ventilation inhomogeneity (VI) as assumptions underlying the calculation are invalid; an alternate index that has been suggested is Scond.1

Aim To compare these two methods of CDI assessment in CF children

Methods Children with cystic fibrosis (CF; 67) and healthy controls (61) performed multiple breath washout with sulphur hexafluoride measured using mass spectrometry. Scond was calculated from 1.5 to 6 turnovers and Scond* from breath 2 to 3 turnovers.

Results All measures of VI were significantly higher for CF vs control, mean difference: LCI 4.0, Scond 0.054, Scond* 0.081.

In CF, LCI correlated better with Scond* than Scond (See figure: correlation coefficient LCI vs. Scond* 0.75; LCI vs. Scond 0.42). If children with moderate-severe VI (LCI > 11) were excluded there was an improved correlation for both relationships (correlation coefficient LCI vs. Scond 0.83; LCI vs. Scond* 0.86).

An asymptote for the Scond vs LCI relationship was at Scond 0.07 and Scond* 0.13.

Conclusion Scond* quantifies the mechanism of VI in moderate to severe lung disease, but it may reach asymptote in very severe VI.

REFERENCE

SLEEP DISORDERED BREATHING IN CHILDREN WITH SPINA BIFIDA. TIME TO SCREEN?

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Background Spina bifida is associated with sleep disordered breathing (SDB) particularly when associated with Arnold-Chiari malformations. Studies suggest that moderate/severe sleep apnoea is present in up to a third of spina bifida patients (Patel 2015) and yet there are no national guidelines that recommend screening for SDB in children with spina bifida. There is evidence to suggest that many children present late and this can be associated with unnecessary morbidity and even mortality (Kirk 1999).

Aim To assess the prevalence of SDB in children with spina bifida, presenting through clinical presentation alone in the West of Scotland and to explore whether there is a case for screening all children with spina bifida for SDB.

Method The database of the Spina Bifida Association Scotland and clinical records from the regional centre in the Royal Hospital for Children, Glasgow were used to identify all children with spina bifida in the West of Scotland. The level of the spinal lesion, presence of an Arnold-Chiari malformation or ventriculo-peritoneal shunt was established, as was the number who had had sleep studies performed and who had required ventilator support.

Results 108 children were identified; 44/108 (40%) had an Arnold-Chiari malformation (1 type I, 43 type II); 64/108 had lumbar abnormalities, 14/108 lumbosacral, 14/108 thoracolumbar, 9/108 sacral and 4/108 thoracic. 52/108 had a VP shunt at some point. Only 14 children had presented with clinical symptoms that lead to a sleep study being undertaken (snoring 7, apnoeas 7, cough/wheeze 2, restless night 2, morning headache 2). 5 children had mixed central and obstructive apnoeas, 1 obstructive sleep apnoea, 2 hypoventilation. 8 children went on to require non-invasive mask ventilation of these 7/8 had an Arnold-Chiari malformation (p = 0.005), 7/8 had a previous VP shunt (p = 0.02), 5/8 had lumbar abnormalities and 3/8 thoraco-lumbar.

Conclusion Clinical presentation alone only identifies a small proportion of cases of SDB in children with spina bifida, with a high proportion of these requiring intervention. We remain concerned that there are many children with spina bifida with undiagnosed SDB who may benefit from treatment, particularly those with Arnold-Chiari malformations and therefore that screening is indicated.
Introduction  Congenital lung agenesis is a rare abnormality which has been reported to have an estimated incidence of 1 in 15,000 pregnancies and to carry a poor prognosis, especially when associated with congenital heart anomalies. (1) However, precise incidence data has previously been unavailable and clinical management has improved in recent years with an unknown impact on outcomes.

Methods  We used the North of England register of congenital anomalies (NorCAS) cross-referenced with clinical data from our regional paediatric respiratory centre to calculate the first known accurate incidence estimate for this condition. Detailed clinical and outcome data were then collected with informed consent from the families of affected infants.

Results  The incidence of lung agenesis was 0.12 per 100,000 live births (95% confidence interval 0.03–0.31). Four cases were identified with a median age at follow-up of 5 years. Lung agenesis was associated with complex congenital heart disease (complete atrioventricular septal defect with left atrial isomerism) in one case, and with aortic coarctation and atrial septal defect in another. Both these patients had their heart defects successfully repaired. The third patient had a normal heart but musculoskeletal problems, while lung agenesis was the only anomaly in the fourth patient. All four patients are well and not on home oxygen.

Conclusions  We reviewed the course of the four patients with lung agenesis born in the region between 2004 and 2013, and report that medium term outcomes have been good, even when associated with congenital heart disease and other anomalies. This information will provide a useful starting point when counselling parents whose unborn baby has an antenatal diagnosis of lung agenesis. We aim to follow-up these patients to report long-term outcomes as these remain unknown and there is concern about the potential for the development of late onset pulmonary hypertension.

REFERENCE

From oxygen to the ITU

P187 POTENTIAL IMPACT OF NON-ARTERIAL BLOOD GAS SAMPLING ON CLINICAL PRACTICE

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Introduction  Arterial blood gas (ABG) analysis is commonly used to monitor patients with acute respiratory problems but can be painful and associated with potentially serious side-effects. Recently there has been renewed interest in non-arterial forms of blood gas analysis with a suggestion that such techniques could replace >60% of ABGs. (2) It is however difficult to quantify the impact that non-ABG sampling could have as there is little published data on the clinical burden that ABGs represent. We were interested in establishing how many ABGs are routinely done in a real world setting.

Methods  We retrospectively analysed acute admissions (all cause) to a ward based level 2 Respiratory Emergency Care Unit (RECU) at a University Teaching Hospital over a 6 month period. Prospectively we analysed visual analogue pain scores (VAS) and complications including attempt rates.

Results  Over a 6month period (Apr-Oct) 57 of 111 patients admitted to the RECU had complete datasets. A total of 432 ABGs were obtained from this cohort giving an average of 7.6 ABGs (range 2 – 22) per patient per admission and on average patients had 2.3 ABGs each day whilst on RECU.

Overall the mean number of attempts per ABG of the prospectively collected cohort (n = 100) was 1.6 (range 1 – 8; 44% physician obtained). Taking this into account a patient could therefore expect to be stabbed 3.7 times every day or 11 times in total during their admission through the RECU. The most commonly documented complication amongst our patients was pain but average pain scores were relatively low in keeping with other published data (median VAS = 3, IQR 5); no serious complications were reported.

Conclusions  Our data suggests that patients requiring admission to a level 2 respiratory unit experience a high burden of ABG testing during their stay. Using published literature on the potential impact of Non-ABG sampling our data suggests that >500 ABGs could be avoided each year on our acute respiratory unit alone.

REFERENCE

P188 NASAL HIGH FLOW (NHF) – IS IT APPROPRIATELY PRESCRIBED? A RETROSPECTIVE CASE REVIEW OF 93 ADULT PATIENTS REQUIRING NASAL HIGH FLOW OXYGEN WITHIN A DISTRICT GENERAL HOSPITAL

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Background  Nasal high flow (NHF) delivers warmed humidified oxygen up to 65L/min and is commonly used in intensive care units (ICU). In a non-critical care setting it can provide a bridge to ICU for hypoxic patients, and symptom palliation for those unsuitable for escalation.

Evidence to support NHF use in a non-critical care setting is limited. We aimed to review patient selection and clinical outcomes for individuals commenced on NHF over two winter periods, in order to highlight prognostic indicators and develop a clinical guideline.

Method  We performed a retrospective review of 93 cases managed with NHF September-December in 2014 and 2015 in a medium sized DGH. Patients were included regardless of diagnosis and treatment escalation plan. Data was collected on patient demographics, diagnosis, management and clinical outcome. Results described as % (n).

Results  93 patients (mean age 71.2 yrs, SD 14.8). Presenting complaint: Pneumonia 57.0% (53), Aspiration pneumonia 4.3% (4), Pulmonary embolism 4.3% (4), Pulmonary oedema 3.2% (3), Interstitial Lung Disease 3.2% (3), other diagnoses 28.0% (26). Clinical indication: Hypoxia 64.5% (60), hypoxia and work of breathing 14.0% (13), work of breathing alone 7.5% (7) and palliation 14.0% (13).

42 patients (45%) were assessed by ICU, of which 21 patients were deemed appropriate [mean age 65 yrs (40–84)]. 62% (13) were intubated. 81% (17) survived to discharge; 5.9% (1) required LTOT. No significant correlation was present between flow rates and mortality (p = 0.7).