and is an increasing component factor in children. Upper body obesity may directly augment narrowing of the upper airway. Studies have shown that the dimensions of neck and waist circumference predict OSAS severity in adults.

The most common cause of OSAS in children is considered to be narrowing of the upper airway due to adenotonsillar hypertrophy, with more recent data indicating that it may also be related to airway size and structure. During childhood growth and development, the upper airway is a site for progressive alteration, including changes in growth of tissues as well as functional adaptations in neuromuscular tone and ventilatory drive. Within this time, augmented growth of the adenoids or tonsils could predispose children to upper airway obstruction, principally during sleep. It has been proposed in studies that assessed the mechanical properties of the upper airway, including ventilatory responses to resistive loading, that the upper airway of children is more resistant to collapse compared to adults. Conversely, the dysfunctional soft palate, due to relative hypotonia, in young children without concomitant disorders could also be considered as a cause of obstructive sleep apnea, mostly resolving spontaneously with increasing age. There could be additional contributing factors to developing OSAS and therefore anatomical dimensions of tissues may play a more significant role in maintaining airway patency, particularly during different stages of growth and development.

Regardless of the variability of the diagnostic criteria for obstructive SDB, the prevalence is elevated in certain pediatric populations. This includes children with anemia, such as SCD, in which previously reported prevalence rates range from 5% to 79%. There is arguably a three-pronged approach to understanding the pathophysiological factors involved in OSAS by addressing: i) anatomical factors, ii) upper airway collapsibility, and iii) obesity. There have been challenges in exploring these factors and thus identifying the genetic and environmental risk factors associated with SDB in these populations, due to relatively few studies that include contemporaneous controls assessed against the gold standard polysomnography.

**Effect of ethnicity on SDB**

There are few data comparing the prevalence of SDB between ethnic groups, with lack of data in important populations, including people of African origin, who are typically reported in manuscripts as Black. Several of the studies enrolled a relatively high proportion of people of African origin but not all reported any differences in
SDB prevalence across ethnicities. Traeger\textsuperscript{23} included 48\% Black children, Montgomery-Downs\textsuperscript{41} included 23.1\% and 38.3\% Black children in respective age brackets, Rosen\textsuperscript{37} included 41\% Black children, and Bixler\textsuperscript{39} included 13.8\% Black children. However, Verhulst\textsuperscript{24} included 95\% White children while the studies of Uliel\textsuperscript{22} and Moss\textsuperscript{25} involved only White children.

There are potentially important differences in prevalence of SDB in populations of various ethnicities.\textsuperscript{36,37} Rosen reported an increased prevalence of SDB in Black children in her prospective, cross-sectional, population-based study.\textsuperscript{37} Eight hundred and fifty children aged 8–11 years of age, 41\% Black and 46\% pre-term (<36 weeks gestation), were enrolled, with equal numbers of White and Black pre-term children. Home cardiorespiratory polygraphy studies and Child Sleep Questionnaires were collected. Defined as an OAHI $\geq$5 or an OAI $\geq$1, Black children were four to six times more likely to have OSAS (8.7\%) than White children (2.2\%). Regardless of race, OSAS was three to five times more likely in pre-term (7.4\%) than full-term (2.4\%) children. Black children were more likely to snore and tended to have higher body mass index. There are relatively few data in populations from the Indian subcontinent or China, in whom craniofacial anatomy may play a more important role than obesity.\textsuperscript{65}

“Normal” sleep is different in children of different ethnicities,\textsuperscript{37} as well as for different ages,\textsuperscript{23,66} and socio-economic circumstances. However, due to lack of awareness or other limitations, many published studies have not taken into account these factors, making the classification of “normal sleep” in children problematic. When interpreting literature findings for a specific demographic, we must take into account all confounding factors present within that population – including confounders that the authors may not have addressed. This requires particular attention if comparing ethnicities exposed to different environments, eg, Africans living in Africa vs people of African origin living in a Western country. This highlights a significant limitation in our assessment of this topic but highlights the need for further, well-designed studies that take into consideration confounding factors and the reflection of a “true” demographic.

Genetic factors, including underlying conditions, may be responsible in part for the association between ethnicity and increased prevalence of obstructive SDB. For instance, the underlying mechanisms for adenotonsillar hypertrophy and differences in airway size and structure might be different for particular populations and ethnicities that exhibit a higher prevalence of OSAS. Alterations in the dimensions and shape of the airway may occur in people with anemia secondary to hemoglobinopathies\textsuperscript{61} eg, SCD,\textsuperscript{50} and other anemias, including thalassemia,\textsuperscript{63} as well as Down syndrome\textsuperscript{67} and disorders of bone, such as achondroplasia and osteogenesis imperfecta, or of soft tissue, such as mucopolysaccharidoses.\textsuperscript{68}

In order to explore this, we present an overview of some of these conditions, particularly illustrated by an array of anemia-related disorders and their association with obstructive SDB, before focusing upon SCD and the implications involved with this condition.

### Populations with higher prevalence of OSAS

#### Down syndrome

Children with Down syndrome have a high incidence of OSAS,\textsuperscript{69,70} considered to be related to a combination of adenotonsillar hypertrophy and reduction in the size of the airway in relation to midface hypoplasia and a small jaw. There is some evidence for an effect of OSAS on memory consolidation\textsuperscript{71} and attention\textsuperscript{72} in this population. Uong studied a Down syndrome population without OSAS to assess this population for predisposition to OSAS.\textsuperscript{73} They found a smaller mid and lower face skeleton – ie, smaller mental spine to clivus length, hard palate length, and mandible volume in the Down syndrome group. They also found a smaller airway volume in Down syndrome compared to healthy children.\textsuperscript{73} Adenoid and tonsil volumes were smaller in the Down syndrome group, whereas the tongue, soft palate, pterygoid, and parapharyngeal fat pads were of a similar size to healthy children. Hence, the size of the upper airway is proportionate to soft tissue crowding due to these smaller craniofacial dimensions.

#### Conditions with abnormal bone or connective tissue

Other conditions involving abnormal development of bone or connective tissue from an early age, including achondroplasia, Pierre–Robin syndrome, and Prader–Willi syndrome, may be associated with a small airway related to differences in growth of the mandible and maxilla. In addition to the effect of large tonsils and adenoids, these anatomical differences may play a role in the development of central SDB,\textsuperscript{68,74} but the pathophysiology may be complex.\textsuperscript{75,76} Central apneas may be related to pressure on the brainstem and cervical spinal cord related to foramen magnum stenosis. However, there are relatively few large series comparing the anatomy of the upper airway in these patients to matched controls.
Anemias

Iron deficiency, chronic renal failure, and other chronic diseases, such as β-thalassemia and SCD, are common causes of anemia. Defects of red cell stability, deformability, and metabolism cause a wide range of anemia for which the genetic basis is understood. β-thalassemia and SCD affect 400,000 newborns each year, with several million affected worldwide. Hereditary spherocytosis and pyruvate kinase deficiency are less common, but for all anemias, a wide variety of comorbidities are well recognized and are considered to occur via a variety of mechanisms. The possibility that OSAS and/or NOD play a role has received only modest consideration. The association between anemia and SBD, however, is not well understood.

The normal adult hemoglobin (HbA) consists of two α and two β globin chains combined with a heme molecule. Modifications of the normal structure of hemoglobin can occur through point mutations in the globin gene. Heme production requires adequate iron stores. Anemia is defined as a hemoglobin concentration below 10–13 g/dL, dependent on gender, age, race, and with a higher threshold at altitude. It affects 1.6 billion people worldwide and is frequently associated with poor health. Red cell disorders are therefore a significant burden to health services, with complications impacting quality-of-life and employment opportunities. Low hemoglobin, red cell destruction, and disturbed iron homeostasis lead to alterations in tissue oxygenation and chronic damage to the kidneys, brain, lungs, and heart. However, despite the overlap in the range of complications, the possibility that comorbid SDB might play a role in complications of anemia has received relatively little attention.

Iron deficiency anemia

There are very few reports including sleep studies in iron-deficient patients in any age group, although there is a probable association with periodic leg movements. In adults with newly diagnosed OSAS, iron levels were lower and hepcidin levels were higher compared with controls. Treatment of anemia with erythropoietin and/or iron was associated with improvement in SDB in adults with chronic heart failure.

Thalassemia

Homozygous α-thalassemia is typically lethal. In β-thalassemia, erythropoiesis is limited because of a genetically determined (due to many autosomal recessive genes) inability of the bone marrow to synthesize beta globin chains. The condition is common in the Mediterranean. OSAS has been described in a child with β-thalassemia intermedia. Extramedullary hematopoiesis appeared to have obstructed the nasopharynx on the CT scan. Treatment with hydroxyurea and blood transfusion was associated with relief of symptoms after six weeks. In a study of 10 Israeli patients with β-thalassemia major (n=6) or intermedia (n=4), arousals were common and were induced by periodic limb movements. There was no evidence for OSAS. Multiple sleep latency tests demonstrated that there was objective daytime sleepiness. In another study of 120 children with severe β-thalassemia, 19 (15.8%) snored habitually, of whom 16 had PSG. Ten had OSAS on PSG, which was moderate to severe in six. All 10 had adenoidal hypertrophy and eight had tonsillar enlargement. Those with OSAS had higher serum ferritin compared to those without OSAS. Reactive lymphoid hyperplasia was seen in all six patients who had adenotonsillectomy. Thalassemia patients have typical craniofacial characteristics with a relatively wide maxilla and nose. In one study, three features, including two involving the position of the hyoid bone, as well as tongue length, were shorter in patients with β-thalassemia. The middle airway space was also significantly wider and there was a trend for a wider inferior airway space in these patients.

Congenital dyserythropoietic anemia

Congenital dyserythropoietic anemias are rare autosomal recessive disorders of erythropoiesis typically requiring frequent transfusion. Type 1 is common in Bedouin Arabs living in the Negev desert in Israel. One study looking at 10 of these children and adolescents from one extended Bedouin family found that arousals were common in this condition, mainly related to periodic limb movements, but there was little evidence for OSAS. A PubMed search found no papers on OSAS in other rare hereditary anemia, namely Fanconi anemia, Diamond-Blackfan syndrome, hereditary spherocytosis, or pyruvate kinase deficiency.

Sickle cell disease

SCD is a multi-organ disease with multifaceted pathophysiological mechanisms – the simplicity of the genetic mutation that causes SCD belies the complexity of the disease’s pathophysiology. The sickle cell trait, which originated in West and Central Africa centuries ago,
appears to have arisen as an evolutionary adaptation of natural selection on genetic resistance to malaria. From Africa, the gene spread along the Mediterranean, through Persia and India, and across the Atlantic. More recently, with migration, it has spread to Europe. SCD is now the most common genetic condition at birth in England, affecting 1 in every 2000 live births. Pre-conception counseling and invasive prenatal diagnosis are available but not all couples want these options for a condition with a wide spectrum of outcomes. With increased migration rates and successive generations of people living with SCD, there will be an increasing prevalence of SCD in many countries. Considering the morbidity and mortality associated with SCD, a greater understanding of the complications of the disease and the development of new interventions and therapies is paramount in order to treat this population adequately.

SCD is caused by a genetic mutation of a single base pair change, adenine to thymine, in the 6th codon of the β chain, which results in the alteration of one amino acid (glutamine acid to valine) in position 6 of the β chain of hemoglobin, producing hemoglobin S (HbS; sickle hemoglobin). SCD is an autosomal recessive genetic condition. “Sickle cell anemia” is the term used when both β globin chains exhibit the abnormal HbS (ie, homozygous for sickle hemoglobin (HbSS)). These patients tend to have a worse prognosis and are often symptomatic. The expression “sickle cell disease” includes any genotype where one abnormal HbS β globin chain is inherited with a different abnormal β globin chain that commonly interacts with it. There are many identified hemoglobin variants, but the most common are HbC or Hb β-thalassemia (β− or β0), HbO Arab, HbD Punjab, and HbS Antilles. An abnormal HbS beta globin chain plus a normal HbA beta globin chain constitutes “sickle cell trait” (HbAS). Heterozygotes, ie, those with sickle cell trait, typically do not exhibit symptoms and may not be aware they carry the gene. Hbβ0-thalassemia may be as severe as HbSS, whereas HbSC and Hbβ+ thalassemia tend to be milder. With increasing global migration, there is an increase in combination traits, such as HbβSth, HbE/ath, for which the full range of phenotypic traits is yet to be determined.

When sickle red blood cells (RBCs) become deoxygenated, the HbS becomes insoluble and produces polymers that aggregate into tubular fibers. These enlarge and deform the RBCs, giving the RBC the characteristic sickle shape. Upon restoration of normal oxygen tension, the cells do not necessarily return to normal shape and are left with reduced solubility and elasticity. In addition, RBCs in SCD are typically stiffer compared with normal RBCs. In SCD, RBCs may become dehydrated, which increases the propensity of HbS to polymerize when deoxygenated.

There are three broad ways that HbS contributes to the pathophysiology of the disease. Firstly, the deformed shape and rigidity can lead to vascular obstruction and consequently ischemia. Secondly, the membrane damaged RBCs lead to increased adherence to, and damage of, the vascular endothelium and thus promote a proliferation of white blood cells, cytokines, growth factors, and coagulation proteins, which together increase vascular obstruction. Thirdly, membrane damage causes hemolysis – RBC breakdown. In a healthy person, an RBC survives 90–120 days in the circulation. However, due to the abnormal structure, sickle red blood cells only survive 10–20 days until they are sequestered by the spleen, thus resulting in hemolysis.

The cardinal features of SCD are chronic hemolytic anemia and complications related to recurrent vaso-occlusion. One of the strongest precipitants of vaso-occlusion (or polymerization of sickle hemoglobin) is oxygen desaturation, which has been associated with increased risk of a number of complications, including central nervous system events. Oxygen desaturation in SCD is in part due to the presence of dysaemoglobins such as carboxyhemoglobin and methemoglobin, which are increased during intravascular hemolysis but fail to transport oxygen. SDB/OSAS is often comorbid with SCD, potentially exposing patients to additional hypoxia. Many adults and children with SCD experience at least some degree of nocturnal oxyhaemoglobin desaturation, which could be as a consequence of OSAS; other potential causes include pulmonary disease, shunting at cardiac or pulmonary level or the decreased oxygen affinity for hemoglobin S. There is a good case for diagnosing obstructive SDB and NOD in SCD as evidence is accumulating that upper airway obstruction and the degree of oxygen desaturation affects cognitive function as well as the rate of complications, including stroke.

Pathophysiology of sleep-disordered breathing in sickle cell anemia

Our knowledge of the pathophysiology of SDB and NOD in the SCD population is limited. Hypotheses, which are not mutually exclusive, include the following:
(i) hypoventilation due to chronic lung disease,
(ii) presence of OSAS,
(iii) left ventricular diastolic dysfunction,
(iv) pulmonary hypertension,
(v) shunting at cardiac or pulmonary level,
(vi) presence of dyshemoglobins

The most common cause of OSAS in children with SCD is adenotonsillar hypertrophy, the same primary cause identified for the general pediatric population. However, the underlying mechanisms for adenotonsillar hypertrophy and differences in airway size and structure might be different for SCD with OSAS compared with non-SCD children with OSAS, hence the higher prevalence in this specific pediatric SCD population. For instance, in SCD, adenoid and tonsillar hypertrophy might be as a consequence of a compensatory mechanism for functional asplenia, or as a consequence of upper respiratory infections due to limited opsonization of bacterial pathogens. Differences in craniofacial development due to atypical growth trajectories or extramedullary hematopoiesis should be considered. This includes variants of maxillary structure in the vertical, horizontal, and transverse planes; poor lip seal; and imbalance in the function of the orofacial muscles, particularly the mandibular elevator and depressor muscles, which may lead to upper airway obstruction and mouth breathing.

There are two crucial questions related to OSAS in children with SCD that are yet to be resolved:

(i) What factors differ between children with SCD with and without OSAS, after matching for adenotonsillar size?
(ii) When similar groups of children undergo adenotonsillectomy, why is the outcome better in some children than others?

Literature exploring OSAS in adults with SCD is markedly limited in comparison to OSAS in children with SCD. Sharma and Whitesell conducted small studies evaluating SDB in adults with SCD suggesting prevalences of 44% and 50%. Due to improvements in genetic screening and clinical management, the trend in life expectancy for SCD is increasing. Our knowledge of SDB in an adult SCD population, therefore, needs to be broadened to appropriately treat this novel population; extrapolating data from childhood studies is inadequate.

There may be other contributory factors within the upper airway to SDB in the SCD population that could be targeted for potential therapy. For example, central sleep apnea (CSA), or central sleep apnea syndrome (CSAS), distinct from obstructive SDB, is a sleep-related disorder which is also associated with a reduction in blood oxygen saturation. In CSA, the effort to breathe is diminished or absent, typically for 10–30 s either intermittently or in cycles. There are very few data on CSA or CSAS in SCD and the possibility that coincidental Chiari malformation plays a role has received little attention. Further discussion of this is beyond the scope of this review but warrants further research.

Regarding sleep studies in children with SCD, Strauss et al reported a novel study on the prevalence of SDB in Black children with and without SCD, measuring PSG outcomes in 36 subjects with SCD (aged 6.9±4.3 years) and 36 presumably Black control subjects (aged 6.6±3.4 years). It was part of a more comprehensive investigation on the prevalence, contributory mechanisms, and pulmonary and vascular consequences of oxygen desaturation in children with SCD at the Children’s Hospital of Philadelphia (CHOP). Strauss et al identified a greater prevalence of OSAS in SCD (7/36; 19.4%) compared to healthy children (0/2; 0%), with increased arousals (13.7±4.7 events/hr vs 10.8±3.8 events/hr, \( P<0.05 \)), lower arterial oxygen saturation nadir (84.3%±12.3% vs 91.2%±4.2%, \( p<0.05 \)), and increased peak end-tidal \( \mathrm{CO}_2 \) (53.4±8.5 mm Hg vs 42.3 ±5.3 mm Hg, \( P<0.001 \)) in the SCD group. In contrast to the data from Rosen’s population-based study, but in line with the previous data from CHOP, none of these, presumably Black, control children had OSAS.

The Sleep Asthma cohort (SAC) is an ongoing longitudinal study of children with SCD aged 4–18 years, resident in the United States of America or the United Kingdom, who have undergone PSG regardless of symptoms related to sleep. A cross-sectional analysis of baseline data was collected as part of the SAC study from eligible patients aged 4–18 years who were HbSS or compound heterozygotes for HbSβ-thalassaemia zero (HbSβ⁰). The study was designed to evaluate the contribution of sleep abnormalities and asthma to SCD-related morbidity. Of 243 children (median age 10 years, 50% male, 99% Black, 95% HbSS), OSAS defined by obstructive AHI at cut points of \( \geq 1 \) or \( \geq 5 \), was present in 100 (41%) or 25 (10%) children, respectively. In multivariable analyses, habitual snoring and lower daytime oxygen saturation were risk factors for OSAS.

OSAS therefore appears to be four to six times more common in Black children than White children and is particularly prevalent in those with SCD.
available, in addition to adenotonsillar hypertrophy, differences in facial features and upper airway anatomical structure probably play a role in the prevalence of OSAS.

Anatomy of the airway
Anatomical dimensions were compared between ethnic groups in a study that found increased tongue area and soft palate length in adult African-Americans. OSAS may be more closely related to these soft tissue dimensions in Black adults, who have generalized leptoprosopic facial shape (longer facial height with less depth) while White adults with OSAS tend to have brachycephaly (shorter, wider skull and pharynx dimensions). Investigating anatomical variation of the upper airway and related structures is therefore a fundamental area of OSAS research in all populations.

We will now review studies that have explored structural components of the upper airway using imaging, primarily MRI, as well as other clinical methods. Again, we need to be mindful when interpreting these studies as the literature on SDB in SCD rarely includes matched control children of African origin without SCD. Additionally, the majority of PSG studies of children with SCD have selected patients with symptoms. Socioeconomic and environmental factors should also be taken into account, including air pollution, smoking, and poor nutrition, as well as the genetic influences which apply to all populations.

Anatomical variations of the upper airway and associated structures
Imaging the upper airway in normal growth and development
There are relatively few studies that explore developmental changes of the upper airway in normally developing children and they exhibit some contradictory findings. Arens et al found that the lower face skeleton had a strong linear correlation with age and height of their healthy subjects aged 1–11 years. The measurements for mental spine to clivus length and intermandibular length delineated the mid and lower face skeletal dimensions. As the mid and lower face skeletal dimensions increased in size with increasing age and/or height, the size of the upper airway tissues within this demarcated space also grew at a constant proportion. Therefore, no age-related growth velocities for any tissue contributing the airway were found; this included adenoidal and tonsillar tissues, as well as the size of the tongue, soft palate, tonsils, parapharyngeal fat pads, and pterygoids. Additionally, nasopharyngeal airway dimension grew at a constant proportion to skeletal growth. A study with similar outcomes for this age group by Vogler et al (2000), using comparable methodology, also found that the adenoid pad grew at a constant proportion to age, up to 10 years, with the largest size reached aged 7–10 years. Their study, however, continued beyond this age and subsequently the growth velocity diminished until 60 years old.

In contrast to these findings are studies by Fujioka et al and Jeans. Fujioka et al found that adenoid:nasopharyngeal (AN) ratio, demarcated on lateral skull radiograph, did exhibit some age-related growth velocity during childhood. The AN ratio typically increased from 1.5 months to 1 year 3 months, reaching its highest value at 4 years 6 months, after which it gradually decreased until 12 years 6 months and subsequently decreased dramatically at 15 years 6 months. The study also proposed that linear measurements of the adenoids and nasopharyngeal airway that generate an AN ratio greater than 0.8 were related to a subjective assessment of subjects with enlarged adenoids. These findings were corroborated by Jeans et al using lateral neck cephalometry. Although they found a constant growth velocity of the total nasopharynx area between 3 and 11 years, they noted a mild reduction in the area of the nasopharyngeal airway between 3 and 5 years that occurred simultaneously with a mild increased growth velocity of the soft tissues. Thereafter, nasopharyngeal airway continued to grow linearly with no further increase in soft tissue within the nasopharyngeal distribution. This AN ratio measurement could warrant further research, particularly with the advent and accessibility of superior imaging modalities compared to the standard plain radiograph that was used originally.

While there are some discrepancies, these studies generally show a similar trend of constant proportion of growth of the upper airway and its structures up to 10 years old. It is interesting to note the reduction of nasopharyngeal airway with simultaneous increase in soft tissues at age 3–5 years, considering that OSAS is commonly diagnosed around a similar age. This may indicate a mechanism that predisposes to OSAS – a narrowed nasopharynx due to overgrowth of soft tissues, such as the adenoids. Hence, making the AN airway measurement is important.

It would be interesting to see whether the growth trajectory for SCD children matches the growth trajectory for normal, healthy children. Any apparent age-related growth velocities in SCD could provide further information for risk stratifying for SDB. Imaging of the anatomical dimensions of the upper airway in developing children and young adults may provide insight into structural growth changes linked with SDB and...
specific populations—focusing upon the aforementioned regions of interest, including the nasopharyngeal airway, AN airway ratio, and mid to lower face skeletal dimensions, eg, mental spine to clivus length. The imaging modality that is widely available and most appropriate in reproducing three-dimensional innate images of soft tissues, such as those comprising the upper airway structure, is MRI. This generates a medium for comparison of upper airway anatomical measurements between patient groups.

Employing MRI to assess the structure of the upper airway in OSAS populations

The advent of MRI has led to greater understanding of growth and development of the upper airway during childhood. MRI provides an innately scaled, three-dimensional image of the tissues of the upper airway. Advantages include the avoiding of ionizing radiation in a pediatric population and the superior resolution for soft tissues compared to radiographs.\(^2\) There are limitations including the comparatively long acquisition time of an MRI scan compared with a lateral neck X-ray or CT scan, which may precipitate motion artifact. Additionally, some studies used sedation or anesthesia as part of their protocol in young children, which could potentially lead to diminution of muscle tone impacting on airway measurements.

A literature search using the terms “obstructive sleep apnea”, “adenoids”, “airway”, and “MRI” was performed to explore methods used for measuring regions of interest in the upper airway in pediatric OSAS populations.

Table 2 compares the anatomical landmarks used by each study to demarcate the upper airway with particular regions of interest, and Table 3 describes the key features of each of these studies. The previously mentioned imaging studies have been included for comparison.

MRI of the upper airway: comparison between healthy controls against OSAS populations

Several studies, as illustrated in Tables 2 and 3, found upper airway anatomical differences between OSAS and non-OSAS populations. The volume/area/size of the upper airway was smaller in subjects with OSAS,\(^47\)–\(^50\) and in adults, the volume of the lateral pharyngeal walls was larger in OSAS.\(^51\) The adenoids, tonsils,\(^47,49,50,123,124\) retropharyngeal nodes,\(^49,50,123\) deep cervical nodes,\(^50,123\) and parapharyngeal fat pads\(^49\) were larger in OSAS. The soft palate\(^47\) and the total soft tissues were found to be larger in OSAS.\(^51\) The volume of the mandible was similar in the control and OSAS groups in some studies,\(^47\) but smaller in the OSAS group in others. Regarding the tongue in patients with OSAS compared with controls, it was found to be either similar in children\(^47\) or larger in adults.\(^51\) Abdominal visceral fat was greater in OSAS.\(^49\)

In a study using respiratory-gated MRI under sedation, Arens et al found a smaller upper airway cross-sectional area, particularly during inspiration, in children with OSAS.\(^125\) Airway narrowing occurred during inspiration without evidence of complete airway collapse, while airway dilatation occurred during expiration. The magnitude of the fluctuations in cross-sectional area during tidal breathing was significantly greater in OSAS compared to healthy children.

Patini conducted a systematic review\(^126\) to assess the effectiveness of MRI in evaluating upper airway structures in children with OSAS and, after strict exclusion parameters were applied (initially 1170 titles for possible inclusion), a meta-analysis was performed on three of the studies of patients with OSAS and controls: Arens,\(^47\) Arens,\(^48\) and Cappabianca.\(^124\) The studies showed differences in minimum retropharyngeal cross-sectional area, nasopharyngeal airway, combined upper airway volume, tonsillar, and adenoid cross-sectional and volumetric indices. In addition, in the latter study, midsagittal cross-sectional area of the soft palate was higher, mandibular volume was lower, and the vertical position of the hyoid bone was significantly lower in the OSAS group compared with controls.\(^124\) There were also differences in the position of the maxilla and the mandible, in both cases in the sense of retroposition.\(^124\) The results suggested that MRI could be a useful tool in pediatric OSAS populations for calculating the total volume of the upper airway structures and thus enabling diagnostic examination and clinical management in practice, as it allows assessment of regions of interest that appear to contribute to obstruction—lumen, soft, and skeletal tissue.

MRI of the upper airway—comparison between healthy controls against SCD populations and association with OSAS

To our knowledge, there is only one MRI study that evaluates the upper airway in children specifically with SCD with comparison to OSAS. Strauss et al found that children with SCD tended to have a smaller upper airway and larger adenoids, retropharyngeal nodes, and deep cervical nodes.\(^50\) Only lymphoid tissues were measured in this study. In terms of sleep studies, as previously referenced, Strauss et al identified a greater prevalence of OSAS in SCD (19%) compared to healthy children (0%).\(^50\) They found that in children with
SCD, higher AHI, lower arterial oxygen saturation nadir, increased peak end-tidal CO$_2$, and increased arousals correlated with upper airway lymphoid tissue size.

In a clinical study including subjects with SCD, Salles et al found a positive correlation between height/age z score and cervical circumference. The presence of nocturnal oxygen desaturation was associated with cervical circumference and abdominal circumference. There was a negative correlation between oxygen desaturation and maxillary intermolar distance and mandibular intermolar distance, ie, in SCD subjects that exhibited nocturnal oxygen desaturation, smaller maxilla, and mandible measurements were noted (measured using cephalometry, not MRI).

In summary, children with OSAS typically tended to have smaller sized airways and larger lymphoid and soft tissues compared to children without OSAS. Determining the proportion of growth of the upper airway and related structures for children with anemia such as SCD could potentially provide novel targets for therapies, particularly regarding the size of the lower facial skeleton.

Hypotheses stemming from the literature: considerations for potential treatment targets for SDB

The findings in the literature suggest it is plausible that people with OSAS have smaller sized airways due to smaller lower facial skeleton (eg, smaller mandible) and thus exhibit overcrowding within a defined area – rather than purely having enlarged adenoids and tonsils.

**AN ratio**

It is also possible that it is the relationship between the size of the adenoid to the nasopharyngeal airway, ie, the AN ratio, that is most important. Adenotonsillectomy is based on the assumption that hypertrophic adenoids and tonsils are the cause of OSAS and thus removal should alleviate symptoms. Differences in AN ratio may suggest why adenotonsillectomy is ineffective in some patients. For instance, if the adenoids are removed but this has little effect on the AN ratio, this may be because, even though the adenoids appeared hypertrophic, the nasopharynx was sufficiently large enough to combat this; hence there was no therapeutic effect from targeting this one factor alone.

Thus, an alternative treatment option could aim to target a combination of factors associated with OSAS, as highlighted in these studies: a triad of lymphoid tissue hypertrophy, airway volume distribution, and lower facial skeleton.

**Lymphoid tissue hypertrophy**

Children with OSAS show increased nasal and oropharyngeal inflammation with increased local and systemic inflammatory markers and pro-inflammatory cytokines which enhance lymphoid tissue proliferation. Novel therapies to combat these effects include anti-
Table 3 Describes these studies found

<table>
<thead>
<tr>
<th>Paper</th>
<th>Subjects</th>
<th>Measurement method and software</th>
<th>Outcome</th>
<th>Statistics used</th>
<th>Figures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fujioka et al 1979. Radiographic eva-</td>
<td>1398 infants and children (1</td>
<td>Lateral radiographs. Adenoid (linear measurement from maximal convexity along inferior margin of</td>
<td>An adenoidal-nasopharyngeal ratio greater than 0.8 was present in 94% subjectively judged to have enlarged</td>
<td>Information not available</td>
<td>Figure S1</td>
</tr>
<tr>
<td>luation of adenoidal size in children:</td>
<td>month–16 yrs)</td>
<td>adenoid shadow to straight part of anterior margin of basiocciput), Nasopharynx (linear</td>
<td>adenoids</td>
<td></td>
<td></td>
</tr>
<tr>
<td>adenoid nasopharyngeal ratio</td>
<td></td>
<td>measurement from posterior superior edge of hard palate to antero-inferior edge of sphenobasioccipital synchondrosis. When synchondrosis is not clearly visualized, the site of crossing postero-inferior margin of lateral pterygoid plates and floor of bony nasopharynx)</td>
<td></td>
<td>Figure S2</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arens et al 2001. MRI of upper airway</td>
<td>18 children with OSAS (age 4.8±2.1</td>
<td>1.5 Tesla Siemens vision system. Axial and sagittal T1- and T2-weighted images with 3-mm slice thickness and 1 NEX. Slices spanning from base of orbital cavities to epiglottis. Axial = T1-weighted image, maximal tonsil cross-sectional area level, cross section measurements of oropharyngeal airway (anterior border soft palate/tongue, lateral border tonsils, posterior border pharyngeal constrictor muscle), tonsils, pterygoids, parapharyngeal fat pads. Also, linear measurements across transverse line centre of tonsils (inter tonsillar width, bilateral tonsillar width, bilateral fat pad width, bilateral pterygoid width, intermandibular distance). Other axial levels - largest adenoid cross section, maxilla width, internal distance between mandibular heads. Sagittal = T1-weighted midsagittal, cross-sectional area of nasopharyngeal airway (anterior border vomer, posterior border adenoid, inferior border horizontal line above hard and soft palate), adenoid, soft palate, tongue, mandible, hard palate. Also, linear measurements along oblique line of mental spine, centre soft palate and clivus (ie, tongue oblique, soft palate oblique, airway oblique, adenoid oblique, mental spine-dibius oblique). Length - hard palate (anterior nasal spine to end palate bone) Volumetric measurements = adjacent axial slices. Combined upper airway (nasopharynx and oropharynx), tongue, soft palate, mandible (T1-weighted), tonsils, adenoid (T2 - better resolution lymphoid tissues)</td>
<td>Two-tailed unpaired t test, Wilcoxon rank test, or chi-square test. The Pearson correlation used to assess linear correlation of volume percent difference of each OSAS-control pair and AHI</td>
<td></td>
<td></td>
</tr>
<tr>
<td>structure of children with OSAS</td>
<td>yrs); 18 matched</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(Continued)
Table 3 (Continued).

<table>
<thead>
<tr>
<th>Paper</th>
<th>Subjects</th>
<th>Measurement method and software</th>
<th>Outcome</th>
<th>Statistics used</th>
<th>Figures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uong 2001. MR imaging of upper airway in children with down syndrome</td>
<td>11 down syndrome children without OSA (age 3.2 yrs±1.4 yrs); 14 controls (age 3.3±1.1 yrs)</td>
<td>1.5-T Siemens vision system, VIDA software. Sequential T1 and T2-weighted spin-echo axial and sagittal. Axial retropalatal axial TI-weighted image at level of maximal tonsillar cross-sectional area they measured tonsils, pterygoids, parapharyngeal fat pads, airway cross section. Other axial images – largest adenoid area, maximum maxilla width, internal distance between mandibular heads. Sagittal: midsagittal T1-weighted image they measured cross-sectional areas of adenoid, soft palate, tongue, mandible, hard palate, combined nasopharyngeal and oropharyngeal airway. Length of hard palate (nasal spine to end palatine bone). Mandible size (mental spine and the clivus through soft palate centroid). Volumetric measurements – adjacent axial slices. Adenoid, tonsils, (both T2), tongue, soft palate, mandible, total nasal/oral pharyngeal airway (rest T1)</td>
<td>In Down syndrome group found smaller airway volume, mid and lower face skeleton. Shorter mental spine-clivus distance, hard palate length and mandible volume. Adenoid and tonsil volume smaller. Tongue, soft palate, pterygoid, and parapharyngeal fat pads similar to controls</td>
<td>Two-tailed unpaired t test, Wilcoxon rank test, or chi-square test</td>
<td>Figure S3</td>
</tr>
<tr>
<td>Arens et al 2002. Linear dimensions of the upper airway structure during development</td>
<td>92 normal children</td>
<td>1.5 Tesla Siemens vision system. Sequential T1-weighted images. VIDA software. Sagittal = midsagittal T1-weighted, lower face sagittal skeletal growth by measuring mental spine-clivus length (distance between mental spine – ie, point of insertion of geniohyoid muscle – and clivus passing through soft palate centroid. Linear measurements – tongue oblique width, soft palate oblique width, nasopharyngeal airway oblique width, adenoid oblique width. Axial = lower face skeletal growth was determined by intermandibular length (between medial aspects of both mandibular rami along transverse line passing through tonsillar centroid) at level of maximal tonsillar cross section. Linear measurements included tonsillar width, bilateral tonsillar width, bilateral fat pad width, bilateral pterygoid width. Also, oropharyngeal width (maximal oropharyngeal width on a line parallel to intermandibular line)</td>
<td>In midsagittal plane, mental spine-clivus distance related linearly to age. Tongue, soft palate, nasopharyngeal airway, adenoid increased with and maintained constant proportion to mental spine-clivus distance. Linear relationship for mandibular growth growth measured along intermandibular line on axial plane and age. Intertonsillar, tonsils, parapharyngeal fat pads, pterygoid widths maintained constant proportion to intermandibular width with age. Lower face skeleton grows linearly along sagittal and axial planes from 1st to 11th year. Soft tissues, incl tonsils and adenoids, surrounding upper airway grow proportionally to skeletal structures</td>
<td>Linear regression analysis</td>
<td>Figure S4</td>
</tr>
</tbody>
</table>

(Continued)
Table 3 (Continued).

<table>
<thead>
<tr>
<th>Paper</th>
<th>Subjects</th>
<th>Measurement method and software</th>
<th>Outcome</th>
<th>Statistics used</th>
<th>Figures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arens et al 2003. Upper airway size analysis by MRI of children with OSAS</td>
<td>20 children with OSAS (age 3.7±1.4); 20 controls</td>
<td>1.5 Tesla Siemens vision system. Sequential T2-weighted spin echo axial sections obtained spanning from orbital cavity to larynx. Images transferred to SUN workstation. 3DVIEWNX software (fuzzy connectedness segmentation). 1. Upper airway centerline = bounded by upper nasopharynx (posterior edge vomer), lower oropharynx (superior part epiglottis). Airway regions adjacent to adenoid, tonsils, overlap measured along centerline. 2. Upper airway cross-sectional area = at planes orthogonal to centreline. Mean and minimal cross section of total airway; mean cross section adjacent to adenoid, tonsils and overlap between the two. 3. Upper airway volume = centerline length and mean cross-sectional area. Volumes for 10 consecutive segments at 10% increments of centerline were computed.</td>
<td>In OSAS group mean and minimal cross-sectional area of total airway smaller. Upper airway in OSS is most restricted where adenoid and tonsils overlap</td>
<td>Two-tailed unpaired t test, Wilcoxon rank test, or chi-square test</td>
<td>Figure S5</td>
</tr>
<tr>
<td>Schwab et al 2003. Identification of upper airway anatomic risk factors for OSAS with</td>
<td>48 subjects with OSAS; 48 controls</td>
<td>1.5 Tesla (online supp). Volumetric measurements = lateral pharyngeal walls (retropalatal and retroglossal); soft palate; tongue (genioglossus, geniohyoid, hyoglossus, myohyoid, digastric, myohyoideus muscles); parapharyngeal fat pads; total</td>
<td>In OSAS group, volume of lateral pharyngeal walls, tongue, total soft tissues larger. Increased risk of OSAS the larger the volume of the tongue, lateral pharyngeal walls and total soft tissue</td>
<td>Chi-square tests and t tests. Multiple logistic regression models used to obtain adjusted odds ratios and 95% CI for effect of 1 SD change in size of airway and soft tissue measurement.</td>
<td>Figure S6</td>
</tr>
<tr>
<td>Arens et al 2005. Changes in upper airway size during tidal breathing in children with OSAS</td>
<td>10 OSAS children (age 4.3 ±2.3 yrs); 10 matched controls</td>
<td>1.5 Tesla Siemens Sonata system using 2D trueFISP sequence. VIDA software. Axial images. Respiratory gating performed. Data for 40 mm axial from epiglottis to upper nasopharynx obtained from 10×4 mm slices using 10 successive breaths in tidal breathing (Vt) – 10,30,50,70,90% of inspiration and of expiration.</td>
<td>Subjects with OSAS – smaller upper airway cross-sectional area; airway narrowing during inspiration but without airway collapse; airway dilatation during expiration; greater fluctuations in airway area during tidal breathing.</td>
<td>Two-tailed unpaired t test, Wilcoxon rank test, or Chi-square test</td>
<td>Figure S7</td>
</tr>
<tr>
<td>Schwab et al 2006. Family aggregation of upper airway soft tissue structures in normal subjects and pts with sleep apnea</td>
<td>55 sleep apnea probands; 55 proband siblings</td>
<td>1.5 Tesla MRI scanner spin echo axial and sagittal images. 3D volumes of lateral pharyngeal walls (retropalatal and retroglossal regions); soft palate; tongue (genioglossus, geniohyoid, hyoglossus, myohyoid, digastric, myohyoideus muscles); parapharyngeal fat pads; soft tissue.</td>
<td>Volume of lateral pharyngeal walls, tongue, total soft tissue demonstrated heritability. Upper airway larger in normal subjects. Tongue, lateral parapharyngeal fat pads, lateral parapharyngeal walls larger in patients with apnea</td>
<td>Mixed model analysis of variance ANOVA; odds ratio</td>
<td>Figure S8</td>
</tr>
</tbody>
</table>
Table 3 (Continued).

<table>
<thead>
<tr>
<th>Paper</th>
<th>Subjects</th>
<th>Measurement method and software</th>
<th>Outcome</th>
<th>Statistics used</th>
<th>Figures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arens et al 2011. Upper airway structure and body fat composition in obese children with OSAS</td>
<td>22 obese children with OSAS (12.5±2.8 yrs; BMI z score 2.3±0.3); 22 obese controls</td>
<td>16 channel Philips 3.0 Tesla Achieva Quasar TX scanner. Amira software. <strong>Volumentric measurements</strong> – Upper airway = nasopharynx (superior to soft palate and continues anteriorly, through choanae, with nasal cavities); oropharynx (region between soft palate and larynx, posterior 1/3rd of tongue as anterior border); hypopharynx (posterior to larynx, includes pyriform recesses and vallum); Lymphoid tissue = (adenoid, combined palatine tonsils, combined retropharyngeal nodes – between internal carotid arteries from base skull to hyoid bone – and deep cervical lymph nodes – internal jugular vein from base skull to hyoid bone). Tongue (inc. genioglossus and geniohyoid muscles). Soft palate. Mandible. Head/neck = parapharyngeal and subcutaneous fat.</td>
<td>Size of lymphoid tissue correlated with OSAS severity but BMI z score did not have modifier effect on lymphoid tissue. OSAS subjects had increased size parapharyngeal fat pads, abdominal visceral fat (did not correlate with OSAS severity)</td>
<td>Two-tailed paired t tests, Wilcoxon signed rank tests or McNemar’s tests. Pearson correlations derived between AHI and BMI z score. Mixed effects regression models using AHI as dependant variable and conditional regression models with OSAS vs non OSAS as dependant variable</td>
<td>Figure S99</td>
</tr>
<tr>
<td>Cappabianca et al 2012. Magnetic resonance imaging in the evaluation of anatomical risk factors for pediatric obstructive sleep apnea – hypogonsal: a pilot study</td>
<td>80 Caucasian children aged 4–15 years, of which 40 had been diagnosed with OSAS</td>
<td>Retrospective analysis MRI head and neck between March 2010–2012, 1.5T magnet Symphony, Siemens. <strong>Axial T1-weighted images</strong>: cross-sectional area oropharyngeal airway, bounded anteriorly by soft palate or tongue, laterally by tonsils, posteriorly by pharyngeal constrictor muscles; intermandibular distance, maximum distance between the mandibular rami; maxillary width, maximum transverse diameter of superior jaw. <strong>Axial T2-weighted images</strong>: largest cross-sectional area of adenoids, tonsils, pterygoids, and parapharyngeal fat pads. <strong>Mid sagittal T1-weighted image</strong>: cross-sectional area of nasopharyngeal airway, bounded anteriorly by vomer, posteriorly by adenoid, and inferiorty by horizontal line above the hard and soft palate; the cross-sectional area of adenoids, soft palate, tongue, mandible, and hard palate. Other: distances from a transverse line perpendicular to the floor passing through the hyoid (hy, the most cranially located point of the hyoid bone) to nasion (hy-n), to sella (hy-s) and to supramentale (hy-B) and the angles sella– nasion– subspinale (SNA), sella– nasion– supramental (SNB) and subspinale– nasion– supramental (ANB). <strong>Adjacent axial slices to determine volumes</strong>: combined upper airway (nasopharynx and oropharynx), adenoid, tonsils, tongue, soft palate, and mandible. <strong>Volumentric measurements</strong> (except tonsils and adenoid) from axial T1-weighted slices from base of orbital cavities to epiglottis. T2-weighted images for adenoid and tonsils due to better resolution of lymphoid tissue. Tonsillar volume = volume of right and left tonsils</td>
<td>The total upper airway volume of children with OSAS (1.4±0.7 cm³) was significantly smaller (p&lt;0.01) compared to controls (1.6±1.1 cm³); soft palate volume was significantly larger (p&lt;0.01) in OSAS group (3.9±1.3 cm³) compared to controls (3.1±1.4 cm³); adenoid and tonsils were significantly larger in children with OSAS compared to controls (p&lt;0.01) with mean adenoid volume in children with OSAS 9.1±1.8 cm³ vs 6.3±2.1 cm³ in controls and mean tonsillar volume in OSAS 9.2±1.5 cm³ vs 6.5±1.7 cm³ in controls. Interestingly, similarly to Arens (2001) they also noted a smaller mandibular volume (p=0.05) in the OSAS group (22.2±2.23) compared to control group (25.4±2.4 cm³)</td>
<td>Student’s t test</td>
<td>Figure S10124</td>
</tr>
</tbody>
</table>
Table 3 (Continued).

<table>
<thead>
<tr>
<th>Paper</th>
<th>Subjects</th>
<th>Measurement method and software</th>
<th>Outcome</th>
<th>Statistics used</th>
<th>Figures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Strauss et al 2012. Upper airway lymphoid tissue size in children with sickle cell disease</td>
<td>36 children with SCD (aged 6.9±4.3 yrs); 36 controls</td>
<td>1.5 Tesla Siemens vision system. Axial and sagittal T1- and T-2 weighted images 3 mm slice and 1 NEX, from orbital cavity to larynx. Annonymised, converted to DICOM format. AMIRA software. Airway = nasopharynx (superior to level of soft palate and continuous anteriorly, through choanae, with the nasal cavities); oropharynx (between level of soft palate and larynx, communicating anteriorly with oral cavity, posterior 1/3rd of tongue as anterior border); hypopharynx (posteroalateral to larynx, communicating with cavity of larynx through auditus, incl pyriform recessed and vallecculae). Lymphoid tissues = adenoid, combined palatine tonsils, combined retropharyngeal nodes – lymph nodes between internal carotid arteries from base of skull to hyoid bone – and combined deep cervical lymph nodes – internal jugular vein from base skull to hyoid bone</td>
<td>Children with SCD – smaller upper airway with larger adenoid, retropharyngeal nodes, deep cervical nodes. AHI correlated positively with upper airway lymphoid tissues size</td>
<td>Two-tailed unpaired t tests and chi-square test. Pearson correlations between AHI and upper airway lymphoid tissues.</td>
<td>Figure S11</td>
</tr>
<tr>
<td>Nanadalike et al 2012. Adenotonsillectomy in obese children with OSAS: MRI findings</td>
<td>27 obese children with OSAS (age 13±2.3 yrs; BMI z score 2.5±0.3)</td>
<td>AMIRA software. Philips 3.0 Tesla Achieva scanner with 16 channel surface array coil. Images anonymised converted to DICOM. Sagittal: airway – nasopharynx (superior boundary is base of skull – basi-sphenoid and basi-occiput; inferior boundary upper soft palate; anterior boundary nasal cavities, choana orifice, posterior nasal septum; posterior boundary pharyngobasilar fascia, superior pharyngeal constrictors). Oropharynx (superior boundary soft palate; inferior boundary upper epi-glottis – vallecula fossa – and root of tongue; anterior boundary oral cavity, posterior and lateral boundary superior and middle pharyngeal constrictors). Lymphoid tissues (adenoid, combined palatine tonsils, lingual tonsil, combined retropharyngeal nodes – located between internal carotid arteries from skull base to hyoid bone – plus deep cervical lymph nodes level II – located internal jugular vein from skull base to hyoid bone). Tongue (inc genioglossus and geniohyoid muscles). Soft palate. Mandible. Head and neck subcutaneous fat</td>
<td>Pts followed up 6.1±3.6 mo after AT. AT improved AHI. AT increased volume of nasopharynx and oropharynx, reduced tonsils, no effect on adenoid, lingual tonsil, retropharyngeal nodes. Increase volume soft palate and tongue</td>
<td>Wilcoxon signed rank test for paired data</td>
<td>Figure S12</td>
</tr>
<tr>
<td>Parikh et al 2013. Deep cervical lymph node hypertrophy: a new paradigm in the understanding of pediatric obstructive sleep apnea</td>
<td>70 children with OSAS (mean age 7.47 yrs; BMI 23.63 kg/m²) and 76 healthy matched controls (mean 8.00 yrs; BMI 20.87 kg/m²)</td>
<td>1.5 Tesla (Siemens Vision System, Iselin, NJ). Four regions of lymphoid tissue: tonsil, adenoid, retropharyngeal nodes (between internal carotid arteries from the skull base to the hyoid bone), and the upper jugular lymph nodes (along the internal jugular vein from the skull base to the hyoid bone)</td>
<td>Children with OSAS have larger volumes of deep cervical lymph nodes and adenotonsillar tissue than normal controls</td>
<td>Chi-square analysis and Student’s t test were used to compare demographics and lymph node volumes. Fisher’s exact test and chi-square analysis were used to compare sleep data</td>
<td>Figure S13</td>
</tr>
<tr>
<td>Paper</td>
<td>Subjects</td>
<td>Measurement method and software</td>
<td>Outcome</td>
<td>Statistics used</td>
<td>Figures</td>
</tr>
<tr>
<td>-------</td>
<td>----------</td>
<td>---------------------------------</td>
<td>---------</td>
<td>----------------</td>
<td>---------</td>
</tr>
<tr>
<td>Salles et al 2014. Association between morphometric variables and nocturnal desaturation in sickle cell anemia</td>
<td>85 children with SCD (9±4 yrs; BMI z score −0.4 in apneic group and −1.0 in non-apneic group)</td>
<td>An oral cavity assessment in the Frankfort position with tongue in relaxed position and with a mouth opening angle of 20°C to the mandibular condyle. A 20°C fixed-aperture compass was used, which was placed on the topography of the temporomandibular joint, the tip of its upper leg aligned with the upper central incisors, to yield desired mouth opening. Another compass was used to obtain measurements of maxillary intermolar distance (size of maxilla); and mandibular intermolar distance (mandibular size) which were then transposed to a ruler. The overjet, cervical circumference and abdominal circumference were measured with a ruler/tape</td>
<td>A positive correlation was found between height/age z-score and cervical circumference. Nocturnal desaturation was associated with cervical circumference and abdominal circumference. A negative correlation between desaturation and maxillary intermolar distance and mandibular intermolar distance</td>
<td>Student's t test for independent samples or the Mann–Whitney test to compare two means. Spearman’s test to test the correlation between variable.</td>
<td>No imaging available as measurements taken via clinical method</td>
</tr>
<tr>
<td>Schwab et al 2015. Understanding the Anatomic Basis for Obstructive Sleep Apnea Syndrome in Adolescents</td>
<td>Adolescents 12–16 years from CHOP study. 49 obese with OSA; 38 obese and 50 lean controls</td>
<td>Upper airway MRI was performed using a 3T scanner (Magnetom Sonata; Siemens). Amira 4.1.2 image analysis software. Sagittal: airway volume, cross-sectional area, minimum airway area in the retropalatal (RP), retroglossal (RG), and nasopharyngeal (NP) regions; minimum anteroposterior airway width, minimum lateral airway width in the RP and RG regions. Airway length: distance between palatal plane and parallel plane through base of epiglottis. Volumetric analysis of upper airway soft tissue structures on axial T1-weighted MRI scans: soft palate, tongue genioglossus muscle, other tongue (geniohyoid, hyoglossus, myohyoid, digastric, and mylohyoid) muscles, parapharyngeal fat pads, lateral pharyngeal walls (including tonsils), pterygoid muscle, epiglottis, sum of soft tissue volumes. Axial T2-weighted for tonsils (right and left combined) and adenoid as better resolution of lymphoid tissue</td>
<td>Obese with OSAS had increased adenotonsillar tissue and smaller nasopharyngeal airway compared with all controls. The size of other upper airway soft tissue structures (volume of the tongue, parapharyngeal fat pads, lateral walls, and soft palate) was similar between both groups</td>
<td>Adjusted analysis of covariance was used to compare the three groups with a sub-domain-specific, Bonferroni-corrected level of significance</td>
<td>Figure S14</td>
</tr>
<tr>
<td>Tong et al 2016. MR Image Analytics to Characterize the Upper Airway Structure in Obese Children with Obstructive Sleep Apnea Syndrome</td>
<td>30 children 8–17 yrs (15 obese with OSAS; 15 obese controls)</td>
<td>Philips Achieva 3T machine. Axial T2-weighted and sagittal T1-and T2-weighted sequences. &quot;Object&quot; measurements: skin, pharynx, nasopharynx, oropharynx, mandible, fat pad, adenoid, tonsils, tongue, soft palate. 3D region enclosed by the boundary surface was considered to represent the &quot;object.&quot;</td>
<td>OSAS group: increase in linear size, surface area, volume of adenoid, tonsils, fat pad. Fat pad and oropharynx become less round/more complex in shape in OSAS</td>
<td>( \phi = U/Q ). Logistic regression based on a sigmoid model</td>
<td>Figure S15</td>
</tr>
</tbody>
</table>
inflammatory agents, such as leukotriene receptor antagonists. Montelukast, a leukotriene receptor antagonist, has been used in trials and has shown a significant reduction in adenoid size and respiratory-related sleep disturbances.\(^{128}\) Leukotriene receptors LT1-R and LT2-R showed increased protein expression within adenoid tissues and there were also increased levels of LTB4, LTC4, LTD4, and LTE4 within adenoid tissue.\(^{128,129}\) However, similarly to adenotonsillecctomy, there is the possibility of adenoid regrowth. Studies combining adenotonsillecctomy and Montelukast as a postsurgical agent look promising.\(^{129}\)

**Volume of airway**

Arens et al showed that regional analysis of the upper airway in children with OSAS was most restricted where adenoid and tonsils overlap.\(^{40}\) However, segmental analysis revealed that the upper airway was restricted throughout the initial two-thirds of its length and that the narrowing was not in a discrete region adjacent to the adenoids or tonsils, but rather in a continuous fashion along both. Documenting the distribution of the volume throughout the airway, rather than just the value of the volume, could be important in understanding this disease process in both healthy populations and those with underlying diseases.

**Lower facial skeleton**

**Craniofacial differences**

Arens et al and Cappabianca found that the mandible was smaller in children with OSAS.\(^{47,124}\) Populations that exhibit craniofacial abnormalities, with a higher incidence of OSAS, include Down syndrome, Treacher Collins syndrome, Crouzon syndrome, Apert syndrome, and Pierre Robin syndrome.\(^{64}\) Uong et al found that Down syndrome subjects were more likely to have smaller mental spine to clivus lengths.\(^{73}\) Therefore, exploring these anatomical measures as potential targets for further research could be warranted. Although it can be studied using MRI, the lower facial skeleton has received relatively little attention in populations at risk of OSAS. It would be interesting to ascertain whether there are any differences in craniofacial morphology owing to extramedullary hematopoiesis in chronic anemias and to determine whether growth of the mandible and maxilla matches the trajectory in comparison to healthy, typically developing children – ideally with sequential MRI into adulthood.

**Lower facial skeletons in SCD**

Salles et al showed smaller maxilla and mandible measurements, measured using cephalometry, in SCD subjects who exhibited nocturnal oxygen desaturation.\(^{116}\) Studies have shown craniofacial differences in the SCD population exhibiting mandibular retrusion, maxillary protrusion with elongated vertical development, and generalized pronounced protrusion of the midface. Skeletal immaturity and impedance of growth alongside angular bony deformities are commonly seen in patients with SCD.\(^{130}\) Anoxic events potentially precipitate premature closure of the epiphysis, whereas craniofacial structure abnormalities could be due to extramedullary hematopoiesis.\(^{131}\) Studies have shown that patients with SCD have reduced z-scores for weight for age, height, elbow breadth, skinfold thickness, and mid-upper arm circumference, that are indicative of global growth deficit. Interestingly, studies have reported lower levels of vitamin D in this population, which were associated with morbidity. Vitamin D supplements might promote growth to combat these anatomical defects.

If genetic disposition is also explored, there could be interventions based on this. A study by Arun et al evaluated MYO1H gene polymorphisms and haplotypes as risk factors for mandibular retragnathism and identified a single-nucleotide polymorphism rs3825393 to be associated with mandibular retragnathism.\(^{132}\) Previous studies have shown that there are distinct combinations of myosin heavy chain isoforms in the masseter muscles and that these muscle combinations may have mutual effects on bones. SCD patients may have a genetic predisposition for mandibular retragnathism influencing their susceptibility to develop OSAS.

Techniques including mandibular advancement or mandibular distraction osteogenesis for treatment of retron-positioned mandibles or mandibular hypoplasia have produced promising results in children with OSAS with improvement in flow limitation.\(^{133}\) These oral jaw repositioning devices are relatively new and not commonly used in the pediatric population. In addition to adenotonsillecctomy, alternative methods for treatment include nasal corticosteroid spray or Montelukast, and continuous or bi-level positive airway pressure (CPAP; BiPAP). Exceptionally rare cases may require tracheostomy.

**SDB and complications of SCD – implications for treatment?**

SDB and NOD have been associated with increased incidence of vaso-occlusive crisis, stroke, neurocognitive deficits, nocturnal enuresis, cardiovascular, and autonomic dysfunction in SCD.\(^{93,97,103,134–136}\) However, there are few studies looking at mechanisms which might lead to
Stroke and OSAS/NOD in SCD as an example
The prevalence of cerebral infarction, both overt and covert in people with SCD, is much greater than compared to the healthy population. Case reports in the 1980s suggested that OSAS was a risk factor for stroke in SCD. A cohort study showed that nocturnal hypoxemia predicted central nervous system events. Interestingly, stroke appears to be more common in those with a longer history of upper airway obstruction. A laboratory study by Setty et al reported that the release of cellular mediators in hypoxemia, and the relationship between anemia and oxyhemoglobin desaturation, suggested that cerebral vasculopathy and stroke might occur through hypoxia-mediated pathways. In patients, high cerebral blood flow on transcranial Doppler appears to be associated with low daytime oxygen saturation.

Abnormal magnetic resonance angiography was associated with NOD in another study. Huang et al highlighted that children with SCD without OSAS have more prominent upper airway reflexes than children with SCD without OSAS. Reduced upper airway neuromuscular reflexes may contribute to the increased upper airway collapsibility in SCD children with OSAS. The study speculated that cerebrovascular disease in children with SCD may predispose to abnormal regulation of upper airway muscles during sleep, therefore increasing the risk of OSAS in this population. The question is: does cerebrovascular disease predispose to OSAS or does OSAS-inducing hypoxia lead to cerebrovascular disease?

Conclusion
SDB is more common in Black ethnicities, and even more prevalent in those with SCD. SDB has significant implications for health leading to chronic and debilitating diseases. There appear to be structural differences in the airway between those with and without OSAS, as well as between ethnicities. OSAS populations tend to have smaller airways, larger lymphoid tissues, and smaller lower facial skeletons demarcated by measurements of the mandible and mental spine to clivus. There appears to be an increased prevalence of OSAS in SCD populations with hypoxia as a key manifestation which appears to increase the risk of complications, including stroke. Understanding a potential relationship between the AN ratio, lymphoid hypertrophy, airway volume distribution, and lower craniofacial structure could potentially help to stratify those children most at risk of OSAS. Combatting any structural defects with appropriate interventions could reduce the duration spent in hypoxemia and consequently reduce the risk of comorbidities in this population, therefore warranting early treatment interventions.

Abbreviation list
AN, adenoid to nasopharynx ratio; AASM, American Academy of Sleep Medicine; AHI, Apnea/Hypopnea Index; BiPAP, bi-level positive airway pressure; BMI, body mass index; CAI, central apnea index; CHOP, Children’s hospital of Philadelphia; CPAP, continuous positive airway pressure; CSA, central sleep apnea; CSAS, central sleep apnea syndrome; CT, computed tomography; ECG, electrocardiography; EEG, electroencephalography; EMG, electromyography; EOG, electro-oculography; ETpCO₂, end-tidal partial pressure carbon dioxide; HbA, normal adult haemoglobin; HbAS, sickle cell trait haemoglobin; HbSS, homozygous sickle cell haemoglobin; MRI, magnetic resonance imaging; NOD, nocturnal oxygen desaturation; Non-REM, non rapid eye movement sleep; OAHI, Obstructive Apnea/Hypopnea Index; OSAS, obstructive sleep apnea syndrome; pCO₂, partial pressure carbon dioxide; PSG, polysomnography; RBC, red blood cell; REM, rapid eye movement sleep; RIP, respiratory inductance plethysmography; SAC, sickle Sleep Asthma Cohort study; SaO₂, arterial oxygen saturation; SCA, sickle cell anaemia (HbSS); SCD, sickle cell disease (HbSS, HbSC, HbSßthalassemia); SDB, sleep disordered breathing; SpO₂, pulse oximeter oxygen saturation.

Acknowledgments
Dr Kirkham’s research is supported by Action Medical Research grant GN2509.
Disclosure
The authors report no conflicts of interest in this work.

References
34. Franklin KA, Lindberg E. Obstructive sleep apnea is a common disorder in the population-a review on the epidemiology of sleep apnea. J Thorac Dis. 2015;7(8):1311–1322. doi:10.3978/j.issn.2072-1439.2015.06.11


Supplementary materials

Figure S1 (A) Adenoidal measurements. “A” represents distance from point of maximal convexity, along inferior margin of adenoid shadow to line drawn along straight part of anterior margin of basiocciput. “A” is measured along line perpendicular from point A to its intersection with B. (B) Nasopharyngeal measurement. “N” is distance between posterior superior edge of hard palate, and D, anteroinferior edge of sphenobasiosseal synchondrosis. When synchondrosis is not clearly visualised, point D can be determined as site of crossing posteroinferior margin of lateral pterygoid plates P and floor of bony nasopharynx.


Figure S2 (A) Anatomical outlines of an axial T1 image at the level of the maximal tonsillar area of a control Subject. Transverse black arrow represents the intermandibular distance. (B) Anatomical outlines of a midsagittal T1 image of a control subject. Oblique black arrow represents the mental spine-clivus oblique distance.


Figure S3 (A) T1-weighted axial image at the retropalated level (right) and segmented regions of interest (left) of a control subject. (B) Midsagittal T1-weighted image with mental spine-clivus distance shown (right) and segmented regions of interest (left) of a control subject.

Figure S4 (A) Mid-sagittal T1-weighted image with mental spine–clivus oblique line shown (white line). Linear measurements were obtained along this line. (B) Axial T1-weighted image at the level of maximal tonsillar cross-sectional area. Note the transverse intermandibular line (white line) passing the center of each tonsil. Linear measurements were obtained along this axis. Maximal oropharyngeal width was obtained on this plane (dotted white line).


Figure S5 Left: three-dimensional display of an upper airway and its centerline in a subject with OSAS. Top right: gray-level two-dimensional scene of the cross-section of the airway orthogonal to the centerline at marker locations: A = adenoid; A + T = adenoid and tonsil overlap; T = tonsils; E = epiglottis. Bottom right: plot of the cross-sectional area function.


Figure S6 (A) Midsagittal magnetic resonance image (MRI) of a normal subject, demonstrating the upper airway regions: retropalatal (RP) — from the level of the hard palate to the caudal margin of the soft palate; and retroglossal (RG) — from the caudal margin of the soft palate to the base of the epiglottis. Soft palate, tongue, airway, mandible, and subcutaneous fat are denoted with arrows. Fat is bright (white) on an MRI. (B) Representative three dimensional upper airway volume in a patient with sleep apnea (left) and in a normal subject (right). Note that the upper airway volume is smaller in the RP region than in the RG region in both subjects and that the length of the total airway and the individual regions (RP/RG) is not equivalent between the apneic and normal subjects. Airway volume is smaller in the RP region in this apneic subject compared with the control subject; airway volume is similar in the RG region in this apneic subject compared with the control subject.


Figure S8 Volumetric reconstructions from a series of 3-mm contiguous axial magnetic resonance (MR) images of the mandible (gray), tongue (orange/rust), soft palate (pink/purple), lateral parapharyngeal fat pads (yellow), and lateral/posterior pharyngeal walls (green) in a normal subject (top panel) and in a patient with sleep apnea (bottom panel). The upper airway is larger in the normal subject than in the patient with apnea. In addition, the tongue, lateral parapharyngeal fat pads, and lateral pharyngeal walls are larger in the patient with apnea. Note: Reprinted with permission of the American Thoracic Society. Copyright © 2020 American Thoracic Society. Schwab RJ, Pasirstein M, Kaplan L, Pierson R, Mackley A, Hachadoorian R, et al. 2006. Family aggregation of upper airway soft tissue structures in normal subjects and patients with sleep apnea. Am J Respir Crit Care Med. 173. 453–63. Official journal of the American Thoracic Society.

Figure S10 (A) CSA oropharyngeal airway. (B) Combined upper airway volume, T1 weighted axial scan. (C) Transverse line perpendicular to the floor passing through the hyoide to nasion, to sella to supramentale. (D) Midsagittal palatal CSA.


Figure S11 (A) Left 4 images. Mid sagittal (top) and axial T2-weighted images at the level of the nasopharynx (bottom), of a patient before (left) and after (right) adenotonsillectomy. Solid white arrows indicate adenoid tissue (gray) and dotted arrows indicate nasopharyngeal airway. Note residual adenoidal tissue and larger airway after adenoidectomy. In this example, a 29% reduction in adenoid volume was noted on MRI. (B) Right 2 images. Axial T2-weighted images at the level of the oropharynx of a patient before (left) and after (right) adenotonsillectomy. Solid white arrows indicate tonsillar tissue (gray) and dotted arrows indicate oropharyngeal airway. Note the larger airway and small left residual tonsil tissue after tonsillectomy.

Note: Reprinted from CHEST, 142(1), Strauss T, Sin S, Marcus CL, Mason TBA, McDonough JM, Allen JL, et al., Upper airway lymphoid tissue size in children with sickle cell disease, 94-100, Copyright 2012, with permission from Elsevier.

Figure S12 MRI volumetric analysis of lymphoid tissue volumes in an OSA patient (green-upper jugular lymph nodes, orange-tonsil tissue, red-retropharyngeal lymph nodes, magenta-adenoid tissue). (A) Three dimensional reconstruction of lymphoid tissue using Amira® software. (B) Axial T2-weighted DICOM image with lymphoid tissue tracings. (C) Coronal T2-weighted DICOM image with lymphoid tissue tracings. (D) Sagittal T2-weighted DICOM image with lymphoid tissue tracings.


Figure S13 Upper airway, soft tissues, and mandibular reconstructions. Head and neck surface rendering with three-dimensional reconstructions of the upper airway, soft tissues, and mandible of a subject with obstructive sleep apnea syndrome in various views: lateral (top left), anterior oblique (top right), superior oblique (bottom left) and posterior (bottom right). Airway (light blue), tongue (brown), mandible (white), soft palate (blue), tonsils (yellow), adenoid (magenta), retropharyngeal nodes (red), and deep cervical nodes (green).

**Figure S14** Anatomical definitions of upper airway landmarks on midsagittal and axial slices with 3D reconstruction of regions of interest.


**Figure S15** 3D surface renditions of objects: np (nasopharynx), op (oropharynx), mn (mandible), tn (tonsils), tl & tR (left and right tonsil), fp (fat pad), ad (adenoid), tg (tongue).