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Key paediatric messages from Amsterdam

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ABSTRACT The Paediatric Assembly of the European Respiratory Society (ERS) maintained its high profile at the 2015 ERS International Congress in Amsterdam. There were symposia on preschool wheeze, respiratory sounds and cystic fibrosis; an educational skills workshop on paediatric respiratory resuscitation; a hot topic session on risk factors and early origins of respiratory diseases; a meet the expert session on paediatric lung function test reference values; and the annual paediatric grand round. In this report the Chairs of the Paediatric Assembly’s Groups highlight the key messages from the abstracts presented at the Congress.

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Early life risk factors for wheezing and asthma

There is increasing interest in exposure during pregnancy and even prior to conception on the risk of childhood respiratory diseases [1]. LODGE et al. [2] investigated the association between grand-maternal smoking at 10–12 weeks of pregnancy and asthma risk in grandchildren using data from Swedish registries. Children had a 6–22% increased risk of asthma during the first 6 years of life if their grandmothers smoked during early pregnancy, independent of maternal smoking. These findings support possible epigenetic transmission of risk from environmental exposures in previous generations. In a cross-sectional study, PESCE et al. [3] reported that children exposed to maternal vaginitis in pregnancy were more likely to have had bronchitis, pneumonia and wheezing in the first 2 years of life, and current asthma at 6 years of age. However, as the researchers themselves acknowledge, these data should be confirmed in a prospective study in order to avoid recall bias and misclassification. Use of paracetamol in pregnancy has been repeatedly reported to be associated with wheezing and asthma in the first years of life [4], but the problem of confounding by indication has also been raised [5]. MAGNUS et al. [6] explored the issue in the Norwegian Mother and Child Cohort and concluded that although there is evidence of confounding by indication this does not fully explain the association between paracetamol use only during pregnancy and asthma at 3 and 7 years of age in the offspring. As for early post-natal factors associated with wheezing and asthma, living on farms has been shown to protect children from asthma and allergies. A major factor involved in this effect is the consumption of unprocessed cow’s milk [7]. Within the Pasture Study Group, which includes children living in rural areas in five European countries, samples of the milk usually consumed were taken at the age of 4.5 years [8], while doctor-diagnosed asthma was parent-reported at age 6 years. The risk of asthma was reduced when consuming raw farm milk as compared with shop milk. Part of the effect was attributable to the content of omega-3 fatty acids, which could counterbalance the synthesis of pro-inflammatory leukotrienes and prostaglandins. Accumulating evidence suggests an association between early growth patterns and wheezing in infancy. Within the NINFEA birth cohort, PAPOVIC et al. [9], using a novel modelling approach (SITAR) that allows for mutual adjustment of different aspects of growth, showed that larger size and faster growth over the first 18 months of life, but not the timing of peak weight velocity, were independently associated with an increased risk of wheezing at 18 months of age.

Asthma and allergy: genetics and environment

The asthma risk locus 17q21 modifies the effects of smoking and early viral infections on subsequent asthma [10]. LOSS et al. [11] of the Pasture Study Group followed children in rural areas of several countries in Europe from birth until 6 years of age. Single nucleotide polymorphisms (SNPs) in the genes ORMEL and GSDMB at locus 17q21 were assessed in cord blood. Exposures to siblings or to animal sheds affected wheezing largely in carriers of known asthma risk alleles. LOSS et al. [11] conclude that the 17q21 locus might be related both to environmental susceptibility and to progression from wheeze to asthma. The associations between traffic-related air pollution (TRAP), asthma and allergies in children are heterogeneous; possibly due to genetic polymorphisms that modify the response to oxidative stress [12]. BOWATTE et al. [13] investigated if variants in the glutathione pathway alter associations between early life TRAP exposure and childhood asthma and allergies in a high risk birth cohort of children in Australia. Baseline TRAP exposure was not associated with any outcome. Carriers of GSTM1null and GSTT1null had an increased risk of current asthma at 7 years and current eczema at 18 years of age. GSTT1null carriers exposed to TRAP in early life had an increased risk of wheeze and asthma at 12 years of age when compared with TRAP exposed carriers of wild-type GSTT1. Polymorphisms of CD14 are associated with a lower risk of allergic disease in children, but little is known about the effects in adult life. LAU et al. [14] aimed to examine if the association between CD14 polymorphisms and allergic sensitisation in middle age was modified by sibling exposure in early life in participants from the Tasmanian Longitudinal Health Study. Exposure to siblings at 6 months, 2 years and 4 years was associated with a significantly reduced risk of allergic sensitisation. They showed evidence of CD14 SNPs and sibling interaction for allergic sensitisation. The gene–environment interaction was strongest in the first 6 months of life, suggesting that this may be a critical period for altering the risk of allergy.

European Respiratory Society task force on primary ciliary dyskinesia and beyond

Published data on primary ciliary dyskinesia (PCD), a rare disease leading to chronic airways inflammation and infection, are scarce and often based on small case series. A consensus statement of a European Respiratory Society (ERS) task force published in 2009 [15] concluded that the evidence base for diagnosis and treatment of PCD is poor and more research is needed. Nasal nitric oxide (nNO) is currently recommended as a screening tool. PANAGIOTIS et al. [16] evaluated the diagnostic performance of nNO for detection of PCD by examining 12 published studies and concluded that it has high overall diagnostic accuracy. However, COLLINS et al. [17] using data from prospective referrals to the PCD service at University
Hospital of Southampton, where the incidence of PCD is 11%, underlined that the very high negative predictive value of the test (99.1%) makes nNO a good rule-out test, but the low positive predictive value (26.3%) makes it a poor test for general screening. They calculated that screening the general population would reduce the positive predictive value to 0.04%. Within the European project BESTCILIA, HALBREICH et al. [18] compared the lung function (forced expiratory volume in 1 s (FEV1)) of patients with PCD to the Global lung function initiative (GLI) 2012 reference values and to published data from UK patients with cystic fibrosis (CF) [19]. FEV1 was significantly reduced compared with normal reference values. As for the comparison with CF data, FEV1 was similarly low in children (e.g. at age 6–9 years patients with PCD had a FEV1 of 87% predicted, while in CF patients FEV1 was 90%); while in adults aged 18–21 years FEV1 was better in PCD (76% predicted, 95% CI 73–80%) than in CF (66% predicted, 95% CI 65–68%).

Infant lung function

Objectively assessing lung function in infants remains one of the most challenging aspects of respiratory physiology. For the most part measurements are performed following sedation, use complex equipment and require highly trained staff [20, 21]. A recent survey on the clinical utility of infant lung function testing shows that while many centres were using lung function tests in infants to aid in clinical decision making there were many barriers to practical use [22]. One of these barriers was the availability of appropriate reference values for the infant version of spirometry (the raised volume rapid thoracic compression technique (RVRTC)). LUM et al. [23] reported reference values for modern commercial RVRTC equipment (Jaeger BabyBody; Care Fusion, San Diego, CA, USA) derived from data in 367 infants over 607 visits in four centres. Significant differences to the previously published RVRTC reference equations were noted [24], suggesting that users of the Jaeger BabyBody RVRTC equipment should adopt these new reference equations for their research and clinical RVRTC work.

Much of our understanding of lung growth and development has been derived from longitudinal measurements of lung function in birth cohorts [25, 26]. However, the repeated measurement of infant lung function tests is difficult. KONALIK et al. [27] reported the feasibility of obtaining repeated multiple breath washout (MBW), body plethysmography and RVRTC data (in that order) over the first 2 years of life in the CHILD birth cohort. Of the three tests, MBW measurements had the highest feasibility at any one visit (range: 79–98%), with body plethysmography (70–83%) and then RVRTC (49–56%) being successively lower. In the 191 infants that attended more than one visit, longitudinal data was obtained in 63% of MBW attempts compared with 50% of body plethysmography and 33% of RVRTC assessments. These data suggest that feasibility will be influenced by test order, further highlighting the importance of consideration of the primary outcome in assessments of lung function in infancy. In a unique study, GRAY et al. [28] reported longitudinal measurements of infant lung function in the Drakenstein birth cohort. These investigators assessed lung function in unsedated infants at two time-points (6 weeks and 1 year). Of the 507 eligible infants, data was obtained in 472 (93%) at 6 weeks and 377 (74%) at 1 year for at least one lung function test. Paired tidal breathing or MBW outcomes were obtained in 234 and 209 infants, respectively, representing over 50% of the tested infants. These results demonstrate for the first time that unsedated measurements of some forms of infant lung function tests can be obtained longitudinally and may assist in allowing more wide spread integration of lung function testing in this age group.

Lung function in older children

PFEILGER et al. [29] compared changes in spirometry and multiple breath nitrogen washout after cold dry air challenge. In those children who had a positive challenge test the decrease in FEV1 and the increase in lung clearance index (LCI) correlated well. SEMD (ventilation heterogeneity in conducting airways) was a predictor of airway hyperresponsiveness (AHR); exhaled nitric oxide fraction (FeNO) seemed to predict airway AHR as well. This might be helpful in assessing AHR in young children. DE LEEUW et al. [30] showed that most children with asthma have normal spirometry, even if they have severe asthma. LCI had better discriminating power between severe and mild-to-moderate asthma; however, the vast majority of children had a normal LCI, irrespective of asthma severity, which may limit the clinical utility of MBW techniques in asthma. Two research groups reported data on a new lung function technique, structured light plethysmography (SLP) [31–33]. SLP is a noninvasive, noncontact test that does not require cooperation by the patient. It records real-time images of chest wall movement, using a grid of light on the chest and abdomen of the patient, which is imaged by two cameras. With this technique data on breathing pattern, the tidal breathing flow/volume graph, inspiratory or expiratory flow limitation, and regional subdivision may be assessed. A group from Verona, Italy showed that with SLP expiratory flow limitation could be detected in children with an acute asthma attack versus children with stable asthma [31]. Researchers from the UK assessed changes in tidal breathing parameters in children with acute asthma and found significant differences before and after bronchodilation [32]. The same group collected normative data on several parameters of SLP in healthy children [33]. This technique seems feasible in.
likely (OR 2.02, 95% CI 1.26–3.24) following either adenotonsillectomy, adenoidectomy or no treatment. Untreated children were twice as likely to have normal MBW outcomes than those with CF demonstrated altered lung clearance (LCI) and convection dependant ventilation distribution (Scond), while children with CF also had altered acinar ventilation when compared with healthy children. Two studies reported longitudinal tracking of LCI and spirometry in children with CF, with both demonstrating that measurements of lung clearance were more likely to deteriorate over time than spirometry [36, 37]. While Jayasuriya et al. [38] demonstrated that LCI was more likely to be abnormal (1458%) out of 24 children) compared with FEV1 (521%) out of 24) in children following bone marrow transplant, further reinforcing the importance of using the most appropriate test for each specific lung condition. Several studies in children born preterm highlighted these concepts with varying degrees of pathophysiology reported. Bar-Yissay et al. [39] reported spirometry and forced oscillatory mechanics before and after bronchodilator responsiveness in preterm children with and without bronchopulmonary dysplasia (BPD). As reported previously [40], preterm children exhibited reversible obstruction; however, the differences did not completely return to normal with lung function remaining lower in preterm compared with term born children. In contrast to other data presented at the Congress (discussed later), there were no differences in lung function between those children with and without BPD, which may be attributed to sample size and/or differences in cohort characteristics. O’Dea et al. [41] reported that while the majority of children (aged 9–11 years) born very preterm (<32 weeks completed gestation) had altered lung structure on computed tomography (CT) this did not influence peak exercise capacity in these children, which was not different to that of term born controls. It was, however, noted that dynamic flow limitation during exercise was common in preterm children with BPD suggesting an altered response to maximal exercise unrelated to exercise capacity. The same children were followed longitudinally with paired lung function at 4–7 and 9–11 years [42]. As is commonly reported, preterm children with and without BPD had lower spirometry that term controls [40]. Spirometry was noted to decline over time only in preterm children with BPD with their mean±SD FEV1 reducing by −0.47±0.92 (p<0.02) z-scores. The number of children with obstructive lung function (FEV1/forced vital capacity less than the lower limit of normal) increased from 32 to 52%. The significance of these changes is not clear, but suggests an active disease process in preterm children with BPD that requires detailed investigation.

LCI measured by MBW has already been shown to be a sensitive marker of CF lung disease, which is now being used as a surrogate outcome measure in clinical trials [43], as a surveillance tool to monitor structural lung disease [44], and as a prediction tool for pulmonary exacerbations in children with CF [45]. However, LCI response to therapy for pulmonary exacerbations is heterogeneous in CF patients, the overall effect size is small and the results are often discordant with FEV1 [46]. A group from Toronto presented their data on whether LCI can track disease progression: they observed an association between LCI and functional residual capacity from MBW [51, 52].

Sleep monitoring

The pathogenesis and treatment of sleep disorders in children are generally well understood; however, the optimal treatment of obstructive sleep apnoea (OSA) and the potential impact of comorbidities is less certain [53, 54]. Domany et al. [55] examined the rate of treatment failure in 659 children with moderate-to-severe OSA. Children were assessed for residual OSA using a screening questionnaire following either adenotonsillectomy, adenoidectomy or no treatment. Untreated children were twice as likely (OR 2.02, 95% CI 1.26–3.51) to have ongoing OSA compared with children treated with
adenotonsillectomy, while there were no differences between adenotonsillectomy and adenoïdectomy suggesting that adenoïdectomy may offer additional treatment pathways for paediatric OSA. While laboratory based sleep studies remain the gold standard for the diagnosis of sleep disorders, the cost and availability remain barriers to their use. Options that allow the screening of children to minimise the requirement for polysomnography (PSG) are urgently needed. Van Eyck et al. [56] and Weber et al. [57] reported the potential of screening in children with OSA. Van Eyck et al. [56] assessed the ability to identify OSA in 130 obese children using nocturnal oximetry when compared with full PSG studies, and reported poor sensitivity (58%) and specificity (88%) of oximetry concluding that oximetry alone was insufficient to identify OSA in obese children. Weber et al. [57] assessed the role of unattended PSG in 135 children over a 3-year period. Study failure was highest in the first year (52%) and dramatically decreased to 15% in the third year. The primary cause of failure was loss of sensor, which occurred more commonly in younger children. While these data highlight the importance of training in these forms of studies, this study confirms the benefits of unattended PSG in subgroups of patients. While the assessment and treatment of airway inflammation is common in children with asthma, its role in children with sleep disorders is less clear [58]. Studies presented at the 2015 ERS International Congress reported increased 8-isoprostane [59] and leukotriene B4 [60] in children with OSA. Supino et al. [59] assessed 8-isoprostane from urinary and/or exhaled breath condensate samples and matched PSG in 84 children with sleep disordered breathing. Children with moderate-to-severe OSA had higher 8-isoprostane (median (interquartile range) 52.1 (39.2–100.6) pg·mL$^{-1}$) than children with primary snoring only (31 (18.8–41.9) pg·mL$^{-1}$; p<0.005) with significant associations between 8-isoprostane and the apnoea–hypopnoea index (AHI) in children with OSA. In contrast, Alexopoulos et al. [60] found no relationships between leukotriene B4 and the AHI ($r$=-0.11; $p=0.26$) or desaturation index ($r$=-0.13; $p=0.21$) in 104 children with confirmed sleep disordered breathing. However, children with high leukotriene B4 (defined as >75th centile of leukotriene B4 levels in healthy controls) were over three times more likely (adjusted OR 3.19, 95% CI 1.16–8.74; $p=0.02$) to have tonsillar hypertrophy suggesting a link with upper airway inflammation and some aspects of sleep pathogenesis in children.

**Infection and inflammation in CF**

Early diagnosis of pulmonary infection with *Pseudomonas aeruginosa* is necessary to enable timely eradication therapy. However, this is particularly challenging in those who are unable to expectorate sputum. Specific *P. aeruginosa* quorum sensing signal molecules have potential as biomarkers to aid diagnosis of infection. In a prospective, multicentre observational study over 3 years, UK researchers found a high sensitivity and specificity of such a molecule (2-heptyl-4-hydroxyquinoline (HHQJ)) in plasma (86% sensitivity and 86% specificity, respectively) and urine (79%, 71%) compared with conventional microbiological culture [61]. In a longitudinal analysis of 45 children without *P. aeruginosa* infection at baseline, plasma HHQ levels were significantly associated with subsequent positive culture during the following year. The presence of fungi in the Airways of patients with CF is common, but the pathogenic role of most fungi is unclear. The most prevalent species are *Aspergillus* and *Candida*. Metagenomic studies have opened a new dimension, and can identify a large population of fungal microbiota in CF, which are usually not seen in cultures. In a Spanish, prospective, observational cohort study, metagenomic investigations revealed 82% had fungi in contrast to none in culture [62]. In adults, metagenomic screening increased the detection rate of fungi from 30% in culture to 70%; in most cases the fungi were *Candida* (38%) and *Aspergillus* (11%). Patients with newly acquired fungal isolates had worse lung function when compared with persistent culture-negative patients, suggesting that newly acquired isolates impact on disease progression. A research group from Qatar reported a high prevalence of *Candida dubliniensis* (65%) in the sputum of their CF patients. However, the clinical significance of their observation remains unclear [63]. A retrospective data analysis of the UK registry in 2013 revealed an association of inhaled tobramycin and colistin use with increased risk of both non-*Aspergillus* and *Aspergillus* fungal isolation, with odds ratios of 1.33 and 1.40, respectively. In contrast to tobramycin use (with an OR 1.34), there was no statistically significant association between colistin use and *Aspergillus* isolation [64]. Allergic bronchopulmonary aspergillosis (ABPA) causes a high degree of morbidity due to its symptomatology, and the side-effects of long-term corticosteroid therapy. A research team from Germany presented data on the influence of domestic pets on ABPA acquisition. Pet owners showed a higher occurrence of clinical proven ABPA (23%) compared with non-pet owners (8%), which was still highly significant after adjustment for other variables in a multiple logistic regression analysis [65]. The Tel-Aviv CF Centre presented their experience with *Nocardia* infection among 200 CF patients. Eight (4%) had at least one sputum sample positive for *Nocardia*; half of whom had a mild course or no symptoms, however, the other half developed severe lung disease, for which they recommend an early aggressive multidrug therapy [66].

Inflammatory markers are a potentially useful tool for monitoring lung disease in CF, but could only be used routinely if they can be measured noninvasively. Researchers from the Australian AREST-CF study
group measured metabolites from BAL using mass spectrometry, and found an association between decreased levels of adenosine and its metabolite inosine as well as oxidised glutathione and CF lung disease [67]. These data provide new insights into adenosine signalling pathways involved in airway inflammatory responses, and have identified a number of metabolites associated with early disease that could potentially serve as noninvasive biomarkers and therapeutic targets in early CF. In a study from the Netherlands, CF exacerbations were best predicted by a set of nine volatile organic compounds (VOCs), provided that the time interval between breath sampling and exacerbations was not longer than 7 days [68]. VOCs can also be emitted by P. aeruginosa and Aspergillus fumigatus cultures, and their fingerprints are time dependent and differ between mono-cultures and co-cultures, which complicates the in vivo determination [69]. Nitric oxide plays an important role in the regulation of airway calibre and host defence against certain pathogens. In CF airways, nitric oxide is reduced and may contribute to CF lung disease. The mechanisms resulting in reduced airway nitric oxide formation in CF are not completely understood, but recent studies suggest a direct link between cystic fibrosis transmembrane conductance regulator (CFTR) dysfunction and low nitric oxide production from nitric oxide synthases. Changes in FeNO may serve as a biomarker of restored CFTR function in the CF lower airway during CFTR modulator treatment [70]. The effect of ivacaftor on airway nitric oxide has been assessed for the first time by a Canadian research team. They measured FeNO before and 4 weeks after initiation of ivacaftor therapy [71, 72]. In a total of 15 patients, pulmonary function improved significantly and mean ± SD FeNO increased from 8.5 ± 5.0 to 16.2 ± 15.5 ppb. The effect was more pronounced in paediatric compared with adult patients, but there was no linear correlation between changes in FeNO, pulmonary function or sweat chloride concentration.

A French group presented their research on histidinylated polylsines (pLK) and suggest pLK30-His8 as a promising new alternative anti-inflammatory agent or mucolytic, as it is able to compact DNA and liquefy sputum of CF patients [73]. The Catalonia CF newborn screening programme presented their results on cascade carrier testing of relatives when a positive case was discovered [74]. All parents were identified as heterozygous carriers of the corresponding mutation, with the exception of three parents who had, in addition to the corresponding mutation, another CFTR mutation in trans. These three parents all had pathological sweat tests (chloride 65–75 mmol·L⁻¹) and mild symptoms, although they had normal lung function. The authors believe this emphasises the benefits of genetic testing of parents for potential new CF diagnoses, as well as informing genetic counselling for families. However, there are potential drawbacks such as discovering non-paternity, which is a recognised issue in most countries.

There is ongoing controversy about the routine use of bronchodilators in CF. The effect of long-term anticholinergic tiotropium bromide therapy in patients with CF has never been investigated. In a retrospective study of 172 patients, subjects treated with tiotropium bromide for 24 months showed a significant difference in mean annual change of FEV₁ [75]. However, this difference was small (0.43±4.4% versus 2.2±5.18% in controls), and tiotropium bromide therapy was not associated with a lower exacerbation risk. Nonetheless clinical experience shows this can be useful in patients with intractable wheezing not responding to usual combination therapy with inhaled corticosteroids and a long-acting β₂-agonist.

**Neonatology and intensive care**

In a very recent Cochrane review, elective use of high frequency oscillatory ventilation (HFOV) was shown to reduce the risk of BPD; however, lung mechanics during HFOV is still incompletely understood [76]. To elucidate this, two studies from the same research group evaluated the use of resonant frequency in infants and demonstrated for the first time that non-uniform chest wall displacement decreased with increasing oscillation frequency, but asynchronies became more evident [77, 78]. In a study by Bhat et al. [79], lung mechanics during volume-targeted ventilation were compared with pressure-limited ventilation in a group of term or near-term infants and found to be associated with fewer episodes of hypocarbia, although not a reduction in the time to successful extubation. This is very similar to what has been previously reported by the same group of researchers regarding preterm infants suggesting that, regardless of gestational age, an important benefit of volume-targeted ventilation is reduction of the risk for over-ventilation [80]. In a very elegant experimental study, Bourke et al. [81] could visualise the response to methacholine in a mouse model of BPD and using phase-contrast microscopy demonstrated an excessive contraction in the small airways of lungs exposed to hyperoxia and perinatal inflammation. The susceptibility to AHR in children with BPD is a clinical problem, with twice as many of former BPD patients being asthmatic at school age [82]. This model holds the potential for further investigations of the underlying mechanisms to reveal new targets for treatment of airway hyperreactivity in BPD.

Newborn infants, and particularly those born preterm, are at risk for apnoea due to immature autoregulation of respiration; however, several factors may play a role and the following studies have contributed to increased knowledge about the mechanisms behind apnoea in the newborn period. Rossor et al. [83] have demonstrated an increased ventilatory response to hypercarbia in preterm infants receiving...
caffeine therapy. Contrary to this, term infants of smoking and substance abusing mothers exhibited a dampened ventilatory response to hypercarbia, possibly contributing to an increased risk for sudden infant death syndrome [84]. Finally, with the use of synchronised PSG and pH intraluminal impedance no causal relationship between gastro-oesophageal reflux and apnoea could be shown in a mixed group of term and preterm infants [85]. Caffeine therapy for apnoea of prematurity has been shown to reduce the rates of cerebral palsy and cognitive delay; however, no long-term effects on survival without disability or sleep apnoea have been found [86].

Evidence is now accumulating regarding the use of high-flow nasal cannula (HFNC) in the newborn period. In a systematic review conducted by CHAKRABORTY et al. [87], data from a total of 1112 preterm infants enrolled in randomised or quasi-randomised clinical trials were analysed showing similar efficacy to other modes of noninvasive ventilation. As previously reported there was a significant reduction in the incidence of nasal trauma with HFNC, but also a trend towards lower rates of air leaks and mortality. A majority of the infants were born moderately to late preterm suggesting that, at least in this group of more mature infants, the safety profile with less adverse effects is an advantage of HFNC. Interestingly, this is in line with the preference of nursing staff according to a recent study in which nurses favoured nasal continuous positive airway pressure (CPAP) as the choice of respiratory support post-extubation in the most preterm infants, but preferred HFNC in infants ≥28 weeks gestation [88].

In very preterm infants, one of the most commonly used indications for HFNC in clinical practice is weaning from CPAP in infants with evolving or established BPD. In a small cross-over study performed around 2 weeks of age in preterm infants with an average gestational age of 27 weeks, SHETTY et al. [89] reported no significant advantage with regard to work of breathing, thoracoabdominal asynchrony or oxygen saturations with HFNC compared with CPAP. Evidence from YODER et al. [90] suggests that HFNC is comparable to nasal CPAP in preventing extubation failure, but data on various other indications for HFNC use is lacking. HFNC as the primary mode of respiratory support in preterm infants will be tested in the HIPSTER study, a randomised trial designed for noninferiority, and others studies are on the way [91].

Infection and immunity
Respiratory viral infections remain a major cause of morbidity and mortality worldwide. The importance of neutrophils in the inflammatory response to respiratory viruses is now well known. While their role in contributing to the severity of symptoms appears well established it is still unclear how important they are in viral clearance. SAINT et al. [92] presented data from in vitro gene microarray experiments suggesting that when neutrophils are exposed to the respiratory syncytial virus (RSV) a number of antiviral pathways are activated. They concluded that this supported the suggestion that neutrophils play an important active role in the innate antiviral response. FONSECA et al. [93] presented data from a rodent model of RSV infection in which, as expected, the acute illness was characterised by an intense airways neutrophilia that rapidly resolved. However, following these animals to 21 days post-infection there was persistence of viral replication in airways dendritic cells although no virus was identifiable in BAL, suggesting that the virus may be able to persist at low levels in these cells in vivo as well as in vitro. The same group also presented evidence to support the suggestion that the virus may be acquired in utero [94]. Their study found evidence of viral RNA in 60% of human cord blood monocytes and they speculated that in utero exposure may have an impact on the host’s ability to mount an effective long-term memory response. Admissions to hospital with acute bronchiolitis continue to place a considerable burden on health systems. BOYD et al. [95] undertook a health economics assessment of changing the criteria for taking infants out of oxygen using a threshold saturation of 90% in air rather than 94% and suggested that this would result in a saving of approximately EUR 356 per patient without any compromise in safety. This does of course only apply to infants whose saturation in air falls below 94% so cannot be extrapolated to all admissions, and it is also important to remember this was undertaken in patients in whom the saturation data was being used to inform when oxygen therapy should be discontinued, that is they were in the recovery phase of the illness. Hence the results should not be extrapolated to the concept that any infant with saturations >90% early in the course of the illness can be safely discharged. A Russian group presented data suggesting that the use of recombinant α2-interferon administered once daily to children aged <3 years admitted with respiratory viral infections resulted in more rapid resolution of inflammation and clearance of virus [96].

Prevention remains the goal of much of this research, but to date, there is no imminent prospect of an effective vaccine entering routine practice. A large Canadian study retrospectively reviewed outcomes for more than 19000 infants who had been prescribed palivizumab for any reason enrolled over a 9-year period. They found that, when compared with the 60% who received all five doses, health outcomes in those who were less compliant were worse [97]. The increase in hospitalisation was relatively modest, but duration of hospitalisation if admitted was markedly increased (7.8 versus 16.6 days). Effectively addressing the issues underlying low rates of compliance remains one of the great challenges across the whole of healthcare. A second large study from
the same group found no significant benefit could be identified among infants with CF given palivizumab when compared with infants with CF who did not receive the humanised antibodies [98].

The study by Korten et al. [99] addressing the impact of rhinovirus infections on the nasopharyngeal microbiota exemplified the potential of modern sequencing techniques to transform our understanding of the interactions between the human host and microbiota occupying different niches within and on the body, and how these interactions drive important components of health and disease. By sampling the nasopharynx biweekly they confirmed that many rhinovirus infections were largely asymptomatic. Their data indicates that symptomatic infections appear to be associated with lower bacterial diversity and higher bacterial density, in large part supporting some earlier studies using conventional microbiology indicating that density of bacterial pathogens is an important determinant of the severity of nasal symptoms in those with an apparent viral “cold”. As in other studies, they found a degree of resilience within the microbiome although the change might persist for up to 3 weeks. Frequent symptomatic infections did appear able to produce more long lasting changes.

Studies addressing the potential for host factors to impact on these interactions included studies addressing the structure and function of cilia. Hirst et al. [100] demonstrated electron microscopy changes in some cilia from some patients with conditions included under the umbrella term ciliopathies (genetic disorders of non-motile cilia) (figure 1). A multinational team used immunofluorescence to explore functional correlates of genetic mutations in patients with PCD and proposed this approach would be valuable in diagnosis of patients with loss-of-function mutations and missense variants [101]. Another condition associated with significant respiratory morbidity is ataxia telangiectasia. Devaney et al. [102] highlighted the delay in diagnosis of ataxia telangiectasia, although they were unable to identify an impact of this delay on respiratory status. The potential of targeting host mediators in order to prevent disease progression was again raised in a presentation in which the authors noted that ceramides detected in tracheal aspirates from premature infants varied between those who developed chronic lung disease and those who did not [103]. A group from London presented data suggesting that aspiration was relatively common in apparently normal children with chronic or recurrent respiratory problems [104]. Interestingly, there was apparently no difference in the pattern of symptoms or severity in those in whom aspiration was judged to occur as compared with those in whom it was considered that aspiration was not occurring.

FIGURE 1 Cystic cilia tip from Sensenbrenner’s ciliopathy. Reproduced from [100] with permission from the author.
Other groups addressed a range of infectious diseases. A group from Korea again highlighted the importance of considering *Chlamydia trachomatis* as a cause of acute respiratory illness in infancy, with the organism being identified in 8.3% of a very large cohort of infants with acute bronchiolitis [105]. The potential use of novel technologies in directing management was raised by an exploratory study in which MALDI-TOF mass spectrometry was used to characterise protein composition associated with parapneumonic effusions [106]. The challenges of early diagnosis and instituting appropriate therapy for hydatid disease even in areas in which it is endemic was again raised, with the authors noting that 18% of subjects required lobectomy [107]. Research into pulmonary infections and host responses is entering an exciting period as the power of "omics" technologies such as gene sequencing (16S rRNA and the more complex "shotgun" sequencing) metabolomics and computational biology are brought into this area.

**Viruses and asthma and the airway microbiome**

The role of viruses in asthma pathogenesis and asthma symptoms remains a hot topic. In a symposium on virus-driven and host-defence immune responses in acute virus-induced asthma James Gern (University of Wisconsin, Madison, WI, USA) argued that allergic inflammation inhibits interferon responses to virus infections, with rhinovirus C being the most prominent example. This was in line with a recent review by Lambecht *et al.* [108] on the immunology of asthma. In this review, reduced interferon production after exposure to viral agents, due to interference with eosinophil-derived transforming growth factor-β and upregulation of chitinases and chitinase-like proteins, was suggested to be related to increased susceptibility to viruses in patients with asthma. According to James Gern, this decreased response may be attenuated by omalizumab. He showed data that omalizumab reduces rhinovirus-induced asthma attacks during the peak season and speculated that enhanced interferon-α production by omalizumab reduces viral replication, thereby mitigating disease severity and exacerbation risk [109]. Whether inhaled interferons may reduce or mitigate viral exacerbations in asthma remains to be seen.

Peter Le Soeuf (The University of Western Australia, Crawley, Australia) showed that viral infections are associated with an altered airway microbiome, which may lead to a higher risk of viral infection and asthma. Michael Cox (Imperial College, London, UK) showed that the airway microbiome is an equilibrium that can be influenced by internal and external factors. Diseases may cause selection of the microbiome and asthma severity seems to be associated with changes in the microbiome. However, in a small number of infants diagnosed with wheeze, no significant differences were found in the airway microbiome between acute attacks and during short-term follow-up. However, rhinovirus infections did change the bacterial copy numbers in patients with chronic obstructive pulmonary disease and, not surprisingly, smoking alters the airway microbiome. Studies on the influence of the airway microbiome on asthma development are scarce. A small study in children suggested that microbiome characteristics modulate host inflammatory and immune systems in patients with asthma [110]. TEO *et al.* [111] studied the nasopharyngeal microbiome during the first year of life in 234 infants and looked at both viral and bacterial communities. Early asymptomatic colonisation with *Streptococcus* strongly predicted asthma, and not surprisingly antibiotics disrupted the microbiome. The authors suggest that the nasopharyngeal microbiome might offer preventive strategies to asthma development by targeting pathogenic bacteria. Whether manipulation of the airway microbiome may offer new opportunities to interfere with airways and lung diseases remains to be shown, but is a highly exciting area.

**E-health in paediatric asthma**

Two multicentre studies from the Netherlands assessed the effect of a text alert to improve adherence to inhaled medication and of a virtual asthma clinic, respectively [112, 113]. With real-time medication monitoring coupled with text reminders VASBINDER *et al.* [112] were able to improve adherence to treatment; however, this did not result in better asthma control or asthma-related quality of life. Results were similar in the UK, where an intervention using electronic monitoring with feedback (figure 2) did not improve asthma control as assessed by the primary outcome, the Asthma Control Questionnaire (figures 3 and 4), although the UK, where an intervention using electronic monitoring with feedback (figure 2) did not improve asthma control or asthma-related quality of life. Results were similar in the so-called virtual asthma clinic used by VAN DEN WINGAART *et al.* [113]. While reducing clinic visits by 50% similar asthma control could be maintained by offering patients and their parents web-based contacts. VOOREND-VAN BERGEN *et al.* [116] showed that a somewhat similar approach was successful in the so-called virtual asthma clinic used by VAN DEN WINGAART *et al.* [113]. While reducing clinic visits by 50% similar asthma control could be maintained by offering patients and their parents web-based contacts. VOOREND-VAN BERGEN *et al.* [116] showed that a somewhat similar approach was
measured and shared with healthcare providers, although the social acceptability of Smartinhalers needs attention [117]. Although systematic reviews did not show an overall effect of e-health interventions on asthma outcomes, careful analyses of studies that did show an effect can be helpful in determining which interventions may be useful, in which populations, how often and how extensively [118].

**Visualising the airway and lung**

**Antón-Pacheco et al.** [119] presented the long-term outcome and the quality of life of a group of children with severe airway malacia who had undergone metallic stenting. Seven out of 23 patients had their stents in place for more than 5 years. 13 metallic stents, one tracheal and 12 bronchial, were bronchoscopically placed at median age of 5 months. Complications were fewer than expected. The impact on the quality of life was considered as mild to moderate. **Krivec et al.** [120] also presented their experience with therapeutic interventions in the airways during flexible bronchoscopy. In 37 patients (median age 2.8 years, range 0.1–17.0 years) they performed 55 procedures, including large mucous plugs and bronchial cast removals (39 procedures), foreign body extractions (nine procedures) and other airway occlusion resolutions (one large laryngeal granulation, two subglottic membranes, one tracheal membrane and three bronchial membranes). No serious complications were reported.

![FIGURE 2 Electronic adherence monitor (Smartinhaler; Adherium, Auckland, New Zealand). Image courtesy of Mark L. Everard.](image)

**FIGURE 2** Electronic adherence monitor (Smartinhaler; Adherium, Auckland, New Zealand). Image courtesy of Mark L. Everard.

**FIGURE 3** Adherence over a year for the intervention (Smartinhaler feedback group; adherence=70% for the whole 12 months) and control (normal care; adherence=49% for the whole 12 months) groups in a) the morning and b) the afternoon. The boxes represent the interquartile range and the horizontal lines indicate the median. Data from [114] with permission from the author.
In tracheomalacia, DOUROS et al. [121] studied the use of helical CT scans in 21 children with brassy cough or, as they call it, “vibrating” cough (accompanied by a sense of vibration in the chest wall). All children underwent fibre-optic bronchoscopy and tracheomalacia was confirmed in 19 children, who were further investigated with helical CT. The cross-sectional area ratio of the trachea at the level of maximum end-expiration collapse during end-expiration and end-inspiration (CSR), determined the basis for the helical CT diagnosis. Four children who underwent helical CT scans for reasons unrelated to tracheomalacia served as controls. Mean (95% CI) CSR was 0.59 (0.54–0.65) and 0.89 (0.78–0.98) in patients and controls, respectively. Helical CT diagnosed tracheomalacia in four (21%), 13 (68%) and 17 (89%) patients, if the criterion used was CSR $\leq 0.5$, CSR $\leq 0.6$ and CSR $\leq 0.7$, respectively. All the controls had CSR >0.7. The authors proposed that CSR $\leq 0.7$ would be an appropriate threshold for tracheomalacia in children.

In a retrospective study, FASSEEH et al. [122] reviewed 97 infants (78.4% females) who were diagnosed with subglottic haemangioma at Alexandria University Hospitals in Egypt from 2002 to 2012. The mean age of onset and diagnosis were 2.8±1.1 and 5.6±1.7 months, respectively. It was noted that 87.6% of the cases had biphasic stridor, 68% had grade 3 airway obstruction and 35.1% were associated with cutaneous haemangiomas. Most of the cases received oral propranolol (52.6%), some received systemic steroids (30.9%) and a few received both oral steroids and propranolol (16.5%). In the steroids group, 40% of the cases showed a rapid response, 36.7% showed a slow response and 23.3% of cases did not respond to treatment. The best response rate was noted in the propranolol group (98.0%) with a rapid or even very rapid response to treatment without reported side-effects. In an interesting Spanish multicentre retrospective study (involving 11 centres), MORENO et al. [123] presented data regarding the anatomical and clinical characteristics of tracheal bronchus. 93 patients with tracheal bronchus diagnosed by bronchoscopy (81.6%) and/or chest CT were included. All of them were right sided and 66% were in the lower third of the trachea. Broncho-obstructive episodes (60.2%) were the predominant clinical manifestation reported, whereas 44% had pneumonia. Children with tracheal bronchus were associated with heart disease (39.7%), tracheomalacia (35.5%), gastro-oesophageal reflux (31%) and Down syndrome (10.5%). Most of them (73%) had a favourable outcome.

WALKER et al. [124] investigated whether physician’s interpretation of gross findings in children with persistent respiratory symptoms correlated with the information obtained during bronchoscopies they underwent regarding the appropriate antibiotic prescription. They retrospectively audited the bronchoscopic findings, BAL cell count and culture, and decision to commence antibiotics over a 3-month period in one paediatric centre. 22 patients aged 1–15 years underwent bronchoscopy and BAL. The commonest indication was persistent cough. These results gave a sensitivity of 84% and specificity of 57% for the physician decision to initiate antibiotics based on macroscopic bronchoscopy findings. The authors concluded that experienced physician decisions to initiate antibiotics based on bronchoscopy findings have good sensitivity and result in the majority of patients receiving antibiotics appropriately.

Exercise-induced asthma is considered the most common cause of exertional dyspnoea, but other causes such as cardiovascular or pulmonary diseases, poor fitness, hyperventilation and vocal cord dysfunction are also implicated. However, a significant proportion of symptomatic adolescents do not fit into these categories with symptoms of dysfunctional breathing limiting physical activity. BENZRAHI et al. [125] presented data on 59 adolescents with exercise-induced dysfunctional breathing other than exercise-induced asthma. 45 patients showed dysfunctional breathing, but did not meet the criteria of hyperventilation or vocal cord dysfunction. They were considered to have exercise-induced dyspnoea. Frequently there was a history of competitive sport. Typically dyspnoea was accompanied by significant lactic acidosis and oxygen desaturation, as in exercise-induced arterial hypoxaemia. Patients showed a typical breathing pattern, and usually showed a good response to physiotherapy aiming at modification of the breathing pattern.
The clinical utility of routine CT chest scanning is still debated, as CT scanning in young children often requires sedation and the biological effects of radiation later in life are uncertain. Some still advocate for routine annual surveillance, while most believe it should be restricted to clinical need on an individual basis. Due to technical progress, the scanning time can be reduced to 0.4 s and radiation exposure can be reduced further. This makes CT scanning possible during normal breathing (CT-b) without anaesthesia, and has been investigated in infants with and without CF in the first year of life. All CT-b scans showed good concordance between the two scores for air trapping (intraclass correlation coefficient=0.98). Infants with CF showed a significant increase in air trapping score expressed as a z-score at 10 weeks and 1 year of age [126]. Due to concerns over the radiation burden of CT scans, there is still an interest in the use of magnetic resonance imaging (MRI) for long term monitoring of lung disease. Researchers from Heidelberg investigated the correlation between MBW and MRI chest scores in children with CF and healthy subjects. Morphological, perfusion and global MRI chest scores were evaluated in stable CF children and were positively correlated with a higher LCI z-score; the concordance between abnormal findings in MBW and MRI was above 70% [127]. Both had worsening scores during an exacerbation, and significantly improved after antibiotic therapy. However, use of MRI scanning in routine clinical practice seems unlikely given the time, expense and the fact that sedation or general anaesthesia are required for younger patients.

References

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