DEVELOPMENT OF A BREATHELESSNESS CLINIC FOR PATIENTS WITH CHRONIC OBSTRUCTIVE PULMONARY DISEASE (COPD)

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Introduction: Patients with chronic obstructive pulmonary disease (COPD) are often breathless despite optimal medical care and pulmonary rehabilitation. Breathlessness frequently becomes more disabling as COPD advances and may precipitate hospitalisation. Proactively managing breathlessness may improve patient outcomes.

Aim: To describe the development of a multidisciplinary (MD) non-pharmacological Breathlessness Clinic for patients with COPD.

Methods: A literature review of evidence-based non-pharmacological treatments for breathlessness was undertaken. Two randomised controlled trials (RCT) reported on MD clinics in England focusing on the non-pharmacological interventions for breathlessness. A travel grant enabled visits to the clinics, to meet with clinicians, identify assessment parameters, treatment options and outcome measures. On return to Australia, a framework for the clinic was refined and a range of resources developed including a Breathlessness DVD and a relaxation CD, and the acquisition of affordable and effective handheld fans and pedometers. In order to evaluate the service, an RCT was designed with an eight-week wait-list control to ensure evaluation was planned, standardised and rigorous.

Results: The clinic began accepting referrals in May 2016 with a team consisting of a respiratory physician, specialist respiratory nurses, physiotherapists, occupational therapist, dietitian and clinical psychologist. The target population are patients with documented COPD who remain breathless despite standard care. The primary outcome for the RCT is the Chronic Respiratory Questionnaire mastery of breathlessness subscale. Secondary outcomes include breathlessness intensity and unpleasantness and confidence managing breathlessness measured by a numerical rating scale.

Conclusion: The development of a non-pharmacological Breathlessness Clinic for patients with COPD has resulted in an exciting option for patients experiencing refractory breathlessness. The RCT will objectively assess the benefit of this intervention for the patients. Patients remaining breathless may be offered additional psychological and/or pharmacological treatment.

Grant Support: Judith Meppem Travel Scholarship 2015

Declaration of interest: Nil
PRESENTATION RE DEVELOPMENT OF NEWLY ESTABLISHED INTEGRATED RESPIRATORY NURSING SERVICE IN CANTERBURY

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Introduction/Aim: The Canterbury District Health Board aims to provide services which are "better, sooner, and more convenient" for the people of Canterbury.

Historically Canterbury has had three specialty nursing services offering assessment and education to respiratory patients in the community. This has led to confusion as to the role of each nursing service and to which service primary health and hospital staff should refer. In some cases, this has led to duplication of referrals and service, with at times a referral sent to all three nursing services.

The aim of the project was to integrate three community nursing services, Cardio-Respiratory Integrated Specialist Service (Christchurch Hospital), the Community Respiratory Service (formally known as Canterbury Initiative) and CanBreathe (local Asthma Society).

• To offer a coordinated approach to patient education on COPD and Asthma and streamline access across services.
• To stop duplication of referrals across services.
• To support primary health to manage complex respiratory patients within their community.

Methods: A small working group of five was set up, with representation from the three respiratory nursing services. They have developed an algorithm outlining the referral pathway to the appropriate nursing service via a single point of entry. The patient intervention takes place in the community, which may include development of management plans and referrals to other health providers. There is increased collaboration with General Practice through education and joint community visits. The group has identified the need for a shared electronic platform to increase visibility of patient outcomes. Healthpathways provides information regarding the Integrated Respiratory Nursing Service, and can be accessed from Primary and Secondary Care.

Results: The project has:
• Simplified the referral process with a single point of entry.
• Stopped duplication of referrals and service provision.
• Increased collaboration between the community nursing providers and general practice.

Conclusion: Development of an Integrated Respiratory Nursing Service now provides a “better, sooner, and more convenient” intervention for the people of Canterbury.

Grant Support: Nil

TRANSFORMING RESPIRATORY CARE IN HAWKE’S BAY (NZ): ENGAGING PATIENTS AS INFORMED PARTNERS IN THEIR OWN HEALTH CARE

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Introduction/Aim: Management of respiratory conditions has historically been by secondary care responding to the 20% of acute hospital presentations.

Chronic Obstructive Pulmonary Disease is the 4th leading cause of death in New Zealand.

New Zealand has the 2nd highest prevalence of asthma and associated high death rates; 6 times higher for Pacific, 5 times higher for Maori.

We aimed to form an evidence based pathway of integrated care for patients diagnosed with or suspected to have respiratory disease.

Methods: Early intervention in primary care:
• Developing competent workforce
• Sector boundaries removed

Service – Practice Nurse Led – performing spirometry across 18 practices.

Spirometry training with annual accreditation – Respiratory Scientist Mentors for practice nurses – RCNS and BHB

Information technology:- Diagnostic support and care planning tool leading to joint care planning.

Priorities:
• Patient directed education enabling good understanding of their condition
• Empowering patients to be active partners in their health and well-being
• Ongoing long term monitoring and support

Results:
• Obvious growth in patient knowledge and improved quality of life
• Improved service quality in primary care leading to greater equity of care
• Decrease in secondary service referrals from 658(2012) to 28(2015) for diagnostics
• Decrease in bed days by 740 days (2014–2015)
• Decrease in length of stay, decrease in emergency presentations/admissions
• Significant engagement from high deprivation areas and indigenous people
• Improved health literacy

Conclusion: Additional funding for primary care has removed the barrier of cost of accessing early intervention and provided protected time for patient education.

Diagnostic software has assisted health practitioners to standardise best practice intervention ensuring accurate diagnosis, and progression to evidence based care planning, optimising medication, and improving quality of life.

Change management drove itself once staff recognised the benefits to all concerned but particularly for patients.

Patient quote: ‘This has changed my life and given me back my independence….I didn’t think I would ever feel this good ever again’.
NURSES CONFIDENCE AND COMPETENCE IN PROVIDING INHALED MEDICATION EDUCATION TO PATIENTS

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Introduction: Chronic obstructive pulmonary disease (COPD) is a common respiratory disorder. Inhaled medications are widely used to control symptoms and improve the quality of life of patients. Studies have shown that up to 93% of patients with COPD are unable to use their inhaler devices correctly, resulting in inadequate drug deposition. This figure could be higher since the influx of new devices over the past 18 months.

Aim: To assess the confidence and competence of nurses working on a respiratory ward in delivering inhaler device education to patients.

Methods: Subjects were given a questionnaire evaluating confidence in providing inhaler device education to patients. Subjects were asked if supervision of patients taking inhaler medication is required and how often it should be done. Subjects then undertook a competency assessment reviewing the correct steps in using each of the 8 different inhaler devices.

Results: 25 subjects (85% female, 6 yrs (±7.9) experience in nursing) were enrolled in the study. 96% of subjects reported that patients should be supervised when taking inhaler medications and rated their confidence in inhaler device knowledge as ‘not confident’ (16%), ‘moderate’ (76%) or ‘extremely high’ (8%). No subject could correctly perform all the steps of use for all inhalers. When the steps for all devices were tallied, the average total score of correct steps was 47% (SD ± 17%). There was no correlation between total correct steps and years in nursing, or confidence levels.

Conclusion: Despite confidence, competence and knowledge of the correct steps of inhaler use required to teach inhaler technique to patients is poor amongst a group of respiratory ward nurses. There is a need to provide an ongoing educational intervention to improve the inhaler device competence of nurses working in respiratory medicine so that patients receive education regarding correct inhaler use to optimise disease management.

RESPIRATORY NURSES’ KNOWLEDGE AND PRACTICE BEHAVIOURS IN COPD-RELATED ADVANCE CARE PLANNING: PRELIMINARY FINDINGS FROM A CROSSSECTIONAL SURVEY STUDY

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Introduction: Chronic obstructive pulmonary disease (COPD) is a progressive illness that leads to significant morbidity and mortality. Despite high symptom burden, COPD patients have limited access to palliative care, and many are unaware of their prognosis. Advance care planning (ACP) is crucial for patients to make their wishes known if they are unable to speak for themselves in the future. However, ACP is not routine in practice, and many health professionals are reluctant to initiate end-of-life conversations.

Aims: To explore respiratory nurses’ knowledge and practices regarding ACP for patients with COPD.

Methods: ANZ respiratory nurses were invited to participate in an online survey via email listservs and snowballing. Questions covered validated ACP-related knowledge and practice indicators¹. Data were analysed using descriptive statistics, with thematic analysis of free response questions.

Results: Eighty-nine respiratory nurses participated. Between 64–86 (72–97%) correctly answered 8/10 key knowledge indicators, but 57 (64%) were unsure regarding inter-state transferability of advance directives (AD) and 38 (42%) incorrectly thought that a formal AD template was necessary to be legally binding. While 54–85 (61–95%) agreed/strongly agreed with 21/23 ACP practice indicators, 22 (25%) were ambivalent regarding personal confidence in communicating ‘bad news’ and only 45 (50%) routinely initiated ACP. Seventy-four (83%) agreed/strongly agreed that ACP was part of their professional role, but 22 (25%) were ambivalent as to whether ACP was the physician’s responsibility and or indeed their own (26, 29%). Thematic analysis identified a need for time, clear guidance and training to be consistent concerns.

Conclusions: While respiratory nurses are well placed to commence ACP with COPD patients, many were uncertain of ACP legislation and unsure which members of the clinical team should engage in ACP discussions. Variation in confidence to undertake ACP suggests the need for further training to help nurses support patients in articulating their wishes.

Status of research at time of abstract submission: Data collection still ongoing.

Grant Support: Nil


Declarations: The authors have no conflicts of interest to declare.

With thanks to: TSANZ (State and special interest groups), RCN, PCNA, NZNO, LFA, PCV, SVHA, ADHB, SAFGP, and all those clinicians, that facilitated circulation of this survey, and to Dr Tracy Smith for involvement in study design.
IMPROVING PATIENT FLOW AND REDUCING BED BLOCK DURING INFLUENZA SEASON IN A TERTIARY HOSPITAL
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Introduction/Aim: Seasonal influenza places increased demand on the emergency department and hospital resources. Limited single room capacity means that patients often remain in the emergency department >24 hours or there is suboptimal ward placement increasing the risk of transmission between both patients and staff.

Methods: This year, patient and clinical decision making was improved due to development of an Influenza flowchart based on current best practice guidelines.1,2 A local clinical business rule was developed and resources disseminated widely. Once single room capacity was exhausted, patients meeting case definition were cohorted in the respiratory ward, a viral swab taken and oseltamivir commenced.3 Cohort rooms on other wards were co-opted depending on demand.

Results: Improved placement of influenza patients
During June to September 2016 a total of 298 patients were identified as testing positive for influenza A or B. Overall 85% of patients who tested positive, n=252 required admission. Despite high number of influenza admissions 84.5% of patients were isolated correctly (n=213). 25% of total admissions were placed on the respiratory ward with at least 43% of these patients cohorted in a four bedded room (n=28/64).

Decreased length of stay
A four-fold reduction in the number of patients who stayed in ED for more than 24 hours was apparent over the peak winter month of August despite a higher number of presentations to ED overall.

Improved awareness of influenza symptoms and clinical decisions in line with matrix
Despite a 75% increase in presentations in the peak month of August the increased familiarity with the risk matrix was reflected in improved management of patients. A total of 59.8% of patients (n=178) were given oseltamivir, thus greatly reducing the length of stay in single rooms from 7 to 3 days providing the patient was afebrile. Local hospital data also demonstrated an absence of influenza outbreaks in areas where the matrix was used correctly.

Conclusion: The cohort flowchart directed care and provided guidance to clinicians to ensure the safe and appropriate placement of patients thus reducing the risk of influenza outbreaks. Cohorting reduced demand on respiratory single rooms and emergency department length of stay remained lower than yearly predictions despite higher presentations in 2016.

Grant Support: Nil

REFERENCES

Asthma and Allergy 1

ASTHMA CONTROL, AIRWAY INFLAMMATION AND GUT MICROBIOME ARE IMPROVED BY SOLUBLE FIBRE SUPPLEMENTATION
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Introduction/Aim: Westernised diets are typically low in fibre, which may be contributing to asthma development and progression. Observational data has shown that a low fibre intake is associated with worse airway inflammation and lung function. To date, there have been no fibre intervention studies in humans with asthma. However in animal models of airway disease, dietary fibre has been shown to modulate the gut microbiome and reduce airway inflammation. This study aimed to investigate the effects of soluble fibre supplementation in asthmatic adults.

Methods: A randomised, placebo controlled 3-way crossover study in 17 stable asthmatics, using 7 days supplementation with inulin (12g/day), inulin+probiotic (multi-strain >25 billion CFU) and placebo. Clinical assessment, induced sputum and faecal collection occurred before and after each treatment. Selected sputum was dispersed and total and differential leucocyte counts performed. Faecal bacteria were determined by fluorescent in situ hybridisation and faecal SCFA by HPLC-MS. Changes in asthma control (ACQ), lung function (FEV1), gut microbiota, SCFAs and induced sputum cell counts were analysed using Wilcoxon signed–rank and Spearman’s correlations.

Results: ACQ improved (-0.35(-0.50,-0.13) (median(IQR)); p=0.006) and sputum eosinophil% decreased (-1.0(-2.5,0.0), p=0.006) following the inulin intervention only. Significant changes in bacterial taxon relative abundance were seen; Bifidobacterium increased following inulin+probiotic supplementation with a trend following inulin supplementation. Anaerostipes increased following both the inulin and inulin+probiotic supplements. An unidentified Erysipelotrichaceae taxon and Roseburia decreased following inulin supplementation only. Total faecal SCFA and acetate showed a trend to increase following inulin supplementation and changes in SCFA were correlated with changes in FEV1 (Rs=0.53, p<0.001; Rs=0.51, p=0.001), sputum eosinophils (Rs=−0.39, p=0.019; Rs=−0.34, p=0.048) and Erysipelotrichaceae (Rs=−0.40, p=0.013; Rs=−0.35, p=0.030).

Conclusion: Short term inulin supplementation beneficially alters gut microbiome, improves asthma control and airway inflammation in asthma. Soluble fibre supplementation warrants further investigation as a potential non-pharmacological addition to asthma management strategies.

Grant Support: John Hunter Hospital Charitable Trust
NEW MEASURES OF ADHERENCE TO INHALED THERAPY IN PATIENTS WITH ASTHMA

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Introduction: Poor adherence is an important contributor to inadequate asthma control. Electronic monitors provide an objective measure of inhaler adherence. Adherence is often expressed as the mean proportion of prescribed puffs taken during the study period, but this may not adequately reflect variations in use over time. This work aims to develop new metrics to examine patterns in both dose and time to better quantify adherence.

Methods: This is a retrospective analysis of previously published data (Foster et al, JACI 2014) comparing subjects with uncontrolled asthma who received inhaler reminders and usage feedback (IRF), designed to improve preventer adherence versus no reminder or feedback (non-IRF). The adherence of all participants was measured by electronic monitoring. Daily dose records were divided into AM and PM (half-days). We constructed curves based on the proportion of prescribed puffs taken (dose-based measure) and whether or not any puff was taken (time-based measure) over time, and calculated the difference in the area under the actual and expected cumulative curves (AUC). Mann–Whitney test was used to assess differences in these measures between groups.

Results: We examined data from 105 asthma patients with Asthma Control Test (mean ±SD) of 14.6 ±3.4 and age of 41 ±15 years. IRF patients took more doses on average than non-IRF patients in the first two months (104.9 ±89.4 vs 64 ±40.4% of prescribed dose, p<0.0001). They also had shorter average gaps between doses (2.5 ±3.4 vs 4.4 ±6.9 half-day, p<0.01). Dose-based and time-based AUC were higher in IRF-compared to non-IRF-patients (1.7 ±62.2 vs −33.2 ±37.8%, and −21.9 ±23.9 vs −44.8 ±28.2% of expected AUC, p<0.0001, respectively).

Conclusion: We have partitioned adherence into dose-based and time-based measures. Patients differed both in amount of medication taken as well as gaps between doses. Further study will determine the relationship between these patterns and aspects of symptom control and future risk.

Grant Support: Nil

“I’M A PERSON WITH ASTHMA...NOT THE ASTHMA FIRST.”:
PERSONAL EXPERIENCES OF SEVERE ASTHMA: A REVIEW

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Introduction/Aim: 5-10% of the asthma population have severe asthma, which is diagnosed when good asthma control is not achieved with high doses of inhaled treatment, or, good control can be achieved only with high dose inhaled treatment. Previous studies have largely focused on the clinical aspects of severe asthma, and there is very little exploring the personal experience of living with the condition. Our aim was to review the qualitative literature examining the personal experiences of adults living with severe asthma.

Methods: We conducted a systematic review to identify qualitative English-language publications about the personal experience of adults living with severe asthma, excluding studies in which patients were pre-selected by their participation in other research. A meta-ethnographic method and comparative thematic analysis approach were adopted.

Results: From 574 articles, five studies met the inclusion criteria. The topics of the included papers were notably medically focussed. There was a paucity of literature on the physical burden of asthma symptoms. Our analysis revealed that the side effects of medications, particularly oral corticosteroids, communication with health care providers, physical and mental burden, and effects on relationships, work and leisure played a dominant role in these experiences. Feeling excluded from decision-making and being left uninformed about the condition were common experiences.

Conclusion: To our knowledge, this is the first systematic review of the qualitative literature regarding people’s experiences of living with severe asthma. Together, these findings suggest the experience of severe asthma poses challenges to personal autonomy and that people with severe asthma make efforts to re-claim control over their lives. These findings provide important signposts for a deeper exploration of this experience. Our findings suggest there is a paucity of studies exploring the personal experience of living with severe asthma and there is clearly much more to be understood.

Support: CRE for Severe Asthma
IMPROVING ASTHMA EMERGENCY DEPARTMENT DISCHARGE PROCESSES

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Introduction/Aim: Repeated asthma related emergency department (ED) presentations are associated with an increased risk of life threatening asthma. This study aimed to determine if a streamlined ED discharge process would improve level of asthma control and asthma self-management behaviours, encourage patients to follow up with their GP and reduce asthma related re-presentations to EDs.

Methods: We used a controlled cohort study design to evaluate the Asthma ED Discharge Protocol intervention. Patients presenting to the intervention site were provided with a spacer, an education resource (Asthma Control Pack), written prompt to visit GP, comprehensive education and referral to 1800 ASTHMA for follow-up at 1, 5 and 16 weeks post discharge. Control site patients received usual care and referral to 1800 ASTHMA for follow-up at 5 and 16 weeks. Data on behavioural and health outcomes was collected for each cohort via telephone follow-up.

Results: Demographic characteristics at five week contact point:

<table>
<thead>
<tr>
<th></th>
<th>Intervention</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of participants</td>
<td>24</td>
<td>16</td>
</tr>
<tr>
<td>Female</td>
<td>75% (18)</td>
<td>88% (14)</td>
</tr>
<tr>
<td>Average age (mean s.d.)</td>
<td>48 (15.7)</td>
<td>47 (19.6)</td>
</tr>
</tbody>
</table>

Rapid and significant improvements in level of asthma control and delivery device accuracy are seen when a systematic and formalised discharge process is implemented. Results at five weeks post discharge:

<table>
<thead>
<tr>
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<th>Intervention</th>
<th>Control</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asthma Control Score 20+</td>
<td>79% (19/24)</td>
<td>19% (3/16)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Correct delivery technique</td>
<td>96% (23/24)</td>
<td>44% (7/16)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Using spacer</td>
<td>100% (24/24)</td>
<td>94% (15/16)</td>
<td>1.00</td>
</tr>
<tr>
<td>Visited GP</td>
<td>75% (18/24)</td>
<td>56% (9/16)</td>
<td>0.22</td>
</tr>
<tr>
<td>Received asthma action plan from GP</td>
<td>22% (4/18)</td>
<td>33% (3/9)</td>
<td>0.53</td>
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</tbody>
</table>

The cost of delivering the Asthma ED Discharge Protocol was $96 per participant, representing a cost of $282 per patient with well controlled asthma at five weeks.

Conclusion: Formalising asthma related ED discharge processes shows potential in enhancing longer term asthma management. Patients who received this intervention were four times more likely to have well controlled asthma and twice as likely to use their delivery device correctly.

Grant Support: Australian Centre for Health Services Innovation (AusHSI)
AZITHROMYCIN REDUCES EXACERBATIONS IN ADULTS WITH PERSISTENT SYMPTOMATIC EOSINOPHILIC ASTHMA

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Introduction/Aim: Acute exacerbations of asthma cause a significant global burden of illness. Macrolide antibiotics may treat persistent asthma.

Methods: We performed a randomized double-blind placebo controlled trial to determine whether oral azithromycin decreases the frequency of severe asthma exacerbations in adults with symptomatic asthma despite current use of inhaled corticosteroid (ICS) and long-acting bronchodilator (LABD), and who had no hearing impairment or prolongation of the corrected QT interval.

Results: We screened 582 patients and randomized 420 to receive azithromycin 500mg (213 patients) or placebo (207 patients) 3 times per week for 48 weeks. Azithromycin reduced severe asthma exacerbations (0.61/patient-year) compared with placebo (1.07/patient-year; incidence rate ratio [IRR] 0.59; 95% confidence interval [CI] 0.42, 0.83; p=0.002). In eosinophilic asthma (EA), azithromycin reduced severe asthma exacerbations (IRR 0.40, CI 0.23, 0.69) and reduced the proportion of patients experiencing at least one severe exacerbation (placebo 44.7%, azithromycin 26.7%, p=0.010). Azithromycin reduced exacerbations in EA defined using blood eosinophils (IRR 0.44, CI 0.23, 0.83), but not in the non-eosinophilic asthma phenotype (IRR 0.92, CI 0.62, 1.37). Diarrhoea was more common in azithromycin treated patients (34% vs 19%, p=0.001). Azithromycin treated patients had fewer respiratory infections (20% vs 36%, p<0.001).

Conclusion: Adults with persistent symptomatic eosinophilic asthma despite ICS/LABD experience fewer severe asthma exacerbations when treated with the addition of oral azithromycin for 48 weeks. With azithromycin, there is an increase in gastrointestinal adverse effects but fewer respiratory infections. Azithromycin can be added to ICS/LABD therapy in persistent eosinophilic asthma to reduce severe asthma exacerbations.

Grant Support: National Health and Medical Research Council of Australia

Declaration of interest: None.

ANZCTR No: 12609000197235
PULMONARY REHABILITATION GUIDELINES FOR AUSTRALIA AND NEW ZEALAND

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Introduction/Aim: The aim of the Pulmonary Rehabilitation Guidelines (Guidelines) is to provide evidence-based recommendations for the practice of pulmonary rehabilitation (PR) specific to Australian and New Zealand (NZ) healthcare contexts.

Methods: A writing group of 28 experts in PR (10 in the lead group), was selected from expressions of interest. The Guideline methodology adhered to the Appraisal of Guidelines for Research and Evaluation (AGREE) II criteria. Nine key questions were constructed in accordance with the Population, Intervention, Comparator, Outcome (PICO) format and reviewed by a COPD consumer group for appropriateness. Systematic reviews were undertaken for each question and recommendations made with the strength of each recommendation based on the Gradings of Recommendations, Assessment, Development and Evaluation (GRADE) criteria. The Guidelines were externally reviewed by a panel of experts (Australian and NZ respiratory physicians, physiotherapists).

Results: In brief, the Guideline panel recommended that people with mild to severe COPD should undergo PR to improve quality of life and exercise capacity and to reduce hospital admissions; that PR could be offered in hospital gyms, community centres or at home, and could be provided irrespective of the availability of a structured education program; that PR should be offered to people with bronchiectasis, intermittent lung disease and pulmonary hypertension, with the latter in specialised centres. Due to insufficient evidence, the Guideline panel was unable to make recommendations relating to: PR program length beyond 8 weeks; optimal model for maintenance after PR; use of supplemental oxygen during exercise training. The Guideline document discussed the need for culturally appropriate PR programs for Indigenous people with COPD in Australia and NZ to reduce the gap in health outcomes.

Conclusion: The Australian and New Zealand Pulmonary Rehabilitation Guidelines present an evaluation of the evidence for nine PICO questions, with recommendations to provide guidance for clinicians and policy makers.

Grant Support: The Lung Foundation Australia funded the face-to-face meetings and teleconferences and provided administrative support.

UNDERSTANDING BARRIERS AND FACILITATORS TO PULMONARY REHABILITATION REFERRAL, UPTAKE AND PARTICIPATION USING THE THEORETICAL DOMAINS FRAMEWORK: A SYSTEMATIC REVIEW

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Introduction/Aim: Strong evidence supports pulmonary rehabilitation (PR) for people with COPD, yet <5% of those eligible receive PR annually. The Theoretical Domains Framework (TDF) is an integrative framework to explain issues around implementation of best practice evidence in healthcare settings. The TDF was used to understand the constructs that influence referral to, attendance at, and completion of PR by people with COPD.

Methods: A systematic review of studies (qualitative or quantitative) reporting data relating to referral, uptake and/or participation in PR. Extracted data were mapped to one of the 14 domains of the TDF in order to understand the nature of barriers and facilitators.

Participants: Individuals aged >18 years with a diagnosis of COPD; or health professionals working with people with COPD.

Results: A total of 6969 references were screened with 48 studies included and 323 relevant items mapped to the TDF. The most frequently represented domain was ‘Environment’ (34/48 included studies, 34% of mapped items) which included waiting time, burden of illness, travel, transport and health system resources. Other frequently represented domains were ‘Knowledge’ (18/48 studies, eg clinician knowledge of referral processes, patient understanding of rehabilitation content) and ‘Beliefs about consequences’ (16/48 studies, eg beliefs regarding role and safety of exercise, expectations of rehabilitation outcomes). Barriers to referral, uptake or participation represented 61% (n=191) of all items mapped to the TDF. All domains of the TDF were represented, however items were least frequently coded to the domains of ‘Optimism’ and ‘Memory’. Methodological quality of included studies was fair (mean quality score 9/12, SD 2).

Conclusion: The benefits of PR for people with COPD are well established. This review highlights the complex interaction between environment, knowledge, attitudes and behaviours influencing uptake and participation in PR. Overcoming challenges associated with the personal and/or healthcare system environment will be imperative to improving access and uptake of PR.

Grant Support: Nil
ANTIBIOTICS FOR PERSISTENT COUGH OR WHEEZE FOLLOWING ACUTE BRONCHIOLITIS IN CHILDREN (REVIEW)

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Introduction/Aim: Bronchiolitis is a significant health burden globally. It is a clinically diagnosed condition, characterised by tachypnoea; crackles and/or wheeze in children <2 years. Although typically self-limiting, persistent symptoms may continue beyond the acute phase (<14 days), increasing the burden of disease and risk of re-hospitalisation. This review aims to determine the efficacy of antibiotics, compared to a control, to reduce persistent respiratory symptoms following acute bronchiolitis (within six months of illness).

Methods: The previous review was updated in August 2016, using the following databases: Cochrane Airways Group Register of Trials; Cochrane Central Register of Controlled Trials (CENTRAL); MEDLINE; EMBASE; Clinicaltrials.gov; WHO Trial Portal; and the Australian and New Zealand Clinical Trials Registry.

Two review authors independently assessed articles against predetermined selection criteria: randomised controlled trials (RCTs), comparing antibiotics with controls, in the post-acute phase of bronchiolitis (>14 days), in children < 2 years of age diagnosed with bronchiolitis.

Results: Of 344 papers retrieved, only 2 studies1,2 involving 249 children were included. Using an intention-to-treat analysis at 6 months, no significant difference was observed between treatment groups for the primary outcomes: number not cured at follow up (OR 0.69; 95% CI 0.37 – 1.28.), or the number re-hospitalised with respiratory illness (OR 1.19; 95% CI 0.67 – 2.12). There was no significant difference for any of the secondary outcomes at 6 months: recurrent wheeze (OR 0.78; 95% CI 0.35 – 1.73) or bacterial resistance (OR 0.78; 95% CI 0.31 – 1.94).

Conclusion: There is currently no evidence to support the use of antibiotic treatment at the point of illness to prevent post-acute bronchiolitis symptoms. However, there were only 2 studies and children were randomised at the point of bronchiolitis. No studies have randomised children at the point of symptom persistence confirming the need for further RCTs to inform clinical practice.

Key Words: Bronchiolitis, Indigenous, antibiotics, persistent symptoms, respiratory

Nomination for New Investigator Award: No
Grant Support: Nil
Conflict of Interest: No conflicts of interest to declare


LUNG CANCER

TLG INCREASES THE SPECIFICITY OF PET-CT ANALYSIS OF THE MEDIASTINUM IN NSCLC

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Introduction/Aim: Endobronchial ultrasound-guided transbronchial needle aspirate (EBUS-TBNA) and positron emission tomography-computed tomography (PET-CT) are crucial in mediastinal staging of non-small cell lung cancer (NSCLC). However, the specificity of PET-CT in assessing the malignancy status of a mediastinal or hilar lymph node is only 40-79%. Total lesion glycolysis (TLG), defined as lesion volume multiplied by lesion SUVmean, has prognostic value in NSCLC, but to our knowledge has not been used to predict the malignancy status of lymph nodes. Our aim was to determine if the TLG can improve the specificity of PET-CT analysis in predicting the malignancy status of mediastinal and hilar lymph nodes in NSCLC.

Methods: A retrospective study of patients with NSCLC who underwent EBUS-TBNA and PET-CT between January 2012 and December 2014 at Westmead Hospital was performed. A blinded, experienced nuclear medicine physician measured the PET-CT TLG of lymph nodes sampled via EBUS-TBNA. The malignancy status of each lymph node was determined by the EBUS-TBNA histopathology, or the results of surgical sampling when this was performed.

Results: We identified 40 patients who met study criteria. A total of 39 mediastinal and hilar lymph nodes were analysed. Using a cut-off TLG of 5000, the sensitivity of PET-CT was 87% and the specificity was 92%. A receiver operating characteristic curve generated showed good accuracy of TLG, with an area under the curve of 0.92.

Conclusion/discussion:: The TLG may be a helpful tool in determining the malignancy potential of mediastinal and hilar lymph nodes in NSCLC, in particular by increasing the specificity of PET-CT analysis. This in turn would aid the avoidance of unnecessary EBUS-TBNA procedures, and the associated costs and delays to surgery. Prospective studies are required to further evaluate this novel tool.

Grant Support: Nil
PREOPERATIVE EXERCISE TRAINING FOR PATIENTS WITH NON-SMALL CELL LUNG CANCER: A COCHRANE SYSTEMATIC REVIEW.

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Aim: To determine the effect of preoperative exercise training on postoperative outcomes including risk of developing a postoperative pulmonary complication, postoperative duration of intercostal catheter and length of hospital stay in adults scheduled to undergo lung resection for non-small cell lung cancer (NSCLC).

Methods: We searched the Cochrane Central Register of Controlled Trials, PubMed, EMBASE, PEDro and SciELO (The Scientific Electronic Library Online) up to March 2016. Randomised controlled trials (RCTs) were included in which study participants who were scheduled to undergo lung resection for NSCLC were allocated to receive either preoperative exercise training or no exercise training. The two review authors independently screened and assessed studies for inclusion. Risk of bias was assessed using the Cochrane seven evidence-based domain table Meta-analyses were conducted where possible.

Results: Five RCTs involving 167 participants were identified. Overall, the risk of bias in the included studies was high. Pooled data from three studies demonstrated that preoperative exercise training reduced the risk of developing a postoperative pulmonary complication by 70% (RR 0.30; 95% CI 0.14 to 0.66). Compared to the control group, the number of days patients in the intervention group needed intercostal catheters was lower (MD −3.33 days; 95% CI −5.35 to −1.30 days) (pooled data from two studies), and postoperative length of hospital stay was also lower in the intervention group (MD −4.34 days; 95% CI −5.65 to −3.03 days) (pooled data from three studies).

Conclusion: Preoperative exercise training appears to reduce the risk of developing a postoperative pulmonary complication, the duration of intercostal catheter and postoperative length of hospital stay in people undergoing lung resection for NSCLC. The findings of this review should be interpreted with caution due to disparities between the studies, methodological limitations, risk of bias and small sample sizes. This systematic review emphasises the need for larger RCTs.

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Declaration of interest: Nil.
Nomination for awards: Physiotherapy.

IS IT LUNG CANCER?: EVALUATING THE PERFORMANCE OF A “LUNG LESION FOR INVESTIGATION” SERVICE AT A NEW SECONDARY HOSPITAL

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Introduction/Aim: Access to, and quality of cancer services are important determinants of patient outcomes from respiratory malignancy. Existing Australian data is limited to retrospective analyses of patients with proven malignancy; these data do not fully describe the journey of all patients with suspected respiratory malignancy. A prospective registry was developed to inform service requirements and benchmark quality.

Methods: Patients referred to St John of God Midland Hospital, a general secondary hospital, for assessment of possible respiratory malignancy were prospectively included. Analyses of clinical data were descriptive.

Results: Between November 2015 and September 2016, 107 patients were referred. 69 were referred by general practitioners and 79 were seen in outpatient clinics. Referral triggers were symptoms with radiological abnormality (76/109), incidental findings (24/107) or abnormality on CT scans performed for lung cancer screening (7/107). The commonest abnormalities were mass (25/107), and pulmonary nodules (single, 23/107; multiple, 15/107). Invasive diagnostic investigations (undertaken in 64/107) established a tissue diagnosis of malignancy in 54, of which 39 were lung cancer. CT surveillance was recommended for 39; four patients were not investigated further. Median time from outpatient referral to first review was eight days (range 0–190; 58/79 of outpatient were seen within 14 days. Patients who required endobronchial ultrasound at a tertiary centre waited a median of 34 days (n=6, range 25–98) from first review to referral for treatment compared to a median of 15 days (n=46, range 0–115) for those who did not.

Conclusion: This pilot registry describes the wide variation in diagnostic pathways taken when investigating possible lung cancer. In addition to providing prompt outpatient assessment, lung cancer services are needed for inpatients presenting acutely unwell. The waiting times for outpatient assessment and diagnostic testing was short for the majority of patients, but there were delays for some patients highlighting the need for ongoing service improvement.

Grant Support: Nil
PLEURX CATHETER DRAINAGE SYSTEMS ARE A SAFE, EFFECTIVE AND ECONOMICAL STRATEGY FOR THE MANAGEMENT OF MALIGNANT PLEURAL EFFUSION AND ASCITES. A SINGLE CENTER EXPERIENCE.

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Introduction/Aim: Malignant pleural effusion (MPE) and malignant ascites (MA) contribute to significant breathlessness for patients with advanced cancer. Managing this fluid has required frequent admission for repeated needle thoracocentesis or pleurodesis. Since 2008, we have added indwelling tunnelled catheters to the management options available to clinicians within our LHD.

Methods: This project is a retrospective review of our experience with PleurX catheters inserted for management of MPE and MA between 2008–2016.

Results: All patients reported immediate improvement in breathlessness.

Two hundred and ninety-three PleurX catheters were inserted. The indication was MPE in 176 and MA in 111. Bilateral pleural PleurX were inserted in 17 patients. Pleural and peritoneal catheters were inserted simultaneously in 3 patients.

A technical success rate of 100% was achieved. Drains remained in situ for a mean of 64.0 days (range 2–392 days). Almost all (87%) of patients died with the PleurX catheter insitu. Pleurodesis was achieved in 6.1% (n=18) of patients and PleurX was removed without fluid accumulation. The remaining patients 6.9% remain alive with the PleurX insitu.

PleurX were inserted in patients MPE and MA due to NSCLC 20.5% (n=60), breast 19.8% (n=58), gastric/esophageal 7.8% (n=23), pancreatic 7.5% (n=22), ovarian 7.5% (n=22), colorectal 6.1% (n=18), adenocarcinoma of unknown origin 4.1% (n=12), mesothelioma 3.8% (n=11), hepatocellular carcinoma 2.7% (n=8), SCLC 1.7% (n=5), melanoma 1.7% (n=5), bladder 1.7% (n=5), cholangiocarcinoma 1.4% (n=4), renal cell carcinoma 1% (n=3), angiosarcoma 0.7% (n=2), endometrial 0.7% (n=2), and alcoholic cirrhosis 1.4% (n=4), others/unknown 9.9% (n=29).

Complications from drain insertion included; wound infection 2.4% (n=7), blocked catheter 1.7% (n=5), fell out/cuff exposed 1.7% (n=5), leakage from drain site 0.7% (n=2), bowel injury 0.3% (n=1).

The cost per patient on average for the entire duration of pleural and peritoneal PleurX was $1759 and $984 respectively (ongoing hardware cost and insertion cost). The average number of pleural drainage bottles and peritoneal drainage bags per patient was twenty-nine and twenty-two respectively.

Conclusion: This study demonstrates the safe use of tunnelled PleurX catheters in patients who present with MPE or MA. Once inserted, subsequent drainage of re-accumulated effusion or ascites can be managed in the comfort of the patients home. We believe the use of a PleurX catheter should be considered as a first line approach in the management of refractory MPE and MA.

Grant Support: Nil

No conflict of interest to disclose.
MDT V NON-MDT CARE IN LUNG CANCER: DOES CO-MORBIDITY STATUS HAVE AN IMPACT?
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Introduction/Aim: Multidisciplinary team (MDT) care affects lung cancer outcomes including treatment receipt1,2 and survival1,3. This project analyses lung cancer outcomes according to MDT care in an Australian dataset from 2006–2012 aiming to adjust outcomes for co-morbidity status (Charlson co-morbidity index and Colinet simplified co-morbidity score).

Methods: A consecutive cohort of cases diagnosed between 1 January 2006 and 31 December 2012 were identified through the local Clinical Cancer Registry (ClinCR). Histopathology, TNM stage, ECOG status and survival were analysed according to MDT care. The Charlson co-morbidity index has been calculated via ICD coding. Colinet simplified co-morbidity scoring will be completed via validated items in institutional medical records.

Results: 1022 cases of primary lung carcinoma were identified; 296 (29%) were presented at MDT (“MDT”), 726 (71%) were not (“non-MDT”). SCLC comprised 33/296 (11%) of MDT and 121/726 (17%) of non-MDT cases. TNM data showed higher rates of staged (vs unstaged) cases for the MDT group compared with non-MDT group (97% vs 84.4%, p<0.001). Stage IV patients were half as likely to be presented at MDT than other stages (22% stage IV vs 45% stage I-II combined). ECOG status was better recorded for MDT than non-MDT (72% vs 49.6%, p<0.001). Survival analysis, adjusted for age, disease stage and ECOG status, showed an overall 52% increase in the relative probability of death for non-MDT vs MDT cases (HR 1.52 [95%CI 1.25 – 1.85]; P<0.0001). Stage IV patients were half as likely to be presented at MDT than other stages (22% stage IV vs 45% stage I-II combined). ECOG status was better recorded for MDT than non-MDT (72% vs 49.6%, p<0.001). Survival analysis, adjusted for age, disease stage and ECOG status, showed an overall 52% increase in the relative probability of death for non-MDT vs MDT cases (HR 1.52 [95%CI 1.25 – 1.85]; P<0.0001). An initial data extract comprising ICD coding for a subset of 264 patients included 168 (63%) MDT cases and 96 (36%) non-MDT cases, most of whom had a co-morbidity score of 2 or less.

Discussion/Conclusion: In an analysis adjusted for age, TNM stage and ECOG status, MDT care is associated with improved survival. This project seeks to update this analysis with adjustment for co-morbidity status according to two previously validated measures.

Grant Support: Nil

A SIT-TO-STAND TEST IS USEFUL TO ASSESS EXERCISE OXYGENATION IN PATIENTS WITH INTERSTITIAL LUNG DISEASE
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Introduction/Aim: Exercise desaturation on six-minute walk test (6MWT) is associated with poorer outcomes in patients with interstitial lung disease (ILD). The 60s sit-to-stand test (STST) is faster and requires less space than 6MWT, however its utility in ILD patients has not been assessed. The aim was to compare oxygen desaturation between the 6MWT and STST in ILD patients.

Methods: Consecutive subjects were recruited from RPAH ILD clinic. Patients performed 6MWT & STST in randomised order with 30mins recovery. 6MWT was performed according to ATS/ERS standards. STST was performed by a single operator, requiring the patient to sit and stand out of a chair as many times as possible in 60s whilst oxygen saturations (SpO2) and heart rate (HR) were monitored. Nadir SpO2 was defined as the lowest SpO2 recorded during the exercise or recovery period. Significant desaturation on 6MWT was defined as nadir SpO2<88%.

Results: 50 subjects were studied (mean age 63.4±11.7 years; 30 males). The underlying cause of ILD was idiopathic pulmonary fibrosis (48%), connective tissue disease (38%), and other (14%). Mean FVC was 72.58±16.3%, DLCO 57.46±16.2%, and resting SpO2 98.64±1.4%.

Median 6MWD was 500m (range 300-817m) and nSTS was 26 (range 18–66). Patients desaturated more on 6MWT than STST (mean nadir SpO2 90.34±6.7% vs 93.28±5.4%, p<0.0001). There was a strong correlation between nadir SpO2 on 6MWT and STST (Spearman’s r = 0.88, p<0.0001). The area under the receiver operating curve for significant desaturation on 6MWT using the nadir SpO2 on STST was 0.96 (95%CI 0.91–1.0). Nadir SpO2 <94% on STST gave a sensitivity of 100% and specificity of 87.1% for significant desaturation on 6MWT.

Conclusion: In an ILD cohort, nadir SpO2 on STST and 6MWT correlated strongly. The STST is suitable in an office setting, and nadir SpO2 on STST of ≥94% appears to rule out significant desaturation on 6MWT.

Grant Support: Nil
TO-025
VASCULAR REMODELLING IS REVERSED FOLLOWING BMPR2-EXPRESSING ENDOTHELIAL PROGENITOR CELL THERAPY IN A MCT-INDUCED PAH RAT MODEL
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Introduction/Aim: Pulmonary arterial hypertension (PAH) is caused by pulmonary vascular remodelling. Reduced expression of the bone morphogenetic protein receptor type-2 (BMPR2) is causally linked to PAH. Previously, we have augmented endothelial progenitor cells (EPCs) to over-express BMPR2 and transplanted them in a monocrotaline (MCT)-induced PAH rat model resulting in an amelioration of the disease. We now assess the effects of our BMPR2-EPC and EPC only therapy on vascular remodelling in this MCT-induced PAH rat model.

Methods: Rats (n=8) were injected with MCT, and then at day 10, rats were intravenously injected with EPCs only, AdBMPR2 transfected EPCs, AdTrackLuc transfected EPCs or uninjected. After a further 8–10 days, PAH was assessed and lungs were extracted and processed into paraffin blocks. Immunohistochemical analysis on vessels 50 μm or less was performed with α-smooth muscle actin (α-SMA) Ki67 and proliferating cell nuclear antigen (PCNA).

Results: After 10 days PAH was attenuated as shown by a significant reduction in RVSP, mPAP and Fulton Index in the BMPR2-transduced EPC group compared to disease only, EPCs only and irrelevant virus-transduced EPC groups. BMPR2-EPC treated animals a significant reduction in vessel muscularisation (31.88%) vessel thickness (24.87%) and a significant reduction in cellular proliferation as shown by PCNA (35.77%) and Ki67 (66.86%) staining compare to disease only.

Conclusion: Amelioration of PAH can be achieved using BMPR2 modified EPCs. There’s a significant reduction in distal vessel muscularisation and cellular proliferation following EPCs and BMPR2-EPCs treatment, with a greater effect seen in the BMPR2-EPCs group.

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TO-026
REDUCED GCR, HDAC2 AND HSP90 IN PRO-INFLAMMATORY LYMPHOCYTES IN THE SMALL AIRWAYS DURING BOS
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Introduction: Immunosuppression therapy following lung transplantation fails to prevent bronchiolitis obliterans syndrome (BOS), primarily a disease of the small airways, which we have shown is associated with lack of suppression of pro-inflammatory lymphocytes. We have also recently shown reduced glucocorticoid receptor (GCR), histone deacetylase 2 (HDAC2) and heat shock protein 90 (Hsp90) in these pro-inflammatory lymphocytes following transplant. We hypothesize that these lymphocytes target the small airways during BOS.

Methods: Blood, bronchoalveolar lavage, proximal and distal airway bronchial brushings were collected from patients with BOS (n=5), stable lung transplant patients (n=18) and healthy aged-matched controls (n=10). Intracellular pro-inflammatory cytokines and expression of GCR, HDAC2 and Hsp90 were measured in lymphocytes subsets following culture using flow cytometry.

Results: BOS was associated with an increase in CD8 T, NKT-like and NK cells in the distal airways compared with stable patients and controls. There was an increase in IFNγ, TNFα and reduced GCR, HDAC2 and Hsp90 expression in these lymphocyte subsets in patients with BOS compared with stable patients and controls. There was a correlation between the percentage of CD8+ T cells expressing HDAC2, GCR and Hsp90 from distal brushings with FEV1 (HDAC2:R=−.676 , p=.031; GCR: R=−.632 , p=.039).

Conclusions: BOS is associated with increased pro-inflammatory CD8+ T, NKT-like and NK cells in the distal airways. Treatments that increase GCR, HDAC2 and Hsp90 in these lymphocyte subsets may reduce graft rejection.

Grant Support: RAH Respiratory Clinical Trials
Declaration of interest: No declaration of interest
**MATRIX METALLOPROTEINASE-7 (MMP7) MAY BE A BIOMARKER OF EARLY CHRONIC LUNG REJECTION**

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**AIM:** Lung transplantation is the best option for patients with end-stage lung diseases. Chronic rejection is a major complication post-operation. More than half of all lung transplant recipients experience chronic rejection but early in the course of disease few symptoms are clinically apparent. The aim of this study was to investigate the use of MMP7 as a biomarker of early chronic lung rejection.

**METHODS:** Forty adult lung transplantation patients at The Alfred had blood collected and lung function measured at 4 time points: pre-transplantation, 3-, 12- and 24 months post-transplant. Measurements of plasma MMP7 were performed using sandwich immunoassay. Clinical variables including age, BMI, smoking history and history of acute graft rejection were also recorded. Univariate and multivariate analyses assessed the clinical variables associated with plasma MMP7 levels. The utility of MMP7 as a biomarker of disease progression was assessed by receiver-operating characteristic curve analysis. Disease progression was defined as a greater than 5% fall in percent predicted Forced Vital Capacity in 1 second (FEV1%) between the 12 and 24 month time points.

**RESULTS:** Seventeen of 40 patients experienced a decline in lung function between 12 and 24 months post-operation. Patients who experienced disease progression had higher levels of plasma MMP7 at 12 months post-transplant than those who remained stable (mean stable 0.6 pg/mL vs mean progressed 1.1 pg/mL; p<0.001). Increased levels of MMP7 predicted disease progression (area under the curve = 0.8, p=0.003).

**CONCLUSION:** There are no biomarkers able to detect disease progression of chronic rejection before patients experience lung function decline. MMP7 is a biomarker of disease progression in idiopathic pulmonary fibrosis, a fibrotic lung disease with shared mechanisms of pathogenesis to chronic rejection. Elevated MMP7 levels may be detectable prior to clinical manifestation of lung function decline. MMP7 levels may be a biomarker of disease progression in chronic rejection.

**Grant Support:** Nil

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**DEAD SPACE VENTILATION IS LINKED TO EXERCISE CAPACITY AND SURVIVAL IN DISTAL CHRONIC THROMBOEMBOLIC PULMONARY HYPERTENSION**

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**Introduction:** Cardiopulmonary exercise testing (CPET) is frequently used for the evaluation of patients with pulmonary hypertension (PH). Patients with non-operative distal CTEPH represent a unique subgroup of PH where microvascular disease resembling pulmonary arterial hypertension (PAH) may predominate and efficacious medical therapy is now available. However, little is known regarding the detailed CPET profile of distal CTEPH, and whether ventilation and gas exchange responses are different to PAH.

**Methods:** Forty-nine consecutive patients with non-operable distal CTEPH according to multidisciplinary team assessment and 45 matched PAH patients underwent CPET and right heart catheterization. Patients were followed up for median of 3.2 years.

**Results:** Pulmonary haemodynamics were similar in distal CTEPH and PAH groups but patients with distal CTEPH achieved lower percentage predicted peak oxygen consumption (59±13 vs 66±14%, p<0.05). At peak exercise, higher physiological deadspace fraction (Vd/VT) (0.45±0.07 vs 0.35±0.07, p<0.0001) and larger arterial-to-end tidal carbon dioxide gradient (9±3 vs 5±3mmHg, p<0.0001) were observed in distal CTEPH compared with PAH. Ventilatory efficiency expressed as VE/VCO2 slope was also more impaired in distal CTEPH (52.2±10.1 vs. 43.8±8.4 L/min/L/min, p<0.0001). In only the distal CTEPH group, higher Vd/VT correlated with lower peak oxygen consumption (r = −0.46, p = 0.003) and was associated with worse survival.

**Conclusions:** Compared with PAH, a distinct pattern of pathophysiology is observed in distal CTEPH which is characterized by increased deadspace ventilation resulting in worse ventilatory efficiency and greater impairment of exercise capacity. In distal CTEPH, deadspace ventilation correlates with exercise capacity and may have potential prognostic relevance.

**Conflict of Interest:** Nil
HYPOXEMIA

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Aim: Patients with interstitial lung disease (ILD) frequently experience intermittent hypoxaemia (IH) during exercise and sleep. Chronic IH is associated with increased oxidative stress and maladaptive cardiovascular consequences, including pulmonary hypertension (PH). We aimed to compare plasma markers of oxidative stress [protein carbonyl (PC) and thiol concentrations] in two ILD cohorts who were exposed to the intervention of oxygen versus medical air.

Methods: Two previously described ILD patient cohorts with either significant sleep-related hypoxia (>10% of sleep time with SpO2<90%), or exercise-induced desaturation (SpO2<88% during 6-minute walk test, 6MWT) participated in randomised, double-blinded, sham-controlled, cross-over studies. Paired plasma samples were taken following both supplemental oxygen and medical air during exercise and sleep. Plasma samples were analysed for PC and thiol concentrations using the DTNB assay with the 96-well plate method.

Results: Eleven Sleep-ILD patients and 14 Exercise-ILD patients were studied (combined population: 11 females; mean age 66.9±9.1 yrs; BMI 30.9±5.8; FVC 71.4±16.8%; DLCO 49.1±11.4%, nadir 6MWT SpO2 84.5±4.8%, RVSP 40.3±13.7mmHg, %sleep time<90% 30.8±17.9 (n=11)). Baseline plasma PC concentrations (on air) were elevated, ranging between 0.12-0.90 nmol/mg, (normal 0.1 nmol/mg), and baseline thiol concentrations were depleted (range 68.0-256.3 μmol/L; normal: 400–600 μmol/L). Lower thiol concentrations were associated with lower nadir air oxygen (R=0.82, p=0.002), and increased RVSP (R=0.62, p=0.03). With the application of oxygen during sleep and exercise, plasma PC levels were not significantly reduced, compared with air. Anti-oxidant thiol levels however, were significantly improved with oxygen in both sleep and exercise cohorts [Sleep: 131.0±28.0 μmol/L (air), 173.8±50.7 μmol/L (oxygen), p=0.03.; Exercise: 171.4±52.4μmol/L (air), 213.9±70.5 μmol/L (oxygen), p=0.002].

Conclusion: ILD patients with IH during sleep or exercise have an increased burden of oxidative stress. Lower anti-oxidant thiol levels are associated with more severe sleep hypoxia and PH. Supplemental oxygen improves anti-oxidant capacity in ILD cohorts.

Grant Support: Primary Care

OXGEN ATTENUATES OXIDATIVE STRESS IN ILD PATIENTS WITH NOCTURNAL AND EXERCISE-INDUCED HYPOXEMIA

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Aim: Patients with interstitial lung disease (ILD) frequently experience intermittent hypoxaemia (IH) during exercise and sleep. Chronic IH is associated with increased oxidative stress and maladaptive cardiovascular consequences, including pulmonary hypertension (PH). We aimed to compare plasma markers of oxidative stress [protein carbonyl (PC) and thiol concentrations] in two ILD cohorts who were exposed to the intervention of oxygen versus medical air.

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Conclusion: ILD patients with IH during sleep or exercise have an increased burden of oxidative stress. Lower anti-oxidant thiol levels are associated with more severe sleep hypoxia and PH. Supplemental oxygen improves anti-oxidant capacity in ILD cohorts.

Grant Support: Primary Care

HIGH RATES OF RESPIRATORY SYMPTOMS AND AIRWAYS DISEASE IN MENTAL HEALTH INPATIENTS IN A TERTIARY CENTRE

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Introduction: People with severe mental illness (SMI) have a lower life expectancy than the general population. A significant proportion of this excess mortality is due to a higher prevalence of cardiac and metabolic disease. Less is known of the prevalence of respiratory disease in this group. This cross sectional, observational study aimed to assess the prevalence of respiratory disease and symptoms in patients admitted to an inpatient mental health unit.

Methods: 82 inpatients had a structured interview and questionnaire completed. The questionnaire included self-reported diagnoses of common diseases and screening questions designed to detect respiratory disease and sleep disordered breathing. Spirometry was performed on the basis of symptoms and smoking status. Access to primary care was assessed including surrogate markers for comprehensive care such as vaccination status.

Results: Patients reported high rates of respiratory symptoms including wheeze, dyspnoea and cough. Productive cough was significantly associated with tobacco use (p<0.005). 52% of patients reported daily tobacco use and 13% used cannabis at least monthly. Ten patients (17%) had spirometry consistent with COPD of whom 6 did not have a formal diagnosis of COPD previously. Symptoms suggestive of sleep disordered breathing were common with 9 patients (11%) reporting witnessed apnoeic episodes while sleeping. Vaccination rates were low for both pneumococcal and influenza vaccine.

Conclusions: People with SMI have high rates of respiratory symptoms with a high prevalence of COPD on spirometry. Half of the COPD cases were not previously diagnosed suggesting a hidden burden of respiratory disease in patients with SMI.

Key words: mental illness, COPD, smoking.

Support: Prince Charles Foundation Novice Researchers Grant

TO-029

TO-030
A QUALITATIVE STUDY USING INNOVATIVE TECHNOLOGY VIA A SMARTPHONE APPLICATION TO PROVIDE DEMONSTRATIVE INHALER TECHNIQUE EDUCATION FOR ASTHMA PATIENTS
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Introduction/Aim: Evidence suggests more than 75% of asthma patients fail to use their inhalers correctly resulting in frequent exacerbations. Sub-optimal education for health professionals and subsequently patients, may be a contributing factor. Innovative technology (‘augmented reality’), accessed via smartphone applications, can function as an educational tool to address these issues. Therefore the aim of this study is to obtain health professional, asthma patient and key community stakeholder perspectives on the feasibility of augmented reality technology to improve asthma inhaler technique education.

Methods: A patient information poster displaying images of the 22 asthma inhaler devices was produced based on existing evidence based recommendations and resources. Augmented reality technology was then utilised to provide a video demonstration of correct inhaler technique for each device, made accessible via a free smartphone application. Twenty-one semi-structured, one-on-one interviews with health professionals, asthma patients and key community stakeholders were conducted between August and September 2016. Data was analysed thematically using the Triandis model of interpersonal behaviour.

Results: Interviews with asthma patients highlighted perceived competence with inhaler technique. However, health professionals and key community stakeholders identified that this perception was misguided and that poor inhaler technique is a facilitating condition for incorrect inhaler use and sub-optimal disease management. Delivering inhaler technique education using augmented reality was favoured by all participants, particularly around ease of use with the ability to visually display various inhaler techniques for each device. However, all participants identified some barriers, particularly for use among the elderly.

Conclusion: Augmented reality may be a novel means to address poor inhaler technique among certain cohorts of asthma patients and serve as a prompt for health professionals to initiate review of inhaler devices. A randomised controlled trial design is needed to evaluate efficacy of this technology for use in the clinical care setting.

Grant Support: The Hospital Research Foundation; COI: None

THE DEBILITATING BURDENS OF LIVING WITH SEVERE ASTHMA
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Introduction/Aim: People with severe refractory asthma (SRA) likely face a sizeable burden, beyond that of people with milder disease. We aimed to explore the little-known experiences of people living with SRA.

Methods: Participants identified as SRA via the Australasian Severe Asthma Network and/or specialist asthma clinics across seven Australian states were invited for telephone interview. Semi-structured interviews were conducted consecutively until no new themes emerged. Interviews were recorded, transcribed and thematically analysed.

Results: Most of the 25 participants (aged 23–81 yrs; 68% female) experienced daily symptoms. Key themes were: ‘The body as a hindrance’: SRA placed wide-ranging limits on life from daily chores to career, relationships and family life; ‘Alone with asthma’: Interviewees felt alone in understanding the experience of asthma; they suffered emotional distress including frustration, hopelessness and/or depression, but formal emotional support services were lacking; ‘Striving to adapt’: Participants expressed varying degrees of adjustment to their diagnosis, from denial to full acceptance; ‘Concerns and experience with treatments’: Most interviewees accepted the need for medicines but were concerned about oral corticosteroid side-effects and disliked reliance on treatment, which they felt took over their life; ‘Day-to-day medical care’: Participants reported need for more accessible, knowledgeable GPs, and better communication with GPs/specialists; ‘Acute care’: Exacerbations were frightening for participants/family yet they avoided emergency department presentation to avoid the disruption that hospitalisations placed on daily lives; ‘Support needs’: practical and emotional support varied from none at all to provision by willing/fatigued family. Participants felt compelled to ‘push through’ due to parenting/financial responsibility.

Conclusion: SRA imposes long-term, debilitating burdens and should be considered differently to milder asthma. There is an urgent need to improve support services and primary care for SRA patients, and assistance for their families.

Grant Support: Funded by Asthma Australia with independent research grants from Boehringer Ingelheim, GlaxoSmithKline and Novartis
QUITTING EXPERIENCES AND SMOKING CESSION PREFERENCES OF SMOKERS IN AUSTRALIAN GENERAL PRACTICES


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Introduction/Aim: Approximately one in five patients attending general practice is a daily smoker. Information on previous quitting experiences and smoking cessation preferences of smokers attending general practice is necessary to inform effective interventions.

Methods: Current/ex-smokers (aged ≥40 years, ≥10 pack years) were recruited from general practices across Melbourne for the RADICALS® study – a cluster randomised controlled trial of an interdisciplinary model of care to reduce the burden of smoking and lung disease in Australian primary care. Patients completed an exhaled carbon monoxide (CO) test to confirm smoking status and a structured questionnaire during a face-to-face interview.

Results: Of the 653 participants identified across 38 general practice clinics in Melbourne, 456 (70%) were current smokers; 425 (93%) of whom were daily smokers. The median exhaled CO reading was 22 ppm (IQR 14–29). On a scale of 1 (low) to 10 (high), the self-reported motivation (median 6; IQR 4–8) and confidence to quit (median 5; IQR 3–7) were modest among current smokers. Of the current smokers, 237 (52%) had attempted quitting in the previous 12 months; 156 (66%) of whom reported two or more attempts in the past year. The most common pharmacological treatments used were nicotine replacement therapy (128, 54%) and vareniclne (72, 30%). The most popular non-pharmacological treatment was hypnotherapy (40, 17%). Approximately 40% of smokers would use medications to aid future quit attempts. E-cigarettes have been used by 21 (9%) current smokers with at least one quit attempt in the previous year, and would be considered to help future quit attempts by 132 (29%) current smokers.

Conclusion: Many smokers in primary care consider and have used non-evidence-based smoking cessation aids. Healthcare professionals should use motivational counselling and emphasise evidence-based treatments when providing support to smokers.

Grant Support: NHMRC, Cyril Tonkin Scholarship, Boehringer Ingelheim, Lung Foundation of Australia, Eastern Melbourne PHN

DIAGNOSIS OF COPD IN AUSTRALIAN GENERAL PRACTICES: EXPERIENCE FROM THE RADICALS® TRIAL


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Introduction/Aim: Diagnosis and management of COPD are underpinned by strong scientific evidence and local guidelines are available to guide decision making. However, previous studies have suggested under-diagnosis and misdiagnosis of COPD in primary care. We describe our experience in case-finding from the RADICALS® (Review of Airway Dys-function and Interdisciplinary Community-based care in Adult Long-term Smokers) trial in Australian general practices.

Methods: Current/ex-smokers (aged ≥40 years, ≥10 pack years, including those with an existing diagnosis of COPD) have been recruited from 38 general practices across Melbourne. Following case-finding using the COPD-6® (Vitalograph Inc., Kansas, USA), those with FEV1/FEV6 ≤0.75 underwent spirometry. COPD diagnosis was made based on spirometry (FEV1/FVC < 0.7), COPD assessment test (CAT) score and modified Medical Research Council (mMRC) dyspnoea scale grade.

Results: Of 990 participants, 201 (20%) individuals had an existing diagnosis of COPD. Spirometry referrals have been made for 396 individuals (40%). Spirometry has been performed in 335, of whom 238 (71%) have been confirmed as having COPD. Only 105 (52%) of those currently managed as having COPD had spirometric confirmation. COPD-6® (FEV1/FEV6 ≤0.75) testing had a positive predictive value of 70% for COPD.

Conclusion: Case-finding of at risk patients using handheld devices measuring FEV1/FEV6, and subsequent spirometry testing revealed continuing under-diagnosis of COPD. Diagnostic spirometry is essential in avoiding misdiagnosis of COPD in Australian primary care.

Grant Support: NHMRC, Cyril Tonkin Scholarship, Boehringer Ingelheim, Lung Foundation of Australia, Eastern Melbourne PHN
FEASIBILITY OF A PEER-LED ASTHMA AND SMOKING PREVENTION PROGRAM (ASPP) IN AUSTRALIAN SCHOOLS WITH HIGH INDIGENOUS YOUTH

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Introduction/Aim: The high global burden of asthma and tobacco smoking among Indigenous people may potentially be reduced by appropriate interventions that target prevention of tobacco smoke uptake and improved asthma management. We undertook a pilot study in two Darwin schools with a high proportion of Indigenous youth to determine the feasibility of an innovative, peer-led, school-based education program called the Asthma and Smoking Prevention Project (ASPP). A subset of children with reported persistent respiratory symptoms were also clinically evaluated to determine the lower airway inflammatory profile and optimise asthma management.

Methods: The ASPP is founded on an evidence-based program to improve asthma management and prevent the uptake of tobacco smoking. The program uses a student-centred approach in which senior students (Peer Leaders) deliver the ASPP to Grade 7 students using activities, videos and games. Students completed questionnaires related to asthma and smoking at baseline and 3-months after program delivery. Students with respiratory symptoms at 3 months were invited for a comprehensive clinical evaluation and tests including sputum induction.

Results: The ASPP was well received. Of the 203 students involved, 56 (28%) were Indigenous and 70% completed baseline and follow-up questionnaires. Self-reported asthma was high (19%), 10% of students reported smoking and 63% reported exposure to tobacco at home. Of the 22 students who were clinically evaluated, 41% were Indigenous. Clinically important airway inflammation was high; 23% had Fractional Exhaled Nitric Oxide levels ≥35ppb, 88% had airway neutrophilia (>15%) and 29% had airway eosinophilia (>2.5%). Optimisation of medication and management was required in 59% of students.

Conclusion: Our study demonstrates the feasibility of implementing a school-based prevention and intervention program for at-risk groups. The high prevalence of clinically important airway inflammation and sub-optimal asthma management highlights the need for a community-based study on persistent respiratory symptoms in adolescents to reduce the burden of chronic lung disease particularly for Indigenous Australians.

Grant Support

RESPONSE OF AIRWAY EPITHELIAL CELLS TO RHINOVIRUS IN AN ALLERGIC CYTOKINE ENVIRONMENT

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Introduction/Aim: Rhinovirus is the most common trigger of acute exacerbations of allergic asthma. The airway epithelium is the target of rhinovirus infection, and is an important source of pro-inflammatory and anti-viral mediators. Airway epithelial cells (AECs) from patients with allergic asthma exhibit a pro-inflammatory phenotype. We aimed to characterise how the Th2 cytokine environment in allergic asthma alters the response of AECs to rhinovirus and to investigate the signalling pathways involved.

Methods: Cells of the well-differentiated human airway epithelial line, Calu-3, were cultured with or without the Th2 cytokines IL-4 and IL-13 for 48 hours, then stimulated with Poly I:C (TLR3 agonist) or imiquimod (TLR7 agonist), or infected with rhinovirus16. Quantitative real-time PCR and nCounter assays were used to characterise changes in expression of mRNA for pro-inflammatory and anti-viral mediators as well as viral pattern-recognition molecules.

Results: In the allergic cytokine environment, enhanced expression of mRNA for the pro-inflammatory cytokines IL6 and IL8 was induced by Poly I:C and imiquimod, while expression of anti-viral response genes IFIT1, STAT1, IRF7 and IFIH1 was induced only by Poly I:C. Following rhinovirus infection, Calu-3 cells pre-treated with Th2 cytokines exhibited significantly higher expression of IL6, IL8, CXCL10, CCL5, IL32 and CFB. Expression of IFNB1 and IFNL2/3 was either unchanged or modestly increased in cells pre-treated with Th2 cytokines. These alterations were accompanied by increased expression of the pattern recognition receptor genes TLR3, IFIH1, DDX58, and also of ICAM1, the cell surface receptor for rhinovirus.

Conclusion: Th2 cytokines appear to promote increased production of pro-inflammatory mediators following rhinovirus infection. Furthermore, any impairment of anti-viral host defence observed in allergic asthmatics is unlikely to be due to the effects of Th2 cytokines. Increased viral entry, together with enhanced signalling via TLR-3, MDA-5, and RIG-I, may explain the exaggerated inflammatory response to rhinovirus infection observed in patients with allergic asthma.

Conflict of Interest: No
Grant Support: Nil
SEX DEPENDENT EFFECTS ON AIRWAY RESPONSIVENESS OF MATERNAL HYPOXIA-INDUCED INTRAUTERINE GROWTH RESTRICTED JUVENILE MICE

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Aim: Previous studies have shown that intrauterine growth restriction (IUGR) is associated with asthma in children, however the reason is unknown. In early childhood, the prevalence of asthma is greater in males than females. The aim of this study was to investigate the impact of maternal hypoxia-induced IUGR on airway hyperresponsiveness in male and female juvenile mice.

Methods: Pregnant female BALB/c mice were housed under hypoxic conditions (10.5% O2) from gestational days E11-E17.5 (IUGR group; gestation period = 21 days). Following hypoxic exposure, mice were returned to a normoxic environment (21% O2). A second group of pregnant mice were housed under normoxic conditions throughout pregnancy (Control group). Weights of offspring were recorded until 2 weeks of age at which point responsiveness to methacholine and thoracic gas volume (TGV) were assessed.

Results: The IUGR offspring were lighter at birth compared with Controls, but not at 2 weeks of age. There were no differences in snout-vent length or abdominal circumference between groups at 2 weeks of age. At 2 weeks, airway resistance after methacholine challenge was greater in male IUGR offspring compared with Controls, but not in female IUGR offspring compared with Controls. In contrast, while there was no difference in peripheral lung mechanics between male IUGR offspring and Controls, IUGR female offspring had increased tissue damping and elastance after methacholine challenge, compared with Controls. There was no difference in TGV between groups.

Conclusion: Maternal hypoxia induced IUGR offspring were smaller at birth but exhibited ‘catch up’ growth by 2 weeks of age. The physiologically consequences of IUGR in the juvenile period were sex dependent: airway hyperresponsiveness in male offspring, and impaired peripheral mechanics in female offspring. Sexual dimorphism in the response to IUGR may contribute to differences in the prevalence of asthma between males and females in early childhood.

Grant Support: NHMRC, Asthma Foundation WA.

Declaration of Interest Statement: None.
MEASURING BRONCHIAL HYPERRESPONSIVENESS USING A MORE SOPHISTICATED BIOSTATISTICAL METHOD: THE LINEAR MIXED EFFECTS MODEL

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Introduction/Aim: Bronchial hyperresponsiveness (BHR) is typically measured using a dichotomous variable or the log transformed dose response slope (logDRS), both of which lead to loss of information. Furthermore, regression of the logDRS does not consider the initial FEV1 and the resulting estimates are difficult to interpret. In contrast, linear mixed models (LMM), a standard biostatistical tool that appears to have never been used for BHR, overcome these limitations, using all the data collected and producing more intuitive estimates. We aimed to test the utility of modelling BHR using a LMM on associations with known risk factors. Regression of the logDRS was used as a comparison.

Methods: In 2010–2012, a subsample (n=687) of the 1961-born Tasmanian Longitudinal Health Study (TABS) cohort were assessed by questionnaire, spirometry and methacholine challenge. Repeated FEV1 measurements were modelled using a dichotomous variable or the log transformed dose (Female=1). BHR was associated with sex, height and former smoking were not.

Results: In both models current smoking, asthma status, and age were associated with BHR, sex, height and former smoking were not.

<table>
<thead>
<tr>
<th>Predictor</th>
<th>β-coefficient (95%CI)</th>
<th>p-value</th>
<th>β-coefficient (95%CI)</th>
<th>p-value</th>
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</thead>
<tbody>
<tr>
<td>Sex (Female=1)</td>
<td>−2.54 (−10.4,5.36)</td>
<td>0.53</td>
<td>0.14 (−0.11,0.39)</td>
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<tr>
<td>Smoking</td>
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<td></td>
</tr>
<tr>
<td>-Former</td>
<td>0.68 (−5.15,6.51)</td>
<td>0.82</td>
<td>0.01 (−0.17,0.19)</td>
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<tr>
<td>-Current</td>
<td>−14.1 (−22.0,−6.31)</td>
<td>&lt;0.001</td>
<td>0.06 (0.016,0.50)</td>
<td>0.037</td>
</tr>
<tr>
<td>Asthma</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-Remitted</td>
<td>−13.3 (−19.8,−6.83)</td>
<td>&lt;0.001</td>
<td>0.43 (0.23,0.64)</td>
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<tr>
<td>-Current</td>
<td>−36.9 (−44.1,−29.6)</td>
<td>&lt;0.001</td>
<td>1.16 (0.95,1.37)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Height</td>
<td>−0.17 (−0.63,0.29)</td>
<td>0.46</td>
<td>−0.01 (−0.02,0.01)</td>
<td>0.21</td>
</tr>
<tr>
<td>Age</td>
<td>−10.0 (−14.5,−5.52)</td>
<td>&lt;0.001</td>
<td>0.26 (0.12,0.40)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Conclusion: A LMM can be used to assess risk factors for BHR. By incorporating all data, it is likely to be more sensitive and provide more statistical power to identify risk factors.

Grant Support: NHMRC, Clifford Craig Medical Research Trust

MECHANICAL FORCES ATTENUATE ANTI-VIRAL IMMUNITY IN PRIMARY HUMAN AIRWAY EPITHELIAL CELLS FROM ASTHMATIC DONORS

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Introduction/Aim: Bronchoconstriction (BC) is a major physiological event in asthma, leading to clinical symptoms and the generation of mechanical force within the airway. Asthma is characterised by periods of good health, interrupted by exacerbations. Most asthma exacerbations are caused by respiratory viruses, including rhinovirus (RV). During a viral exacerbation of asthma, BC and viral infection occurs simultaneously; these factors may interact in the airway. This study aims to investigate the role of mechanical forces on innate anti-viral immunity during RV infection in primary human airway epithelial cells.

Methods: Primary airway epithelial cells from asthmatic donors (n=5) were collected at bronchoscopy and air liquid interface (ALI) cultures established. After 21 days, cell were infected with RV1B at a multiplicity of infection of 0.001 for 6 hours. Infected and control cells were then exposed to apical compression (30cm H2O pressure) using 5% CO2 in air for 10 minutes every hour for 96 hours. Cells were harvested at 24-hour intervals. Samples were analysed for viral and interferon (IFN) RNA, and IFN, transforming growth factor-β (TGF-β) and mucus proteins.

Results: Compression induced the release of TGF-β protein across all time points (At 96 hours post infection, control(Mean±SEM): 302±120pg/ml, compressed:1726±909pg/ml). Compressed cells produced less type-I and type-III IFNs following RV1B infection (At 96 hours, IFNb virus alone(Mean±SEM):173±89pg/ml, compression-virus 58±17pg/ml and INFl virus alone: 603±280 (SEM) pg/ml, compression+ virus 125±35pg/ml. Compression also induced an increase in viral replication.

Conclusion: We demonstrate that mechanical forces similar to those induced during bronchoconstriction in vivo induce production of TGF-β and impair innate anti-viral immunity in primary human airway epithelial cells. This is the first time that mechanical forces have been shown to impact on innate immunity and the data have direct relevance to human disease.

Grant Support: University of Newcastle post graduate scholarship, The Singleton Foundation.

Conflict of Interest: Nil
POTENTIAL PROTECTIVE ROLE OF microRNA-22 AGAINST AIRWAY REMODELLING

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Introduction/Aim: The epithelium of asthmatics is differentially dysregulated, which may allow repair but not proper regeneration after noxious stimuli (e.g. viruses and allergens) leading to further remodelling. The influenza virus H1N1 is one of the major causes of asthma exacerbations. Differentiation specific microRNAs; e.g. miR-22, may be key epigenetic factors involved in aberrant epithelial cell differentiation of asthmatics upon exposure to H1N1. We hypothesise that the impaired miR-22 expression in epithelium of asthmatics after H1N1 infection may be the link between abnormal differentiation and innate immune dysregulation in these cells. Our aims are to determine the role of miR-22 in primary bronchial epithelial cells of severe asthmatics by determining its expression in these cells. Our aims are to determine the role of miR-22 in primary bronchial epithelial cells of severe asthmatics by determining its expression in these cells.

Methods: Primary bronchial epithelial cells (pBEC) from severe asthmatic and non-asthmatic subjects were cultured under air-liquid-interface (ALI) condition. Cells were incubated with H1N1 (MOI 5). miRNAs and mRNA were isolated using RNAeasy mini kit with some modifications and subjected to Tagman miR-22 and mRNA assays.

Results: pBEC from non-asthmatics and asthmatics infected with H1N1 showed the same viral titers. In cells from non-asthmatics, miR-22 expression decreased 6h after infection but then increased and recovered to basal levels by 24h post infection. In cells from asthmatics, miR-22 expression remained unchanged during infection. Expression of the miR-22 targets, CD147 (MMP inducer) and HDAC4 (epigenetic regulator) mRNA were decreased during infection in cells from non-asthmatics. However in cells from asthmatics, CD147 expression increased during infection and HDAC4 remained unchanged.

Conclusion: Despite similar level of infection, miR-22 expression differs in epithelial cells from asthmatics and non-asthmatics during H1N1 infection. The reduction of CD147 and HDAC4 during H1N1 infection in cells from non-asthmatics may represent a self-defence mechanism against further cellular remodelling.

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Grant Support

ACUTE INHIBITION OF NOTCH SIGNALLING ABLATES MUC5AC PRODUCTION IN HUMAN AIRWAY EPITHELIAL CELLS FROM ASTHMATIC, NON-ASTHMATIC AND COPD DONORS.

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Introduction: Mucus overproduction in patients with asthma and COPD is linked to increased hospitalisations and morbidity. The Notch family of receptor proteins regulate airway mucus production via differentiation mechanisms that direct cells towards secretory or ciliated morphology. However little is known about Notch’s role during mucus production, following differentiation.

Aim/Hypothesis: To examine the impact of inhibiting Notch signalling on mucus production in fully differentiated primary bronchial epithelial cells (pBECs) from control subjects as well as patients with asthma or COPD. We hypothesize that Notch inhibition will downregulate mucus production.

Methods: pBECs from 4–5 donors of each cohort were grown at air-liquid interface (ALI) culture for 25 days to promote multicellular differentiation. At this time, cells were treated with dibenzazepine (DBZ), a potent inhibitor of Notch signalling for a further 96h during which, apical lining fluid was collected every 24h for assessment of Muc5AC release. At the completion of the experiment, samples were also collected for protein, mRNA and histological analysis.

Results: DBZ treatment significantly reduced MUC5AC expression and release in all phenotypes as assessed by qPCR, ELISA and immunofluorescence. Western blotting/qPCR revealed significant reduction of NOTCH3 intracellular domain (NICD3) and Notch3 mRNA in pBEC from all donor phenotypes. Reduced expression of NICD1 was also observed, but was restricted to cells from non-asthmatic following treatment. Finally, the goblet cell marker protein CLCA1 was unchanged across all treatments.

Conclusion: Notch inhibition reduced MUC5AC expression and secretion from differentiated pBECs, independent of goblet cell number. This reduction is NOTCH3 dependent and occurs in pBECs from control, asthmatics and patients with COPD, suggesting that Notch regulates MUC5AC production independent of secretory cell differentiation. The decrease of NICD1 in pBEC from control donors suggests additional regulation of MUC5AC production that may be compromised in cells from COPD or asthma patients.

Grant Support: NHMRC project grant #1064405.
CIGARETTE SMOKE-INDUCED INFLAMMATION IN MACROPHAGES: INVOLVEMENT OF THE SPHINGOSINE-1-PHOSPHATE (S1P) SIGNALLING
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Introduction/Aim: There have been conflicting reports on the possible role of NLRP3 inflammasome activation in IL-1-driven inflammation in COPD. We hypothesized that in COPD and/or in response to cigarette smoke, there is increased activation of the NLRP3 and/or alternative AIM2 inflammasomes, regulated via sphingosine-1-phosphate (S1P) signalling.

Methods: Human primary alveolar macrophages and THP-1-derived macrophages were analysed for release of IL-1β (ELISA) and its cleavage-activation in response to cigarette smoke extract (10%, 24h). Expression and localization of IL-1β, NLRP3, AIM2, ASC (pro-caspase-1 recruiter), and cleaved caspase-1 were assessed by immunofluorescence/confocal microscopy and/or Western blot. The effects of the caspase-1 inhibitor ZYVAD-fmk or NLRP3 antagonist glyburide, and treatment with S1P or the S1P receptor regulator FTY720, with or without exposure to cigarette smoke extract were investigated.

Results: In alveolar macrophages, cigarette smoke induced increased intracellular expression of IL-1β (~15%, p<0.05 on pooled data from 3 donors), with particulate staining for cleaved IL-1β increased both inside and outside the cell (5 fold, p<0.001 on pooled data). These particles were partially co-localized with particulate NLRP3 or AIM2 which were also increased. In THP-1 macrophages, cigarette smoke induced similar NLRP3/cleaved IL-1β complexes, which were inhibited by 20μM ZYVAD-fmk or 20μM glyburide (p<0.05). At 10nM, S1P or FTY720 significantly protected THP-1 macrophages from cigarette smoke-induced activation of IL-1β.

Conclusion: IL-1β-driven inflammation in cigarette smoke-exposed macrophages is initiated by both NLRP3 and AIM2 inflammasomes, the former regulated by S1P signalling system, which suggests novel therapeutic targets in COPD.

EVALUATING ANTI-CD20 THERAPY FOR STAT3-MEDIATED LUNG FIBROSIS
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Introduction: The STAT3 signalling pathway has recently been implicated in the pathogenesis of Idiopathic Pulmonary Fibrosis (IPF).

Hypothesis: We hypothesise that the pro-fibrotic effects of STAT3 involve B cell-mediated immune regulation.

Methods: We have analysed immune cell composition in human lung biopsy tissue and examined the effect of B cell depletion on bleomycin-induced lung fibrosis in vivo.

Results: A trend towards increased B-cell activating factor, APRIL and CXCL13 are observed in IPF patient serum versus age match controls. In addition we observed an increase in the number of mature B cells in the lungs of IPF patients. Genetic depletion of B cells in gp130757F; Stat3−/−/− attenuated bleomycin-induced fibrosis. The therapeutic potential of depleting follicular B cells using anti-CD20 treatment was assessed. Mice were given two 100 μg doses of anti-CD20 antibody (provided by Genentech Inc USA), or IgG2a isotype control i.p. 7 days prior to and 7 days after bleomycin, and the extent of fibrosis measured 21 days after the last dose. FACs analysis of blood taken on days 0, 7 and 28 days post-bleomycin-treatment revealed an almost complete depletion of CD19+ and B220+ B cells. However, the extent of fibrosis, assessed using micro-CT imaging and HPLC analysis of hydroxyproline levels, was not significantly different between treatment groups.

Conclusion: Although antibody depletion of follicular B cells had no effect on bleomycin-induced fibrosis, residual B cells remained in the lung of these mice. Current studies are analysing B cell subsets in fibrotic lung tissue from mice and IPF patients.

Grant Support: This work is funded by NHMRC Project Grant GNT1067511 and a British Lung Foundation Priming Grant.
INTERACTION OF DIETARY FATTY ACIDS WITH OBESITY-INDUCED CYTOKINES IN PRIMARY PULMONARY FIBROBLASTS

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Introduction: Obesity is an important risk factor for developing severe asthma. Dietary fatty acids, increased in the serum of obese people, activate systemic innate immune responses. Furthermore, we have shown that a single high fat meal increases airway inflammation and impairs bronchodilator recovery in asthma.

Aim: The aim of this study was to investigate whether dietary fatty acids directly cause inflammation and/or synergise with obesity-induced cytokines in human pulmonary fibroblasts and to elucidate underlying mechanisms.

Methods: Primary human pulmonary fibroblasts were incubated with BSA-conjugated fatty acids for 24hr, before stimulation with TNF-α for another 24hr. IL-6 and CXCL8 release was measured using ELISA. IL-6 acts as a general marker for inflammation and CXCL8 is a potent neutrophilic chemoattractant. The following fatty acids were used: arachidonic acid (AA, ω-6), eicosapentaenoic acid (EPA, ω-3), and palmitic acid (PA, saturated fatty acid) at increasing concentrations of 1, 10 and 100μM.

Results: AA induced substantial IL-6 (n=11, P<0.05) and CXCL8 (n=9, P<0.0001) release by pulmonary fibroblasts. EPA and PA did not induce CXCL8 or IL-6 release. Stimulation with the combination of AA and TNF-α resulted in greater IL-6 (n=9, P < 0.0001) and CXCL8 (n=8, P <0.0001) release than AA alone. The effect of the combination AA/TNF-α on IL-6 release was greater than the sum of the individual effect of AA and TNF-α, indicating synergy.

Conclusion: These findings suggest that dietary fatty acids are important modulators of inflammatory responses and that there is an interaction between arachidonic acid and TNF-α, resulting in a synergistic inflammatory response in pulmonary fibroblasts. This could indicate that obese asthmatics compared to lean individuals, are more prone to airway inflammation after a high fat meal.

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NOVEL ROLE OF INFLAMMASOMES IN THE MOLECULAR PATHOGENESIS OF EMPHYSEMA

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Introduction/Aim: Innate immune responses within the lung mucosa, for instance against cigarette smoke (CS), depend on the activation of pattern recognition receptors via multiprotein complexes called inflammasomes, which facilitate the maturation and release of the pro-inflammatory cytokines interleukin (IL)-1β and IL-18. In recent years a plethora of data from clinical studies and mouse disease models have demonstrated that excessive inflammasome activation promotes several diseases. In that regard, emphysema a major debilitating components of COPD, has shown a link between IL-1β and IL-18 and disease. Despite these observations, a causal role for specific inflammasomes in emphysema is unknown. Therefore we aim to reveal a novel functional link between inflammasome activation and the pathogenesis of emphysema.

Methods: To identify whether inflammasomes promote emphysema, we have utilised a genetic model (gp130–/–) for spontaneous emphysema and an acute (4 days) CS-induced model, both characterised by elevated alveolar type II cell (ATII) apoptosis that is driven by the cytokine IL-6.

Results: Here, we demonstrate that lung tissues from emphysema patients, as well as from spontaneous (gp130–/– mice) and CS-induced emphysema mouse models, are characterized by excessive productions of the gene encoding AIM2 which forms a cytoplasmic DNA-sensing inflammasome and IL-1β (but not IL-18) protein. Furthermore, genetic blockade of Aim2 in gp130–/– mice prevents the development of emphysema by suppressing augmented ATII cytokinemia. A positive correlation also exists between elevated IL6 and Aim2 mRNA levels, and Aim2 mRNA and IL-1β protein expression levels, in emphysema patients.

Conclusion: Collectively, we define for the first time that hyper-activation of the endogenous IL-6/Aim2/IL-1β axis in the lung augments ATII apoptosis, which in turn causes emphysema. Finally, we believe our study also has the potential to aid future therapeutic strategies to target inflammasomes (AIM2) alone or in combination with IL-6 in human emphysema.

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STEREOID RESISTANT CD8+CD28nullNKT-LIKE PRO-INFLAMMATORY CYTOTOXIC CELLS IN COPD
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Introduction: NKT-like cells represent a bridge between innate and adaptive immunity. We have reported increased numbers of steroid resistant cytotoxic/pro-inflammatory senescent CD28nullNKT-like cells in BAL in COPD associated with increased cytopoietic to bronchial epithelial cells. These cells express drug efflux pump Pgp-1 and loss of glucocorticoid receptor (GCR). Recently, loss of histone deacetylase 2 (HDAC2) and heat shock protein 90 (Hsp90) have been reported from CD8+ T cells from COPD patients and we hypothesized these molecules may be decreased in CD28nullNKT-like cells (particularly the CD8+ subset).

Method: Blood was collected from a group of COPD patients and aged matched controls and expression of CD28, Pgp-1, GCR, HDAC2, Hsp90 and pro-inflammatory cytokines determined in CD8+ and CD8-NKT-like cells in the presence of 1μM prednisolone, low-dose cyclosporine A (binds to GCR-Hsp90 complex) and heat shock protein 90 (Hsp90) have been reported from CD8+ T cells from COPD patients and we hypothesized these molecules may be decreased in CD28nullNKT-like cells (particularly the CD8+ subset).

Results: Loss of GCR, HDAC2 and Hsp90 (but not Pgp-1) expression was identified from CD28nullCD8+NKT-like cells compared with CD28+ counterparts. Loss of GCR, HDAC2 and Hsp90 was associated with increased production of IFNγ and TNFα and increased steroid resistance. Uregulation of GCR, HDAC2 and Hsp90 was noted in the presence of prednisolone + low dose cyclosporine A and increased HDAC2 was noted in the presence of prednisolone + cyclosporine A.

Conclusion: Steroid resistance in pro-inflammatory CD28nullCD8+NKT-like cells is associated with multiple mechanisms. Combination prednisolone, low-dose cyclosporine A and theophylline treatment therapy inhibits pro-inflammatory cytokine production from these cells and may reduce systemic inflammation in COPD.

Grant Support: NHMRC

Declaration of Interest: No declaration of interest

EFFECT OF TIOTROPIUM AND OLODATEROL, ALONE AND WITH EXERCISE TRAINING, ON EXERCISE ENDURANCE IN COPD
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Introduction/Aim: Physical deconditioning is common in patients with COPD, limiting exercise tolerance. PHYSACTO® (NCT02085161) tested the effects of bronchodilators alone or with exercise training (ExT), combined with a standardised physical activity self-management behaviour-modification (BM) programme, on exercise endurance time (EET) in patients with COPD.

Methods: A 12-week randomised, partially double-blind, placebo (P)-controlled, parallel-group trial at 34 sites in Australia, New Zealand, USA, Canada and Europe. Interventions (all with 12-week BM): P; tiotropium (T) 5μg; T + olodaterol (T+O) 5/5μg; T+O 5/5μg with 8 weeks’ ExT (T+O 5/5μg + ExT). EET (log transformed) during an endurance shuttle-walk test (ESWT) to symptom limitation was assessed after 8 weeks (primary end point) and 12 weeks.

Results: 303 patients (200 men) were randomised and treated (full analysis set n=274). Mean post-bronchodilator FEV1 was 1.59 L (56.7% predicted), EET significantly increased with T+O 5/5μg and T+O 5/5μg + ExT versus P at 8 weeks (Figure); 13 patients reached test termination criteria (20 minutes) without symptom limitation (P, n=0; T, n=3; T+O, n=2; T+O + ExT, n=8) at 8 weeks. No safety concerns were identified.

Conclusion: T+O 5/5μg, alone and combined with ExT, improved EET during ESWT compared to P in moderate to severe COPD.
Grant Support: The study was funded by Boehringer Ingelheim

Declaration of interest statement: PF has received in the past 5 years honoraria for educational and advisory board involvement and/or received conference attendance support for the following: Global Initiative for COPD (GOLD), Improvement Foundation, Lung Foundation Australia, and Remedy Healthcare; AstraZeneca, Boehringer Ingelheim, CSL-Behring, GlaxoSmithKline, Menarini, MundiPharma, and Novartis. TT is the Principal Investigator of PROactive project and received consultancy fees from Boehringer Ingelheim, Novartis, and Bayer. JB received research funding via the Research Institute of the McGill University Health Centre from AstraZeneca, Boehringer Ingelheim, GlaxoSmithKline, Merck, Novartis, Nycomed, Pfizer and Theratechneologies; and has served on speakers, consultation panels and/or advisory boards for these pharmaceutical companies. FM is a consultant for Boehringer Ingelheim and served on advisory boards for Boehringer Ingelheim, GlaxoSmithKline, and Pfizer and has received payment for lectures including service on speaker bureaus from Boehringer Ingelheim, GlaxoSmithKline, Nycomed and Pfizer. NL is an employee of Evidera and works with a variety of companies and organizations and expressly prohibited from receiving any payment or honoraria directly from these organizations for services rendered. DE, DDS, LK and AH are employees of Boehringer Ingelheim. WJ served on advisory boards and/or speaker bureaus for Astra Zeneca, Boehringer Ingelheim, Novartis and GSK. KLL received grants/research support from GSK, consulting fees from Schering-Plough and Merck Frosst and served on speaker bureaus/honoraria for GSK, Astra-Zeneca, Pfizer, Merck Frosst, Air Liquide and Health International.

Declaration of interest statement: Peter Frith has received in the past 5 years honoraria for educational and advisory board involvement and/or received conference attendance support for the following: Global Initiative for COPD (GOLD), Improvement Foundation, Lung Foundation Australia, and Remedy Healthcare; AstraZeneca, Boehringer Ingelheim, CSL-Behring, GlaxoSmithKline, Menarini, MundiPharma, and Novartis.

RESPONSIVENESS OF TWOMINUTE WALK DISTANCE AFTER PULMONARY REHABILITATION IN CHRONIC OBSTRUCTIVE PULMONARY DISEASE

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Aim: Field exercise tests (e.g. six minute walk distance [6MWD]) are important measures of functional exercise capacity in people with COPD. Shorter tests such as the two-minute walk distance may offer advantages in some populations, but lack information about responsiveness to change. This study examined reliability, validity and responsiveness of the two-minute walk test in people with COPD attending pulmonary rehabilitation (PR).

Methods: At pre-PR assessment, study participants completed a two-minute walk test twice in addition to usual measures (6MWD, Chronic Respiratory Questionnaire). At post-PR assessment following a standard PR program, measures were repeated and global rating of change scores obtained (patient and therapist). Pre-post program change scores were examined for correlations with change in two-minute walk distance and used (where r>0.3) to estimate the minimal importance difference through anchor-based methods. Distribution-based estimates based on standard error of measurement were examined. Test-retest reliability (ICC, Bland Altman agreement) and validity (Pearson correlation with 6MWD) were reported.

Results: Pre-program assessment was conducted in 59 participants (68±10 years, FEV1%pred=48±20%). Test-retest reliability of two-minute walk distance was high (ICC=0.985) with mean difference between trials of 2.4m (95%CI 0.7 to 4.0, p=0.006). Two and six-minute walk distance was highly correlated (r=0.87, p<0.01). Post-program assessment was completed in 36 patients (69±8 years, FEV1%pred=52±21%). Anchored against clinically meaningful change in 6MWD, the minimal important difference in two-minute walk distance was 3.5m (area under curve=0.82, 95%CI 0.67 to 0.97) and agreed with the distribution-based estimate of 3.4m. Two-minute walk distance and 6MWD did not significantly improve post-program in this sample, in contrast to QOL measures.

Conclusion: Improvement in two-minute walk distance of at least 3.5m following a PR program corresponded to a clinically meaningful change. Due to a learning effect between first and second trials a practice two-minute walk test is recommended.

Grant Support: Nil
NO ADDITIONAL BENEFIT FOR 6MWT OR HADs IN COMBINING COGNITIVE BEHAVIOURAL THERAPY AND PULMONARY REHABILITATION

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Introduction/Aim: Controlled trials of psychological interventions in people with chronic obstructive pulmonary disease (COPD) generally target generalised anxiety and depression. This randomised controlled trial aimed to determine whether adding a cognitive behavioural therapy program for the sensation of breathlessness (CBT-BREVE) to a comprehensive pulmonary rehabilitation program (CPRP) significantly improved health outcomes beyond those achieved with a CPRP alone.

Method: Eight week CPRP cycles at the Repatriation General Hospital were block randomised to include CBT (BREVE) or social group (CPR) interventions. People with COPD and at least moderate airflow obstruction (FEV1 < 80% pred, FEV1/FVC < 70%, GOLD Grade ≥ 2) were eligible for inclusion. Primary (six minute walk test – 6MWT, Hospital Anxiety and Depression scale –HADs) and secondary outcomes (Multidimensional Dyspnoea Profile -MDP, Chronic Respiratory Questionnaire- CRQ, sedentary and physical activity (accelerometry -Actigraph GT3X+) and symptom scores) were assessed one month before and one, six and 12 months post intervention. Participant feedback was sought after completion of the 12 month assessment. Differences in clinical outcomes between groups (intention to treat) were assessed across all four assessments via Chi2 test.

Results: 101 participants met all inclusion criteria with no significant (p<0.001) differences between subjects eligible for participation but declining (n=66 GOLD grade ≥ 2) and those participating in the trial (n=101 mean age 70.1 ± SD 8.5, 54 males, FEV1 % pred 47.7 ± 16.3). Preliminary analysis indicates no statistical or clinically significant differences between groups at any assessment point for 6MWT or HADs. Analysis of secondary outcomes is ongoing. Participant feedback was uniformly positive for both forms of intervention.

Conclusion: Despite overwhelmingly positive feedback for the CBT program, combining CPR with a CBT program specific for sensation of breathlessness did not result in greater improvements in functional exercise capacity, anxiety or depression, beyond those achieved with standard CPR.

Key Words: Pulmonary rehabilitation, chronic obstructive pulmonary disease, cognitive behavioural therapy

Grant Support: National Health and Medical Research Council Project Grant (# 1010309) Clinical Trial registration: ANZCTR12611000292976

ENHANCEMENT OF TRAINING RESPONSE FOLLOWING PULMONARY REHABILITATION WITH DOWNHILL WALKING: A RANDOMISED CONTROLLED TRIAL

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Introduction/Aim: Downhill walking (DW) enhances quadriceps contractile muscle force (CMF) with less symptoms than level walking (LW) in patients with COPD. This study sought to determine the effectiveness of DW compared to LW as a 12-week comprehensive pulmonary rehabilitation program in patients with COPD.

Methods: 39 COPD patients (62±9 yrs; FEV1 49±17%pred) were randomised to PR with DW or LW. Exercise capacity (6-minute walk test, 6MWT [primary outcome]; cycle endurance test), muscle force and quality of life were assessed before and after program completion. Training responses and the proportion of patients exceeding the 30m minimally important difference (MID) for 6MWT were compared between groups via Chi2 test.

Results: The magnitude of improvement was similar across all outcomes in both groups (Table 1). However, a significantly greater proportion of patients in DW exceeded the 6MWT MID compared to LW (94% vs 65%, p=0.03).

Table 1. Training responses in LW and DW.

<table>
<thead>
<tr>
<th>Outcome</th>
<th>LW (n=17)</th>
<th>DW (n=22)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>∆ 6-minute walk test (metres)</td>
<td>+67 [22–103]</td>
<td>+62 [45–98]</td>
<td>0.52</td>
</tr>
<tr>
<td>∆ Cycle endurance test (seconds)</td>
<td>+260 [55–665]</td>
<td>+740 [85–885]</td>
<td>0.16</td>
</tr>
<tr>
<td>∆ Muscle Force (NM)</td>
<td>+29 [13–48]</td>
<td>+17 [5–26]</td>
<td>0.20</td>
</tr>
<tr>
<td>∆ CRQ (total points)</td>
<td>+12 [10–22]</td>
<td>+20 [10–27]</td>
<td>0.47</td>
</tr>
</tbody>
</table>

Conclusion: These results support the use of downhill walking as a useful adjunct in pulmonary rehabilitation programs for patients with COPD.

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THE EFFECT ON HRQoL OF ONGOING FEEDBACK DURING A MAINTENANCE WALKING PROGRAM: AN RCT

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Introduction/Aim: This study evaluated the effect on health-related quality of life (HRQoL), endurance exercise capacity and daily physical activity of the addition of ongoing feedback (Intervention Group [IG]) versus no feedback (Control Group [CG]), to a 12-month unsupervised maintenance walking program which followed a supervised walking training program in people with COPD.

Methods: Participants were randomised at baseline to the IG or CG. Both groups completed the same 2-month supervised, walking training program followed by a 12-month unsupervised maintenance walking program during which the IG received ongoing feedback (telephone calls, biofeedback and progressive goal setting) and the CG received no feedback.

Results: Seventy-five of 95 (79%) participants who entered the main maintenance program [mean (SD); age 69 (8) yrs.; FEV1 43% predicted (15)] completed the study. There were no between-group differences in HRQoL, endurance exercise capacity or daily physical activity levels on completion of the 12-month maintenance program when compared to baseline measures (St George's Respiratory Questionnaire [SGRQ] total score mean difference (MD): 2 point, 95% CI −4 to 8; Endurance Shuttle Walk Test [ESWT] time: MD −96 seconds, 95% CI −253 to 61; Daily steps: MD 757 steps, 95% CI −103 to 1617) or compared to the 2-month assessment (following supervised training) (SGRQ total score MD: 4 points, 95% CI −2 to 10; ESWT time: MD −54 seconds, 95% CI −254 to 137; Daily steps: MD 617 steps 95% CI −435 to 1669).

Conclusion: This study demonstrated that following a 2-month supervised walking training program, ongoing feedback was no more effective than no feedback in maintaining HRQoL, endurance exercise capacity or daily physical activity levels in people with COPD during a 12-month unsupervised maintenance walking program.

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PARTICIPANTS WHO DO NOT ACHIEVE THE MID IN THE 6MWT MAY STILL HAVE A TRAINING-RESPONSE

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Introduction/Aim: The six minute walk test (6MWT) is a clinical tool widely used for evaluating functional exercise capacity. The primary outcome of the 6MWT is the six minute walk distance (6MWD). Current practice recognises the use of the minimally important difference (MID) as a key index of a response to pulmonary rehabilitation (PR). However, the 6MWT also provides information on dyspnoea, heart rate (HR) and degree of desaturation (nadir SpO2) achieved during the test. This information is typically not considered when evaluating patient outcome to PR. Whilst an individual may not achieve the MID, it is possible other indices from the 6MWT may reflect improvement. The aim of this study was to compare changes in peak dyspnoea, HR and nadir SpO2 following PR in participants that did not have a clinically significant change (non-MID) in their 6MWD (−30m≤MID≤30m) with a group that did achieve the MID (>30m).

Methods: Data was retrospectively analysed from subjects that completed a 6MWT both pre and post PR over a 2 year period. The 6MWT was measured prior to and immediately following an 8 wk, twice weekly PR program. Dyspnoea (0–10 Borg scale) and HR were recorded immediately at the end of the 6MWT while nadir SpO2 was measured during the test.

Results: 81 subjects did not achieve the MID (6MWD: 3 ± 17m) whereas 91 subjects achieved the MID (6MWD: 74 ± 42m). For the non-MID group, following PR subjects reported a lower dyspnoea (=0.7 ± 1.7, p=0.01) and lower nadir SpO2 (=1.3 ± 4.4%, p=0.01) whereas HR remained unchanged (=2.6 ± 13.5 beats.min−1, p=0.12). For the MID group, nadir SpO2 (=1.6 ± 4.7%, p<0.01) was lower following PR and there was a tendency for dyspnoea (=0.4 ± 1.9, p=0.053) to be lower. End exercise HR for the MID group remained unchanged (=0.2 ± 12.3 beats. min−1, p=0.9).

Conclusion: These data demonstrate the benefit of using all the data obtained from the 6MWT to evaluate the overall response to PR. Individuals who may not show a clinical improvement in 6MWD may be less breathless during the 6MWT suggesting an improved response to exercise. Whilst the 6MWD remains the primary outcome from the 6MWT, other exercise related indices from this test should be considered when evaluating overall response to an intervention.

Grant Support: Nil
TO-054

PATIENTS WITH X-LINKED RETINITIS PIGMENTOSA HAVE ALTERED AIRWAY CILIARY FUNCTION AND STRUCTURE

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Introduction:Aim: Cilia are tentacle-like extensions from cell bodies, which are motile or immotile depending on their function. The airways contain motile cilia to move mucus, whereas cilia in the eyes act as immotile sensory antennas. Ciliopathies are diseases caused by developmental defects in motile or immotile cilia. Relationships between motile ciliary diseases and immotile ciliary diseases have not been studied. Case reports have emerged that infer patients exist with a phenotype consistent with both diseases. We hypothesised that patients with a pre-diagnosed immotile ciliopathy – Retinitis Pigmentosa – will also have altered motile cilia

Methods: The inferior nasal turbinate was brushed for samples of airway epithelium from 12 patients (range 8–73 years) with Retinitis Pigmentosa. Epithelial strips of each patient were examined under high-speed videomicroscopy to capture ciliary waveform and beat frequency. Epithelium was also processed for electron microscopy and cilia were photographed. Ciliary structure and orientation was quantified from these photographs. A Mann–Whitney U test was used to analyse the difference in ciliary orientation between patients with and without Retinitis Pigmentosa.

Results: Ciliary beat frequency of patients with Retinitis Pigmentosa was normal (mean 8Hz). Ciliary waveform was uncoordinated in video playback. Patients with Retinitis Pigmentosa had periodic waveform segments of increased intensity and frequency compared to background breath sounds, mostly spanning expiration for a period of 0.03–1.2 seconds at frequencies of 100–1050Hz, and occasionally spanning shorter inspiratory segments. Recordings of patients with crackles revealed brief (6–20 millisecond) discontinuous sounds with a distinguishing waveform identifiable within them.

Conclusion: Digital breath sound analysis may be more sensitive than manual auscultation in detecting wheeze in our study. Patients with clinically-described wheeze had periodic waveform segments of increased intensity and frequency compared to background breath sounds, mostly spanning expiration for a period of 0.03–1.2 seconds at frequencies of 100–1050Hz, and occasionally spanning shorter inspiratory segments. Recordings of patients with crackles revealed brief (6–20 millisecond) discontinuous sounds with a distinguishing waveform identifiable within them.

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TO-055

DETECTION AND DEFINITION OF ABNORMAL PAEDIATRIC BREATH SOUNDS USING DIGITAL STETHOSCOPIES COMPARED TO STANDARD AUSCULTATION

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Introduction:Aim: The audio characteristics of abnormal paediatric breath sounds are not well-defined and digital techniques to capture them have not been widely investigated. Our study aimed to define the audio-logical features of abnormal paediatric breath sounds objectively and compare the ability of digital stethoscopes to detect them against auscultation using standard bell-and-diaphragm stethoscopes.

Methods: Twenty children with normal breath sounds, generalized wheeze or crackles, and some with a diagnosis of cystic fibrosis were auscultated by a paediatric consultant and digitally recorded using both the Littman™ 3200 Digital Electronic Stethoscope and a Clinicloud™ Digital Stethoscope. We used spectrographic analysis and recording playback to detect abnormal breath sounds and define their audio waveform characteristics.

Results: Digital stethoscopes were more sensitive than standard auscultation in detecting wheeze in our study. Patients with clinically-described wheeze had periodic waveform segments of increased intensity and frequency compared to background breath sounds, mostly spanning expiration for a period of 0.03–1.2 seconds at frequencies of 100–1050Hz, and occasionally spanning shorter inspiratory segments. Recordings of patients with crackles revealed brief (6–20 millisecond) discontinuous sounds with a distinguishing waveform identifiable within them.

Conclusion: Digital breath sound analysis may be more sensitive than manual auscultation in detecting breath sound abnormalities in children, with potential applications for improved diagnosis, data sharing and disease monitoring. Further research and development is needed.

Grant Support: Nil
THE EFFECT OF PARTICLE SIZE DELIVERED BY PRESSURIZED METER DOSE INHALERS (pMDIs) IN ASTHMATIC ADOLESCENTS

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Introduction/Aim: Spacer device use is currently recommended with aerosolised corticosteroids to reduce mouth and throat deposition produced by larger particle generating inhalers. Anecdotally, spacers are seldom used as prescribed, which has particularly been observed in the adolescent age group at Princess Margaret Hospital for Children in Perth. The recommendation to use a spacer may not be necessary with new asthma medications producing smaller particle sized aerosols. We aimed to assess the effect of particle size in pMDIs with and without a spacer.

Methods: Fourteen adolescents aged 13–17 years with mild stable asthma were randomised to use a pMDI with or without a spacer in a randomised cross-over study. Radiolabelled (Tc99m) corticosteroids of different mass median aerodynamic diameters (fluticasone propionate 3.5 μm (Flixotide®), and beclometasone dipropionate 1.1μm (QVAR®)) were inhaled using correct technique, as assessed by a Clinical Nurse Specialist, and dose of drug quantified immediately after with a 2D gamma camera scan. Radiation detected was quantified per region of interest accounting for attenuation by body tissues. Drug deposition in the lungs and orogastric regions was compared in adolescents inhaling the same drug with and without a spacer using a Wilcoxon matched pairs signed-rank test.

Results: Fourteen adolescents completed two visits each, with and without a spacer. We did not observe a significant difference in the QVAR group lung deposition with or without spacer (median 32.9 vs 23.0, p=0.47) or the Flixotide group lung deposition with and without spacer (median 38.3 vs 34.1, p=0.08). However, we may not have had the power to detect differences given the small sample size in the dataset.

Conclusion: Preliminary data suggests QVAR (p=0.47) may be used without a spacer in adolescents, provided inhalation technique is assessed by an appropriately trained clinical professional. We would still recommend that Flixotide be used with a spacer.

Grant Support: Princess Margaret Hospital Foundation

EFFECTS OF IN UTERO SMOKE EXPOSURE ON LUNG FUNCTION FROM INFANCY TO ADULTHOOD AND THE CONTRIBUTION OF GLUTATHIONE S-TRANSFERASE POLYMORPHISMS

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Introduction/Aim: In utero smoke exposure is associated with abnormal lung function, wheeze and asthma from infancy and throughout childhood. The negative influence of in utero smoke exposure is modified by genetic polymorphisms in genes important in detoxifying tobacco by-products, including the Glutathione S-transferase (GST) gene, a respiratory anti-oxidant. The longterm effects of in utero exposure on respiratory health into adulthood are unclear.

Methods: The Perth Infant Asthma follow up study is a longitudinal birth cohort of 253 subjects recruited antenatally from a general population with lung function carried out 1, 6,12 months, 6,11.18 and 24 years of age. Antenatal smoking history was collected from both parents at recruitment. DNA was collected at 6 and 11 years on a subgroup of 180 subjects.

Results: Either parent smoking during the pregnancy was associated with wheeze (OR=2.5, 95% CI 1.01- 6, p=0.048) and asthma (OR 2.7, 95% CI 1.2-5.9, p=0.012; n=123) in the offspring at 6 years of age, but not thereafter. Maternal smoking during the pregnancy was associated with reduced lung function in the offspring at 1 month (mean VmaxFRC of 90ml/sec (SD) versus 104ml/sec ; p=0.034; n=242) and 6 years of age (mean FEV1% predicted 99% 15 (SD) versus 105% 15, p=0.039; n=110) but not at later assessments.

Amongst those children exposed to in utero tobacco smoke (n=32), the GSTM1 homozygote null genotype was associated with significantly lower lung function at 6 years of age compared to those with GSTM1 non-null genotype (mean FEV1% predicted 93.5 15 (SD) versus 104.6; p=0.03).

Conclusion: In utero tobacco smoke exposure negatively impacts lung function and respiratory symptoms up to 6 years of age, but this effect does not persist into adulthood. Polymorphisms in anti-oxidant genes may contribute to worse lung function in children who were exposed to in utero tobacco smoke.

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SPECTRUM OF LUNG DAMAGE IN PRIMARY CILIARY DYSKINESIA AS SEEN ON COMPUTED TOMOGRAPHY
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Introduction/Aim: Primary Ciliary Dyskinesia (PCD) and Cystic Fibrosis (CF) are both inherited, progressive, incurable, respiratory conditions causing significant morbidity and mortality. In the lungs, damage is reflected as a spectrum of structural changes that can be detected on computed tomography (CT). To monitor disease progression and guide therapy, scoring systems for CT scans have been developed in CF, which quantify the extent and severity of changes seen. Most research examining structural lung changes on CT scans use scoring systems derived from a CF cohort. All studies to date describing lung damage in PCD use these CF-derived tools. This assumes lung damage in the two conditions is identical, which potentially results in a failure to identify PCD-specific changes. Our study addresses this assumption

Methods: We retrospectively analysed of 58 CT scans from 40 adult and paediatric patients with PCD. The following abnormalities found in CF were scored according to presence and extent: bronchiectasis, bronchial wall thickening, atelectasis, mucous plugging, and air trapping. In addition, an experienced respiratory radiologist reviewed all scans to look for the presence of any abnormalities that differ from the above.

Results: Bronchial wall thickening was the most common abnormality, and air trapping the least common. All abnormalities were present significantly more often in the middle and lower lobes compared to the upper lobes, with all p values <0.001. When present, all abnormalities were significantly more extensive in the middle and lower lobes compared to the upper lobes. Bronchiectasis, mucus plugging, atelectasis (p=0.005) and air trapping (p=0.005) were all present significantly more often in the PCD cohort than the respective CF cohorts. The PCD-unique changes identified were dextrocardia, extensive tree-in-bud patterns, whole lobe atelectasis, and interlobar and interlobular septal thickening.

Conclusion: Significant structural changes were seen on CT scans in patients with PCD, which were not consistent with those previously described in patients with CF. Our findings illustrate the need for development of a PCD-specific scoring system, which can function as a tool for the objective assessment of disease status, progression and efficacy of therapy.

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THE VENTILATORY RESPONSE TO HYPOXIA REMAINS BLUNTED IN PRETERM INFANTS WITH BRONCHOPULMONARY DYSPLASIA
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Introduction/Aim: The early postnatal environment in which very preterm infants develop is relatively hypoxic compared to in utero. Additionally, very preterm infants often require breathing support, including supplemental oxygen (O2); neonatal exposure to > 28 d O2 is an NICHD diagnostic criterion for bronchopulmonary dysplasia (BPD). Neonatal supplemental O2 may alter chemoreceptor “resetting”. Therefore, we aimed to quantify the hypoxic ventilatory response (HVR) in very preterm infants (<32 w gestation) during the neonatal period and determine if neonatal supplemental O2 has lasting effects on the HVR at 12–15 m corrected postnatal age (PNA).

Methods: The HVR was evaluated in very preterm infants at 36 w postmenstrual age (PMA) and 12–15 m corrected PNA. Breathing variables, including tidal volume (VT), respiratory rate (f), minute ventilation (Ve) and inspiratory flow (Vi) were measured under normoxic (21% O2) and hypoxic (14% O2) conditions. Hypoxia induced change in breathing variables was assessed using Wilcoxon Matched-Pair Signed-Rank test.

Results: Preterm infants (n=28; 17 with BPD) had no HVR at 36 w PMA (p>0.05). In contrast, the HVR was present at 12–15 m PNA (n=13; 3 with BPD), manifest during hypoxia as 1) increased VT (mean difference 2.0 mL/kg; 95% CI 1.2, 2.9 mL/kg; p=0.001); 2) increased Ve (28.3; 12.7, 44.0 mL/min/kg; p=0.002) and 3) increased Vi/VT (52.8; 41.2, 64.3 mL/s; p=0.001). Low birth weight z-score (Spearman Rho= 0.622; p=0.031) and increased duration of supplemental O2 (-0.613; p=0.034) were associated with smaller increases in VT during hypoxia. Preterm infants with BPD had less change in VT (p=0.048) and Ve (p=0.024) during hypoxia at 12–15 m PNA compared to preterm infants without BPD.

Conclusion: Infants born very preterm have absent ventilatory response to hypoxia in the neonatal period. Persistence of a blunted HVR beyond the first year of life in infants with BPD warrants further investigation.

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**PRAGMA-CF: A QUANTITATIVE MEASURE OF STRUCTURAL LUNG DISEASE IN YOUNG CHILDREN WITH CYSTIC FIBROSIS.**

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**Introduction/Aim:** Chest computed tomography (CT) is the gold standard for detecting structural lung disease in cystic fibrosis (CF), however there has been a lack of outcome measures appropriate for young children. The aim of this study was to develop and validate a quantitative CT-based measure of structural lung disease in children aged below 6 years. We hypothesised that this outcome is biologically relevant and suitable for use as a clinical trial outcome.

**Methods:** All patients undergoing annual surveillance CT and bronchoalveolar lavage (BAL) were included. To assess structural lung disease extent, the PRAGMA-CF method was developed: a grid was overlaid on CT slices and annotated hierarchically for the presence of bronchiectasis, bronchial wall thickening, and mucous plugging (inspiratory scan) or trapped air (expiratory scan). Overall disease (%Dis) and trapped air (%TA) were expressed as the proportions of cells with disease (respectively) compared to healthy.

Thirty scans were randomly selected for rescoring to assess repeatability (intraclass correlation coefficient, ICC). Linear mixed model analysis was used to compare CT outcomes to the presence of neutrophil elastase (NE) and infection from BAL.

**Results:** 683 scans from 256 patients were included in the final analysis. ICCs to assess repeatability were above 0.8 (grade of excellent). CT outcomes were associated with both the presence and history of NE and infection. Sample size calculations showed that multicentre clinical trials of intervention can be performed with around 100 patients.

**Discussion:** PRAGMA-CF is a sensitive tool to monitor structural lung disease in CF. It is highly repeatable, biologically plausible (associated with clinical markers of lung disease), and is suitable for use as an outcome measure in multicentre clinical trials. This study provides both a rationale and a means to test interventions aimed at preventing structural lung disease in young children with CF.

**IDENTIFICATION OF A PLASMA microRNA PROFILE IN PULMONARY TUBERCULOSIS PATIENTS THAT IS MODULATED BY ANTI-MICROBIAL THERAPY**

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**Introduction/Aim:** microRNA expression profiles are of interest as a biomarker of tuberculosis (TB). The effect of anti-TB therapy on miRNA profiles in pulmonary TB patients is unknown and was examined in this study.

**Methods:** Plasma miRNA levels for 175 miRNAs were compared in 20 TB patients and matched healthy controls from China. 87 miRNAs were differentially regulated between the two groups. Ten of these were analysed in a test cohort of 100 pulmonary TB patients sampled prior to the commencement of antibiotic therapy and at one, two and six months during treatment.

**Results:** Six miRNA were differentially expressed in the test group of TB patients. miRs -29a and -99b were up-regulated, whilst miRs -26a, -146a and -652 were down-regulated. A combination of 4 miRNA distinguished pulmonary TB from healthy controls with a sensitivity of 84%, with an AUC of 0.941. Within one month of treatment, significant modulation of miRs -29a, -99b, -26a and 146a was seen in successfully treated patients, although not all miRNAs had returned to baseline at completion of treatment.

**Conclusion:** Six miRNA were differentially expressed in the test group of TB patients. miRs -29a and -99b were up-regulated, whilst miRs -21, -26a, -146a and -652 were down-regulated. A combination of 4 miRNA distinguished pulmonary TB from healthy controls with a sensitivity of 90% and a specificity of 84%, with an AUC of 0.941. Within one month of treatment, significant modulation of miRs -29a, -99b, -26a and 146a was seen in successfully treated patients, although not all miRNAs had returned to baseline at completion of treatment.

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ZINC DYSHOMEOSTASIS AND AUTOPHAGY AS CRITICAL DETERMINANTS FOR AIRWAY EPITHELIAL DYSFUNCTION IN COPD
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Introduction/Aim: Airway inflammation and cigarette smoke in COPD induces autophagy in bronchial airway epithelial cells (bAEC). Zinc (Zn) deficiency is also a stimulus for autophagic dysfunction. We hypothesised Zn deficiency potentiates COPD, and with cigarette smoke exposure, causes autophagic insufficiency and epithelial fragility.

Our aims were to determine whether cigarette smoke exposure produces Zn dyshomeostasis and autophagic dysfunction, and if this impairs the epithelia barrier, and induces the production of a COPD-related pro-inflammatory phenotype.

Methods: Emphysematous mouse lung tissues were assessed via immunofluorescence for free-Zn. Air-liquid interface (ALI) cultures of human bAECs were exposed to Zn depletion, TNFα/IFNγ, and cigarette smoke extract (CSE10%). End points included western analysis for Zn transporters ZIP1 and ZIP2, autophagy/apoptosis regulators e.g. Light chain-3-II, Sequestosome, Beclin and X-linked inhibitor of apoptosis (XIAP), and tight junction proteins Claudin-1 and ZO-1, transmission electron microscopy to visualise autophagy, barrier function assessment including permeability of microscopy to visualise autophagy, barrier function assessment including permeability of fluorescein tracers, and ELISA for RANTES and Thymic stromal lymphopoietin (TSLP).

Results: Lung tissues from cigarette smoke-exposed mice showed significantly reduced free-Zn in the epithelium. Further, human bAECs stimulated with TNFα/IFNγ, in ZnDep/CSE10% showed an increase in ZIP1 (p=0.001) and decrease in ZIP2 (p=0.007), and increased autophagy with reduced capacity to degrade cellular debris. Autophagic dysfunction associated with the induction of apoptosis via the reduction of Beclin and XIAP proteins (both p<0.05). Reduced epithelial barrier function was evidenced by decreases in Claudin-1 and ZO-1 protein, reduced electrical impedance (3.5-fold, p<0.05), and increased permeability. A significant increase in RANTES and TSLP was observed for ZnDep, with/without CSE10%.

Conclusion: We demonstrate cigarette smoke depletes free-Zn from the airway epithelium, and Zn deficiency is a significant potentiator of autophagic and apoptotic dysregulation, the production of potent AEC-derived inflammatory mediators, and epithelial fragility. In combination with cigarette smoke exposure, Zn dyshomeostasis may be a major determinant for the pathogenesis of the COPD phenotype.


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IDENTIFYING A NOVEL THERAPEUTIC STRATEGY FOR ASTHMA: TARGETING AIRWAY EPITHELIAL CELL RESTITUTION
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Introduction/Aims: Recent advances have implicated the airway epithelium as a key driver in asthma pathogenesis, partly due to its dysregulated response to damage. Novel therapies focusing on protecting and repairing vulnerable airways, particularly in early life, could transform asthma treatment by preventing disease development/progression. (1)To analyse primary airway epithelial cells (pAEC) migration patterns post-wounding in vitro. (2)To identify mechanisms and potential therapeutic targets enhancing wound repair processes.

Methods: pAEC were obtained from non-asthmatic and asthmatic paediatric airways. Scratch wounds were performed on pAEC monolayers to assess repair and imaged every 30 minutes (IncuCyte ZOOM®; Essen Bioscience). Migration trajectories of leading-edge pAEC were analysed using ImageJ. Integrin gene and protein expression were investigated by qPCR and In-Cell®Western, respectively. Total RNA was sequenced(Illumina Hi-Seq2500) and differential gene-expression analysis was performed(DESeq2 and Ingenuity Systems(QIAGEN)).

Results: Response to wounding in asthmatic children was deficient and lacked specificity with significantly lower mean migration distance(non-asthma and asthma; 256.9±7.1 and 152.3±8.6μm(mean±SEM)), velocity(0.42±0.02 and 0.22±0.01μm/min), directionality (91.3±0.1% and 61.1±0.1%) and forward migration index(95.3±0.1 and 64.6±0.1%). A major regulator of cell migration, i.e. integrin α5β1, was investigated in pAEC. Lower gene(5, 5.9-fold, p=0.0001; j1, 1.5-fold, p<0.05), and protein(5, 2.8-fold, p<0.05; j1, 3.1-fold, p<0.05) levels in pAEC from asthmatic children(n=11) compared to control(n=23). RNA-seq analysis identified 1,153 differentially expressed genes in asthmatic children(n=6) relative to control(n=4) with overrepresented gene ontologies related to integrins and extracellular matrix. Drug database screening identified several clinically safe drugs that are now being examined for drug repurposing potential to restore integrin expression and aid wound repair.

Conclusion: These novel experiments demonstrate abnormal migration behaviour of asthmatic airway epithelium post-wounding. Some mechanisms controlling this disease phenomenon were identified like decreased integrin α5β1, and multiple transcriptional mechanisms were dysregulated in astmas, some of which are targetable by existing drugs. Supporting airway epithelial repair and barrier integrity may be a novel therapeutic avenue for asthma.
Introduction/Aim: Rhinovirus infection triggers acute exacerbations of asthma. Expression of IL-33, an instructive cytokine of type 2 inflammation, is upregulated during experimental rhinovirus infection of asthmatic subjects and correlates with the production of type-2 cytokines and eosinophilic inflammation. Using a novel model of virus and allergen exposure, we sought to determine whether anti-IL-33 therapy attenuates a rhinovirus-induced asthma exacerbation in mice.

Methods: To simulate the synergistic effects of virus infection and allergen exposure on asthma susceptibility, mice were exposed to low dose pneumonia virus of mouse (PVM; 1 pfu) and low dose (1 μg) cockroach antigen (CRE) in early life and again in later life. Four weeks after the final CRE exposure, mice were inoculated with rhinovirus (RV-1B, TCID50 5x10^6). Anti-IL-33 or dexamethasone was administered intraperitoneally twice/week between the CRE and RV challenge, then daily until euthanasia.

Results: Both early-life and later-life exposures to PVM and CRE were necessary for disease onset and progression. IL-33 levels were elevated immediately following the final CRE exposure and persisted in the airways until the time of RV-1B challenge. Mice co-exposed to PVM/CRE, but not CRE or PVM alone, presented with eosinophilic inflammation, increased numbers of type 2 innate lymphoid cells, mucous hypersecretion and elevated IL-13 levels following RV infection. Treatment with anti-IL-33 or dexamethasone attenuated the RV-1B-induced type 2 inflammation but had no effect on mucous production. Critically, anti-IL-33, but not dexamethasone, promoted the expression of antiviral cytokines, accelerating RV-1B viral clearance.

Conclusion: Both anti-IL-33 and dexamethasone suppress the magnitude of type 2 inflammation during an rhinovirus-induced acute exacerbation, however anti-IL-33 has the added benefit of boosting antiviral immunity and lowering viral burden.

Grant Support: Pfizer Inc, and The Australian Infectious Disease Research Excellence Award, The University of Queensland.
SELF-PERCEIVED BURDEN AS A BARRIER TO
COMMUNICATION AT THE END OF LIFE IN COPD

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Introduction/Aim: The end of life (EOL) trajectory in Chronic Obstructive Pulmonary Disease (COPD) is known to complicate prognostication, potentially preventing timely transition to palliative care, thus the primary responsibility for care provision remains with the family of the person with COPD. We aimed to explore the experiences of older people living with Stage IV COPD in a rural community, and how COPD influenced their relationships and identity.

Methods: Eleven older people with Stage IV COPD living in regional NSW undertook narrative interviews, discussing their ‘COPD journey’. The interviews were transcribed, then analysed and interpreted in accordance with a hermeneutic phenomenological methodology.

Results: Spousal care was generally well accepted and expected by married participants as an extension of the marital role. However, when adult children were involved in care or decision making, participants expressed guilt and frustration about perceived and actual burden placed on them. This self-perceived burden, grounded in their own experiences caring for loved ones, had flow on effects to their uptake of coping and treatment strategies, and Advance Care Planning (ACP) as well as creating communication barriers between each of the parties. Parents aimed to protect their adult children from distress or impacting on their lives, by under-communicating need and disease specific information. Paradoxically, many participants went on to design their ACP to refuse treatment rather than extend any impact on their children’s lives, contrary to their children’s ACP requests of them.

Conclusion: With the protection of adult children being a primary determinant of end of life decision making, it is critical to establish more formal and effective communication pathways between family and health professionals to facilitate a ‘good death’ within the family group.

Grant Support: Charles Sturt University Faculty of Science Fee Waiver for Higher Degree Research Students

ASTHMA AND PHYSICAL ACTIVITY IN CHILDHOOD: ARE THEY LONGITUDINALLY RELATED?

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Aim: There is increasing interest in the role of physical activity (PA) in the development of asthma but the potential for asthma to influence PA has hindered the progress in understanding this given the lack of relevant longitudinal data in children. The primary aim of this study was to explore whether having asthma leads to lower PA levels, and/ or whether lower PA levels leads to more asthma in children and adolescents between the ages of 8 and 14 years.

Methods: This analysis was conducted in 4983 children who participated in the Longitudinal Study of Australian Children (LSAC) between the ages of 6 and 14. Data on asthma and PA were collected via questionnaires and Time Use Diaries biennially since the children were 4 years old. Bi-directionality of this relationship was investigated using a cross-lagged model for PA and asthma implemented using structural equations modelling. Incident-current asthma was defined as doctor’s diagnosis since previous wave and either use of asthma medications or current wheeze in the past 12 months. PA was time in minutes spent doing medium to vigorous physical activities in a day. Included confounders were gender, older siblings, socioeconomic z-score and body mass index (BMI).

Results: Effect estimates for each regression suggested that there was no association between PA and asthma at any age and in either direction; the adjusted regression coefficients for the effect of asthma on subsequent PA were: 0.98 (0.90, 1.07), 0.93 (0.85, 1.01), 1.07 (0.97, 1.17) and 0.99 (0.88, 1.11). Similarly, the adjusted log of OR for the effect of PA on asthma was 0.99 (0.99, 1.00) at each age.

Conclusions: These results suggest that PA at any of the observed ages does not prospectively predict incident-current asthma in this longitudinal analysis. Similarly, incident-current asthma does not predict future PA in the observed ages.

Grant Support: N/A

Declaration of conflict of interest: The authors report no conflict of interest. The authors alone are responsible for the content and writing of the paper.

Keywords: asthma, adolescents, children, physical activity
BLOOD EOSINOPHILS AND SERUM IgE PREDICT RESPONSE TO OMALIZUMAB IN PATIENTS WITH SEVERE ALLERGIC ASTHMA: INNOVATE TRIAL POST-hoc ANALYSIS

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Introduction: Response to biologic therapy differs among severe allergic asthma (SAA) patients. Biomarker-guided therapy may help to identify subgroups who demonstrate greater clinical benefits from omalizumab than the overall SAA population.

Methods: A post-hoc analysis of INNOVATE trial data (Humbert et al. Allergy 2005) was performed to determine the impact of treatment on exacerbation rates and health-related quality-of-life (AQLQ) of SAA patients stratified by peripheral blood eosinophils per mcI (EOS) and serum IgE (IU/ml). Negative binomial regressions were utilized within biomarker strata to estimate response to omalizumab vs placebo, controlling for potential confounders.

Results: An exacerbation reduction was observed among omalizumab-treated patients vs placebo in EOS-high strata: EOS≥150/IgE>75 stratum (n=231): 0.59, p<0.0001. The benefit on AQLQ exceeded the minimal clinically-important difference (≥0.5) and demonstrated statistically significant improvements in both EOS-high strata: EOS≥150 (n=24) 0.55, p=0.0008; EOS≥150 (n=310) 0.53, p=0.001. The benefit on AQLQ was most apparent in the EOS≥150/IgE≥75 stratum (n=231): 0.59, p<0.0001.

Conclusion: This analysis is among the first to examine a combination of biomarkers to assess response to omalizumab in SAA patients. They suggest that subgroups with a combination of increased IgE and eosinophils may experience a greater clinical benefit. However, caution must be used in interpreting these results given their post-hoc nature.

Grant Support: This analysis was supported by Novartis Pharma AG and Genentech.

INCREASED EXPRESSION OF IL-27 IN NEUTROPHILIC ASThma

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Introduction/Aim: Asthma is a heterogeneous inflammatory disorder consisting of multiple endotypes, those with neutrophilic asthma (NA) have innate immune activation and tend to be resistant to corticosteroid therapy. We have shown that steroid-resistant asthma can involve IL-27, a monocyte/macrophage-derived innate cytokine speculated to be involved in TH1 responses. The aim of this study was to examine the expression of IL-27 in asthma inflammatory phenotypes and determine relationship between IL-27 and other immune factors involved in asthma pathogenesis. We hypothesised that sputum gene expression of IL-27 would be increased in participants with NA compared with other asthma phenotypes.

Methods: Induced sputum samples from 80 non-smoking adults with stable asthma were assessed for total and differential cell counts along with gene expression for IL-27R, IL-27p28, IL-27EBI3, IFN-γ and TNF-α. Inflammatory phenotype was determined using a single sputum samples we eosinophilic asthma (EA) when eosinophils ≥2% and neutrophils <61%, neutrophilic asthma (NA) when neutrophils ≥61% eosinophils <2% and paucigranulocytic asthma (PA) when eosinophils <2% and neutrophils <61%.

Results: The participants had a mean age of 55 years, 37 (42%) were males and 63 (79%) were prescribed inhaled corticosteroids. FEV1 %predicted was lowest in NA (mean 63%) and significantly lower than those with PA (80%; p=0.009). Sputum gene expression of both IL-27 subunits (p28 and EBI3), IFN-γ and TNF-α were significantly increased in NA compared with EA. Furthermore, the expression of IL-27EBI3 in NA was significantly (p<0.05) higher than PA participants. IL-27R mRNA level was significantly negatively correlated with degree of airflow obstruction (FEV1/FVC%) (r=0.24, p<0.05), while gene expression of the subunit, IL-27EBI3 mRNA, was positively associated with sputum total cell count (r=0.315, p<0.05), neutrophil% (r=0.486, p<0.05) and age (r=0.223, p<0.05).

Conclusion: Elevated gene expression of IL-27 suggests this may be an important cytokine in the pathogenesis of NA.

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RISK FACTORS FOR VOCAL CORD DYSFUNCTION IN A DIFFICULT ASTHMA POPULATION

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Introduction/Aim: Vocal cord dysfunction (VCD) is characterized by inappropriate adduction of the vocal cords giving rise to symptoms of dyspnea, voice change, wheezing and throat tightness. VCD can be misdiagnosed as asthma, resulting in delayed or inappropriate treatment. A high proportion of patients with difficult-to-treat asthma are reported to also have co-existing VCD.

Methods: One hundred and seventeen consecutive difficult asthma patients referred by respiratory specialists underwent the Alfred assessment protocol. Vocal cord dysfunction was diagnosed either by: an ENT specialist, or clinically, based on characteristic symptoms, and supported by a positive Pittsburgh VCD index or VCD-Questionnaire. Univariate analyses were performed to identify clinical factors associated with vocal cord dysfunction. Predictors were compared using unpaired t-test or chi-square test where appropriate. Predictors with a p-value \( \leq 0.25 \) were then included in a multivariate logistic regression model. Each predictor was assessed for confounding and significance. Continuous variables were assessed in mean (SD).

Results: Forty (34.2\%) of 117 difficult asthma patients had VCD. The majority (n=33) of these also had asthma demonstrable by variable airflow obstruction. Patients with VCD were more likely to: be female (OR 2.45, 95\%CI 1.03-5.84, p=0.04); have poorer asthma control and quality of life (ACT 13\%±5 vs. 15\%±5, p=0.023; AQLQ 3.59\%±1.41 vs 4.61\%±1.38, p=0.001); have more frequent exacerbations [3\%2–5 vs 2 (0–2) over six months, p=0.003], and: be less severely obstructed (FEV1/FVC ratio 68\%±13 vs. 61\%±16, p=0.028). On multivariate logistic regression, independent predictors for VCD were FEV1/FVC and AQLQ.

Conclusion: VCD was present in a third of our patients with difficult asthma. VCD was associated with female sex and poorer asthma outcomes despite better lung function. Improved lung function and poorer quality of life were both independent predictors of VCD. Our findings highlight the importance of identifying and addressing VCD in this challenging patient group.

Grant Support: Nil

ASThma GENe Signatures expression in bronchial biopsy

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Introduction/Aim: The expression of TH-2 high (CLCA1, POSTN, and SERPINB2) genotype has been shown to predict response to inhaled steroids in mild asthma and a 6 genes signature (6GS) in sputum cells (CLC, CPA3, DNASEI1L3, IL1B, ALPL, and CXCR2) distinguishes between asthma inflammatory phenotypes. The aim was to determine gene expression of both signatures in endobronchial biopsies from adults with asthma and related to asthma severity and airway inflammatory phenotypes.

Methods: Non-smoking adults with asthma had current respiratory symptoms, and evidence of variable airflow obstruction. Asthma severity was defined according to GINA. Inflammatory phenotypes using BAL cell count. The 9 genes were analyzed by qPCR using extracted RNA.

Results: Biopsies were evaluated from: severe asthma (n=40), mild/moderate asthma (n=32), with no differences in age, sex, atopy or smoking history between the groups. Gene expression of SERPINB2 was significantly increased in severe compared with mild/moderate asthma (p=0.022) and inversely associated with FEV1\% predicted in participants with asthma (r=–0.337, p=0.005).

Gene expression CLC and SERPINB2 were significantly increased in eosinophilic asthma (n=20) compared with paucigranulocytic asthma (n=21, p=0.003 and p=0.0009 respectively), while ALPL was increased in eosinophilic compared with mixed-granulocytic asthma (n=13, p=0.0056). Only gene expression for CLC was associated with asthma BAL eosinophils \( \% r=0.325, p=0.006 \). Gene expression for IL-1\( \beta \) was increased in neutrophilic (n=18) compared with paucigranulocytic asthma (p=0.006) and was significantly associated with BAL neutrophils \( \% r=0.234, p=0.0496 \).

Conclusion: SERPINB2 expression in bronchial biopsy is associated with severe asthma, while CLC and IL-1\( \beta \) gene expression are associated with BAL eosinophilia and neutrophilia respectively.

Grant Support: John Hunter Hospital Charitable Trust 2016.
MACROPHAGES REGULATE THE DEVELOPMENT OF RSV INDUCED ASTHMA EXACERBATIONS

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Introduction/Aim: Viral respiratory infections trigger severe exacerbations of asthma, worsen disease symptoms and impair lung function. To investigate the mechanisms underlying viral exacerbation, we established a mouse model of respiratory syncytial virus (RSV)-induced exacerbation after allergen sensitisation and challenge.

Method: We investigated the effects of RSV infection on the induction of asthma exacerbation in a mouse model. Airway hyper-responsiveness (AHR) was assessed by flinvent. In some experiments, pulmonary macrophages were depleted by 2-Chloroadenosine; TNFα and MCP-1 were neutralized by monoclonal antibodies.

Results: RSV infection of OVA-sensitised/challenged BALB/c mice resulted in significantly increased AHR and macrophage and neutrophil infiltration. Exacerbation was accompanied by increased levels of inflammatory cytokines (including TNFα, MCP-1, and KC) compared to uninfected OVA-treated mice or OVA-treate mice exposed to UV-inactivated RSV. Dexamethasone treatment completely inhibited all features of allergic disease including AHR and eosinophil infiltration in uninfected OVA-sensitised/challenged mice. Conversely, dexamethasone treatment following RSV-induced exacerbation only partially suppressed AHR, and failed to dampen macrophage and neutrophil infiltration or inflammatory cytokine production (TNFα, MCP-1, and KC). This mimics clinical observations in patients with exacerbations, which is associated with increased neutrophils and often poorly responds to corticosteroid treatment. Interestingly, we also observed increased TNFα levels in sputum samples from neutrophilic asthmatic patients. While RSV-induced exacerbation was resistant to steroid treatment, inhibition of TNFα and MCP-1 function or depletion of macrophages suppressed features of disease including AHR, macrophage and neutrophil infiltration.

Conclusion: Our findings highlight critical roles for macrophages and inflammatory cytokines (including TNFα and MCP-1) in viral-induced exacerbation of asthma and suggest examination of these pathways as novel therapeutic approaches for disease management.

Key Words: RSV, macrophage, steroid resistance, airway hyper-responsiveness, asthma exacerbation, mouse model

Nomination for New Investigator Award, N/A

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INVESTIGATING THE NON-ION CHANNEL EFFECTS OF CFTR MODULATORS ON INNATE IMMUNE VIRAL RESPONSES

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Introduction & Aims: Airway epithelial cells (AEC) from CF lungs respond to viral infection with poor innate immune responses. The contribution of defective CF transmembrane conductance regulator (CFTR) channel function is not known. We investigated whether CFTR correcting drugs could improve responses by CF AEC to human rhinovirus (HRV).

Methods: Primary AEC obtained from 11 children with CF (five Class II, six Class III mutations) and six healthy controls by bronchial brushing, were expanded in vitro. Monolayers were stimulated with purified HRV1b at multiplicity of infection 12.5 for 24 hours, with or without 72 hours treatment by CFTR therapies lumacaftor and/or ivacaftor. RNA was collected and sequenced on the Illumina HiSeq 2500 platform. Supernatant was assessed by ELISA for interferons (IFN) and inflammatory cytokines.

Results: Data were interpreted as ratio of treatment response to unstimulated AEC. In healthy AEC, HRV1b stimulation resulted in increased IL-8 and RANTES, but not IL-6, IFNβ, IFNα1 or IFNα2. Responses to HRV1b by healthy AEC did not significantly change in the presence of CFTR modulators. In contrast, AEC of both CF genotypes generated an IL-8 and an IL-6 response to HRV1b (p<0.05) Interestingly, Il-6 response by CF AEC was no longer significant following CFTR modulator treatment. Combined CFTR therapy for Class II CF was also found to reduce RANTES production (p<0.05). The IFN response to HRV1b was mixed, with minimal levels detected following stimulation except for IFNα1, which was elevated in CF AEC supernatant (p<0.05). This was no longer significant after CFTR modulator therapy for both Class II and II (p<0.05).

Conclusion: CFTR therapies had a counter-intuitive effect on responses by CF AEC to HRV1b. We are currently analysing RNAseq data to compare innate immune gene networks between treatments and phenotypes.

MOLECULAR AND PROTEOMIC APPROACH TO INVESTIGATE ABNORMAL IRON HOMEOSTASIS IN THE CYSTIC FIBROSIS AIRWAY

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Introduction/Aim: Iron homeostasis appears to be abnormal in the lungs of individuals with the genetic disease cystic fibrosis (CF). Disruption of iron homeostasis in the lung may promote bacterial infection and generate oxidative stress. We therefore tested the hypothesis that iron homeostasis in airway epithelial cells (AEC) from CF patients is abnormal and that this results in oxidative stress.

Methods: Cultured AEC from a CF patient with the F508del/W1282X mutations (immortalized IB3-1 cell line) were exposed to ferric ammonium citrate (FAC; 50 & 200 μg/mL) and the response of iron regulatory pathways assessed by reverse transcription polymerase chain reaction (RT-PCR) and proteomics. Interleukin (IL)-6 levels were also assessed by enzyme-linked immune-assay in the AEC culture supernatant to determine whether iron exposure resulted in a pro-inflammatory response.

Results: IB3-1 CF AEC demonstrated a significant increase in the expression of transferrin receptor (TFR)-1 (iron uptake) at baseline (pre-iron challenge) compared to C38 AEC. The IB3-1 AEC also demonstrated a significant increase in the expression of the iron exporter ferroportin (FPN) by at baseline compared to C38 AEC. Following FAC challenge, the expression of TFR-1 by RT-PCR was significantly reduced in both the IB3-1 and C38 AEC, but expression of FPN remained essentially unchanged. Interleukin-6 levels were significantly increased in the IB-3 CFTR mutant compared to C38 AEC supernatants at baseline and both AEC lines demonstrated a significant increase in IL-6 production following FAC challenge. Preliminary pathway analyses from the proteomic data reveal significant differential expression of proteins involved in oxidative stress and mitochondrial stress responses in the IB3-1 compared to C38 AEC.

Discussion: The airway epithelium in CF demonstrates abnormalities of iron regulation, exhibits a pro-inflammatory phenotype and activation of cell pathways involved in oxidative stress.

Key Words: Iron, cystic fibrosis, airway epithelial cells

Grant Support: NHMRC and TPCH Foundation.

STRUCTURAL DETERMINANTS OF LONG TERM FUNCTIONAL OUTCOMES IN CHILDREN WITH CYSTIC FIBROSIS

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Introduction/Aim: Chest computed tomography (CT) is a potential surrogate endpoint for children with cystic fibrosis (CF). The relationship between structural abnormalities at an early age and long term functional outcomes has not been previously described. We aimed to determine whether CT scores in children under 6 years are predictive of later lung function.

Methods: Volumetric and limited slice CT scans were obtained annually from children enrolled in AREST CF program aged 0 to 6 years. Follow-up spirometry measurements were available for some of the AREST CF cohort from ages 5 to 15. Cox regression and mixed effects models were used to determine at what age and to what extent are CF-CT scores predictive of future lung function decline. Results are presented adjusted for intrinsic disease severity (Homzygous D508 mutation, meconium ileus presence and gender), time elapsed between CT scan and spirometry and test centre.

Results: 681 spirometry measurements were available (mean age 9.3±2.1 years), Total CF-CT score at 5–6 years (n=171) that was greater than the median was significantly associated with shortened time before FEV1 falls below −1.645 Z scores (hazard ratio 3.16 (1.25, 7.99) p=0.015). The extent of mucus plugging and trapped air at ages 5–6 years were strongly associated with FEV1 Z scores between ages of 5–15 (−0.17 (−0.26, −0.07) p=0.001 and −0.09 (−0.14, −0.04) p=0.001 respectively). Predictive power of CT scores in young children is consistently stronger the more time elapses between initial CT scan and spirometry (mucus plugging score at 5–6 explains only 3% of variation in FEV1 Z score at the same age but 37% of variation 5 years later).

Conclusion: Non-bronchiectatic lung abnormalities in young children with CF are important markers of future disease progression. Structural abnormalities are predictive of future FEV1 trajectories and represent promising endpoints in early intervention trials.

Grant Support: NHMRC and USA CF Foundation

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FACE MASKS AND COUGH ETIQUETTE REDUCE COUGH GENERATED BIOAEROSOLS CONTAINING PSEUDOMONAS AERUGINOSA IN PATIENTS WITH CYSTIC FIBROSIS

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Introduction/Aim: The CF Foundation (USA) recently updated their infection control guidelines to include the recommendation that patients with cystic fibrosis (CF) wear surgical face masks in communal areas of health facilities to reduce transmission of aerosolised pathogens. The aim of this study was to investigate the effectiveness of face masks and cough etiquette techniques for reducing viable bioaerosols in patients with CF and chronic P. aeruginosa infection.

Methods: Subjects were enrolled from the Adult CF Centre, The Prince Charles Hospital, Brisbane and positioned in a validated, aerosol-sampling device and performed activities for 5 minutes each: 1. talking; 2. talking with surgical mask; 3. unmasked coughing; 4. coughing with surgical mask; 5. coughing with N95 mask; 6. coughing with hand covering mouth. A 6-stage Andersen Cascade Impactor, positioned at 2 metres, collected and sized viable aerosols. Quantitative sputum and aerosol cultures were performed. Subjects rated their comfort levels with the masks.

Results: 25 (15 male) adults, mean (SD) age of 31.2 (7.7) years and FEV1 of 50.7 (17.4) % predicted were enrolled. 76% subjects produced viable P. aeruginosa during unmasked coughing. Cough etiquette, surgical mask and the N95 reduced infectious cough aerosols of 53.4, 93.8, 94.5% respectively, with the surgical mask rated more comfortable.

Conclusion: Face masks are effective in reducing cough generated infectious droplet nuclei in people with CF, with the surgical mask providing enhanced patient comfort. Cough etiquette provides less protection than masks.

Grant Support: CFF Therapeutics (USA), The Prince Charles Hospital Foundation (Australia)

DIFFERENCES IN THE LOWER AIRWAY MICROBIOTA OF INFANTS WITH AND WITHOUT CYSTIC FIBROSIS

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Introduction/Aim: Infection plays a critical role in the pathogenesis of early cystic fibrosis (CF) lung disease. Differences in the nasopharyngeal microbiota of subjects with CF and healthy controls are apparent from early infancy. This study aimed to compare the composition and structure of the lower airway microbiota of infants below 12 months of age with and without CF.

Methods: Bronchoscopies and bronchoalveolar lavage (BAL) procedures were performed in infants newly diagnosed with CF following newborn screening. Infants undergoing clinically-indicated bronchoscopy were recruited as controls. Quantitative microbiological culture and inflammatory marker (interleukin-8 and neutrophil elastase) measurements were undertaken contemporaneously. 16S ribosomal RNA gene sequencing was conducted on stored samples.

Results: Seventeen infants with CF (median age 2.6 months (interquartile range (IQR) 1.6-4.9 months); male 59%; P.Phe508del homozygous 71%) and nine control infants (median age 5.0 months (IQR 2.9-8.2 months); male 78%) contributed BAL samples. Diversity was reduced in CF compared with control BAL samples (median Shannon diversity index 1.3 (IQR 0.9-2.0) and 2.0 (IQR 1.4-2.3) respectively). Firmicutes was the most prevalent phylum in both groups, accounting for 76.9% and 49% of total reads respectively, however Staphylococcus was predominant in CF samples only (39.8% total reads). The next most prevalent genera in the CF samples were Streptococcus (15.0%), Haemophilus (9.9%), Granulicatella (9.2%) and Gemella (6.7%). In contrast, the five most abundant genera in the control samples were Streptococcus (30.0%), Fusobacterium (12.8%), Neisseria (12.4%), Gemella (6.3%) and Veillonella (6.1%). Staphylococcus and Haemophilus each accounted for 0.4% of total reads.

Conclusion: There are substantial perturbations in the structure and composition of the lower airway microbiota of infants with CF in the first months of life, most notably increased prevalence of Staphylococcus and reduced diversity. Larger longitudinal studies are required to determine the clinical significance of these findings.

Grant Support: KBF is supported by the TSANZ/Vertex CF Paediatric Clinical Fellowship, the Australian Cystic Fibrosis Research Trust
Postgraduate Studentship and the Royal Children’s Hospital Cystic Fibrosis Research Trust. 16S rRNA gene sequencing was funded by grants from the Murdoch Children’s Research Institute “65km for CF” and the RCH CF Research Trust. TWF, SCR, GAS and KMW were supported by the National Institutes of Health (NIH) grant, HL116211 and National Health and Medical Research Council award, NHMRC1043768. These bodies had no role in the study design, data analysis, interpretation or reporting of results.

Declaration of interests: There are no conflicts of interest.

THE RELATIONSHIP BETWEEN EXERCISE CAPACITY AND HOSPITALISATION IN ADULT CYSTIC FIBROSIS

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Introduction/Aim: The six minute walk test (6MWT) is an objective tool used to assess functional exercise capacity in cystic fibrosis (CF). However, the relationship between 6MWT outcomes and need for hospitalisation for treatment of pulmonary exacerbation (PE) has not been studied. Therefore, the primary aim was to determine the relationship between six minute walk distance (6MWD), subjective breathlessness (Modified Borg score) and oxygen saturation (SpO2) during the 6MWT, and frequency and duration of hospitalisation in adults with CF.

Methods: Retrospective cohort study of all adult CF patients treated at The Prince Charles Hospital, Brisbane, who completed a 6MWT in 2014 or 2015. 6MWD, maximum Modified Borg score and nadir SpO2 during the 6MWT were correlated with the number of hospital admissions and total hospital days of treatment for PE’s during the same calendar year.

Results: A 6MWT was performed in 94 subjects [mean age 29.8±9.3yrs, FEV1 56±22% predicted, 6MWD 590±114m, median (IQR) hospital admissions 2 (1–4), median (IQR) hospital days 18.5days (8.3–57.5days)]. Reduced 6MWD significantly correlated with number of yearly hospital admissions and total yearly hospital days for treatment of PE’s. However, strongest correlations were seen between maximum Modified Borg score, nadir SpO2 and total hospital days (Table 1).

<table>
<thead>
<tr>
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<th>6MWD</th>
<th>Maximum Modified Borg score</th>
<th>Nadir SpO2</th>
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<tr>
<td></td>
<td>P-</td>
<td>Correlation value</td>
<td>P-</td>
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<tr>
<td>Yearly hospital</td>
<td>-0.254&lt;0.02</td>
<td>0.546</td>
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<td>admissions</td>
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<tr>
<td>Total yearly</td>
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<td>0.591</td>
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<td>hospital days</td>
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Conclusion: Maximum perceived breathlessness and oxygen desaturation more strongly correlated with hospital utilisation than distance walked, suggesting that subjective exertional dyspnoea and exercise-induced hypoxaemia may be important predictors of exacerbation frequency.

Grant Support: Nil

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OCCUPATIONAL LUNG DISEASES REMAIN UNDERREPORTED AND UNDERCOMPENSATED IN AUSTRALIA

THEODORA AHILAS

Introduction/Aim: Whilst much of the focus on the legal claims in this arena has been on asbestos, the re-emergence of coal workers pneumoconiosis in Queensland over the past 18 months has justifiably showcased lung diseases caused by toxic exposures at work.

The Senate Select Committee on Health stated in The Fifth Interim Report (April 2016) “...a litany of regulator failure and regulatory capture, industry indifference and incompetence, inconsistent risk mitigation and patchy and sometimes compromised health monitoring in Australia” was responsible for the return of a disease that was considered eradicated over 30 years ago.

Theodora Ahilas will present on the rights to compensation for workers with occupational lung diseases.

Theodora will discuss legal entitlements that flow from the diagnosis of an occupational lung disease. The different legal and statutory entitlements that follow in the different States together with the complexity of assessing these claims particularly in the presence of co-morbidities.

LIFETIME OCCUPATIONAL EXPOSURES IN A POPULATION-BASED SAMPLE OF OLDER AUSTRALIAN ADULTS.


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Introduction/Aim: Little is known about how many Australians have been exposed to potentially hazardous agents in workplaces during their lifetime. Assessment of exposure status using only current job data can miss earlier exposures with potential long-term effects, and does not account for changes to occupational exposures over time.

This analysis determines proportions of a national population-based sample of Australian adults ever-exposed to potentially hazardous agents during their working life.

Methods: Individuals were recruited by telephone to participate in a study as controls to a national sample of Idiopathic Pulmonary Fibrosis (IPF) cases in Australia; individually matched 2:1 for age, sex and State. Consenting participants completed a telephone interview collecting data including demographics, self-reported environmental exposures and detailed occupational history. Each reported job was coded to the Finnish Job Exposure Matrix (FinJEM) to assign associated exposures during the relevant time period. Participants were considered exposed to an agent in a job if the probability of exposure was >25%. Data from each job in participants’ occupational history were collated to assess lifetime exposure status. Descriptive statistics were conducted to assess the proportions ever-exposed.

Results: 902 participants completed the interview (625 males and 277 females, mean±SD age 70.8±8.4), reporting a total of 2670 jobs. The FinJEM analysis of ever-exposure produced the following Results

Respirable dust 36.5%; Animal dust 13.6%; Iron 13.0%; Plant dust 13.0%; Asbestos 12.5%; Environmental tobacco smoke 11.0%; Moulds 10.3%; Quartz dust (Silica) 9.9%. Analysis of self-reported occupational exposure data yielded higher proportions: Dusty environment 50.6%; Gases/fumes/chemicals 47.0%; Asbestos 33.7%; Silica 19.5%. As expected, significantly more males were exposed than females for all agents (p<0.05).

Conclusion: A considerable number of older Australians have been exposed to potentially harmful agents within their working life. Data about these lifetime occupational exposures are required to examine potential occupational risk factors for diseases such as IPF.

Grant Support: National Health and Medical Research Council (Project Grant #1106601)

Declaration of interest statement: MJA holds investigator initiated grants for unrelated research from Pfizer and Boehringer-Ingelheim. The other authors declare that they have no competing interests.
Introduction: The development and progression of chronic lung diseases such as bronchiectasis involve recurrent episodes of severe bacterial respiratory infections. Inhaled environmental dust exposure can exacerbate immune responses to respiratory infections in mice. Whether dust exposure leads to more common and severe bacterial infections in the human airway is not yet known.

Aim: We aimed to determine the impact of community-sampled geogenic dust PM10 (particulate matter <10μm diameter) on immortalized human airway epithelial cells (NuLi-1). In particular, we aimed to determine its effects on non-typeable Haemophilus influenzae (NTHi) infection.

Methods: Geogenic dust was collected from remote towns in Western Australia and PM10 was extracted. NuLi-1 cells were exposed to PM10 (10μg/mL in PBS) in vitro for 24h before the addition of live NTHi-86 (MOI 10:1) for 3h. Trypan blue staining was used to determine cell viability, and bacterial infection (attachment and invasion) was determined using a gentamicin survival assay. Epithelial release of IL-6 and IL-8 was assessed using a bead based immunoassay.

Results: Pre-treatment of NuLi-1 cells with dust PM10 significantly increased the ability of NTHi to attach to and invade cells (p<0.001 and p=0.013, respectively). After dust exposure, the IL-6 response was increased (p=0.043) and the IL-8 response was suppressed (p=0.041).

Conclusion: Geogenic dust preparations increased NTHi infection of human airway cells and may contribute to more common and severe respiratory infections. This has important implications for lung health in individuals living in arid environments, such as those in remote Australia, who are exposed to high loads of geogenic dust.

Grant Support: BrightSpark Foundation, Raine Medical Research Foundation

Conflict of Interest: None

Introduction/Aim: Australian Aboriginal children have a much higher incidence of infectious diseases than their non-Aboriginal counterparts. Many Aboriginal communities are exposed to high levels of ambient dust from a geogenic (earth-derived) origin. We have previously reported an association between self-reported dust exposure and poor health in remote Aboriginal communities. The aim of this study was to examine the association between self-reported dust exposure and infectious diseases in Aboriginal children.

Methods: We collected data on self-reported dust levels and dust suppression programs from communities surveyed during the 2004 Environmental Health Needs Survey (EHNS) of remote Aboriginal communities in Western Australia. Health data on infectious diseases from hospitalisation records (2000–2002) and the Western Australia Aboriginal Child Health Survey (WAACHS) were linked to the EHNS. Odds ratios for the association between the health outcomes and exposure were calculated using multivariate logistic regression models within a multi-level framework to take into account the hierarchical structure of the data.

Results: Dust exposure levels were not associated with the prevalence of infectious diseases in the children (p > 0.09 for all comparisons). Lack of a dust suppression program was associated with an increased risk of having chronic ear infections (OR 1.70 95%CI [1.02-2.83], p = 0.04). Higher levels of dust exposure were associated with an increased risk of being hospitalised for upper (OR 1.79 95%CI [1.06-2.02]) and lower respiratory tract infections. Lack of a dust suppression program was associated with an increased risk of being hospitalised for skin infections (OR 2.52 95%CI [1.06-5.89]; 95%CI [1.24-2.76]) and gastrointestinal disease (OR 1.74 95%CI [1.02-2.96]).

Conclusion: While acknowledging that dust exposure in this instance was self-reported, our data suggest that exposure to high levels of geogenic dust may be contributing to the increased prevalence and severity of infectious disease in Aboriginal children.

Grant Support: Nil
TO-082

RESURGENCE OF COAL WORKER’S PNEUMOCONIOSIS IN QLD: PRELIMINARY RESULTS FROM A CLINICAL, OCCUPATIONAL AND RADIOLOGICAL REVIEW

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Aim: To identify the radiological pattern and severity of Queensland cases of coal worker’s pneumoconiosis and to correlate with clinical symptoms and occupational history.

Method: Retrospective review of CT and plain radiograph findings in twenty coal miner’s with radiological and clinical diagnosis of coal worker’s pneumoconiosis. Radiological grade of disease based on the ILO classification has been correlated with clinical symptoms, length of occupational exposure and job roles. Statistical analysis performed with Fisher exact test for categorical data and Wilcoxon test for continuous data.

Results: Preliminary results highlight the difficulties in diagnosis of coal worker’s pneumoconiosis. Although data collection is ongoing, the radiological grade in the majority of cases to date has been low grade (ILO 1). In addition, this case series highlights that in the early stages coal workers presents with soft or ground glass nodules that are low density and therefore extremely difficult to detect on plain radiograph. There has been no case of progressive massive fibrosis from coal exposure alone.

Clinical and occupational history is predictive of diagnosis with most patients having a symptomatic cough and a strong occupational history of dust exposure.

Conclusion: The resurgence of coal worker’s pneumoconiosis in Queensland has been in workers with occupational histories of heavy dust exposure who are usually symptomatic. Radiographic changes are often subtle, particularly on plain radiograph.

Grant Support: Nil

TO-083

EFFECT OF IN UTERO EXPOSURE TO CEILING PARTICLES ON POST-NATAL LUNG FUNCTION

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Introduction/Aims: Assessing the community health impacts of short-term particle exposure events (e.g. fires) can be difficult. One approach involves collecting particle samples from the roof-space of exposed dwellings as a proxy for particles that were generated during the exposure event. In follow up studies from a 2014 fire, we collected particles from ceilings of houses at different distances from the particle source. Using a mouse model, we aimed to determine whether in utero exposure to these particles alters post-natal lung function.

Methods: C57BL/6 mouse dams were intranasally exposed to ceiling particles (100μg in 50μL saline), carbon black particles (control) or saline alone at gestation day (E)13.5, E15.5 and E17.5. At two weeks of age, lung volume (TGV) and lung mechanics (airway resistance; Raw; tissue damping, G; tissue elastance, H) were measured in anaesthetised and tracheostomised mice using plethysmography and the forced oscillation technique (FOT) respectively.

Results: There was no effect of in utero exposure to ceiling dust or carbon particles on TGV in 2-week-old mice (female, P = 0.59; male, P = 0.73). Similarly, there was no effect of exposure to particles on Raw (female, P = 0.18; male, P = 0.88) or G (female, P = 0.46; male, P = 0.10). However, while there was no effect in females (P = 0.35), male mice exposed to ceiling dust (P = 0.04), but not carbon (P = 0.22) had higher H. The magnitude of the response was not associated with distance from the particle source.

Conclusion: In utero exposure to particles of ceiling dust increased lung stiffness, but only in male mice. These data suggest that exposure to ceiling dust may impact on lung growth in a sex specific manner. These responses appeared to be unrelated to the relative contamination of the ceiling dust with particles from a combustion source.

Grant Support: UTas Research Enhancement Grant Scheme; CAR CRE

Competing interests: The authors declare that they have no competing interests.
THE AUSTRALASIAN BRONCHIECTASIS REGISTRY – EARLY STEPS IN MAPPING THE IMPACT OF BRONCHIECTASIS IN AUSTRALIA AND NEW ZEALAND

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Introduction/Aim: Despite a growing worldwide recognition of bronchiectasis as a cause of substantial morbidity and healthcare utilisation, little is known about the prevalence or burden of disease for Australians and New Zealanders. The Australasian Bronchiectasis Registry (ABR) was established to fill this gap and to promote international collaborative research.

Methods: A comprehensive Registry has been developed by the Australasian Bronchiectasis Consortium with the support of Lung Foundation Australia (LFA), the European (EMBARC) and US Bronchiectasis Registries, with common data fields to ensure interoperability and future collaboration, and some unique data-fields to capture information specific to Australia and New Zealand.

Results: As the first step towards a regional standardised registry, the project has been initiated at 15 hospitals, including adult, paediatric, metropolitan and regional sites. Enrolment commenced at the pilot site (Concord Hospital) in March 2016 and as of October 2016 is being undertaken at 10 sites, with the remainder awaiting research governance approvals. To date, 261 patients have been enrolled; females 166 (64%), mean age 64.7 ± 19.4 years.

Conclusion: The ABR is a secure, web-based platform to capture the first comprehensive longitudinal data set of patients with bronchiectasis in Australia and New Zealand. Unique features include the use of opt-out consent, collection of paediatric data and linkage to PBS/Medicare data which will allow for the first time, a topographic map of bronchiectasis in Australia and New Zealand.

Grant Support: The ABR is an initiative LFA and supported by COPD Foundation, EMBARC, Bayer, Insmed, Aradigm, and Pfizer.
THE ROLE OF BRONCHOALVEOLAR LAVAGE (BAL) IN THE DIAGNOSIS OF OPPORTUNISTIC INFECTION IN IMMUNOCOMPROMISED HAEMATOLOGY PATIENTS
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Introduction: Immunocompromised haematology patients are at an increased risk of developing invasive pulmonary infections with significant morbidity and mortality. Where pulmonary sepsis is suspected, our usual practice is to perform chest imaging and refer for fibreoptic bronchoscopy (FOB). We evaluated the diagnostic yield of BAL, the impact of timing and complication rate in a single centre cohort.

Methods: We performed a retrospective study of patients with haematological conditions undergoing FOB between 31/8/2013 to 31/8/2016 by searching The Alfred Bronchoscopy Database and reviewing electronic medical records.

Results: Eighty-one patients underwent bronchoscopy. The average age was 53 years (range 19–78 years). Bronchoscopy was performed within 24 hours of HRCT in 22% and within 72 hours in 54%. Sixty percent of patients returned positive BAL cultures. Fungi were most common (69%), followed by bacterial, viral and malignant causes. Where a radiological diagnosis was suggested, this correlated with BAL findings in 50% of cases. BAL findings were concordant with pre-FOB sputum isolates in 1/20 and 3/10 viral swabs. No major complications were recorded; 16% experienced minor bleeding (<30ml). BAL yield may have been limited by electronic medical records.

Conclusion: BAL has an important role in the diagnosis of invasive pulmonary infections in haematology patients. Our centre demonstrates a high diagnostic yield, prompt performance of the procedure and likely low complication rate.

Declaration of Interests: None to declare.

Grant support: Nil.

COMMUNITY ACQUIRED RESPIRATORY VIRUS DETECTION: COMPARING THE DIAGNOSTIC LABORATORY WITH THE RESEARCH LABORATORY
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Introduction: Pathogen detection by hospital diagnostic microbiology laboratories (HDM) is the standard operating procedure for diagnosis. However, a causative pathogen is often not identified in symptomatic respiratory tract infections. Our study compared pathogen detection in a research laboratory with an HDM to determine differences in testing sensitivity for community acquired respiratory viruses (CARV).

Methods: Samples were derived from a prospective, longitudinal study of lung transplant (LTX) recipients. Swabs of donor and explanted lungs were collected as well as bronchoalveolar lavage (BAL) fluid on post-operative days 1, 7, 21, 42, 63 and 84. BAL sample replicates were sent to the HDM and to a research laboratory where multiple uniplex real-time PCRs were run for human rhinovirus (HRV), respiratory syncytial virus, influenza A (Flu A), influenza B (Flu B), parainfluenza virus (PIV) 1, 2, 3, and human metapneumovirus.

Results: 19 consecutive LTX subjects were recruited (bilateral: heart lung = 18:1) (age 48 ±14 years, mean ±SD) (M=9). Follow up was 59±37 days, range 7–120 days. Using a multiplex PCR, the HDM detected Flu A (n=1) and Flu B (n=1) on donor swabs plus HRV in 7/44 BAL samples. The research laboratory detected Flu A (n=4) in swabs of explanted lungs plus in BAL, HRV (n=7), Flu A (n= 23), Flu B (n = 2) and PIV (n=2). Overall concordance was 19/44, with an unweighted kappa coefficient of 0.113 (SE 0.054).

Conclusion: The real-time PCR used in the research laboratory appeared to have a greater sensitivity for CARV detection compared with the HDM multiplex PCR, perhaps reflecting the greater cycle threshold value allowing detection of lower viral loads in the research laboratory. CARV (particular Flu A) are common early after LTX. The importance of low levels of persistent CARV following LTX is yet to be elucidated.

Grant Support: Nil
PLEURAL PROCALCITONIN IS NOT USEFUL IN DIFFERENTIATING PARAPNEUMONIC FROM OTHER EFFUSIONS

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Introduction/Aim: Pleural effusions are a common complication of pneumonia and chronic diseases. Pleural procalcitonin has been postulated to be useful in differentiating parapneumonic effusions (PPE) from other aetiologies. We aimed to assess the utility of additional pleural procalcitonin measurement to differentiate PPE from non-PPE.

Methods: Consecutive patients with pleural effusion requiring thoracentesis were recruited, and clinical and anthropometric data recorded. Pleural fluid and serum were collected for standard analysis and additional procalcitonin level measurement. PPE were identified as: exudative effusions [Light's criteria (pleural/serum ratio for protein>0.5 or lactate dehydrogenase (LDH) >0.6)] with radiological evidence of a lower respiratory tract infection. Malignant pleural effusions were excluded. Data were compared using a Mann Whitney test or unpaired t-test, correlational analysis via a Spearmans, p <0.05 considered significant.

Results: 38 patients [(63.5±15.8)years; 19 male (mean±SD)] were recruited, 11 (28.8%) with PPE, while 27 (71%) were non-PPE. Pleural: serum ratios were not significantly different (p>0.05) for protein; 0.61 (0.57-0.65) (PPE; median, IQ range) and 0.51(0.36-0.63) (non-PPE), or LDH; 1.36(0.86-2.17) (PPE) and 0.90(0.38-1.35) (non-PPE). Serum procalcitonin was equivalent in PPE and non-PPE [0.2(0.1-0.33)ug/L vs 0.1 (0.1-0.21)ug/L respectively, p=0.07]. Pleural procalcitonin was significantly greater in PPE compared with non-PPE [0.21(0.17-0.41)ug/L and 0.1(0.1-0.15)ug/L respectively, p<0.05]. There were no significant differences between PPE or non-PPE for: serum CRP (40 (26-96)mg/L vs 27.5 (15.5-73.25)mg/L, pleural LDH (258 (112-365)U/L vs 147 (87-300)U/L, or serum WCC (8.3 (5-13)x10^9/L vs (7.6 (4.8-9.0)x10^9/L, all p>0.05. Serum and pleural procalcitonin correlated significantly in both PPE (r=0.99, p<0.0001) and non-PPE (r=0.613, p=0.0007).

Conclusion: Compared with non-PPE, pleural procalcitonin is significantly greater in subjects with PPE, and correlates with serum procalcitonin. However, the difference is small, and unlikely to assist in clinical management of PPE effusions.

Grant Support: Nil

IMPACT OF A SINGLE EDUCATIONAL INTERVENTION ON ANTIBIOTIC PRESCRIBING IN PNEUMONIA AND COPD IS NOT SUSTAINED BEYOND 6 MONTHS

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Aim: The emergence of multi-resistant organisms has been linked to injudicious antibiotic prescribing. We previously showed that a single education session increased adherence to antibiotic guidelines for Chronic Obstructive Pulmonary Disease (COPD) and Community Acquired Pneumonia (CAP). We aimed to measure the longer-term effect of educational intervention.

Methods: Data was collected from all Respiratory admissions to an Australian tertiary hospital for CAP and COPD exacerbations over three consecutive 6 month periods (Period 1: Oct 2013-Mar 2014; Period 2: Apr-Sep 2014; and Period 3: Oct 2014-Mar 2015). Following Period 1, Emergency and Respiratory Department staff attended a one-hour education session. Adherence to guidelines in Period 3 was compared to earlier periods using chi-squared tests for independent proportions.

Results: Period 1 comprised 194 patients (106 COPD, 88 CAP, 69.2±13.5 years, length of hospital stay (LOHS) 8.3±2.9 days, 2 deaths); Period 2: 285 patients (145 COPD, 139 CAP, 68.4±10.5 years, LOHS 8.0±6.5 days, 4 deaths); and Period 3: 196 patients (70 COPD, 126 CAP, 68.0±15.3 years, LOHS 5.8±5.1 days, 3 deaths). In Period 2, appropriate initial antibiotic prescribing for COPD improved from 46% to 71% (p<0.0001), but decreased to 42% in the subsequent 6 months, a significant reduction from Period 2 (p=0.0001) and comparable to pre-intervention rates (p=0.5). Guideline adherence for CAP improved from 31% to 46% immediately post-intervention (p=0.04), but declined towards pre-intervention rates in Period 3 (38.2%, p=0.3). There was a sustained improvement in appropriate change to antibiotics for CAP (13% in Period 1 to 34% in Period 3, p=0.004). Mortality was not significantly different between periods but LOHS was significantly lower in Period 3 for both COPD and CAP.

Conclusion: An educational intervention led to a significant increase in adherence to antibiotic guidelines for COPD and CAP, but this impact waned over time.
ELEVATED NEUTROPHIL–LYMPHOCYTE COUNT RATIO IS ASSOCIATED WITH INCREASED ADVERSE OUTCOME IN INFLUENZA

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Introduction/Aim: The neutrophil-lymphocyte count ratio (NLCR) has been demonstrated to be a useful predictive marker for disease severity. In this retrospective study, we investigated the prognostic impact of this biomarker in patients with influenza infection.

Methods: We retrospectively analysed patients with confirmed influenza A and/or B infection on nasopharyngeal swabs between 1 July 2015 and 31st September 2015. The clinical characteristics, laboratory markers including NLCR and radiological findings were assessed. Influenza complications including pneumonia, respiratory failure, intensive care unit (ICU) admission, length of stay, and 30-days mortality were evaluated. We used NLCR > 5 as a cut off value and the strength of association was reported as odds ratio (OR) and 95% confidence interval (CI).

Results: Total of 320 patients were included in this study. Mean age was 53.82 years. 4.69% (n=15) of the study cohort required ICU admission and inpatient mortality was 3.43% (n=11). The overall mean NLCR was 7.17±7.32 and was significantly higher in patients who developed complications. Patients with NLCR greater than 5 were significantly associated with increased risk of developing pneumonia (OR 3.19, 95% CI = 1.76 to 5.8, p=0.0001), respiratory failure (OR 2.56, 95% CI = 1.33 to 4.91; p=0.005), and had longer length of stay (116.3 hours vs 77.11 hours, p=0.0002). There was no significant association between rate of ICU admission and 30-days mortality with elevated NLCR.

Conclusion: A high NLCR is associated with increased unfavourable outcome in patients with influenza. The utility of NLCR as a predictive marker for pneumonia and respiratory failure in influenza to guide clinical decision making warrants further evaluation.

Grant Support: Nil

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SATURATED FATTY ACIDS, BUT NOT N-6 POLYUNSATURATED FATTY ACIDS OR CARBOHYDRATES, INCREASE AIRWAY INFLAMMATION IN NON-OBESE ASTHMATICS

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Introduction/Aim: We have previously shown that consumption of a high energy, fast food meal, rich in saturated fatty acids (SFA), n-6 polyunsaturated fatty acids (n-6PUFA) and simple carbohydrates (CHO), causes increased airway neutrophilia in asthma. This study aimed to compare the individual effects of each of these macronutrients on airway inflammation in asthma.

Methods: A randomised, crossover study in 23 adults with asthma (n=12 non-obese, n=11 obese) was conducted. Subjects consumed 3 different isocaloric meals (~1800kJ) on 3 separate occasions, with a minimum washout period of 7 days. The meals were: SFA (delivered as double cream and butter, 70% SFA), n-6PUFA (delivered as safflower oil, 72% n-6PUFA) or simple CHO (delivered as glucose confectionary, 100% CHO). Blood and induced sputum were collected at 0 and 4 hours after consumption of the meal for measurement of inflammatory cells. Postprandial changes in sputum cell gene expression were measured by microarray analysis.

Results: Total white blood cell and blood neutrophil counts increased in both the obese and non-obese subjects, following both of the high fat meals, while there was no increase following the high CHO meal. In sputum, neutrophils increased in the non-obese subjects following the high SFA meal (25.8(16.5-48.0)% median(IQR) versus 60.3 (6.0-94.5)%), but there were no changes following the n-6PUFA or CHO meals. There were no postprandial changes in airway cells in the obese subjects. The genes most significantly upregulated in the non-obese subjects following the high SFA meal were NLRP3 and TLR4.

Conclusion: SFA, but not n-6PUFA or CHO, cause an acute increase in airway neutrophilia in non-obese asthmatics, associated with an activation of the NLRP3 inflammasome. The airways of obese subjects appear to be resistant to the inflammatory effects of SFA, possibly due to desensitisation resulting from chronic exposure to high circulating SFA levels.

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HIGH FAT DIET-INDUCED OBESITY PROMOTES STEROID-RESISTANT ASTHMA THROUGH AN NLRP3 INFLAMMASOME-DEPENDENT MECHANISM

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Introduction/Aim: Steroid-resistant asthma is the major unmet need in asthma management. Effective treatments are urgently required, however, their development is hampered by a lack of understanding of the pathogenic mechanisms involved. An increasing body of evidence shows that a high fat diet (HFD) and/or obesity are associated with steroid-resistant asthma, however, the precise mechanisms involved remain unclear. Studies show that HFD-induced obesity causes increased airway hyper-responsiveness (AHR) in mice and that these responses are driven by NLRP3 inflammasome-mediated IL-1β responses. However, whether HFD/obesity affects steroid responsiveness and whether therapeutically inhibiting the NLRP3 inflammasome is effective for the treatment of HFD/obesity-induced experimental asthma is unknown.

Methods: We have developed a novel mouse model of HFD/obesity and combined it with an established model of ovalbumin-induced steroid sensitive, allergic airway disease (AAD). The effects of steroid treatment and the roles and potential for targeting of NLRP3-inflammasome responses in the lung in HFD/obesity-induced AAD were examined using therapeutic treatment with dexamethasone and a potent, highly selective NLRP3 inhibitor, MCC950.

Results: We show HFD results in significant increases in adiposity and weight gain. Significantly, HFD/obesity induces increased eosinophil numbers in the lung and steroid-resistant AHR in both the absence and presence of AAD. We also show that HFD results in an increase in NLRP3 staining in the airway epithelium as well as significant increases in active caspase-1 in whole lung tissue. Significantly, treatment with MCC950 suppressed HFD/obesity-induced steroid-resistant AHR in both the absence and presence of AAD.

Conclusion: We have developed a novel murine model of HFD-induced obesity that induces steroid-resistant AAD. We have also identified a previously unrecognised role for HFD/obesity-induced, NLRP3 inflammasome-mediated responses in the lung the development of steroid-resistant AAD. Thus, HFD/obesity induces increased NLRP3 inflammasome responses in the lung that may be therapeutically targeted in for the treatment of steroid-resistant asthma.

Grant Support NHMRC & HRMI Conflict of interest: None

IMPARED INDUCTION OF SLC26A4 PROMOTES RESPIRATORY ACIDOSIS AND SEVERE, STEROID-INSENSITIVE ASTHMA

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Introduction/Aim: Respiratory acidosis (RA) is associated with airways hyper-responsiveness (AHR) in severe, steroid-insensitive (SSI) asthma and treatment with sodium bicarbonate (NaHCO3) has been used to correct RA and improve lung function in severe asthma. Thus, dysfunction of the mechanisms that govern homeostatic acid-base balance in the airways may induce RA and the cardinal features of SSI asthma. CO2 produced by cellular respiration is hydrated by carbonic anhydrases into carbonic acid (H2CO3) that dissociates into H+ and HCO3-. These are transported in the plasma to the lungs where HCO3- is converted back into H2CO3 and CO2 that are expelled through breathing. Reduced lung function, which occurs in SSR asthma, impairs removal of volatile H2CO3 and CO2, resulting in acid accumulation and increased arterial PaCO2.

Patients with severe asthma often develop complications from increased PaCO2, which skews the PaCO2/HCO3- ratio resulting in increased H+ concentration and reduced pH.

Methods: We developed mouse models of respiratory infection and ovalbumin-induced SSI allergic airways disease (SSIAAD) that are highly representative of human disease. Microarray analyses of the lungs of mice with SSIAAD showed that all three infections decrease the expression of the chloride (Cl-)/HCO3- pump, Slc26a4, which is crucial for secretion of HCO3- into the airway lumen. Thus, we assessed the roles and potential for targeting of Slc26a4 and RA in SSIAAD.

Results: We show that respiratory infections suppress the induction of Slc26a4 in the airway epithelium in AAD. This is associated with increased levels of free H+ ions in bronchoalveolar lavage. Treatment with NaHCO3 during infection-induced SSIAAD suppressed steroid-resistant AHR. In a complementary gain-of-function study, administration of acetazolamide (a carbonic anhydrase inhibitor) in steroid-sensitive AAD, to mimic the effect of decreased Slc26a4 responses, induced steroid resistance of AHR.

Conclusion: We have identified a previously unrecognised role for Slc26a4 in promoting RA and SSIAAD.

Grant Support: NHMRC, HRMI
LIMITED EXACERBATION OF ASTHMA SYMPTOMS BY ELECTRONIC-CIGARETTE AEROSOL EXPOSURE IN A MOUSE MODEL OF ASTHMA

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Introduction/Aim: Electronic cigarettes (“e-cigarettes”) heat and aerosolise a liquid producing an inhalable aerosol often containing nicotine and flavourings. Very little is known about their potential to impact health – however some research indicates they increase airway resistance in healthy and asthmatic individuals and may facilitate asthma attacks. We aimed to assess the effects of e-cigarette use on lung function (early and late phase asthmatic responses; EPR and LPR respectively) in mice with and without existing ovalbumin-induced allergic airways disease.

Methods: We assessed volume and lung function using plethysmography and the forced-oscillation technique in naïve and asthmatic adult female BALB/c mice each minute for 30 minutes after an acute e-cigarette aerosol exposure. We also measured airway hyper-responsiveness (AHR) in a second group of mice 6 hours after 30 minutes of aerosol exposure. Four types of e-cigarette were tested.

Results: For EPR studies, acute e-cigarette exposure had little effect on lung function, except for mice exposed to vegetable-glycerin vapours, which exhibited immediate, temporary impairments in lung function. There was little difference between asthmatic and non-asthmatic mice. For LPR studies, previous e-cigarette exposure did not exacerbate functional asthmatic responses, and there was a protective effect of nicotine free e-cigarette excipient. Until more information is available, caution should be exercised when advocating e-cigarettes as a safe alternative to tobacco smoking, especially in asthmatics.

Conclusion: This study showed that acute e-cigarette aerosol exposure has limited effects on respiratory health in a mouse model of allergic airways disease. E-cigarette aerosol exposure had both negative and protective effects on respiratory health, apparently dependent on the type of e-cigarette excipient. Until more information is available, caution should be exercised when advocating e-cigarettes as a safe alternative to tobacco smoking, especially in asthmatics.

Declaration of interest: None.

Grant Support: Supported by an Asthma Foundation of Western Australia Project Grant.

MULTIDIMENSIONAL ASSESSMENT AND TARGETED THERAPY OF SEVERE PERSISTENT ASTHMA: A RANDOMISED CONTROLLED TRIAL

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Introduction/Aim: Severe asthma is a heterogeneous disease that causes significant disease burden, is often refractory to treatment and is associated with comorbidities and risk-factors that confound management. Multidimensional assessment (MDA) followed by management that is targeted to ‘treatable traits’ has been proposed as a superior treatment approach. We aimed to test the hypothesis that MDA and targeted-therapy for severe asthma would lead to improved control measured by the asthma control questionnaire (ACQ5).

Methods: A randomised controlled trial of MDA and targeted-therapy (Intervention group, IG) versus usual severe asthma care (UC) was conducted. Participants with severe asthma (n=55) underwent a baseline MDA and 16 week follow-up. The intervention involved individualised therapies tailored to treatable traits of the airway, comorbidities, risk-factors and self-management, delivered via case-management. The UC arm received evidence-based treatment within a severe asthma clinic.

Results: Twenty-eight participants were randomised to IG and 27 to UC; 65.5% were female, with a mean (SD) age of 52.2 (13.9) years and post-bronchodilator FEV1 was 73.0% (21.2).

The intervention resulted in significant improvements in asthma control compared to UC; mean difference (SE) in ACQ5 post IG −0.61 (0.18) compared to 0.07 (0.19) for UC, p=0.01. IG also resulted in significant improvements in health-status, mean difference (SE) in Asthma Quality of Life Questionnaire was 1.01 (0.18) compared to −0.01 (0.18) for UC, p=0.001. Sputum eosinophils (%) increased in the UC 8.20 (3.49) compared to −2.52 (3.73) for IG, p=0.04. There was also a significant reduction in IG CRP; −4.90 (1.42) compared to UC 0.61 (1.36), p=0.01.

Conclusion: Management of severe asthma that is based on multidimensional assessment followed by therapy targeted to treatable traits results in significant improvements. This study supports a precision-based medicine approach in severe asthma.

Grant Support: Hunter Medical Research Institute, University of Newcastle, John Hunter Hospital
PHYSICAL INACTIVITY AND SEDENTARY TIME IN SEVERE ASTHMA

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Introduction/Aim: Physical inactivity and sedentary behaviours are likely to be particularly extreme in severe asthma (SA) populations, secondary to a high symptom and comorbidity burden. The degree of inactivity in SA however has not been quantified. We aimed to objectively measure physical activity (PA) and sedentary time (ST) in patients with SA compared to controls, and describe the correlations of these behaviours with clinical outcomes.

Method: A cross-sectional study of adults with physician diagnosed SA (n=65) and healthy controls (HC) (n=33) was conducted. Participants underwent a multidimensional assessment involving measurement of lung function, exercise capacity, airway and systemic inflammation, asthma control and quality of life. PA and ST were measured for 14 consecutive days using a triaxial accelerometer (ActiGraph wGT3X-BT).

Results: The SA group included 33 (50.7%) females, aged (mean±SD) 53.4±15.7 years. The HC group included 26 (76%) females, [median, IQR] age 39 [29–62] yrs. Compared to controls, people with SA spent significantly less minutes/day in moderate and vigorous intensity PA: 22.7 [12.9–47.0] versus 69.0 [58.7–83.6] (p<0.01); and took significantly less minutes/day in moderate and vigorous intensity PA than controls: 195.18 (58.22) versus 166.53 (50.76) (p<0.05). No statistically significant differences in ST were found between the two populations. There were moderate correlations between PA and ST and systemic inflammation, exercise capacity, asthma control and quality of life.

Conclusion: Compared to controls, people with SA perform less moderate intensity activity, despite engaging in more light physical activity. The amount of activity is associated with important disease outcomes. These data show the potential benefits of targeting inactivity as a non-pharmacological strategy in the management of severe asthma.

Key Words: severe asthma, physical activity, sedentary behaviour, correlation.

Nomination for New Investigator Award

Grant Support: John Hunter Hospital Charitable Trust.

LUNG FIBROBLASTS OF IPF PATIENTS DISPLAY SENESCENCE-LIKE FEATURES IN VITRO

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Introduction/Aim: Idiopathic pulmonary fibrosis (IPF), a fatal lung disease of unknown etiology, occurs predominantly in the elderly. The mechanisms that link aging with the persistent accumulation of lung fibroblasts (LFs) in IPF, remain incompletely understood. Fibroblast senescence, characterized by growth arrest, apoptosis resistance and a senescence-associated secretory phenotype (SASP), occurs in aging and may contribute to IPF. The aim of this study was to characterize senescence in human LFs in vitro, and to ascertain the role of mitochondrial dysfunction in the acquisition and/or reinforcement of the senescent phenotype.

Methods: IPF-LFs lines established from lung tissue of IPF patients were compared with LFs of age-matched controls (Ctrl-LFs). Cell proliferation, telomere length, SASP cytokine production, senescent-associated β-galactosidase (SA-β-Gal) activity and levels of p16 and p21 were assessed by cell counting, PCR, ELISA, cytochemistry and immunoblotting respectively. Increased levels of mitochondrial DNA and superoxide, features of mitochondrial dysfunction, were evaluated by PCR and the MitoSox fluorogenic dye respectively. Rotenone was used to disrupt mitochondrial activity.

Results: IPF-LFs displayed slower growth accompanied by decreased levels of cellular protein and telomere length when compared with Ctrl-LFs (p<0.05). Conversely, IPF-LFs were more resistant to H2O2-induced cytotoxicity than Ctrl-LFs (p<0.05). IPF-LFs also exhibited increased SA-β-Gal activity, SASP cytokine production and levels of p16 and p21 (p<0.05). Furthermore, IPF-LFs had lower levels of the mitochondria-driven proteins Bcl-2 and Bax, but higher levels of mitochondrial DNA and superoxide production than Ctrl-LFs. Rotenone (0.6 μM) increased mitochondrial superoxide in Ctrl-LFs, preceding increases in markers of senescence.

Conclusion: Our study suggests that IPF-LFs exhibit senescent like features, including replicative arrest, resistance to oxidative stress-induced cytotoxicity and induction of a SASP. Mitochondrial dysfunction is tightly linked with senescence and is likely to contribute to the healing/repairing malfunction of LFs in IPF.

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NON-ANTIBIOTIC MACROLIDES RESTORE AIRWAY MACROPHAGE PHAGOCYTIC FUNCTION WITH POTENTIAL ANTI-INFLAMMATORY EFFECTS IN CHRONIC LUNG DISEASES

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Introduction/Aim: We reported defective efferocytosis associated with cigarette smoking and/or airways inflammation in chronic lung diseases including chronic obstructive pulmonary disease and severe asthma. We also showed defects in phagocytosis of non-typeable H. influenzae (NTHi), a common colonizer of the lower airway in COPD. These defects could be substantially overcome with low-dose Azithromycin; however, chronic usage may induce bacterial resistance. We investigated two novel macrolides, GS-459755 (2'-desoxy-9-(S)-erythromycin-based 2'-desoxy molecule) and GS-560660 (Azithromycin-based 2'-desoxy molecule) with significantly diminished antibiotic activity against S. aureus, S. pneumonia, M. catarrhalis, and H. influenzae.

Methods: Macrolide effects on efferocytosis, phagocytosis of NTHi, cell viability, receptors involved in recognition of apoptotic cells and/or NTHi (flow cytometry), secreted and cleaved intracellular IL-1β, staining with DAPI (immuno-fluorescence/ confocal microscopy) and NLRP3, were tested on primary alveolar macrophages and THP-1 macrophages ± 10% cigarette smoke extract.

Results: Dose response experiments showed optimal pro-phagocytic effects of GS-459755 and GS-560660 at low concentrations of 0.5-1 μg/mL, comparable to our findings with Azithromycin. Both macrolides significantly improved phagocytosis of apoptotic cells and NTHi (eg, increases in efferocytosis and phagocytosis of NTHi; GS-459755 23% and 22.5% vs NTHi (flow cytometry), secreted and cleaved intracellular IL-1β (CBA, immuno-fluorescence/ confocal microscopy) and NLRP3, were tested on primary alveolar macrophages and THP-1 macrophages ± 10% cigarette smoke extract.

Conclusion: We conclude that GS-459755 and GS-560660 may be useful for reducing airway inflammation in chronic lung diseases without inducing bacterial resistance.

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ANTENATAL VITAMIN C ADMINISTRATION IN SHEEP IS PROTECTIVE AGAINST FETAL HYPOXEMIA-INDUCED OXIDATIVE STRESS AND AIRWAY REMODELLING IN EARLY ADULTHOOD

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Introduction/Aim: Chronic fetal hypoxemia is a common pregnancy complication associated with intrauterine growth restriction (IUGR) and respiratory disease at birth and in later life. The molecular mechanisms that are responsible are not understood. Hypoxia and oxidative stress are linked to poorer cardiovascular outcomes at birth and in later life and antenatal antioxidant administration ameliorates these effects. Here, we investigate the effect of chronic fetal hypoxemia and antenatal administration of the antioxidant, Vitamin C, on molecular regulation of lung structure and function in lambs.

Methods: Chronically catheterised pregnant sheep carrying female singletons were exposed to normoxia (N; n=20) or hypoxia (H; n=18; 10% O2) with either saline (NS; n=11; HS; n=8) or Vitamin C (VC; n=9; HVC; n=10; maternal 200mg/kg i.v. daily) from 105-138d (term, ~145d). Lung tissue was collected from lambs 9 months after birth. Lung expression of genes as markers of oxidative stress (NADPH oxidase 4, NOX-4), airway remodelling (elastin, ELN) & surfactant maturation (surfactant proteins (SP) – A, B, C, D) were quantified by qRT-PCR. Numerical density of SP-B positive cells in lung tissue was determined by point counting. Data were analysed by two-way ANOVA (P<0.05) for treatment (N vs. H) and drug (S vs. VC).

Results: H induced fetal growth restriction but there was no effect of H or VC on body weight or relative lung weight in lambs at 9 months. H increased gene expression of NOX-4 (NS: 0.0029±0.00017 vs HS: 0.0039±0.00056 mean normalised expression (MNE)) & surfactant maturation (surfactant proteins (SP): A, B, C, D) were quantified by qRT-PCR. Numerical density of SP-B positive cells in lung tissue was determined by point counting. Data were analysed by two-way ANOVA (P<0.05) for treatment (N vs. H) and drug (S vs. VC).

Conclusion: Here we show effects of chronic fetal hypoxemia on molecular programming of oxidative stress and airway remodelling in the lung in early adulthood and that maternal antenatal antioxidant treatment is protective, offering insight into mechanism and possible treatment to improve offspring respiratory outcomes.

Grant Support: British Heart Foundation & NHMRC of Australia
ROLE OF INCREASED IRON LEVELS IN THE PATHOGENESIS OF LUNG DISEASE

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Introduction: Iron is essential for many biological processes. Too much or too little iron can result in a wide variety of pathologies. Altered iron levels and/or dysregulated iron homeostasis have been associated with a number of lung diseases, including chronic obstructive pulmonary disease, lung cancer and cystic fibrosis, however, the mechanisms that underpin these associations and whether iron plays a role in the pathogenesis of lung disease are yet to be fully elucidated.

Aim: To determine the effects of systemic iron overloading on iron levels in the lung and to assess the effects of increased iron on lung structure and function.

Methods: Levels of systemic and pulmonary iron and lung structure and function were assessed in transferrin receptor (TFR)2 mutant mice and wild-type (WT) BALB/c mice fed a 2% carbonyl iron diet compared to WT and normal diet controls, respectively. The effects of increased iron loading on murine models of bleomycin-induced fibrosis and house dust mite (HDM)-induced allergic airway disease (AAD) were also assessed.

Results: Excess iron accumulation was observed in the lungs in both the genetic and diet-induced models of iron overloading. Increased iron levels in the lung were associated with emphysema-like alveolar enlargement, small airways collagen deposition, alterations in baseline lung function and increased airways hyper-responsiveness (AHR). Whilst iron overloading in the absence of bleomycin administration results in the generation of fibrosis in the small airways and AHR to similar levels observed with bleomycin-induced fibrosis, iron overloading had minimal additional effects when combined with bleomycin. Iron overloading also resulted in increased eosinophilic inflammation and severe AHR in HDM-induced AAD.

Conclusion: These data show that increased iron levels in the lung results in emphysema and airways fibrosis that corresponds with reduced function. These models will be used to better understand the role of iron in the pathogenesis of lung disease.

Grant Support: NHMRC & HMRI

Conflict of interest: None

DYSREGULATED STAT3 SIGNALING INDUCES AND REINFORCES FIBROBLAST SENESCENCE IN LUNG FIBROBLASTS OF IPF PATIENTS

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Introduction/Aim: Idiopathic pulmonary fibrosis (IPF) is a chronic fibrosing interstitial pneumonia of unknown cause and has a median survival of only 3 years. We have previously shown that dysregulated activation of the transcription factor STAT3 characterizes functionally different subsets of fibroblasts, the presence of which correlates with IPF progression. The question of what drives these phenotypically divergent cells and prolongs their lifespan remains unanswered. Our hypothesis is that activated STAT3, a transcription factor that is increased in fibrotic lung tissue of IPF patients, contributes to senescence via its roles in cytokine expression and mitochondrial dysfunction. The aim of this work is to evaluate whether STAT3 has a role in senescence of lung fibroblasts (LFs) in IPF.

Method: Primary cultures of LFs were established from lung tissue of IPF patients (IPF-LF) and control donors (Ct-LF). Senescence was induced by exposing cells to hydrogen peroxide (H2O2, 150 μM) for 2 hr. STAT3 activation was assessed at different passages or following H2O2 by immunoblotting. Senescence was evaluated by measuring senescence-associated-β-galactosidase (SA-β-Gal) activity, IL-6 production and expression of cell-cycle arrest protein p21. Mitochondrial superoxide production was assessed using the mitosox fluorogenic assay. STAT3 was inhibited with pharmacological inhibitors or by transfection with siRNA.

Results: A higher percentage of IPF-LFs were positive for SA-β-Gal at earlier or equivalent passages than the Ct-LFs, as were levels of p21 and mitochondrial superoxide (p<0.05). At both early (6) and late (15) passages, IPF-LFs (n=4) produced more IL-6 than Ct-LFs (n=3) (p<0.05). Baseline levels of pSTAT3 were higher in IPF-LFs and increased in Ct-LFs following treatment with H2O2. Targeting STAT3 attenuated IL-6 production, levels of p21 and mitochondrial superoxide production.

Conclusion: Our data suggests that IPF-LFs display accelerated senescence in association with increased STAT3 activation and superoxide production. STAT3 contributes to senescence in IPF through regulating mitochondrial activity.

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Key Words
Nomination for New Investigator Award
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FIUBLIN-1C PLAYS CRITICAL ROLES IN LUNG REMODELLING IN IDIOPATHIC PULMONARY FIBROSIS

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Introduction: Idiopathic pulmonary fibrosis (IPF) is a chronic, progressive, and lethal interstitial fibrotic lung disease with poor survival rate after diagnosis. Airway and/or lung remodelling are major features of IPF and associated with accumulation of extracellular matrix (ECM) proteins. Fibrillin-1 (Fbn1), an important ECM protein has four variations (Fbn1a/b/c/d) and stabilizes other ECM proteins. Fbn1c, one of Fbn1 variants, predominates in both humans and mice, and involves in lung fibrosis.

Aims: To determine Fbn1c levels in IPF patients and elucidate the role Fbn1c in pathogenesis of pulmonary fibrosis.

Methods: Fbn1c was measured in IPF patients and bleomycin-induced lung fibrosis model. Fbn1c deficiency (−/−) mice were used to investigate the mechanism of Fbn1c regulated remodelling in lung fibrosis.

Results: Fbn1 was increased in lung tissue of IPF patients and Fbn1c protein was increased bleomycin-induced experimental lung fibrosis in mice. Fbn1c−/− mice had reduced airway remodelling and lung fibrosis, and were protected against lung function impairment after 28 days bleomycin challenge. In experimental lung fibrosis, Fbn1c interacted with fibronectin, peritostin and tenasin-c to stabilise collagen deposition around the small airway and in whole lungs. Fbn1c regulated transforming growth factor-β (TGF-β) activation by binding with latent TGF-β binding protein (LTBP1), and mediated phosphorylated Smad3 level, a downstream molecular of TGF-β pathway. This process also regulated myofibroblasts number and collagen deposition. Interestingly, Fbn1c−/− mice also had less inflammatory cells, and interleukin (IL)-33 in lungs of experimental lung fibrosis than WT mice.

Conclusions: Our data demonstrates a novel role of Fbn1c in lung remodelling in IPF. Inhibition of Fbn1c may be a therapeutic target in IPF.

Grant Support: NHMRC grant, LFA/Lizotte Family Research Award

CHRONIC OBSTRUCTIVE PULMONARY DISEASE 2

HOME-BASED REHABILITATION FOR COPD USING MINIMAL RESOURCES: AN ECONOMIC EVALUATION

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Introduction/Aim: Pulmonary rehabilitation (PR) is a cornerstone of care for COPD. A new model using a home-based programme has shown equivalent short-term clinical outcomes to a centre-based programme for comparable direct intervention costs. This economic evaluation sought to determine and compare the costs and benefits of both intervention models during the 12-month follow-up period.

Methods: Participants with stable COPD were randomly assigned to complete either home or centre-based PR. Complete case analysis was undertaken for participants who completed the Short-form 36 (SF-36v2) questionnaire at the end of PR and 12 months following programme completion, to enable preference-based classification of health status using the SF-6D and calculation of quality-adjusted life years (QALYs). Direct provider costs for healthcare utilisation were collected from hospitals and the Department of Health & Human Services during the 12-month follow-up period.

Results: Complete SF-36 data were available for 59 participants (36 female, mean [SD] age 71 [8] years, FEV1 50 [20] %predicted) who undertook a centre-based programme (74%) and 60 people (33 female, mean age 68 [9] years, FEV1 50 [20] %predicted) who undertook a home-based programme (70%). Participants in the home-based programme had lower costs for COPD-related outpatient appointments (mean difference $415, 95%CI -$644 to $222) and tended to have lower costs for hospital admissions ($4600, 95%CI -$10,331 to $1130). Overall healthcare provider costs tended to be lower in the home-based group ($2450). The difference in QALYs was not significantly different between groups (0.021, 95% CI 0.022 to 0.064).

Conclusion: In the 12 months following PR, healthcare provider costs were equivalent or lower for home-based participants compared to people undertaking a centre-based programme. This group also demonstrated equivalent QALYs. These results support the clinical implementation of this new model of PR.

Grant Support: Lung Foundation Australia, National Health and Medical Research Council

Declaration of interest: None Declared

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ELECTROCARDIOGRAMS AND CHEST RADIOGRAPHS IN PATIENTS WITH ABNORMAL CARDIAC BIOMARKERS DURING EXACERBATIONS OF COPD

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Introduction/Aim: Biochemical evidence of cardiac dysfunction is common in exacerbations of chronic obstructive pulmonary disease (COPD), even in patients without clinically suspected cardiac disorders. We investigated associations between abnormal N-terminal pro-B-type natriuretic peptide (NT-proBNP) and troponin T and abnormalities on electrocardiograms and chest radiographs in patients hospitalised for exacerbations of COPD.

Methods: Electrocardiograms were assessed for tachycardia, atrial fibrillation, right and left ventricular hypertrophy, and ischaemic changes in 389 patients. Chest radiographs were assessed for signs of heart failure in 350 patients. All assessments were done by two independent examiners blinded to cardiac biomarker status. Associations between abnormalities with at least moderate inter-rater agreement and high cardiac biomarkers (NT-proBNP >220pmol/L and troponin T >0.03ug/L or high-sensitivity >50ng/L) were analysed.

Results: High NT-proBNP values were associated with atrial fibrillation (22% vs. 6%), right ventricular hypertrophy (24% vs. 15%), left ventricular hypertrophy (15% vs. 4%), ischaemia (59% vs. 33%) on electrocardiogram and cardiomegaly (42% vs. 20%) on chest radiographs. High troponin T values were associated with tachycardia (65% vs. 41%), right ventricular hypertrophy (26% vs. 15%) and ischaemic changes (60% vs. 36%) on electrocardiogram. None of these tests were very sensitive or specific for biochemical indicators of cardiac dysfunction: the best-performing indicator for NT-proBNP was ischaemic change (sensitivity 65% and 36%) on electrocardiogram and chest radiographs respectively.

Conclusion: Abnormal electrocardiograms and chest radiographs are associated with abnormal cardiac biomarkers and may indicate cardiac disease in patients with exacerbations of COPD. However, electrocardiograms and chest radiographs have poor sensitivity and specificity for diagnosing acute cardiac dysfunction in this setting. Cardiac biomarkers provide additional information about acute cardiac dysfunction in patients with exacerbations of COPD.

Grant Support: Partially supported by the Waikato Medical Research Fund and the Heart Foundation New Zealand

MicroRNA-21 DRIVES EXPERIMENTAL CHRONIC OBSTRUCTIVE PULMONARY DISEASE THROUGH A SATB1/B AXIS

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Introduction/Aim: Chronic obstructive pulmonary disease (COPD) is the 3rd leading cause of chronic morbidity and death worldwide. Cigarette smoking is the major cause of COPD in developed countries. Current therapies have limited efficacy, fail to halt disease progression, and can cause adverse side effects. Effective treatments are urgently required, however, their development is hampered by a lack of understanding of the mechanisms that promote the pathogenesis of disease. The functions of microRNAs (miRs) in health and disease are firmly established and their aberrant expression has been associated with several lung diseases, including COPD. This study aimed to identify miRs that promote the development of COPD and assess the therapeutic potential of targeting the miRs that we identify.

Methods: We performed miR microarray analyses of the lungs of mice exposed to our highly representative, cigarette smoke (CS)-induced model of experimental COPD. We identified miR-21 as one of the highest upregulated miRs. The roles and potential for targeting of CS-induced miR-21 in the lung were examined using treatment with a specific miR-21 inhibitor (antagomir, Ant-21).

Results: Lung miR-21 expression was increased throughout CS exposure in experimental COPD. Reduced lung function in human COPD patients correlated with lung miR-21 expression. Treatment with Ant-21 inhibited CS-induced lung miR-21 expression and suppressed airway inflammation and small airway fibrosis, and improved lung function, in experimental COPD. In silico analyses identified a potential miR-21/SATB1/S100A9/NF-κB axis. SATB1 is a putative miR-21 target that negatively regulates S100A9, a known inducer of NF-κB activity. Significantly, CS exposure decreased lung SATB1 in our model, and Ant-21 treatment restored SATB1 levels and decreased S100A9 expression and NF-κB activity.

Conclusion: We have identified a previously unrecognised pathogenic role for a miR-21/SATB1/S100A9/NF-κB axis in experimental COPD. Our data highlights miR-21 as a novel therapeutic target for the treatment of COPD.

Grant Support: NHMRC

Conflict of interest: None
RELATIONSHIP BETWEEN TLCO, EMPHYSEMA INDEX, FEV1 AND PULMONARY ARTERY DISTENSIBILITY - A SUB-ANALYSIS OF THE 4C COHORT

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Introduction/Aim: Pulmonary hypertension is an important complication of Chronic Obstructive Pulmonary Disease (COPD), however it can be difficult to identify. Reduction in Carbon Monoxide Transfer Factor (TLCO) is a feature of both emphysema and pulmonary hypertension. CT can quantify emphysema via the emphysema index (EI). In addition, pulmonary artery distensibility (PAD) is known to correlate with mean pulmonary artery pressure measured at right heart catheterisation and can be measured simultaneously using ECG-gated CT. The impact of pulmonary artery distensibility on the relationship of FEV1 and Emphysema Index (EI) with TLCO has never been examined. The aim of this study is to determine the effect of PAD, FEV1(%) and EI on TLCO, with the hypothesis that the addition of PAD data would help account for variability in TLCO unexplained by the other two parameters.

Methods: Patients were analysed from the 4C cohort, a prospective group of patients who were recruited over a 1 year period during acute exacerbations of COPD and had subsequent ECG gated CT and pulmonary function tests performed during stable periods. After excluding patients with missing data, significant motion artefact on CT and those unable to perform TLCO, 34 patients were entered into a multiple linear regression model.

Results: TLCO corrected for haemoglobin was entered as the dependent variable and in order FEV1(%), EI and PAD were entered into the regression model as independent variables. Overall, the model was able to explain 62.5% of the variation in TLCO (Corrected R² 0.625). Both FEV1(%) and EI were significantly correlated with TLCO after controlling for the other variables (FEV1(%) r = 0.512 and P<0.001, EI r = -0.431 and p = 0.003) However, after controlling for FEV1pp and EI, PAD was not significantly correlated with TLCO (r = -0.055, p = 0.637)

Conclusion: FEV1(%) was significantly positively correlated and Emphysema Index was significantly negatively correlated with TLCO after controlling for the other variable. The addition of PAD data did not appear to add to the analysis.

Grant Support: Nil

THE PREVALENCE OF REDUCED TRANSFER FACTOR FOR CARBON MONOXIDE (TLCO) IN SMOKERS WITH NORMAL SPIROMETRY

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Introduction/Aim: The prevalence of Chronic Obstructive Pulmonary Disease (COPD) using spirometry criteria has been extensively studied, however little evidence exists for the usefulness of Transfer Factor for carbon monoxide (TLCO) in the diagnosis and classification of COPD. The guidelines for the Global Initiative for Chronic Obstructive Lung Disease (GOLD) currently utilise spirometry predominately to aid a physician-diagnosis of COPD. Since only a relatively small proportion of smokers develop COPD, we aim to assess how TLCO can serve as a useful early biomarker for the detection of lung disease in younger smokers with normal spirometry.

Methods: We performed a cross-sectional analysis of subjects who have had both spirometry and TLCO measured at the Respiratory Laboratory at Monash Lung & Sleep, Monash Health. Subjects included male and females aged 40 to 60 years with normal spirometry and at least 10 pack-years smoking history.

Results: One hundred and six subjects aged between 40 and 60 with normal spirometry and significant smoking histories were identified. Of this cohort, 77 subjects (73% of the group) had a TLCO below their lower limit of normal (using ATS criteria). The cohort was equally divided into 53 males (of whom 33 had a low TLCO) and 53 females (of whom 43 had a low TLCO)

Conclusion: A low transfer factor for carbon monoxide (TLCO) is prevalent amongst younger smokers with normal spirometry. This low TLCO may be an important early indicator of potential lung damage in COPD.

Grant Support: Nil
SELF-MANAGEMENT INTERVENTIONS INCLUDING COPD EXACERBATION ACTION PLANS IMPROVE HOSPITALISATION RATE AND HEALTH-RELATED QUALITY OF LIFE – A COCHRANE REVIEW

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Introduction/Aim: Exacerbation action plans are considered as a key component of a COPD self-management intervention. Studies of these interventions show contradictory results. In this systematic Cochrane review, we have assessed the effectiveness of COPD self-management interventions that include exacerbation action plans compared to usual care.

Methods: After a literature search, full-text articles were independently assessed by two authors. Study inclusion criteria were: randomised controlled trials published from 1995–May 2015; self-management interventions that included a written action plan for COPD exacerbations; and an iterative process between patient and healthcare provider.

Results: Of 1,633 identified records, 22 studies were included with follow-up times between two and 24 months. Statistically significant beneficial effects of self-management were detected on health-related quality of life (St George’s Respiratory Questionnaire total score, mean difference 2.69 (95% CI −4.49 to −0.90), 10 studies, 1,799 participants, I²: 46%) and respiratory-related hospitalisations (OR 0.69 (95% CI 0.51 to 0.94), 14 studies, 3,157 participants, I²: 57%). Furthermore, a borderline significant trend towards fewer all-cause hospitalisations (OR 0.74 (95% CI 0.54 to 1.03), 10 studies, 2,467 participants, I²: 62%) was observed. No effects on all-cause mortality were found (risk difference 0.00 (95% CI −0.02 to 0.03), 16 studies, 3,296 participants, I²: 48%).

Conclusion: Self-management interventions that include a COPD exacerbation action plan are associated with better health-related quality of life and lower probability of respiratory-related hospitalisations. No excess all-cause mortality risk was observed.

Grant Support: Lung Foundation Australia / Cochrane Airways Australia Scholarship 2016, Lung Foundation Netherlands (grant number 3.4.11.061).

Interventional Pulmonary / Bronchology

EBUS TBNA PRE-SURGICAL STAGING HIGHLIGHTS THE IMPORTANCE OF CONFIRMING CYTOLOGICAL NEGATIVITY IN CASES OF TECHNICALLY N3 NSCLC ON PET

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Introduction/Aim: Following radical treatment of non-small cell lung cancer (NSCLC), surveillance is recommended. Endobronchial ultrasound and transbronchial needle aspirate (EBUS-TBNA) has emerged as a bronchoscopic modality for sampling pulmonary lymph nodes detected on surveillance imaging. Previous treatments that patients have received may make reinterpretation of their imaging, and perhaps bronchoscopic lymph node sampling difficult.

Method: We performed an single-centre audit of all patients who underwent EBUS-TBNA to investigate abnormal CT lymph node appearance following radical treatment for NSCLC from 2005 to 2015. A combination of bronchoscopy reports, pathology reports, imaging reports and chart reviews were used to identify patient information.

Results: Overall, 47 cases were identified. The median duration between treatment and re-biopsy was 30.5 months. PET scans were performed in 57.4% of patients and of those the pulmonary lymph nodes were the sole sites of metastases in 74.1% of cases. The nodes sampled were N1 (34.0%), N2 (72.3%) and N3 (2.1%). The frequency of TBNA samples positive for malignancy was 78.7%. The frequency of biopsies positive for malignancy post surgery was 89.2%, post chemoradiation 50% and post radiation alone 33.3%. Patients with samples positive for malignancy went on to have a change in their treatment in 51.4% of cases. Of those, 72.2% received chemotherapy, 38.9% received radiation and 11.1% received surgery.

Conclusion: EBUS-TBNA has a very high yield in detecting regional lymph node recurrence of disease in the post radical treatment setting. Additionally, the regional lymph nodes were frequently the only readily available biopsy target. The yield of positive biopsies was higher in this case series in surgical patients compared to chemoradiotherapy and radiation. EBUS-TBNA allowed for a change in management in a significant number of patients.

Key Words: EBUS-TBNA, endobronchial ultrasound
Nomination for New Investigator Award: N/A
Grant Support: Nil
Declaration of Interest: There were no competing interests in this study.
A NOVEL MANAGEMENT OF DISTAL ACTIVE HAEMOPTYSIS BY AN INTRABRONCHIAL VALVE AND TISSEEL

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Introduction: TISSEEL (combination of synthetic aprotinin, factor VIII and fibrinogen combined with Thrombin) is a fibrin sealant indicated for use as an adjunct to haemostasis in patients undergoing surgery when control of bleeding by conventional surgical techniques is ineffective or impractical. It is widely used in general surgical procedures but remains a non-standard approach in the management of haemoptysis.

There have been recent case reports describing its effective usage in massive haemoptysis where other techniques such as bronchial embolisation, bronchoscopic suctioning or surgical resection have failed or may be ineffective. However, when TISSEEL is used independently endoscopically, migration, dislodgment or even expectoration of the fibrin clot is a common complication that leads to recurrence of haemoptysis.

We present the first ever reported case of successful endobronchial use of TISSEEL in controlling active distal bronchial bleeding. A 67 year old lady suffering from ongoing haemoptysis secondary to recurrent non small cell lung cancer involving lateral segment of left lower lobe (LB9) underwent therapeutic bronchoscopy utilising a therapeutic video scope (Olympus T180) introduced via a rigid bronchoscope which provided secure airway access. After identifying the source of bleeding, 1 ml of TISSEEL was injected proximally on top of the valve. A size 6 Spiration IBV was then inserted to add stability and prevent expectoration of the fibrin clot. A further 1 ml of TISSEEL was then injected proximally on top of the valve. There was excellent seal after the procedure with resolution of bleeding.

Conclusion: The combination of TISSEEL and an endobronchial valve provides a safe and effective therapeutic option for the medium term control of distal bronchial bleeding secondary to malignancy.

Grant Support: Nil

ENDOBRONCHIAL VALVE INSERTION OUTCOMES FOR EMPHYSEMA: A RETROSPECTIVE ANALYSIS IN A LOCAL PUBLIC HEALTH NETWORK.

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Introduction/Aim: Lung volume reduction (LVR) with endobronchial valves (EBV) for severe emphysema has established evidence demonstrating subjective and objective clinical response. In Australia, the majority of these procedures are funded and performed in the private sector. In this audit, we assessed a cohort of patients treated in a public hospital setting after the implementation of a publically funded LVR-EBV program.

Methods: Patients were selected from established criteria by a multidisciplinary team. 10 patients underwent EBV insertion in the first 18 months of this program. One patient who did not have complete post procedure lung function was excluded. The change in residual volume (RV), total lung capacity (TLC) and relative change in FEV1 was calculated for each patient from testing done prior to EBV insertion compared to three month post procedure follow up. Minimally clinically important differences (MCID) were used to assess for objective response. Patients also underwent clinical review to ascertain for any subjective change in wellbeing.

Results: There was a relative increase in mean FEV1 by 16.2% (p<0.05), a decrease in mean residual volume by 600mls (p<0.05) and a decrease in mean TLC by 540 ml s (p<0.05) following EBV insertion. With regards to minimally clinically important differences, 55.6% of the patient cohort had a greater than 430mls volume reduction in their RV whereas 66.7% of the patient cohort had a relative increase in their FEV1 by more than 10%. One patient had a pneumothorax following valve insertion and one of nine patients had valve migration that required repositioning. All of the patients had a subjective response in their first clinical review post valve insertion with one patient no longer requiring home oxygen.

Conclusion: Our audit demonstrated both a subjective and objective benefit of EBV insertion for carefully selected patients with symptomatic emphysema on best current management. Such a program can be successful in a public hospital setting with local support.

Grant Support: Nil
EBUS TBNA SAMPLING OF REGIONAL LYMPH NODES AS A SUBSTITUTE FOR BIOPSY FOR CONCOMITANT PULMONARY NODULES; STRONG SUPPORTIVE EVIDENCE OF PRIMARY LUNG CANCER FROM EBUS TBNA SAMPLE IMMUNOHISTOCHEMISTRY AND PET

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Introduction/Aim: A peripheral lung nodule with hilar or mediastinal nodes is an opportunity for single biopsy (EBUS TBNA alone) to give diagnosis and staging. This avoids subsequent procedures to confirm nodal metastasis.

Methods: A single-centre audit of medical records of all patients who underwent EBUS-TBNA of hilar or mediastinal lymph nodes as a diagnostic procedure for a peripheral lung lesion from 2007 to 2015 at the Royal Brisbane and Women’s Hospital, Brisbane, Australia. Patients with a distant metastasis were excluded.

Results: There were 194 cases of EBUS TBNA, 157 (81%) were diagnostic of malignancy. 30 patients had malignancy diagnosed at a subsequent procedure. No side effects of EBUS TBNA were observed.

Table 1. Method of diagnosis of malignancy

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Cases (N=187)</th>
</tr>
</thead>
<tbody>
<tr>
<td>EBUS TBNA</td>
<td>157 (84%)</td>
</tr>
<tr>
<td>Surgery</td>
<td>10 (5%)</td>
</tr>
<tr>
<td>Endobronchial ultrasound guide sheath</td>
<td>7 (4%)</td>
</tr>
<tr>
<td>CT guided fine needle aspiration</td>
<td>5 (3%)</td>
</tr>
<tr>
<td>Repeat EBUS TBNA</td>
<td>5 (3%)</td>
</tr>
<tr>
<td>Other</td>
<td>3 (1%)</td>
</tr>
</tbody>
</table>

Table 2. Malignancy types diagnosed at EBUS TBNA

<table>
<thead>
<tr>
<th>Subtype</th>
<th>Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary lung adenocarcinoma</td>
<td>61 (39%)</td>
</tr>
<tr>
<td>Squamous cell carcinoma</td>
<td>29 (18%)</td>
</tr>
<tr>
<td>Mesothelioma</td>
<td>2 (1%)</td>
</tr>
<tr>
<td>Small cell lung cancer and neuroendocrine carcinoma</td>
<td>22 (20%)</td>
</tr>
<tr>
<td>Immunohistochemically confirmed metastasis from other sites</td>
<td>6 (4%)</td>
</tr>
<tr>
<td>Non small cell lung carcinoma, not otherwise specified (NSCLC NOS)</td>
<td>27 (18%)</td>
</tr>
</tbody>
</table>

Of the 27 cases of NSCLC NOS, 23 patients underwent a PET scan, all of which were reported to be consistent with a primary lung lesion with nodal metastasis. All malignancy cases were discussed at a multidisciplinary case conference.

Conclusion: In patients who have a peripheral lung nodule and hilar or mediastinal lymph node enlargement, EBUS TBNA is an effective method of providing both diagnosis and staging, avoiding additional procedures. The use of combination pathology and PET gives convincing evidence between the nodal disease and peripheral lung lesion.

Grant Support: Nil

REFERENCE
PRE OPERATIVE EBUS TBNA PRE-SURGICAL STAGING HIGHLIGHTS THE IMPORTANCE OF CONFIRMING CYTOLOGICAL NEGATIVITY IN CASES OF TECHNICALLY N3 DISEASE ON PET SCANS

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Introduction/Aim: Endobronchial ultrasound and transbronchial needle aspirate (EBUS-TBNA) and Positron Emission Tomography (PET) have emerged as common modalities of staging pulmonary lymph nodes in non-small cell lung cancer (NSCLC). The findings of PET scans are often limited by factors such as infection, inflammation, or dust inhalation. As such, bilateral pulmonary lymphadenopathy in patients with known small cell lung cancer is frequently not representative of their true lymph node stage.

Method: We performed an single-centre audit of all patients with known NSCLC who had a PET scan and also underwent EBUS-TBNA for the explicitly stated purpose of pulmonary lymph node staging from 2005–2015. A combination of bronchoscopy reports, pathology reports, imaging reports and chart reviews were used to identify patient information.

Results: Overall, 40 cases were identified; 4 were excluded due to incomplete collection of data. Of these, PET demonstrated 12 cases of low to intermediate grade bilateral lymph node uptake, 22 cases of unilateral lymph node uptake and two cases of no PET uptake. Of cases with bilateral uptake, there were no cases in which EBUS-TBNA sampling was positive for malignancy. Two cases (9%) were subsequently upstaged to positive nodal involvement when they underwent surgery. Of cases with unilateral nodal involvement on PET scan, the frequency of biopsies positive for malignancy on both EBUS-TBNA, and on those who subsequently underwent surgery was 50%. Dust inhalation was seen on biopsy specimens of 41.7% of patients with bilateral PET uptake and 13.6% of patients with unilateral PET uptake.

Conclusion: This case series also adds to the growing body of evidence of dust inhalation associated with PET positivity. Although symmetrical uptake on PET scanning technically represents N3 disease, both EBUS-TBNA and surgery demonstrate a low rate of positive lymph node metastases. This highlights the role of EBUS-TBNA to rule patients in for surgery, rather than be excluded by PET findings alone.

Key Words: EBUS-TBNA, endobronchial ultrasound

Grant Support: Nil

Declaration of Interest: There were no competing interests in this study.

PREDICTING LIFE EXPECTANCY FOR PIRFENIDONE AND BEST SUPPORTIVE CARE IN IPF (ENCORE)

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Introduction/Aim: Clinical trials in IPF are not designed to estimate long-term survival. This analysis used a survival model to predict life expectancy for patients with idiopathic pulmonary fibrosis (IPF) receiving pirfenidone or best supportive care (BSC).

Methods: Life expectancy was estimated by the area under the curve of parametric survival distributions fit to Kaplan-Meier survival data from clinical studies and IPF registries. Kaplan-Meier survival data for pirfenidone were derived from clinical studies (CAPACITY, ASCEND, RECAP). Kaplan-Meier survival data for BSC were obtained from two independent registries of patients with IPF: the Inova Fairfax Hospital database (n=815) and the National Jewish Health Interstitial Lung Disease (NJH-ILD) database (n=321). The best-fitting distributions were chosen by statistical consideration, visual inspection of the fitted curve and by clinical interpretation. To account for differences between patients enrolled in the clinical trials and the registries, covariate adjustment using propensity scores was used.

Results: Mean life expectancy (95% confidence intervals) was calculated as: 8.7 years (7.7, 10.2) with pirfenidone; 5.9 years (5.1, 6.9) with BSC (Inova); and 6.1 years (5.7, 6.5) with BSC (NJH-ILD). Therefore, pirfenidone improved life expectancy relative to BSC by 2.8 years and 2.6 years as measured by the Inova and NJH-ILD registries, respectively.

Conclusion: The survival model suggests that pirfenidone significantly improves life expectancy compared with BSC by almost 3 years in patients with IPF. Although these findings are based on cross-trial comparisons, they provide support for pirfenidone as an effective treatment option for IPF.

Grant Support: Funded by InterMune International AG, which became a wholly owned subsidiary of F. Hoffmann-La Roche Ltd in 2014.
TWENTY-FOUR WEEK DECLINE IN FVC PREDICTS MORTALITY AT WEEK 52 IN THE INPULSIS® TRIALS

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Introduction/Aim: In Phase III INPULSIS® trials, a significantly lower proportion of patients with idiopathic pulmonary fibrosis treated with nintedanib vs placebo had disease progression with absolute FVC decline of ≥5% or ≥10% predicted at week 52. We explored the impact of FVC change over 24 weeks on subsequent FVC decline and mortality.

Methods: Post-hoc descriptive analysis of proportions of patients with absolute FVC declines of <5%, ±5% or ≥10% predicted from baseline to week 24 and changes in FVC% predicted and mortality between weeks 24 and 52 in these groups conducted using pooled data from both INPULSIS® trials.

Results: 1061 patients (nintedanib 638, placebo 423) were included. FVC decline of ≥5% or ≥10% predicted from baseline to week 24 did not predict FVC decline of ≥5% or ≥10% predicted, respectively, from week 24 to 52. The proportion of patients who died between weeks 24 and 52 increased with increasing FVC decline from baseline to week 24. Among patients with FVC declines of ≥5% or ≥10% predicted from baseline to week 24, more patients treated with nintedanib vs placebo had no further decline or an increase in FVC between weeks 24 and 52.

Conclusion: FVC declines of ≥5% or ≥10% predicted in the first 24 weeks did not predict FVC decline but were associated with higher mortality in the following 24 weeks.

<table>
<thead>
<tr>
<th>Absolute FVC decline % predicted from baseline to week 24</th>
<th>Outcome between weeks 24 and 52</th>
<th>Nintedanib (n%)</th>
<th>Placebo (n%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;5% n</td>
<td>444</td>
<td>259</td>
<td></td>
</tr>
<tr>
<td>Increase/no decline in FVC% predicted</td>
<td>155(34.9)</td>
<td>55(21.2)</td>
<td></td>
</tr>
<tr>
<td>FVC decline ≥5% predicted</td>
<td>133(30.0)</td>
<td>110(42.5)</td>
<td></td>
</tr>
<tr>
<td>Death</td>
<td>8(1.8)</td>
<td>6(2.3)</td>
<td></td>
</tr>
<tr>
<td>≥5% n</td>
<td>143</td>
<td>136</td>
<td></td>
</tr>
<tr>
<td>Increase/no decline in FVC% predicted</td>
<td>56(39.2)</td>
<td>47(34.6)</td>
<td></td>
</tr>
<tr>
<td>FVC decline ≥5% predicted</td>
<td>48(33.6)</td>
<td>41(30.1)</td>
<td></td>
</tr>
<tr>
<td>Death</td>
<td>11(7.7)</td>
<td>14(10.3)</td>
<td></td>
</tr>
<tr>
<td>≥10% n</td>
<td>46</td>
<td>53</td>
<td></td>
</tr>
<tr>
<td>Increase/no decline in FVC% predicted</td>
<td>24(52.2)</td>
<td>21(39.6)</td>
<td></td>
</tr>
<tr>
<td>FVC decline ≥10% predicted</td>
<td>9(19.6)</td>
<td>10(18.9)</td>
<td></td>
</tr>
<tr>
<td>Death</td>
<td>5(10.9)</td>
<td>7(13.2)</td>
<td></td>
</tr>
</tbody>
</table>
BIOMARKERS CAN PREDICT DISEASE PROGRESSION IN IDIOPATHIC PULMONARY FIBROSIS: ANALYSIS FROM THE AUSTRALIAN IPF REGISTRY.

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Introduction/Aim: The natural history of patients with idiopathic pulmonary fibrosis (IPF) is highly variable and difficult to predict. With the advent of new treatments that slow disease progression, there is growing interest in the potential of biomarkers as a method of differentiating rapidly progressive from more stable IPF disease.

Methods: The Australian IPF Registry is linked to a biobank where the blood of a sub-population of patients are stored for analysis. Using this IPF we analysed a panel of 11 plasma proteins thought to be important in IPF pathogenesis: ICAM1, OPNp, MMP7, CXCL13, FBLN1, SPA, MUC1, VCAM1, ENRAGE, IL8, CRP. The plasma protein concentrations were measured using the ELISA technique. Multivariate Cox analysis adjusted for baseline demographic features (age, gender, smoking, BMI) and disease severity (FVC%predicted) was used to determine the effect of these biomarkers in predicting mortality and progression free survival (PFS). PFS was defined as death, or fall in FVC≥10% or DLco≥15% from baseline.

Results: 170 participants (Age 68.4±8.2, 74% male, FVC 81.7±19.3) had blood available for analysis. The median concentrations were (ng/mL): ICAM1 298.9, OPNp 384.75, MMP7 2.8, CXCL13 0.045, FBLN1 60772.5, SPA 223.5, MUC1 0.27, VCAM1 1531, ENRAGE 78.9, IL8 0.063, CRP 7261.4. There were too few participants with measurable IL8 and MUC1 for further analysis. Higher concentrations of ENRAGE (HR 1.01, 95%CI 1.00-1.01, p=0.013), SPA (HR1.01, 95%CI 1.00-1.02, p=0.007) and OPNp (HR 1.02, 95%CI 1.00-1.04, p=0.01) were associated with increased mortality whereas higher OPNp (HR 1.01, 95%CI 1.00-1.02, p=0.029) and ICAM1 (HR 1.00, 95%CI 1.00-1.004, p=0.05) levels were associated with shorter progression free survival.

Conclusion: Biomarkers have a potential role in predicting mortality and disease progression in IPF.

Grant Support
**EFFECT OF CONTINUED PIRFENIDONE TREATMENT FOLLOWING ≥15% DECLINE IN 6MWD IN IPF PATIENTS- POOLED ANALYSIS (ENCORE)**

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**Introduction/Aim:** Previous analyses of patients with idiopathic pulmonary fibrosis (IPF) receiving pirfenidone showed that continued treatment following a ≥ 10% absolute or relative decrease in percent predicted forced vital capacity (FVC) or hospitalization during the first 6 months of treatment provided a benefit during the subsequent 6 months. The aim is to further explore the potential benefit of continued pirfenidone treatment in patients with IPF who had a worsening of 6-minute walk distance (6MWD) within the first 6 months of treatment.

**Methods:** A pooled analysis included all patients randomized to pirfenidone 2403 mg/d or placebo in the ASCEND and CAPACITY studies (N = 1247). All patients who had a 6MWD decline ≥ 15% within the first 6 months of treatment were included. The outcomes were assessed during the subsequent 6-month period.

**Results:** A total of 116/623 (18.6%) and 141/624 (22.6%) patients in the pooled pirfenidone and placebo groups, respectively, demonstrated a 6MWD decline ≥ 15% within the first 6 months of treatment. Outcomes during the subsequent 6 months of continued treatment in these patients are shown in the Table.

**Conclusion:** These results suggest that continued treatment with pirfenidone may confer a significant benefit to patients with IPF who experienced a 6MWD decline ≥ 15% within the first 6 months of treatment.

**Grant Support:** Supported by F. Hoffmann-La Roche Ltd.

**Table.** Outcomes during the 6-month period following an initial 6MWD decline ≥ 15% during the first 6 months of treatment

<table>
<thead>
<tr>
<th>Outcome in Subsequent 6 Months</th>
<th>Pirfenidone, n (%)</th>
<th>Placebo, n (%)</th>
<th>Relative Difference</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥ 15% relative decline in percent predicted FVC or death</td>
<td>17 (14.7)</td>
<td>40 (20.6)</td>
<td>-48.3%</td>
<td>0.010</td>
</tr>
<tr>
<td>≥ 10% relative decline in percent predicted FVC</td>
<td>14 (12.1)</td>
<td>25 (17.7)</td>
<td>-33.1%</td>
<td>0.0001</td>
</tr>
<tr>
<td>Death</td>
<td>3 (2.6)</td>
<td>15 (10.6)</td>
<td>-75.7%</td>
<td>0.460</td>
</tr>
<tr>
<td>≥ 15% decline in 6MWD or death</td>
<td>20 (17.2)</td>
<td>52 (36.9)</td>
<td>-63.2%</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>≥ 15% decline in 6MWD</td>
<td>17 (14.7)</td>
<td>37 (26.2)</td>
<td>-44.2%</td>
<td>0.0001</td>
</tr>
<tr>
<td>Death</td>
<td>3 (2.6)</td>
<td>15 (10.6)</td>
<td>-75.7%</td>
<td>0.460</td>
</tr>
<tr>
<td>Hospitalization for any reason</td>
<td>20 (17.2)</td>
<td>27 (19.1)</td>
<td>-10.0%</td>
<td>0.747</td>
</tr>
<tr>
<td>Hospitalization for a respiratory reason</td>
<td>13 (11.2)</td>
<td>18 (12.8)</td>
<td>-12.2%</td>
<td>0.868</td>
</tr>
</tbody>
</table>

**ANNUAL RATE OF FVC DECLINE IN PATIENTS WITH IPF TREATED WITH PIRFENIDONE: POOLED ANALYSIS (ENCORE)**

PAUL W. NOBLE,1 CARLO ALBERA,2 WILLIS CHOU,3 ULRICH COSTABEL,4 BANN-MO DAY,3 IAN GLASPOLE,5 MARILYN K. GLASSBERG,6 LISA LANCASTER,7 DAVID J. LEDERER,8 STEVEN D. NATHAN,9 CARLOS A. PEREIRA,10 JOHN STAUFFER,11 JEFFREY J. SWIGRIS11

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**Introduction/Aim:** Pirfenidone has been shown to decrease the annual rate of decline in forced vital capacity (FVC) volume in patients with idiopathic pulmonary fibrosis (IPF). This analysis explored this effect in various patient subgroups.

**Methods:** Patients randomized to pirfenidone 2403 mg/d or placebo in the CAPACITY or ASCEND studies were included. The annualized rate of decline in FVC volume from baseline through 12 months was estimated using a mixed-effects model, with study, time-by-treatment, age-by-sex and height-by-sex as fixed effects and patients and time-by-patient (slope) as random effects. The annual rate of FVC decline was estimated from the slope within the subgroups, defined by demographics and baseline disease activity measures.

**Results:** A total of 623 patients in the pirfenidone group and 624 in the placebo group were included in the pooled analysis. Overall, the adjusted annual rate (SE) of FVC decline from baseline to 12 months was -132.0 (12.4) mL for pirfenidone vs -223.5 (12.4) mL for placebo, a difference of 91.5 (16.4) mL. The annual rate of FVC decline favoured pirfenidone over placebo across various baseline demographic and lung function subgroups.

**Conclusion:** Patients with IPF treated with pirfenidone, regardless of baseline demographic or lung function, had a significantly lower annual rate of decline in FVC volume vs those treated with placebo after 12 months.

**Grant Support:** Funded by F. Hoffmann-La Roche Ltd.
THE EFFECT OF HIGH-DOSE, HIGH FINE-PARTICLE FRACTION COMBINATION THERAPY ON VENTILATION HETEROGENEITY IN UNCONTROLLED ASTHMA

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Introduction/Aim: Both the large and small airways are important in severe asthma. However, their response to treatment with high-dose, high-fine particle fraction inhaled corticosteroid/long acting beta agonist (ICS/LABA) combination has yet to be established. We aimed to measure the effect of fluticasone/eformoterol on ventilation heterogeneity measured by multiple breath nitrogen washout and on clinical outcomes. In addition, we aimed to determine predictors of improvements in conductive (Scond) and acinar (Sacin) ventilation heterogeneity. We hypothesised that changing to high-dose fluticasone/eformoterol would improve Scond and Sacin, along with clinical improvement in asthma control.

Methods: 21 patients (7 males) with a doctor diagnosis of asthma, were uncontrolled (5 component asthma control questionnaire score (ACQ5)>1.5) and were currently on ICS or ICS/LABA (maximum of 500μg/day fluticasone equivalent) were enrolled. Baseline Scond, Sacin, ACQ5, exhaled nitric oxide (FeNO), airway hyper-responsiveness (AHR) to methacholine (log10 dose response slope (log10DRS)), and spirometry were measured. Patients took 250/10μg Fluticasone/eformoterol via a spacer for 8 weeks before repeat testing. The relationship between changes in Scond and Sacin, and baseline ACQ5, AHR, spirometry and FeNO were examined.

Results: Treatment improved Scond (0.041±0.025 vs 0.030±0.017L−1, p=0.002) and Sacin (0.127±0.079 to 0.104±0.067L−1, p=0.002). There were improvements in ACQ5 (2.11±0.85 to 1.11±0.85, p=0.0003), log10DRS (1.48±0.64 to 0.92±0.45μmol−1, p=0.0001), FeNO (19.71±13.68 to 11.40±4.94ppb, p=0.001), FEV1 (83.61±14.92 to 89.78±14.7%predicted, p=0.007) and FVC (96.95±12.28 to 101.2±12.99%predicted, p=0.04). The improvements in Sacin and Scond were predicted only by baseline Sacin and Scond, respectively (r=0.54, p=0.01 and r=0.73, p=0.0002, respectively).

Conclusion: Fluticasone/eformoterol with high-dose, high-fine particle fraction improved ventilation heterogeneity, symptoms, AHR, airway inflammation and spirometry in people with uncontrolled asthma. Baseline Sacin and Scond were the only predictors for improvements in ventilation heterogeneity.

Grant Support: NHMRC project grant 103701, MundiPharma.
HETEROGENEOUS BRONCHOCONSTRICTION IN ASTHMA ON VENTILATION SINGLE PHOTON EMISSION COMPUTED TOMOGRAPHY/COMPUTED TOMOGRAPHY(VSPECT/CT) DECREASES AFTER TREATMENT

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1Woolcock Institute of Medical Research and The University of Sydney, Australia, 2Division of Cardiovascular Diseases, Mayo Clinic, Rochester Mn, USA

Introduction/Aim: Areas of heterogeneous bronchoconstriction can be seen as low ventilation on functional lung imaging in asthma. A fundamental characteristic of asthma is airway hyperresponsiveness (AHR), which is associated with ventilation heterogeneity. AHR is slow to resolve with anti-inflammatory treatment and the mechanism is poorly understood. The aim of this study was to determine if anti-inflammatory treatment reduces heterogenous bronchoconstriction by measuring low ventilation areas on VSPECT/CT and investigating the role of peripheral airway heterogeneity in this mechanism.

Methods: Twenty-one asthmatic subjects (13 female Mean ± SD Age: 31 ± 13 yrs, FEV1: 83 ± 16% Pred) had baseline and methacholine challenge VSPECT/CT scans, before and after 8 weeks of combined ICS/LABA treatment. Low ventilation was measured by changes in ventilated volume at two thresholds on VSPECT(80/50 ratio). Dose response slope (DRS) was calculated to measure AHR and is reported as logDRS. Multiple breath nitrogen washout was performed at baseline and after treatment to determine peripheral ventilation heterogeneity (Sacin).

Results: Mean (Range) logDRS pre treatment(preT): 1.44 (0.57-2.4) %, umol¹, logDRS post treatment (postT): 0.96 (0.48 − 1.9) %, umol¹, p=0.001; 80/50 ratio preT: 0.05 (-0.01-0.18), 80/50 ratio postT: 0.03 (-0.04-0.18) p=0.02; Sacin preT: 0.128 (0.024-0.375) L⁻¹, Sacin postT: 0.104 (0.019-0.334) L⁻¹, p=0.005. Improvement in 80/50 ratio with treatment correlated with preT logDRS (p=0.036, r=0.46) but not Sacin.

Conclusion: Anti-inflammatory treatment reduces heterogeneous bronchoconstriction measured on functional lung imaging. The reduction in the development of low ventilation areas (after challenge) with treatment is predicted by pre-treatment airway hyperresponsiveness. Peripheral heterogeneity improved with treatment but did not relate to imaging. Bronchoconstriction patterns on imaging are difficult to quantify but potentially could give valuable mechanistic insight for future treatment improvements.

Grant Support: NH&MRC Project Grant 103701 and MundiPharma

EVIDENCE OF PERSISTENT HYPERVENTILATION FOLLOWING HIGH ALTITUDE EXPOSURE

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Introduction/Aim: High altitude exposure results in acute hypoxia and an increase in the ventilatory drive. This hyperventilatory response is characterised by a worsening breathing efficiency (increased Vt/VCO2) and a lower end tidal CO2 (PETCO2) both at rest and during exercise. Whilst the time-course changes in the hyperventilatory response are well-characterised in acclimated climbers, to date there has been little or no examination of the response during ascent and on return from altitude in a large group of non-acclimatised climbers. The aim of the current study was to examine the changes in gas exchange measures in a group of non-acclimatised climbers during the ascent of Mt Kilimanjaro and immediately on return from altitude.

Methods: 27 (age: 44±15, range 22-66yrs) individuals ascended Mt Kilimanjaro during an 11-day climb. Exercise testing (4-minute step test with gas-exchange) was completed on four occasions at the following altitudes: (1) basecamp, 1850m (PB=690 mmHg); (2) 3500m (PB=505 mmHg); (3) 4840m (PB=428 mmHg) and on return to base camp (4) 850m (PB=690 mmHg).

Results: During the ascent subjects became increasingly hypoxic and there was a decrease in the nadir SaO2 (%) 850m: 96±2; 3500m: 82±3; 4840m: 73±4. During exercise breathing efficiency worsened (Vt/VCO2) 850m: 28±2.9; 3500m: 35.8±4.6; 4840m: 50.7±5.8) and PETCO2 fell (PETCO2: mmHg) 850m: 37.1±3.6; 3500m: 28.5±2.6; 4840m: 20.8±1.9) as subjects ascended. On return from altitude (850m) SpO2 normalised (SpO2(%)=97±3), however there was persistent evidence of hyperventilation with both breathing efficiency and PETCO2 remaining abnormal during exercise (Vt/VCO2: 32.8±3.2; PETCO2:mmHg: 31.2±3.0; p<0.01 vs 850m) and at rest.

Conclusion: With high altitude exposure there is an increased ventilatory drive. Following high altitude exposure and once SaO2 normalises, gas exchange remains altered. We hypothesise that this altered gas exchange may be due to a sustained alkalosis with the central chemoreceptors remaining reset to defend a lower PCO2 established during high altitude exposure.
PROXIMAL & PERIPHERAL AIRWAY RESPONSE TO DIRECT & INDIRECT BRONCHIAL PROVOCATION IN UNTREATED ASTHMA

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Introduction/Aim: Spirometry is used to assess the degree of bronchoconstriction during bronchial provocation tests. However, little is known about how specific features of the proximal and peripheral airways respond to different provocation agents. This study aims to determine how the proximal and peripheral airway resistance, ventilation heterogeneity and nitric oxide levels change in the same airways in response to standard direct and indirect provocation agents relative to placebo.

Method: Newly diagnosed (untreated) patients with atopic Asthma were recruited for the study (n=9). All subjects demonstrated a clinically positive test using Spirometry to both direct (Methacholine) and indirect (Mannitol) provocation tests on separate visits, ~7days apart. On another visit, a placebo provocation was also performed. Before and immediately after all provocation tests, the following measures were made; airway resistance using the Forced Oscillation Technique, ventilation heterogeneity using Multiple Breath Nitrogen Washout technique and nitric oxide flux and alveolar concentrations using the Multiple Exhalation Flow technique.

Results: All data expressed as mean ± SEM. *p<0.05 with respect to placebo change.

<table>
<thead>
<tr>
<th>Measure</th>
<th>Units</th>
<th>Placebo</th>
<th>Direct</th>
<th>Indirect</th>
</tr>
</thead>
<tbody>
<tr>
<td>FEV1</td>
<td>L</td>
<td>−0.151±0.036</td>
<td>−0.826±0.081</td>
<td>−0.651±0.075</td>
</tr>
<tr>
<td>Proximal Airways</td>
<td>R16</td>
<td>hPa/(L/sec)</td>
<td>+0.094±0.156</td>
<td>+0.990±0.266</td>
</tr>
<tr>
<td>Scond</td>
<td>1/L</td>
<td>−0.003±0.003</td>
<td>+0.024±0.006</td>
<td>+0.025±0.004</td>
</tr>
<tr>
<td>NO-Flux</td>
<td>nL/sec</td>
<td>−0.394±0.097</td>
<td>−0.654±0.193</td>
<td>−2.075±0.521</td>
</tr>
<tr>
<td>Peripheral Airways</td>
<td>R4-R16</td>
<td>hPa/(L/sec)</td>
<td>+0.261±0.166</td>
<td>+2.703±0.506</td>
</tr>
<tr>
<td>Sacin</td>
<td>1/L</td>
<td>+0.013±0.013</td>
<td>+0.041±0.008</td>
<td>+0.054±0.010</td>
</tr>
<tr>
<td>CNO</td>
<td>ppb</td>
<td>+1.225±0.842</td>
<td>−0.942±0.342</td>
<td>−3.083±1.781</td>
</tr>
</tbody>
</table>

Conclusion: In the proximal airways, both resistance and heterogeneity increased in response to direct and indirect provocation relative to placebo. However, nitric oxide flux decreased only in response to the indirect agent. In the peripheral airways, heterogeneity remained unchanged. Periphera resistance increased in response to both direct and indirect provocation. However, alveolar nitric oxide levels decreased only in response to the indirect agent.


COMPLEX MEDICATION REGIMENS IN CHRONIC OBSTRUCTIVE PULMONARY DISEASE (COPD) ARE ASSOCIATED WITH DISEASE SEVERITY AND COMORBIDITIES

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1Priority Research Centre for Healthy Lungs and Hunter Medical Research Institute (HMRI), The University of Newcastle (UoN), New South Wales, Australia., 2Department of Respiratory and Sleep Medicine, John Hunter Hospital, New South Wales, Australia., 3School of Nursing and Midwifery, UoN, New South Wales, Australia.

Introduction/Aim: COPD patients are often prescribed multiple medications to manage their respiratory disease and comorbidities. This may result in complex medication regimens, which can increase the risks of poor adherence, medication errors and drug-drug interactions. This study aims to explore the medication burden in COPD and its relationship to clinical outcomes.

Methods: A cross-sectional study whereby COPD patients (n=222) were assessed for demographic information, comorbidities, medication use, clinical outcomes and multidimensional indices was performed. Complexity of medication regimens was quantified using the validated Medication Regimen Complexity Index (MRCI).

Results: Participants (58.6% males) had a mean (SD) age of 69.1±8.3 years, post-bronchodilator %predicted FEV1 of 56.5±20.4%. They suffered from a median of 5 comorbidities. Nearly half (47.8%) of the participants were taking ≥9 medications. The mean (SD) total MRCI score was high (25.1±8.9). COPD-specific medication regimens were more complex than those of non-COPD medications (MRCI: 14.7 versus 9, respectively; p<0.0001). Complex dosage formulations contributed the most (57%) to higher MRCI scores of COPD-specific medications while dosing frequency primarily drove the complexity associated with non-COPD medications (67%). Participants in GOLD quadrant D had the highest MRCI score correlated with 6-minute walk distance (r=−0.288; p=0.0003), Saint George Respiratory Questionnaire (r=0.294; p<0.0001) and prior year exacerbation history (r=0.246; p=0.0002). Multiple regression analysis revealed that comorbid cardiovascular, gastrointestinal or metabolic diseases contributed to higher MRCI scores and/or medication counts. Charlson Comorbidity Index and COPD-specific comorbidity test showed the highest degree of correlations with total MRCI score (r=−0.289; p=0.0001 and r=0.326; p<0.0001, respectively).

Conclusion: COPD patients have complex medication regimens that are associated with disease severity and certain comorbidities. Our data support the need to reduce medication complexity in COPD.

Grant Support: NHMRC, Ramaciotti Foundation, Lung Foundation of Australia, UoN, HMRI.
LONG-TERM EFFICACY OF A1-PI THERAPY IN RAPID AND RAPID EXTENSION TRIALS

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Introduction: RAPID (NCT00261833), the largest (N=180) randomized placebo-controlled trial assessing emphysema progression in alpha-1 anti-trypsin deficiency (AATD) completed to date, was followed by the open-label RAPID Extension trial (NCT00670007).

Methods: Subjects in RAPID received alpha-1 proteinase inhibitor (A1-PI, Zemaira; CSL Behring) or placebo for 2 yrs. Eligible subjects (N=140) continued in Extension to receive A1-PI for another 2 yrs. The Early-Start (N=75) and Delayed-Start (n=64) cohorts were defined by their treatment with A1-PI and placebo in RAPID, respectively. Computed tomography (CT) lung density decline rate was measured annually and forced expiratory volume in 1 sec (FEV1) was measured quarterly.

Results: Annual CT lung density decline rate in the first 2 yrs was less by 0.75 g/L/yr in the Early-Start cohort (~1.51 g/L/yr vs ~2.26 g/L/yr, p=0.021 one sided). In the Delayed-Start cohort, lung density decline was reduced to ~1.26 g/L/yr after switching to A1-PI. Changes in FEV1 over 4 yrs correlated significantly with changes in CT lung density.

Conclusions: In RAPID, A1-PI therapy reduced the rate of lung density decline compared to placebo. Over 48 months, the Delayed-Start cohort showed a greater loss of lung density. These data demonstrate a disease-modifying effect of A1-PI therapy, suggesting that early treatment may reduce emphysema progression in AATD patients.

Funding: Trials were funded by CSL Behring.

Abstract Category: Airway Pharmacology and Treatment

Keywords: Treatments; COPD Management

OMALIZUMAB TREATMENT RESPONSE IN A SEVERE ALLERGIC ASTHMA POPULATION WITH OVERLAPPING COPD

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Introduction/Aim: Asthma and chronic obstructive pulmonary disease (COPD) are common airway diseases. Individuals with overlapping asthma and COPD experience increased health impairment and severe disease exacerbations. Efficacious treatment options are required for this population. Omalizumab (anti-IgE) therapy is effective in patients with severe, persistent asthma, but limited data are available on efficacy in populations with overlapping asthma and COPD. In the current study, we sought to assess the effects of omalizumab treatment on this population.

Methods: Data from the Australian Xolair Registry (AXR) was used to compare treatment responses in individuals with asthma-COPD overlap to severe asthma alone. Participants were assessed at baseline and after 6 months of omalizumab treatment. We utilised several different definitions of asthma-COPD overlap. First, we compared participants with a previous doctor diagnosis of COPD to participants with no COPD diagnosis. We then made comparisons based on baseline lung function, comparing participants post-bronchodilator FEV1 <80% predicted to >80%. In the FEV1<80% population, analysis was further stratified based on smoking history.

Results: Omalizumab treatment markedly improved asthma control and health-related quality of life in all populations assessed, based on ACO-S and AQLQ questionnaire scores. Omalizumab treatment did not improve lung function in populations that were enriched for asthma-COPD overlap (diagnosis of COPD or FEV1<80% ever smokers).

Conclusion: Our study suggests that omalizumab improves asthma control and health-related quality of life in individuals with severe allergic asthma and overlapping COPD. These findings provide real-world efficacy data for this patient population and suggest omalizumab is useful in the management of severe asthma with COPD overlap.

Grant Support: The Australian Xolair Registry was supported by Novartis Pharmaceuticals Australia Pty Ltd, as an Investigator-sponsored study. This analysis was supported by funding from the National Health and Medical Research Council Centre of Excellence in Severe Asthma (www.severeasthma.org.au).
INDACATEROL/GLYCOPPYRONIUM (IND/GLY) REDUCES THE RISK OF EXACERBATIONS VERSUS SALMETEROL/FLUTICASONE (SFC) IN MODERATE-TO-VERY SEVERE COPD PATIENTS IRRESPECTIVE OF PRIOR ICS/LABA/LAMA THERAPY: THE FLAME STUDY
JADWIGA A. WEDZICHA1, KAREN MEZZI2, TIM AYERS3, CHAU THACH2, ROBERT FOGELO, FRANCESCO PATALANO6, DONALD BANERJEI3
1Airways Disease Section, National Heart and Lung Institute, Imperial College London, London, United Kingdom; 2Novartis Pharma AG, Basel, Switzerland; 3Novartis Pharmaceuticals Corporation, East Hanover, NJ, United States

Introduction: FLAME was the first study to show superiority of IND/GLY (LABA/LAMA) in reducing risk of exacerbations vs SFC (LABA/ICS) in COPD patients with history of exacerbations. We report results regarding effect of IND/GLY vs SFC on COPD exacerbations in patients with or without prior ICS/LABA/LAMA triple therapy.

Methods: FLAME, a 52-week, randomised, double-blind study, compared once-daily IND/GLY 110/50 μg vs twice-daily SFC 50/500 μg in patients with moderate-to-severe COPD with ≥1 exacerbation in previous year. In this post hoc analysis, time to first COPD exacerbation (mild, moderate or severe) and moderate/severe COPD exacerbation with IND/GLY vs SFC were analysed. Additionally, treatment effects on rates of exacerbations were assessed in these patients.

Results: Of 3354 patients included in this analysis, 1148 (34.2%) were on prior triple therapy. IND/GLY significantly reduced the rate of any exacerbation and moderate/severe exacerbation vs SFC in patients with/without prior triple therapy (Table). IND/GLY numerically reduced the rate of any and moderate/severe COPD exacerbations vs SFC in patients with/without prior triple therapy (table).

Conclusion: IND/GLY significantly reduces risk of exacerbations vs SFC, irrespective of prior triple therapy in exacerbating patients with moderate-to-severe COPD.

Grant Support: The study was funded by Novartis Pharma AG, Basel, Switzerland

BRONCHODILATOR THERAPY AND EXERCISE ADDED TO SELF-MANAGEMENT BEHAVIOUR-MODIFICATION: EFFECTS ON PHYSICAL ACTIVITY IN COPD
PETER FRITH1 ON BEHALF OF THIERRY TROOSTERS2, KIM L LAVOIE3, NANCY LEIDY5, FRANCOIS MALTAIS5, MARIA SEDENO3, WIM JANSSSENS4, ALAN HAMILTON9, DAMIJAN ERZEN10, DOROTHY DE SOUSAA, LAWRENCE KORDUCKI11, JEAN BOURBEAU7
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Introduction/Aim: PHYSACTO® has shown improved exercise capacity in patients with chronic obstructive pulmonary disease (COPD) receiving bronchodilators (BDs) + exercise training (ExT) added to a self-management behaviour-modification programme (SMBM). We explored the influence of BDs+ExT following SMBM on physical activity (PA) and perceived PA-related difficulty and symptoms in PHYSACTO® (NCT02085161).

Methods: A 12 week, randomised, partially double-blind, placebo (P)-controlled, parallel-group trial. Interventions (all with 12 week SMBM): P; tiotropium (T) 5 μg; T + olodaterol (O) 5/5 μg; T + O 5/5 μg + 8 week ExT. A triaxial accelerometer assessed PA (steps/day), the Functional Performance Inventory assessed patient-reported difficulty engaging in PA and the Chronic Respiratory Questionnaire assessed PA-related dyspnoea.

Results: 303 patients (post BD-FEV1: 57 ± 13 % predicted) were randomised and treated (full analysis set n=274). Change in steps/day at week 12 is shown in the Table; no significant gain in PA was seen by adding T+O to SMBM, versus SMBM+P. A triaxial accelerometer assessed PA (steps/day), the Functional Performance Inventory assessed patient-reported difficulty engaging in PA and the Chronic Respiratory Questionnaire assessed PA-related dyspnoea.

Conclusion: In moderate to severe COPD, SMBM increased PA and the addition of T+O to SMBM was associated with reduced difficulty and symptoms with PA when compared to SMBM+P.

Table: Time to first and rate of any (mild, moderate or severe) COPD exacerbation and moderate/severe COPD exacerbation with IND/GLY versus SFC

<table>
<thead>
<tr>
<th>Subgroup</th>
<th>Hazard ratio (95% CI)</th>
<th>Rate ratio (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>With prior ICS/LABA/LAMA therapy (n=1148)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any (mild, moderate or severe) COPD exacerbation</td>
<td>0.88 (0.78, 0.98)</td>
<td>0.88 (0.80, 0.96)</td>
</tr>
<tr>
<td>Moderate/severe COPD exacerbation</td>
<td>0.79 (0.68, 0.93)</td>
<td>0.80 (0.74, 0.91)</td>
</tr>
<tr>
<td>Without prior ICS/LABA/LAMA therapy (n=2206)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Any (mild, moderate or severe) COPD exacerbation</td>
<td>0.84 (0.76, 0.92)</td>
<td>0.85 (0.78, 0.93)</td>
</tr>
<tr>
<td>Moderate/severe COPD exacerbation</td>
<td>0.77 (0.68, 0.88)</td>
<td>0.82 (0.72, 0.92)</td>
</tr>
</tbody>
</table>

CI: confidence interval, COPD: chronic obstructive pulmonary disease, ICS: inhaled corticosteroid, LABA: long-acting β2-agonist, LAMA: long-acting muscarinic antagonist

Steps/day* at week 12

<table>
<thead>
<tr>
<th>Intervention</th>
<th>Adjusted mean (SE)</th>
<th>Adjusted mean change from baseline (SE)</th>
<th>95% CI</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>SMBM+P (n=55)</td>
<td>6517.71 (325.08)</td>
<td>1098.07 (325.08)</td>
<td>457.47, 0.0009</td>
<td>0.0009</td>
</tr>
<tr>
<td>SMBM+T (n=57)</td>
<td>5572.83 (317.98)</td>
<td>153.19 (317.98)</td>
<td>473.43, 0.0634</td>
<td>0.6304</td>
</tr>
<tr>
<td>SMBM+T+O (n=60)</td>
<td>6813.88 (310.22)</td>
<td>556.79 (310.22)</td>
<td>782.92, &lt;0.0001</td>
<td>0.0011</td>
</tr>
<tr>
<td>SMBM+T+O+ExT (n=57)</td>
<td>5976.43 (317.97)</td>
<td>-69.81 (317.97)</td>
<td>1183.39</td>
<td>0.0813</td>
</tr>
</tbody>
</table>

*Steps/day = number of steps per day
Full analysis set. Analysis of covariance model with “treatment” and “baseline” as covariates. Common baseline mean steps/day (SE): 6419.64 (186.15).

*Measured via triaxial accelerometer.

**Grant Support:** The study was funded by Boehringer Ingelheim.

**Declaration of Interest Statement:** PF has received in the past 5 years honoraria for educational and advisory board involvement and/or received conference attendance support for the following: Global Initiative for COPD (GOLD), Improvemnt Foundation, Lung Foundation Australia, and Remedy Healthcare; AstraZeneca, Boehringer Ingelheim, CSL-Behring, GlaxoSmithKline, Menarini, Mundipharma, and Novartis. TT is the Principal Investigator of PROactive project and received speakers/consultancy fees from Boehringer Ingelheim, Novartis, and Bayer. KLL received grants/research support from GSK, consulting fees from Schering-Plough and Merck Frosst and served on speaker bureaus/honoraria for GSK, Astra-Zeneca, Pfizer, Merck Frosst, Air Liquide and Health International. NL is an employee of Evidera and works with a variety of companies and organizations and expressly prohibited from receiving any payment or honoraria directly from these organizations for services rendered. FM is a consultant for Boehringer Ingelheim and served on advisory boards for Boehringer Ingelheim, GlaxoSmithKline, and Pfizer and has received payment for lectures including service on speaker bureaus from Boehringer Ingelheim, GlaxoSmithKline, Nycomed and Pfizer. MS is employed by Respilus, a non-profit organisation that was contracted by the study sponsor to develop the educational component of their training programme. AH, DE, DDS, and LK are employees of Boehringer Ingelheim. JB received research funding via the Research Institute of the McGill University Health Centre from AstraZeneca, Boehringer Ingelheim, GlaxoSmithKline, Merck, Novartis, Nycomed, Pfizer and Theratechnologies; and has served on speakers, consultation panels and/or advisory boards for these pharmaceutical companies.

**EFFECT OF INDACATEROL/GLYCOPYRRONIUM (IND/GLY) VS SALMETEROL/FLUTICASONE (SFC) ON MODERATE OR SEVERE COPD EXACERBATIONS AND LUNG FUNCTION BASED ON BASELINE BLOOD EOSINOPHIL COUNTS: RESULTS FROM THE FLAME STUDY**

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**Introduction:** The FLAME study compared the effect of IND/GLY vs SFC on exacerbations in COPD patients with a risk of exacerbation. Here we present the rate of reduction of moderate or severe exacerbations and lung function changes based on blood eosinophils in moderate-to-very severe COPD patients.

**Methods:** This was a 52-week, multicentre study. Patients with moderate-to-severe COPD, post-bronchodilator FEV1 ≥ 25% and ≤ 60% and a history of ≥ 1 exacerbation in the previous year were randomised (1:1) to IND/GLY (110/50 μg) once daily or SFC (50/500 μg) twice daily. The data were assessed by blood eosinophil cut offs (<150, 150–<300 and ≥300) and percentages (<2% and ≥2%).

**Results:** 3362 patients were randomised to IND/GLY (n=1680) or SFC (n=1682). The annualised rate of moderate or severe exacerbations was significantly lower in IND/GLY-treated vs SFC-treated patients at all eosinophil counts, although not significant for small number of patients with ≥300 cells/μL. The lung function was significantly improved at all the visits with IND/GLY vs SFC irrespective of eosinophil count (Table 1).

**Conclusion:** In patients with high risk of exacerbations, IND/GLY was superior in reducing moderate or severe exacerbations and showed significant improvement in lung function vs SFC independent of blood eosinophil counts.

**Grant Support:** The study was funded by Novartis Pharma AG, Basel, Switzerland

**Table 1. Annualized rate of moderate or severe exacerbations and lung function improvement in IND/GLY vs SFC.**

<table>
<thead>
<tr>
<th>Subgroups by blood eosinophils</th>
<th>Annualized rate of moderate or severe exacerbations</th>
<th>Rate ratio (95% CI), p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;2% n=1279</td>
<td>0.80 (0.68, 0.93), p=0.004</td>
<td>0.85 (0.75, 0.96), p=0.010</td>
</tr>
<tr>
<td>≥2% n=2019</td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt;150 cells/μL, n=1277</td>
<td>0.72 (0.61, 0.84), p=0.001</td>
<td>0.89 (0.76, 1.03), p=0.128</td>
</tr>
<tr>
<td>≥150–&lt;300 cells/μL, n=1286</td>
<td>0.93 (0.76, 1.14), p=0.464</td>
<td></td>
</tr>
<tr>
<td>≥300 cells/μL, n=735</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Change from baseline in pre-dose trough FEV1(L) at post-baseline visit**

<table>
<thead>
<tr>
<th>Subgroups by blood eosinophils</th>
<th>LS Mean Difference (95% CI), p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;2% n=1237</td>
<td>Day 29: 0.093 (0.074, 0.112), p=0.001</td>
</tr>
<tr>
<td></td>
<td>Day 365: 0.072 (0.049, 0.095), p=0.001</td>
</tr>
<tr>
<td>≥2% n=1946</td>
<td>Day 29: 0.058 (0.043, 0.074), p=0.001</td>
</tr>
<tr>
<td></td>
<td>Day 365: 0.056 (0.038, 0.074), p=0.001</td>
</tr>
<tr>
<td>&lt;300 cells/μL, n=2483</td>
<td>Day 29: 0.083 (0.069, 0.096), p=0.001</td>
</tr>
<tr>
<td></td>
<td>Day 365: 0.067 (0.051, 0.083), p=0.001</td>
</tr>
<tr>
<td>≥300 cells/μL, n=700</td>
<td>Day 29: 0.037 (0.011, 0.062), p=0.005</td>
</tr>
<tr>
<td></td>
<td>Day 365: 0.048 (0.018, 0.079), p=0.002</td>
</tr>
</tbody>
</table>

n, number of patients included in the sub group analysis; CI, confidence interval; FEV1, forced expiratory volume at 1 second; LS, Least square; IND/GLY, indacaterol/glycopyrrolonium; SFC, salmeterol/fluticasone.
CHANGES IN LUNG FUNCTION AND FORCED OSCILLATORY TECHNIQUE (FOT) PARAMETERS FOLLOWING BRONCHIAL THERMOPLASTY (BT) IN PATIENTS WITH REFRACTORY ASTHMA

ING AJ1,4, LANGTON D2, PIERUCCI P1,2,4, FARAH CS1,3,4
1Faculty of Medicine and Health Sciences, Macquarie University, 2Monash University, 3Woolcock Institute of Medical Research, 4Sydney Medical School, The University of Sydney

Introduction/Aim: Bronchial Thermoplasty (BT) improves symptom control and reduces exacerbations in patients with refractory asthma. The precise mechanism remains unclear, but consistent improvement in spirometry has not been recorded. We postulate that assessment of lung function during tidal breathing with oscillometry to measure respiratory mechanics may better reflect the clinical improvements reported.

Methods: Patients with refractory asthma (GINA Step 5) had the Asthma Control Questionnaire (ACQ5), spirometry, resistance at 5 Hz (R5) and reactance at 5 Hz (X5) recorded before and 6 weeks after completing BT. One centre measured impedance with the forced oscillation technique (FOT) [TremoFlo C-100, Thorasys Medical Systems, Canada] and the other with impulse oscillometry (IOS) [Sensormedics].

Results: The average age was 60.8 (1.4) years in the FOT group and 61.2 (11.4) years in the IOS group. ACQ5 scores improved in both groups. In the FOT group, there were significant improvements in spirometry and X5; and the change in X5 correlated strongly with changes in vital capacity and weakly with ACQ5.

<table>
<thead>
<tr>
<th>Pre BT</th>
<th>Post BT</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACQ5</td>
<td>2.80</td>
<td>1.90</td>
</tr>
<tr>
<td>(0.75)</td>
<td>(1.10)</td>
<td>(0.91)</td>
</tr>
<tr>
<td>FEV1 (%pred)</td>
<td>51.7</td>
<td>61.7</td>
</tr>
<tr>
<td>(15.9)</td>
<td>(20.9)</td>
<td>(7.2)</td>
</tr>
<tr>
<td>FVC (%pred)</td>
<td>78.5</td>
<td>89.8</td>
</tr>
<tr>
<td>(18.6)</td>
<td>(18.6)</td>
<td>(13.1)</td>
</tr>
<tr>
<td>R5 (cmH2O.s/L)</td>
<td>5.93</td>
<td>5.61</td>
</tr>
<tr>
<td>(1.77)</td>
<td>(1.73)</td>
<td>(2.29)</td>
</tr>
<tr>
<td>X5 (cmH2O.s/L)</td>
<td>-4.53</td>
<td>-3.88</td>
</tr>
<tr>
<td>(4.23)</td>
<td>(3.07)</td>
<td>(1.88)</td>
</tr>
</tbody>
</table>

Conclusion: The improvement in symptom control following BT may, in part, relate to deflation and consequent improvement in lung mechanics during tidal breathing. Differences in measuring impedance with FOT and IOS techniques will be discussed.

Grant Support: University of Sydney Post Graduate Scholarship

5 YEAR SINGLE CENTRE EXPERIENCE OF AIRWAY STENTING

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Introduction: Airway stenting is a therapeutic intervention for the management of complex malignant and benign central airway obstruction. Our institution commenced an airway stenting program 5 years ago. We present an audit of the patient case-mix and outcomes from this service since inception.

Methods: Chart review of stent insertion cases performed at Liverpool Hospital between Dec 2010 to July 2016. Indications, stent selection, peri-procedure and stent-related complications, survival and impact of stent on symptoms were reviewed.

Results: 37 stents (14 silicon; 20 self-expanding metallic, 3 Dynamic Y) were inserted in 32 patients. Stent numbers increased over the first 4 years then stabilised. The case-mix consisted of emergent malignant airway obstruction and outpatients with both benign and malignant pathology. 20 patients were from the Local Health District and 12 were referred from other Health Districts. 11 patients had stents inserted for non-malignant indications. In these patients, post-operative complications included mucous retention, stent migration, obstructing granulation tissue and mucosal flap tissue. Repeat bronchoscopies to address these complications averaged 4 procedures per patient (range 1–12). 21 patients had stents inserted for malignant indications primarily for palliative indications. 17 patients were able to be either transferred back to their referral centre to complete further therapy or were discharged from their acute hospital admission. There were 2 peri-procedural stent related deaths, 1 due to post-insertion hospital acquired pneumonia and another due to central airway laceration resulting in massive surgical and mediastinal emphysema.

Conclusion: Stenting offers symptomatic palliation from central airway obstruction but stent related complications are common, particularly in benign conditions. For malignant airway obstruction, the short term palliative benefits allow the majority of patients to be discharged from acute hospital admission offering scope for further treatments. Further study is required to look at physiologic and QOL improvements. Deaths – TBM massive emphysema lady, GOershal (pneumonia), include massive mediastinal lymphoma patient.

Grant Support: Nil
IDENTIFYING RISK FACTORS THAT PREDICT POOR TOLERANCE IN PNEUMOTHORACES FOLLOWING TRANSTHRORACIC NEEDLE BIOPSY (TTNB) OF LUNG LESIONS.

SHARAN RANDHAWA1, RANJAN SHRESTHA1
1Respiratory Department, Fiona Stanley Hospital, WA

Introduction: The average risk of developing pneumothorax following TTNB of a lung lesion is about 20%. This can be disabling especially in elderly patients with preexisting respiratory disorders and multiple comorbidities, resulting in hospital admissions either for observation or insertion of chest drain. To date there have been no studies directly correlating any risk factors in predicting poor tolerance to iatrogenic pneumothorax.

Aim:
• To identify all patients with pneumothorax post TTNB and record the outcomes of their hospital admissions.
• To identify if age, smoking history, preexisting respiratory disease and reduced FEV1 and or DLCO are risk factors for poor tolerance of pneumothorax and increased length of stay in hospital.

Methods:
• Demographic data of all patients who underwent TTNB (age, gender), primary lung disease, medical histories, smoking status, lung function tests (FEV1, FVC, DLCO, KCO, TLC) and length of hospital stay if complications occurred were retrospectively collected from two centers (Fiona Stanley Hospital and Fremantle Hospital WA) from August 2014 to February 2016.
• Patients who had a pneumothorax following TTNB who were either admitted for observation vs requiring a chest drain were recorded, including duration of hospital admission. Patients requiring a chest drain were defined as having “poor tolerance” to pneumothorax.
• Chi square and t tests were used to determine associations between age, preexisting COPD, smoking history, comorbidities, reduced lung function (FEV1 <50% or DLCO <50%) and “poor tolerance” to pneumothorax.

Results: A total of 154 TTNB procedures were carried out. 65 of the procedures (42.2%) were complicated with a pneumothorax and 13 patients (8.4%) required aspiration or chest drain. Mean length of stay (LOS) in days were 4.17 +/- 8.73 for all procedures.

Of those who had poor tolerance to pneumothorax, the average FEV1 % predicted was 66.5 +/- SD 19.8. Ie both were moderately reduced. 7 patients (58.3%) out of the 13 patients who required a chest drain had COPD. Chi Square testing did not show any statistically significant association between COPD and poor tolerance to pneumothorax (p=0.599).

There was no statistically significant association between older patients >70 years and poor tolerance of pneumothorax (p=0.934) or between reduced lung function and poor tolerance to pneumothorax (p=0.091).

Conclusion: Correlation analysis shows no association between presence of COPD/having reduced lung function/older age and “poor tolerance” to pneumothorax.

This is a negative study however it was limited by its small numbers. Larger studies looking at these risk factors will be crucial to prevent unnecessary lengthy hospital stays and chest drain insertions caused by a routine TTNB procedure.

Nil declaration of interest
BENEFIT OF STANDARD TRANSBRONCHIAL BIOPSIES AFTER ENDOBRONCHIAL ULTRASOUND GUIDED BIOPSIES FOR PERIPHERAL LUNG LESIONS

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Introduction/Aim: Endobronchial Ultrasound Guide Sheath (EBUS-GS) is used to locate and biopsy peripheral lung lesions with sensitivity and specificity of 73% and 100% respectively. (1) There is minimal data regarding a combined conventional transbronchial lung biopsy (cTBLB) plus EBUS-GS biopsy approach in the same patient. We aimed to assess the utility of this approach, and assess complication rates.

Methods: A retrospective analysis of patients who underwent EBUS-GS and cTBLB from August 2012 to October 2016 at the John Hunter Hospital, Newcastle, NSW.

Results: 73 patients were eligible with mean(SD) age 71(7) years. 43 were male. Mean(SD) lesion size was 26(3.5)mm. Of the 73 patients, lesions were successfully localised using EBUS-GS in 59(80%). Of these 59 lesions, 44 were concentric, 12 were eccentric and data was missing for the remaining 3. EBUS-GS biopsies were positive in 29 patients (49%) and definitive atypical pattern was seen in 6 patients (10%) giving a cumulative sensitivity of 60%. Brushings via guide sheath were positive in 20 patients (33%). cTBLB were definitively positive in 28 patients (38%) and an atypical cytology pattern was seen in 3 patients giving a cumulative sensitivity of 42% using non GS guided procedures. Standard brushing were positive in 15 patients (21%).

Four patients (7%) were diagnosed based on cTBLB alone, where EBUS-GS guided investigations were negative. Combining cTBLB and EBUS-GS at the same procedure increased sensitivity from 60% to 68%. There was one small pneumothorax managed conservatively and one episode of bleeding requiring local adrenaline therapy.

Conclusion: EBUS GS provides a higher success rate for diagnosis of peripheral lung lesions than cTBLB, however when EBUS-GS is combined with cTBLB, it may further increase the diagnostic yield.

Grant Support: Nil

Declaration of interest: Nil to declare

REFERENCE


REDDING HOSPITALISATION AND PROCEDURES IN MALIGNANT PLEURAL EFFUSION - RESULTS OF THE AMPLE RANDOMISED TRIAL

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Introduction/Aim: Malignant pleural effusions (MPEs) affect over one million people each year and herald limited prognoses. Talc pleurodesis and indwelling pleural catheter (IPC) are approved treatments with similar quality-of-life benefits in randomised studies to date, resulting in equipoise. Freedom from hospitalization and further interventions are important goals of management. The Australasian Malignant Pleural Effusion randomised trial aimed to determine whether IPCs are more effective than talc pleurodesis in reducing total hospitalization days in the remaining lifespan of MPE patients.

Methods: Open-labelled, randomized controlled trial recruited 146 participants with symptomatic MPE from nine centers in Australia, New Zealand, Singapore and Hong Kong and followed them for 12 months/until death. Participants were randomized (1:1) to IPC or talc pleurodesis, minimized by region (Australasia vs Asia), malignancy (mesothelioma vs others), and trapped lung (vs not). Primary endpoint was total number of days spent in hospital from trial intervention to death, or 12 months. Secondary outcomes included further pleural interventions, patient-reported breathlessness scores, quality-of-life measures, and adverse events.

Results: The IPC group spent significantly fewer days in hospital than the Pleurodesis group: median (IQR) 10.0 (14) vs 12.0 (14) days, p=0.026, which represented 6.2 (13.8)% vs 11.1 (33.7)% of their remaining lifespan, respectively, p<0.01. Mean reduction in hospitalization days was 3.6 days per patient: 12.7 (IPC) vs 16.3 (Pleurodesis) days. Fewer IPC-treated patients required further invasive pleural drainages (4.1% vs 22.5%, 95% CI 7.7-29.2%), improvement in breathlessness and quality-of-life, and adverse event rates were comparable.

Conclusion: IPC management reduces the time MPE patients spend in hospital before death and minimizes the need for further invasive pleural interventions, while providing equivalent symptomatic improvement, when compared with conventional talc pleurodesis.

Grant Support: The Sir Charles Gairdner Research Advisory Committee, Cancer Council of Western Australia and the Dust Disease Board of New South Wales, Australia.

Declaration of Interests: The authors have no conflict of interest.
Introduction/Aim: Mechanical ventilation, a lifesaving therapy for patients with respiratory failure, has been shown to contribute to mortality by inducing inflammation, which leads to multisystem organ failure through systemic effects. Different regions of the lung have been shown to heterogeneously respond to mechanical ventilation, which suggests that ventilator-induced lung injury may vary regionally. However, the impact of mechanical ventilation on regional lung inflammation is unknown. The aim of this study was to assess regional gene expression in response to mechanical ventilation in the healthy lung.

Methods: We ventilated two groups of BALB/c mice (n = 8 per group) for 2 h using protective [low tidal volume with moderate positive endexpiration (PEEP)] or injurious [high tidal volume with zero PEEP] ventilation strategy. mRNA levels of 19 genes of interest in ten different regions of each mouse lung were quantified using qPCR array and compared between the two groups and a free-breathing control group (n=8).

Results: The mRNA levels of ten genes were not differentially expressed between groups (P>0.05). Five genes had significantly different mRNA levels depending on the ventilation strategy (TFN-α, Cxcl-2, fos, IL-6, and Nfe2l2; P<0.01 for all comparisons). Five genes were differentially expressed between lung regions (TFN-α, Cxcl-2, IL-1β, Vim, and Ccl-2; P<0.05 for all comparisons), while two genes had differential regional expression that depended on the ventilation strategy (IL-6, P=0.02 and Ccl-2, P<0.01).

Conclusion: To our knowledge, this is the first demonstration of regional variation in gene expression in response to mechanical ventilation. Our results provide critical insight into the relationship between regional lung response to mechanical ventilation and regional injury. Future studies should be aimed at understanding the link between local tissue stretch and the expression of these injury related genes.

Grant Support: This study is funded by NHMRC grant # 1077905

Conclusion: These preliminary results show that sleep abnormalities are commonly present in subjects with COPD. While diurnal variability was seen both in spirometry and FOT parameters, they were not associated with sleep outcomes. Ongoing work will help us understand the relationship between circadian changes in lung function and sleep quality and symptoms in COPD.

1 OOSTVEEN E ET AL, RESPIRATORY IMPEDANCE IN HEALTHY SUBJECTS: BASELINE VALUES AND BRONCHODILATOR RESPONSE. ERJ. 2013 DEC;42 (6):1513–23

Grants: Menarini Australia

Grant Support
LUNG ULTRASOUND SURFACE WAVE ELASTOGRAPHY – PRELIMINARY MEASUREMENTS IN PATIENTS WITH INTERSTITIAL LUNG DISEASES

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Introduction/Aim: Lung ultrasound surface wave elastography (LUSWE) is a novel technique to assess surface lung tissue elastic properties by measuring surface wave propagation speed using an ultrasound detector system. The surface wave propagation on lung is noninvasively generated by a local and small mechanical actuation on the skin of the chest wall. Since many interstitial diseases predominantly affect the lung periphery, these may be especially suitable for this type of assessment, non-invasively and without radiation exposure.

Methods: 7 healthy controls and 7 patients who had CT chest identified peripherally distributed pulmonary fibrosis were studied. LUSWE was performed by generating small 0.1 second harmonic vibrations at 100 Hz by indenting a 3 mm area in an intercostal space with a handheld shaker and measuring the surface wave propagation on the lung with a ultrasound probe placed 5 mm away in the same intercostal space. Three measurements, at full inspiration, were made in each of 3 locations over each lung.

Results: All measurements below were made at 100 Hz and are in m/sec±SD.

<table>
<thead>
<tr>
<th></th>
<th>Controls</th>
<th>Patients</th>
<th>p (control v patient)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rt upper anterior</td>
<td>2.00±0.26</td>
<td>3.46±0.67</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Rt lateral</td>
<td>1.89±0.31</td>
<td>2.89±0.69</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Rt lower posterior</td>
<td>2.01±0.35</td>
<td>3.25±0.73</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Lt upper anterior</td>
<td>1.93±0.29</td>
<td>3.41±0.76</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Lt lateral</td>
<td>2.01±0.27</td>
<td>2.99±0.69</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Lt lower posterior</td>
<td>2.08±0.28</td>
<td>3.22±0.53</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

Conclusion: These results suggest that LUSWE can identify a difference in wave speed between control and abnormal, peripherally fibrotic, lungs. Its reproducibility, specificity and sensitivity, and ability to detect longitudinal change are yet to be defined but it has the promising advantage of being both non-invasive and radiation-free.

Grant Support: NIH R01HL125234 from the National Heart, Lung, and Blood Institute.
QUALITY OF LIFE AND SYMPTOMS ARE RELATED TO FORCED OSCILLATION MECHANICS DURING HOME-MONITORING IN COPD

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Introduction/Aim: The forced oscillation technique (FOT) is a non-effort dependent and objective measure of airway function. Home monitoring of FOT and its day-to-day variability may offer value in the management of COPD where peak expiratory flow or spirometry have not. This study assessed the relationship between FOT measures obtained from home-monitoring and quality of life and symptoms in COPD.

Methods: 10 COPD patients were recruited from Royal North Shore Hospital, Concord Hospital and the Woolcock Institute of Medical Research. At study enrolment, standard lung function tests were performed. A FOT home-monitoring device (ResTech srl, ResMon Diary device) was installed in each enrolled participant’s home over a period of 8–9 months. Participants were trained to make unsupervised measurements of FOT, to obtain resistance (Rrs), its standard deviation over the past 14 days (SDRrs), and the difference between inspiratory and expiratory reactance (ΔXrs), a measure of expiratory flow limitation. Daily symptoms were assessed by e-diary based on the COPD Assessment Test (CAT) and monthly quality of life via St. George’s Respiratory Questionnaire (SGRQ). The relationship between FOT measures vs SGRQ and CAT were assessed using mixed modelling.

Results: Participants’ mean±SD age was 69.7±10.8 years, with % predFEV1 38.4±5.2 and baseline SGRQ 53.2±16.3. The median(range) adherence was 95.7%(46.6-100%). Rrs and SDRrs were significantly related to SGRQ (fixed effect estimate 4.84±1.04 cmH2O•L−1, p=0.001, estimate 11.3±4.55 cmH2O•L−1, p=0.015, respectively), ΔXrs was significantly related to CAT (estimate −1.47±0.43 cmH2O•L−1, p=0.001).

Conclusion: Airway calibre and its variability in COPD were related to monthly quality of life, encompassing symptoms, activity limitation and psychosocial impact, while expiratory flow limitation related to daily symptom measures. Thus, FOT measures and variability reflect patient-based outcomes, providing supporting evidence for the utility of home monitoring of FOT in COPD.

Conflict of interest No. Grant Support: NHMRC Postgraduate Scholarship, LFA/Boehringer Ingelheim Top Up Research Grant, NHMRC Project Grant #1065938.

Grant Support: NHMRC Postgraduate Scholarship, LFA/Boehringer Ingelheim Top Up Research Grant.

Grant Support

RANDOMISED CONTROLLED TRIAL OF POLYSOMNOGRAPHY TITRATION OF NOCTURNAL NON-INVASIVE VENTILATION

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Introduction/Aim: Non-invasive ventilation (NIV) settings determined during the daytime can produce patient-ventilator asynchrony (PVA) at night, causing sleep disruption and limiting tolerance. The aim of this study was to determine if polysomnography (PSG) titration of NIV results in less PVA and sleep disruption than daytime titration alone.

Methods: Prospective randomised controlled trial in stable, English-speaking, treatment-naïve individuals requiring long-term NIV. Participants were randomly allocated to Control (daytime titration + sham PSG titration) or PSG (daytime titration + PSG titration). Primary outcome was the PVA and arousal indices on PSG at 10 weeks. Secondary outcomes included adherence (change in average daily NIV use), gas exchange (PaCO2, nocturnal oximetry), symptoms (somnolence, sleep quality, fatigue, dyspnoea) and health-related quality of life (HRQoL).

Results: Sixty participants were randomised (n=30 each group). Most participants (88.3%) had a neuromuscular disorder. There was one death, two drop-outs and three participants in each group did not complete the final PSG. PVA was less frequent in the PSG group (PSG 25.7 (12–68) vs. Control 41.0 (28–182), p=0.046) but there was no difference in arousals (PSG 11.4 (9–19) vs. Control 14.6 (11–19), p=0.258). Change in average daily NIV use was not different (PSG +51.8min (7 to 96) vs. Control +3.6min (−33 to 40), p=0.09) although a pre-specified subgroup with poor early adherence (>4hrs/day) increased their use after PSG (PSG +95min (29–161) vs. Control -23min (−86 to 39), p=0.01). PaCO2, somnolence and subjective sleep quality improved to a similar extent, and there were no differences in HRQoL, objective sleep quality or nocturnal gas exchange.

Conclusion: PSG titration of nocturnal NIV is associated with less PVA but not less sleep disruption than daytime titration. PSG titration improved NIV adherence in those who were poor users initially. No other short-term clinical benefits were demonstrated in this stable outpatient population. Further studies may help clarify the role of PSG titration of NIV.

Grant Support: Austin Medical Research Foundation, Institute for Breathing and Sleep, NHMRC Postgraduate Scholarship
TEN YEAR RETROSPECTIVE CLINICAL AUDIT OF TUBERCULOSIS IN FAR NORTH QUEENSLAND

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Introduction/Aim: To document changes in tuberculosis epidemiology in Far North Queensland for comparison with national data and two previous audits of the region.

Methods: Retrospective clinical audit of all cases of tuberculosis notified to the Cairns Chest Clinic from January 2006 to August 2016. Each case was classified as either confirmed or presumed on the basis of available microbiology/histology and evaluated with respect to site of disease, infectivity, demographics, mycobacterial sensitivities and Human Immunodeficiency Virus (HIV) status.

Results: 448 cases were identified, 370 confirmed. There were 307 cases of pulmonary tuberculosis of whom two-thirds (190/307) were smear positive; 154 cases of extra-pulmonary tuberculosis – mostly nodal (65) or pleural (45); and 21 cases of disseminated tuberculosis. Three-quarters (324/448) of cases were identified in the immigrant population – eighty percent (261/324) from Papua New Guinea (PNG). Of the remaining 124 cases, there were forty Torres Strait Islanders and nineteen Aboriginal Australians. Where mycobacterial sensitivities were known, two-thirds were fully sensitive (244/364); 41 mono-resistant; 78 multidrug resistant; and one extensively drug resistant. Rates of HIV co-infection were less than three percent (10/357); HIV status was not known in eighty percent (261/324) from PNG. Of the remaining one-quarter (324/448) of cases were identified in the immigrant population – eighty percent (261/324) from PNG.

Conclusion: Tuberculosis remains a significant public health problem in Far North Queensland. Total case numbers have increased three-fold since 2006. Much of the excess case burden comes from the immigrant population. Although PNG continues to account for the majority, the number of positive notifiable cases has increased fold since 2010. Rates of tuberculosis amongst Aboriginal Australians have fallen in response to policy changes. The proportion of smear-positive cases has also fallen.

Grant Support: Nil

Declaration of interest: None.

AUDIT OF TUBERCULOSIS SCREENING FOR HEALTH CARE WORKERS IN THE HUNTER NEW ENGLAND HEALTH SERVICE

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Aim: All NSW public hospitals are required to screen new health care workers (HCWs) identified as high risk for tuberculosis (TB) with tuberculin skin testing (TST), followed by a chest x-ray (CXR) if their TST is positive. HCWs diagnosed with TB disease or latent TB infection (LTBI) will be referred for further management.

This audit was undertaken to determine the incidence of TB and LTBI among HCWs in the Hunter New England Health Service (HNEHS), and their management following diagnosis.

Methods: An audit of the TB database Clinic Surveillance System was performed. Data from HCWs including students were collected from 2012 to 2015. This included country of birth, TST/Two-step results, TB/LTBI diagnosis, and management options.

Results: 900 high risk HCWs were audited. Most were born in Australia (36%), followed by China (9%), India (9%) and Malaysia (7%). Australian-born HCWs had lower rates of positive TST (16%) compared to those born overseas (54%).

Test | Result | Number of HCWs
--- | --- | ---
TST | Positive | 340
| Negative | 458
| Defaulted | 3
Two-Step | Positive | 25
| Negative | 70
| Defaulted | 4

Of the 365 HCWs who had a positive TST/Two-Step, there was 1 case of TB disease, 356 cases of LTBI, 2 with TST reversion and 6 who defaulted on their CXR.

Management of LTBI | Number of HCWs
--- | ---
Education | 282
Serial CXR | Attended 10
| Defaulted 32
Respiratory referral | QuantiFERON-TB Gold In-Tube (QFT-GIT) | Positive 6
| Negative 21*
| Defaulted 4
Treatment | 7

*included 2 with TST reversion

Conclusion: There was 1 (0.1%) case of TB disease and 356 (40%) cases of LTBI out of 900 high risk HCWs screened in the HNEHS. Only 7 (2%) HCWs received treatment for LTBI while the majority (79%) received education only. There was a high rate of nonattendance for serial CXR follow-up (76%) and a high rate of negative QFT-GIT following a positive TST (78%).

Grant Support: Nil

Declaration of interest: Nil
TRANSPLANTING THE HUMAN RESPIRATORY VIROME
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Introduction: The pulmonary component of the human respiratory virome (a subset of the human microbiome) is transplanted into the recipient at the time of lung transplantation (LTX). We explored the dynamics of virome transplantation and the role of intercurrent community acquired respiratory viruses (CARV).

Methods: A single centre, prospective, longitudinal study with measures of viral load in recipient nasopharyngeal swabs prior to LTX, swabs of explanted lungs, donor lungs prior to implantation and bronchoalveolar lavage (BAL) on post-operative days 1, 7, 21, 42, 63, 84. Samples were processed to isolate nucleic acids, followed by RNA extraction, cDNA synthesis and qPCR for CARV (Human Rhinovirus, Respiratory Syncytial Virus, Influenza A and B, Parainfluenza Virus 1, 2, 3, and Human Metapneumovirus).

Results: 18 consecutive LTX subjects (bilaterial: heart lung = 17:1) (age 49 ± 13 years, mean ± SD) (M: F= 9:9) were recruited. Follow up was 51±36 days, range 2–108 days. The most frequent CARV was Influenza A, followed by influenza B. In 4 individuals, influenza A virus was detected on the swab of the explanted lung at a low viral load, and in 3 of these, Influenza A was also detected in the day 1 BAL at an increased viral load. Multiple patients had Influenza A detected in consecutive BAL from post-operative days 1, 7 and 21, while Influenza B and Parainfluenza 1/2 were detected from day 21. Indication for transplant was not a determinant of viral detection.

Conclusion: Viral load increases in subjects at the time of LTX, likely associated with immunosuppression. CARV appear to be part of the transplant respiratory virome as analysed by BAL, showing progressive expansion of select populations following LTX. The long-term outcomes associated with the ongoing presence of CARV may be elucidated following this longitudinal study.

Grant Support: Nil

PAEDIATRIC TUBERCULOSIS IN TIMOR-LESTE: OPPORTUNITIES FOR IMPROVING RECOGNITION, DIAGNOSIS AND PREVENTION
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Introduction/Aim: Timor-Leste has the highest incidence of TB in Southeast Asia but a relatively low rate of paediatric notifications. Bairo Pite Clinic (BPC) in Dili manages more than a fifth of all TB cases in the country. Improved understanding of TB epidemiology in Timor-Leste will inform quality improvement activities focused on recognition, accurate diagnosis and prevention of TB in children. We aimed to describe the epidemiology and diagnostic features of notified paediatric TB cases at BPC and to compare this to national paediatric TB notification rates.

Methods: Prospective collection of demographic, clinical and laboratory data for all patients diagnosed with TB at BPC between January 2014 and December 2014 inclusive. Comparison is made with national notification data with respect to paediatric notification rates.

Results: In total, 4090 TB cases were notified. Of these, 426 (10%) were children aged <15y; BPC notified 886 cases, of which 110 (12%) were aged <15y. Contact with a known TB case was reported in 39/110 (39%) BPC cases; 15 (14%) were identified through screening household contacts of infectious index cases. Evidence of BCG vaccination was documented in 36/110 (33%); Malnutrition was identified in 39/110 (35%); HIV in 2/110 (2%). Pulmonary TB was diagnosed in 78/110 (71%) BPC cases and 318/426 (75%) paediatric cases nationally. Bacteriological confirmation occurred in 5/78 (6%) pulmonary cases at BPC and 36/318 (11%) cases nationally. In districts other than Dili and Baucau, 29/131 (22%) paediatric pulmonary TB cases were sputum smear positive.

Conclusion: Paediatric TB is common in high TB burden settings such as Timor-Leste but is under-recognised. Malnutrition is an important risk factor; HIV co-infection is rare. Opportunities for prevention include improved BCG coverage and implementation of household contact tracing. Consideration should be given to improved specimen collection and strategic use of PCR to improve diagnostic certainty.

Grant Support: English Family Foundation
AIRWAY EPITHELIAL INNATE IMMUNE RESPONSES TO CORONAVIRUSES
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Introduction/Aim: Despite the recurring emergence of novel pathogenic coronaviruses (CoVs) SARS-CoV (2002) and MERS-CoV (2012), there is a lack of understanding of host immunity and disease mechanisms during respiratory infection with CoVs. In this study, we characterised the innate immune responses of differentiated primary bronchial epithelial cells (pBECs) to infection with related less virulent OC43-CoV and 229E-CoV.

Methods: Human pBECs were obtained from subjects by brushing during bronchoscopy. Cells were grown at the air liquid interface (ALI) until differentiation with cilia beat and mucus production was observed (25–30 days). Cells were infected at a multiplicity of infection (MOI) 0.1 or 1 with OC43-CoV or 229E-CoV for 6, 24, 48, 72, 96 hours and 7 days. Supernatants, total cell protein extracts and RNA were collected at each time point to measure expression of pro-inflammatory- and anti-viral-cytokines, and virus replication.

Results: OC43-CoV and 229E-CoV demonstrated different viral replication kinetics. 229E-CoV replicated earlier and more efficiently, peaking between 48 and 72 hours. This was associated with an activation of the innate host response, with induction of type I interferon (IFN-α), and IFN stimulated genes, CXCL-10 and viperin. In contrast, replication of OC43-CoV peaked later between 96 hours and 7 days. This virus demonstrated attenuated levels of IFN-α, CXCL10 and viperin. We have begun to investigate virulence factors that control epithelial responses to CoV. Sequence analysis of viral proteins (EMBO Needle) revealed only 49.2% identical matches between OC43 and 229E highlighting the potential for variation in host (epithelial)-virus interaction.

Conclusion: This study demonstrated that both OC43 and 229E-CoVs replicated in differentiated pBECs, but they induce a divergent innate immune response potentially linked to their different replication kinetics. Understanding the host-virus interaction for these less virulent coronaviruses will give insight into pathogenic mechanisms underpinning SARS-CoV and MERS-CoV-induced respiratory disease.

Grant Support: University of Newcastle Research Scholarship UNRSC

Declaration of Interest: The authors have no conflict of interest.

INFLUENZA AND BACTERIAL CO-INFECTIONS: THE GOLD COAST HOSPITAL AND HEALTH SERVICE (GCHHS) EXPERIENCE
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Introduction/Aim: Seasonal influenza infections are a leading cause of mortality and morbidity. Literature suggests that bacterial co-infections complicate 20-33% of influenza cases and contribute to a more severe and prolonged course of illness. The challenge in clinical practice remains in the identification of patients who will benefit from empirical antibiotics.

Methods: This is a retrospective audit of patients with influenza infection diagnosed on viral nasopharyngeal swab in the GCHHS between July and September 2015. Demographics, clinical and microbiological data, complications and outcomes were collected. Data on antibiotic prescription was also collected. Patients with and without bacterial co-infection were compared.

Results: 481 patients were diagnosed with Influenza infection on viral nasopharyngeal swab. Bacterial co-infection was present in 52 (11%) patients. The most common co-infecting bacterial pathogen is Staphylococcus aureus (n=13, 22%), Streptococcus pneumoniae (n=11, 18%) and Pseudomonas aeruginosa (10, 17%). Chronic lung disease and COPD were associated with a significantly higher risk of bacterial co-infection (p<0.05). Other co-morbidities examined including active smoking, active malignancy, chronic cardiovascular disease, end-stage renal failure, diabetes mellitus and immunosuppressed states did not achieve statistical significance. 218 (51%) patients who did not have microbiological findings of bacterial co-infection received antibiotics. Prescription of antibiotics in this group of patients resulted in no significant difference in in-hospital mortality (p=0.22). In-hospital mortality, length-of-stay and ICU admission were significantly higher in patients with bacterial co-infection compared with those without, although 30-day mortality was similar between the two groups.

Conclusion: The incidence of bacterial co-infection in our cohort of patients is lower than anticipated. Our findings suggest a probable over-use of antibiotics. A group of patients who may benefit from empirical treatment with antibiotics is those with chronic lung disease and COPD. A prospective study to develop a scoring system could help identify patients who would benefit from empirical antibiotics.

Grant Support: Nil
INCREASED PROPORTION OF SLOW DECLINERS IN PATIENTS WITH ALPHA-1 PROTEASE INHIBITOR (A1-PI) DEFICIENCY FOLLOWING TREATMENT WITH A1-PI

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Introduction: The RAPID trial (NCT00261833) showed that intravenous A1-PI treatment (Zemaira®; n=93) can significantly reduce the mean decline in lung density vs. placebo (n=85). Following RAPID, 140 patients completed a 2-year open-label Extension trial (NCT00670007) in which all patients received active treatment. However, it is unclear what proportion of patients had a fast or slow decline in lung tissue loss and if the proportion of slow decliners was reduced following A1-PI treatment.

Aim: To compare the effect of A1-PI treatment on the proportion of fast vs. slow decliners in patients completing both RAPID and RAPID Extension.

Methods: Adjusted CT scans were analysed and A loss of >2g/L/yr was used as a cut-off to define fast decline. Relative proportions of fast (>2g/L/yr) and slow decliners (≤ 2g/L/yr) were compared using the Fisher’s exact test.

Results: During RAPID, a significantly higher proportion of patients receiving A1-PI were classified as slow decliners vs. placebo (72% vs 50%; p<0.009). When all subjects received A1-PI in the RAPID Extension, the proportion of subjects classified as fast decliners was similar between the arms (67% vs 75%; p=0.352). [Table 1]

<table>
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<tr>
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<tbody>
<tr>
<td>N</td>
<td>75</td>
<td>64</td>
<td>75</td>
<td>64</td>
</tr>
<tr>
<td>Slow n (%)</td>
<td>54 (72)</td>
<td>32 (50)</td>
<td>50 (67)</td>
<td>48 (75)</td>
</tr>
<tr>
<td>Fast n (%)</td>
<td>21 (28)</td>
<td>32 (50)</td>
<td>25 (33)</td>
<td>16 (25)</td>
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<tr>
<td>p-value</td>
<td>0.009</td>
<td></td>
<td>0.352</td>
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</table>

Conclusion: Treatment with A1-PI therapy is associated with a higher proportion of patients with a slow decline in lung tissue loss compared to placebo. After switching to active treatment there is a progressive shift from fast to slow decliners, further demonstrating the benefit of A1-PI treatment.
EVALUATING BLOOD EOSINOPHILS AND EXACERBATION HISTORY TO PREDICT INHALED CORTICOSTEROIDS RESPONSE IN COPD

DIMITAR SAJKOV1 PRESENTING ON BEHALF OF PETER M. A. CALVERLEY2, KAY TETZLAFF3,4, CLAUS VOGELMEIER5, LEONARDO M. FABBRI6, HELGO MAGNUSSEN7, EMIEL FM WOUTERS8, BERND DISSE9, HELEN FINNIGAN9, GUUS M. ASIJEE9, HENRIK WATZ7

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Introduction/Aim: Inhaled corticosteroids (ICS) are used to reduce the rate of chronic obstructive pulmonary disease (COPD) exacerbations. Debate continues over use of blood eosinophils (EOS) to predict ICS response, with some suggesting a cut-off of ≥2%. In the WISDOM study (NCT00975195), this response was driven by patients with higher EOS levels. The rate of chronic obstructive pulmonary disease (COPD) exacerbations. In patients who do not meet these criteria, ICS may not be as effective as is commonly assumed.

Results: High EOS counts (≥400 cells/μL) were associated with increased exacerbations rate after complete ICS withdrawal only in patients with ≥2 prior exacerbations.

Figure: Rate ratios (ICS withdrawal/ICS) for moderate or severe exacerbations by EOS subgroup and exacerbations history

<table>
<thead>
<tr>
<th>Factors</th>
<th>Number of patients</th>
<th>Rate ratios</th>
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<tr>
<td></td>
<td></td>
<td>&lt;2 exacerbations</td>
</tr>
<tr>
<td>Total</td>
<td>1454</td>
<td>1.14</td>
</tr>
<tr>
<td>Baseline eosinophil</td>
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<td></td>
</tr>
<tr>
<td>&lt;150 μL</td>
<td>864</td>
<td>1.11</td>
</tr>
<tr>
<td>≥150 μL</td>
<td>503</td>
<td>1.02</td>
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<tr>
<td>≥300 μL</td>
<td>421</td>
<td>1.19</td>
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<tr>
<td>Baseline eosinophil</td>
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<td></td>
</tr>
<tr>
<td>&lt;300 μL</td>
<td>1121</td>
<td>1.11</td>
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<tr>
<td>≥300 μL</td>
<td>859</td>
<td>1.02</td>
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<tr>
<td>≥500 μL</td>
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<tr>
<td>Baseline eosinophil</td>
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<tr>
<td>≥500 μL</td>
<td>155</td>
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<tr>
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<tr>
<td>≥1500 μL</td>
<td>738</td>
<td>1.00</td>
</tr>
<tr>
<td>≥2000 μL</td>
<td>208</td>
<td>1.25</td>
</tr>
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</table>


Grant Support: The study was funded by Boehringer Ingelheim.

Declaration of Interest Statement: DS has no real or perceived conflict of interest that relates to this presentation. PMAC has advised various pharmaceutical companies including Boehringer Ingelheim, GlaxoSmithKline, Astra Zeneca, Novartis, Nycomed, Chiesi, Almirall and Takeda on the design and conduct of clinical trials and has spoken at meetings supported in whole or part by these companies. KT, BD, HF and GMA are employees of Boehringer Ingelheim. CV has served as an advisor or consultant for AstraZeneca, Boehringer Ingelheim, GlaxoSmithKline, Novartis, Takeda; and has received grants for clinical research from GlaxoSmithKline and Grifols. LMF has consultant arrangement with, is lecturer for, is on the advisory board for, and receives research support from Boehringer Ingelheim and is lecturer and member of the advisory board for Pfizer. HM has relationships with Boehringer Ingelheim, BerlinChemie, Almirall, AstraZeneca, Chiesi, Novartis, Takeda, AB2BIO, Bayer, and Intermune. EFMW reported serving on advisory committees of Nycomed and as a speaker for AstraZeneca, GlaxoSmithKline, and Novartis; he has received research support from AstraZeneca and GlaxoSmithKline. HW has received consultancy fees and travel support from Takeda; has received consultancy fees from Almirall, AstraZeneca, Boehringer Ingelheim, GlaxoSmithKline, and Novartis; has received research support from AstraZeneca and GlaxoSmithKline; has received lecture fees from AstraZeneca, Almirall, Boehringer Ingelheim, BerlinChemie, GlaxoSmithKline, Merck, and Novartis; has received payment for development of educational presentations from Boehringer Ingelheim and BerlinChemie; and has received travel support from AstraZeneca, GlaxoSmithKline, and Novartis.

Conclusion: Withdrawal of ICS increased the rate of exacerbations only in patients with both raised EOS (≥400 cells/μL) and a history of frequent exacerbations. In patients who do not meet these criteria, ICS may not be as effective as is commonly assumed.
REDUCED PULMONARY ARTERY DISTENSIBILITY ON DYNAMIC CT CORRELATES WITH INCREASED EMPHYSEMA AND IMPAIRED RIGHT VENTRICULAR FUNCTION

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Introduction/Aim: Pulmonary hypertension (PH) in COPD is associated with increased disability and reduced survival. Detection of PH in COPD is challenging due to limitations of echocardiography and potential risks associated with right heart catheterization (RHC). The main pulmonary arteries (PA) become progressively less distensible in PH. Previous studies have shown PA distensibility to correlate well with measured mPAP at RHC. We assessed relationships between PA distensibility, cardiac function and emphysema index measured using dynamic ECG-gated MDCT.

Methods: 34 patients with stable COPD (mean[SD] age 65.8[9.7], FEV1% 43.9[22.5], 18 male) underwent dynamic retrospective ECG-gated MDCT. Emphysema was quantified by CT densitometry based upon a density threshold of <-950 Hounsfield Units (HU). The main pulmonary artery cross-sectional area (CSA) was measured at 10% intervals from 0-90% of the R-R interval. PA distensibility (%) was calculated as ((maximum CSA – minimum CSA)/minimum CSA) x 100. Cardiac function parameters were assessed using semi-automated software with manual correction if required. The relationship of PA distensibility with both parameters were assessed using semi-automated software with manual correction if required. The relationship of PA distensibility with both emphysema index and right ventricular ejection fraction (RVEF) was tested via Pearson’s r.

Results: Mean[SD] PA distensibility was low (16.5[5.9]%), PA distensibility was significantly negatively correlated with emphysema index (r=−0.365, p = 0.031) and positively correlated with reduced RVEF (r=0.483, p=0.0025).

Conclusion: Dynamic MDCT permits simultaneous evaluation of emphysema, PA pressure and right ventricular performance. Increased severity of emphysema appears associated with reduced PA distensibility. Impaired PA distensibility appears associated with decreased RVEF. Dynamic MDCT provides a valuable tool for assessment of heart-lung interaction in COPD.

Antimicrobial prescription in patients dying from chronic obstructive pulmonary disease

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Introduction: It is estimated that by 2050 antibiotic resistance will cause 10 million deaths/year, thus outstripping cancer deaths. In patients dying with advanced cancer, dementia, or residing in hospice, antimicrobials are commonly prescribed in the absence of clinical symptoms to support a bacterial infection. The extent and appropriateness of antimicrobial use in patients with chronic obstructive pulmonary disease (COPD) is unknown.

Aim: To review antibiotic prescription at the time of death in patients who died from COPD.

Methods: A retrospective medical record audit was performed for 475 patients who died over twelve years between 2004–2015. Patients were analysed within three groups: Group 1 – radiographic consolidation, Group 2 – elevated inflammatory markers (WCC) or (CRP) with no radiographic consolidation and Group 3 - normal inflammatory markers and no radiographic consolidation.

Results: Two hundred and twenty one patients died from COPD. Median age was 80 years and 136 (60%) were male. Median respiratory number and duration of use were similar in all three groups. 57 (59%) patients used home oxygen and 156 were ex-smokers (70.6%).

Antimicrobials were prescribed to 201 (91%) patients (Table 1). Antibiotic number and duration of use were similar in all three groups. 57 (59%) Group 2 patients received ceftriaxone and 34 (35%) received azithromycin first line, which are recommended treatments for pneumonia but not acute exacerbations of COPD (AECOPD). Similarly 24 (46%) Group 3 patients (without objective evidence of infection) received ceftriaxone and 16 (31%) received azithromycin.

Conclusion: Almost all patients who died from COPD received antimicrobials during their final hospital admission, irrespective of objective evidence suggesting infection. While antibiotics were mainly prescribed according to guidelines in patients with pneumonia, guideline or evidence based prescription of antibiotics occurred less in patients dying with AECOPD or without objective evidence of infection.

Grant Support: Nil
LUNG FUNCTION TRAJECTORIES OVER THE LIFE SPAN, THEIR ASSOCIATED CHILDHOOD FACTORS AND CONSEQUENCES: A POPULATION BASED LONGITUDINAL COHORT

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Introduction/Aim: Individuals with different patterns of lung function growth and decline may have different risks for developing COPD. We investigated trajectories of lung function from childhood to middle age and their associated childhood factors.

Methods: Using 2142 subjects from the Tasmanian Longitudinal Health Study, pre-bronchodilator FEV1, measured at 7, 13, 18, 45, 50 and 53 years, was modelled with Group Based Trajectory Modelling. Logistic regression was used to investigate the relationship between childhood factors and trajectory groups.

Result: The best fit model showed six distinct trajectories (Figure 1). Based on initial lung function at 7 years, lung function growth and decline rates, the trajectories were labelled as “early low, reduced growth, accelerated decline” (7.4%; n = 159), “early normal/high, normal growth, accelerated decline” (5%; n = 107), “early low, normal growth, normal decline” (28.7%; n = 615), “early low, accelerated growth, normal decline” (4.2%, n = 90), “persistently high” (14%; n = 300) and “normal” (40.7%; n = 871). The first three trajectories had increased risk of COPD (post-bronchodilator FEV1/FVC < LLN) at age 53 years compared with the “normal” group. Childhood asthma, bronchitis, pneumonia/pleurisy, maternal asthma and maternal smoking were positively associated with these three trajectories while negative associations were observed for breast feeding and childhood overweight.

Figure 1 Trajectories of FEV1 from 7 to 53 years of age. The six trajectories represent the latent growth patterns of lung function.
Conclusion: This is the first study to develop lung function trajectories from childhood to middle age in a general population. We identified three trajectories for increased risk of COPD: (1) only accelerated lung decline in adulthood, (2) low initial lung function in childhood with normal growth and normal decline, and (3) low initial lung function in childhood with reduced growth and accelerated decline. Important modifiable childhood risk factors for these three trajectories included exposure to maternal smoking in childhood and a lack of exclusive breast feeding in very early life. Strategies to maximize lung function growth from early life as well as preventing accelerated decline are important for COPD prevention.

Grant Support: National Health and Medical Research Council of Australia, Clifford Craig Medical Research Trust of Tasmania; Victorian, Queensland & Tasmanian Asthma Foundations

Declaration of Interest Statement: There is no conflict of interest.

A FAST TRACK CLINIC IMPROVES DIAGNOSIS AND TREATMENT TIMES FOR THOSE INVESTIGATED FOR LUNG CANCER IN RURAL NEW ZEALAND

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1,2General Medicine, 3Clinical analysis, 4Radiology, 5Respiratory Medicine, Whangarei Hospital, Northland, New Zealand

Introduction: In 2009 a Respiratory Fast Track Clinic (RFTC) was introduced successfully by the Southern District Health Board. We sought to improve on this model by incorporating further methods of biopsy in a similar rural population in New Zealand (NZ).

Methods: The RFTC introduction was bi-phasic, initially patients had their CT scan on their first specialist assessment (FSA) (phase 2). After two months biopsy was incorporated (bronchoscopy, CT-guided biopsy, ultrasound-guided biopsy) into the RFTC (phase 3). Patients with suspected lung cancer were identified between December 2015 and May 2016 prior to the RFTC (phase 1) and the time for the diagnostic pathway was measured and compared to those in phases 2 (May to July 2016) and 3 (July to October 2016). Median times were used for statistical analysis as some patients underwent long waits for personal or clinical reasons thereby distorting the data.

Results: 212 were investigated for suspected lung cancer. Prevalence by age was normally distributed peaking 61–70, with preference for lower socio-economic deciles, equal gender and proportional ethnic distributions. Endobronchial ultrasound (EBUS) was the most utilised biopsy method, followed by bronchoscopy.

Time from GP referral to FSA between phase 1–3 improved significantly (p = 0.005). Similarly, time from FSA to diagnosis and treatment improved significantly, median times reducing from 15 to 0 (p < 0.001) and from 37 to 24 days (p = 0.004) respectively between phases 1–3. (Table 1)

Table 1 The effectiveness of the rapid access clinic on pathway timing from referral to commencing treatment

<table>
<thead>
<tr>
<th>Measurement</th>
<th>Average difference in days phase 1 vs phase 3</th>
<th>P=</th>
</tr>
</thead>
<tbody>
<tr>
<td>GP referral to Appointment</td>
<td>–4.5</td>
<td>0.005</td>
</tr>
<tr>
<td>Appointment to Diagnosis</td>
<td>–16.6</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Appointment to Treatment</td>
<td>–13.8</td>
<td>0.004</td>
</tr>
<tr>
<td>GP referral to Treatment</td>
<td>–17</td>
<td>0.74</td>
</tr>
</tbody>
</table>

Discussion: The RTFC significantly shortened time to diagnosis and treatment in our population. To the best of our knowledge, this is the first study demonstrating a reduction in time to treatment for lung cancer patients in NZ in a fast track clinic. A previous study (1), using EBUS as the initial investigation resulted in faster treatment and seemed to improve survival. EBUS could easily be incorporated into the RFTC model and this may confer a survival advantage.


Grant Support: None

Conflicting Interests: None
DIFFQUICK STAINED CYTOLOGY SMEARS PROVIDE IMPROVED MALIGNANT CELL DNA YIELDS FROM LYMPH NODES AT ENDOBRONCHIAL ULTRASOUND

TRANSBRONCHIAL NEEDLE ASPIRATION (EBUS-TBNA)

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Introduction/Aim: We are currently optimising procedural aspects of Endobronchial Ultrasound Transbronchial Needle Aspiration (EBUS-TBNA) with the aim of converging rapid on-site evaluation (ROSE) with Targeted Amplicon Sequencing (TAS) of “hotspot” DNA mutations and RNA translocations to enhance lung cancer diagnosis. Here we explore a comparison of cellularity and DNA yields between DiffQuik stained cytology smears and formalin fixed paraffin embedded (FFPE) cell blocks.

Methods: In a prospective study of 63 consecutive patients with lung malignancy (m/f ratio 1.2, mean 66.6 ± 9.5 years), EBUS-TBNA was performed using 2–5 (median 3) aspirates with an Olympus needle (NA-201SX-4021). Smear and cell block tumour cell abundance were assessed by experienced cytopathologists on a scale from 0 (absent) to 4 (abundant). Genomic DNA was isolated from both smears and blocks, quantified then sequenced using the TruSeq Amplicon Cancer Panel (Illumina, San Diego, CA) on a MiSeq platform. Diff quik slides were digitally scanned before being used for DNA extraction.

Results: Unexpectedly, the overall cellularity was higher in the cytology slides than the cell blocks: 35/63 cases had 3–4 scores for tumour cell abundance compared with 22/64 cell blocks (p < 0.05, Chi-squared test). 39 cases provided adequate DNA for amplicon sequencing (>50 ng). The highest DNA yields from cell blocks and cytology slides were 4.80 μg and 15.96 μg respectively, with mean (standard deviation; SD) DNA yields from cell blocks (439 ng (858 ng)) and cytology slides (1745 ng (3173 ng)) differing significantly (p < 0.001, paired t-test). Extracted DNA was of good quality exhibiting DNA fragment sizes of up to 10Kb on a 1% agarose gel.

Conclusion: DiffQuik stained cytology slides present numerous advantages over FFPE cell blocks as a source of DNA for molecular pathology including increased DNA yield, and an immediate visual confirmation of adequate malignant cell sampling.

Grant Support: Pathology Queensland Study Education and Research Grant.

Conflict of Interest: Nil

ASThma Control During Pregnancy, 17q21 Variants and Childhood-Onset Asthma

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Introduction/Aim: Our aim was to investigate the effect of FeNO-guided versus symptom-guided asthma management during pregnancy on the prevalence of childhood-onset asthma.

Methods: The Growing Into Asthma (GIA) study is a prospective longitudinal birth cohort following up the offspring of asthmatic women involved in a double-blind RCT comparing a treatment algorithm using FeNO in combination with asthma symptoms ascertained with the asthma control questionnaire [ACQ] (FeNO Group) against a treatment algorithm using ACQ only (clinical group) which found a 50% reduction in asthma exacerbations during pregnancy in the FeNO group. 140/179 (78%) consenting children completed the follow-up at 4–6 years of age. Four asthma-associated 17q21 single-nucleotide polymorphisms (SNPs) rs7216389, rs8076131, rs9303277 and rs2290400 were genotyped.

Results: FeNO-guided asthma management during pregnancy significantly reduced the odds ratio for childhood-onset asthma in the offspring (OR 0.45, 95% confidence interval [CI] 0.21–0.95, p = 0.04). Furthermore frequent wheeze in the past 12 months (OR 0.26; CI 0.08–0.81, p = 0.02), wheeze ever (OR 0.49; CI 0.25–0.98, p = 0.04) and recurrent bronchiolitis (OR 0.20; CI 0.06–0.62, p = 0.005), were less common in the FeNO group. Asthma was significantly associated with all four variants at 17q21 single-nucleotide polymorphisms (SNPs) rs7216389, rs8076131, rs9303277 and rs2290400 were genotyped.

Results: We found that FeNO-guided asthma management during pregnancy significantly reduced asthma prevalence in childhood. 17q21 variants interact with bronchiolitis in early life to increase the asthma risk (gene-environment interaction), but there is no evidence of an interaction with FeNO guided asthma management. Thus the prevention strategy may benefit children with and without variants at 17q21.

Grant Support: NHMRC, Hunter Children’s Research Foundation, Hunter Medical Research Institute, John Hunter Hospital Charitable Trust, University of Newcastle Priority Research Centre GrowUpWell