attended a screening visit; 58/129 screen failed (eg, due to deterioration in peak flow, unable to wean off regular asthma medications) leaving 71 randomised (2.5%) of total patients invited. Trial 2: similar picture, completed July 2011, extended by 6 months due to slow recruitment. Target to randomise =80, target to complete =68. Actual completed: 71/8398 (<1%) of those invited.

**Conclusion** Achieving the completion target in randomised controlled trials requires significant administrative support, and the capacity to increase support should difficulties in recruitment be encountered. Closer partnership with primary care practitioners, better access to primary care patient databases and direct contact with potential recruits can overcome this. Loss of potential recruits during the run-in phase needs exploration, and is of significant importance to improve the efficiency of screening to randomisation. Addressing these issues will mean fewer trials are underpowered and hence provide better return for grant awarding bodies.

**P206 PSYCHOLOGICAL COMORBIDITY IN VOCAL CORD DYSFUNCTION**

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E Gregson, S Campbell, S Lilie, R Stacey, J Haines, S Fowler, A Vyas. Royal Preston Hospital, Preston, UK

**Background** Vocal Cord Dysfunction (VCD) is typically reported to affect young females and has been associated with a psychiatric history. We run a multidisciplinary-based service for VCD patients with input from specialist speech and language therapy, physiotherapy and psychology. We investigated the demographics of our patient cohort with VCD, its association with anxiety and depression and whether this affected treatment response.

**Methods** All patients referred for specialist speech and language therapy at the Royal Preston Hospital Airways Clinic between June 2006 and May 2011 with VCD confirmed by endoscopy were included. During routine clinical care data were collected including demographic details and comorbidities. Patients were also asked to complete the Hospital Anxiety and Depression (HAD) questionnaire. Subjective symptomatic improvement was recorded at patient follow-up visits.

**Results** A total of 95 patients were eligible for study inclusion: 75.7% were female with a median age of 53 (17–83) years while men were older at 59 (37–80) years. Medical comorbidities included asthma (56.8%), reflux (47.4%), chronic cough (17.9%), nasal disease (16.8%) and neurological disease (12.6%). A history of confirmed psychiatric disease was noted in 38.9%. In 43 patients who completed HAD scores, moderate or severe anxiety was found in 41.8% and depression in 23.3%. Response to treatment was excellent (67% showing clinical improvement) and this was independent of medical or psychological comorbidity.

**Conclusions** Our data suggest that VCD affects a wide range of patients, in terms of age, gender, comorbidities and HAD scores which do not impact on treatment response, as a result of the multidisciplinary approach and this success is comparable to most asthma therapies when patient compliance and education are accounted for. It challenges many previously held concepts and supports a multidisciplinary approach to treating VCD.

**P207 DOES THE NIJMEGEN CORRELATE TO THE D12 WHEN USED AS AN OUTCOME MEASURE IN PATIENTS WITH BREATHING PATTERN DYSFUNCTION**

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R Johnston, F Shaw, A Menzies Gow, L Flude. Royal Brompton and Harefield NHS Foundation Trust, London, UK

**Introduction** Breathing pattern dysfunction (BPD) in patients with and without respiratory disease is linked to disproportionate dyspnoea. Asthma patients in particular have a high prevalence of BPD. The Nijmegen questionnaire (NQ) is a validated outcome measure in patients with no underlying respiratory pathophysiology, not in patients with organic disease. There is no validated tool that can be used to assess the effectiveness of breathing retraining in patients with BPD. The Dyspnoea-12 (D12) questionnaire provides a global score of breathlessness severity and is validated in respiratory disease, but not in patients with BPD. The aim was to correlate the D12 to the NQ and assess its use as an outcome measure for patients with BPD. Pre and post scores were compared to assess the effectiveness of Physiotherapy breathing retraining on patients with BPD referred to our tertiary centre.

**Method** We included all Physiotherapy referrals to the outpatient department of the Royal Brompton Hospital for BPD between January and July 2011. The D12 and NQ were completed before and after Physiotherapy interventions and respiratory rate was also measured.

**Results** Initial data were available for 26 patients: mean (SD) age 47.6 yrs (13), 16 (62%) female, 13 (50%) had a diagnosis of asthma. Post intervention data were available for 17 (65%) patients. Respiratory rate did not correlate with the NQ or D12 in any of the groups. There was a statistically significant correlation between the NQ and D12 both pre (p=0.0017, R=0.59) and post (p=0.0156, R=0.58) intervention. A statistically significant difference pre and post intervention was shown in the NQ (p=0.0242 mean (SD)=7.35 (12.1)) and the D12 (p=0.0127).

**Conclusion** The data suggests that breathing retraining provides both a clinically effective and statistically significant improvement in symptoms of patients with BPD. There was a significant correlation between the D12 and NQ scores in patients referred with BPD. This highlights that the D12 may be an appropriate outcome measure in this patient population.

**P208 OBESITY AUGMENTS CIRCULATING NEUTROPHIL LEVELS IN ASTHMA**

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M C Pyn, G Davies, C A Thornton, A Bryant, R Jones. Swansea University, Swansea, Wales

**Introduction** Mechanisms underlying the association between obesity and asthma are not well understood. Obesity is characterised by chronic inflammation and adipose tissue, comprising up to 50% pro-inflammatory cells, produces many pro-inflammatory cytokines and hormones (adipokines). Enhanced systemic inflammation might provide the causal link between obesity and asthma. Although there are good mechanistic data that obesity can augment innate immune function and promote immune dysregulation by reducing regulatory T cell (Treg) numbers there is little work in this area in relation to asthma.

**Methods** A case-control study is being conducted examining six groups of pre-menopausal women (n=120): normal weight (BMI <30 kg/m²), overweight (BMI 25–30 kg/m²) and obese individuals (BMI ≥30 kg/m²) with and without asthma. Asthma diagnosis was physician confirmed, and severity graded. Measures of adiposity, lung function and blood were collected during menstruation. Automated haematology analysis was used to quantify major cell types and chemiluminescence to measure whole blood reactive oxygen species generation following stimulation. Flow cytometry was used to examine major lymphocyte subtypes including Treg cells. A number of circulating cytokines and adipokines will be measured on sample collection completion.

**Results** Interim analysis of 36 individuals revealed a significant increase in circulating total leucocytes with increasing BMI which is more pronounced in asthmatics compared with controls (p=0.022). This appears to be due to a significant increase in neutrophils