temperature was lower in 20/30 patients, max decrease 1.3°C (before 37.1°C (36.3–37.9°C) vs. 36.6°C (35.9–37.4°C)).

Conclusions Early screening using an MRI incubator is a relatively safe procedure in clinically stable infants. Use of sedation was not associated with clinically relevant changes, although these findings warrant further investigation.

**O-163 DOES EARLY SCREENING LEAD TO HIGHER PREVALENCE OF PAEDIATRIC DELIRIUM?**

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10.1136/archdischild-2014-307384.230

**Introduction** Early screening of paediatric delirium (PD) allows for early intervention if necessary. The aim of this study was to determine if early screening with the SOS-PD scale led to higher prevalence of PD in ICU patients.

**Methods** A prospective before-after study design was applied in a population of children aged ≥3 months and admitted for ≥48 h to the PICU. In the before-period the prevalence of PD was estimated in terms of the number of children with PD confirmed by the consulting psychiatrist. During the after-period nurses systematically assessed the children with the SOS-PD scale three times a day in addition to the psychiatric consultation (SOS-PD score ≥4).

**Results** 148 and 150 children were included in the before and after period, respectively. The prevalence of PD was 6.1% and 8.7% for the before and after period respectively (see Table). The relative risk of PD with early screening was 1:43 (95% CI 0.63 to 3.23). In 33 patients (22%) the SOS-PD score was ≥4 on one or more occasions. In 14 of these patients, the child psychiatrist was consulted. In the remaining patients the child psychiatrist was not consulted for the following reasons: only once a high score (n = 9), adverse effects of sedatives (n = 4), and underlying disease/motor restlessness (n = 6).

**Conclusions** Systematic early screening of PD resulted in a higher incidence of PD and could contribute to timely start of treatment.

**Allergology**

**O-164 SPECIFIC IMMUNOGLOBULIN E TO ARA H 2 AS PREDICTOR FOR PEANUT ALLERGY IN CHILDREN IN A GENERAL DUTCH HOSPITAL**

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10.1136/archdischild-2014-307384.231

**Background** Specific immunoglobulin E (sIgE) to Ara h 2 is described as a potential factor for diagnosing peanut allergy in children. However for the Dutch children, limited data are available. In this study the diagnostic value of sIgE to Ara h 2 for children in a general non-university hospital is evaluated and compared with the existing data.

**Methods** Data from 137 peanut sensitised children were collected retrospectively. The primary outcome was peanut allergy or tolerance confirmed by food challenges. Different possible predictors, including sIgE to Ara h 2 (n = 52), were identified by multivariate backward stepwise logistic regression analysis. All significant predictors were combined in a formula for prediction of peanut allergy. Different essential cut-off points were obtained by an ROC curve.

**Results** Multivariate analysis resulted in sIgE to Ara h 2 as only predictor for peanut allergy, with a discriminative ability of 0.87 (95% CI, 0.77–0.97). Sensitivity and specificity values of respectively 55% and 95% were found at a sIgE to Ara h 2 cut-off value of 4.25 kU/L. Hundred percent specificity was reached at a cut-off point of 5.61 kU/L. The mean (SD) sIgE to Ara h 2 level for allergic children was 21.49 kU/L (SD 30.65) compared to 1.07 kU/L (1.56) for tolerant children (p = 0.001).

**Conclusions** Specific IgE to Ara h 2 is the best predictor for peanut allergy in sensitised children in a non-university hospital, comparable to previously published data. These results are a step forward to a generalisation to the Dutch children population.
Abstract O-165 Table 1

<table>
<thead>
<tr>
<th>Odds-ratio</th>
<th>Confidence interval (95%)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.19</td>
<td>1.11–9.56</td>
<td>0.03</td>
</tr>
<tr>
<td>1.85</td>
<td>1.03–3.34</td>
<td>0.04</td>
</tr>
<tr>
<td>1.73</td>
<td>1.01–2.94</td>
<td>0.04</td>
</tr>
<tr>
<td>1.92</td>
<td>1.04–3.53</td>
<td>0.04</td>
</tr>
<tr>
<td>1.73</td>
<td>1.01–2.94</td>
<td>0.04</td>
</tr>
<tr>
<td>1.62</td>
<td>0.96–2.71</td>
<td>0.07</td>
</tr>
</tbody>
</table>

Background and aims Recent studies reveal an important link between the intestinal immune system, microbiota, brain and behaviour. Previously we have shown that food allergy in male mice caused behavioural and neurochemical changes. This study aimed to investigate the effects of a dietary intervention with immunomodulatory short-chain galacto – and long-chain fructo-oligosaccharides (scGOS/ICcFOS), Bifidobacterium breve (Bb) and glutamine (Gln) on behavioural impairments in food allergic mice.

Methods Male C3H mice were fed a control, scGOS/ICcFOS/Bb, Gln, or scGOS/ICcFOS/Bb/Gln (comb) diet shortly after weaning and 2 weeks prior to first sensitisation with whey and cholera toxin (CT), or CT alone. Mice were sensitised for 5 weeks and subsequently orally challenged. Spontaneous alternation was examined in a T maze test 2 days after the last sensitisation and a social interaction test was conducted 1 day after oral challenge. Spontaneous alternation was used to measure exploratory behaviour and spatial memory.

Results Supplementation with scGOS/ICcFOS/Bb or Gln partially prevented reduced spontaneous alternation, whereas supplementation with scGOS/ICcFOS/Bb/Gln completely normalised alternation. Both scGOS/ICcFOS/Bb and Gln partially attenuated reduced social behaviour in food allergic mice. No additional effect of the combination was observed on social behaviour. Supplementation with scGOS/ICcFOS/Bb and/or Gln did not reduce allergic sensitisation, measured by whey-specific immunoglobulins.

Conclusions Supplementation with scGOS/ICcFOS/Bb or Gln partially prevented food allergy-induced behavioural impairments and the combination normalised impaired alternation, without changing allergic sensitisation. Therefore, it is of interest to further investigate the effects of dietary supplementation with scGOS/ICcFOS/Bb and Gln on immune-induced behavioural impairments in infants.

Abstract O-167 Figure 1

**O-166** WITHDRAWN

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**O-167a** INFLAMMATORY SUBTYPES IN WHEEZING INFANTS: ASSESSMENT AND IDENTIFICATION USING INDUCED SPUTUM

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Background Patterns of wheezing during early childhood may indicate differences in aetiology and prognosis of respiratory illnesses.

Objectives This study evaluated sputum cytology in infants with recurrent wheezing to classify sputum inflammatory phenotypes and assessed their characterisation over time.

Methods Sputum induction were performed in 890 infants with recurrent wheezing. Samples were classified as eosinophilic (>2.5% eosinophils), neutrophilic (>54% neutrophils), mixed granulocytic (>2.5% eosinophils, >54% neutrophils), or pauci-granulocytic (≤2.5% eosinophils, ≤54% neutrophils). Sputum induction were repeated after 3 months in infants with oral montelukast sodium (4 mg, QN) or nebulizer ICS (Budesonide aerosol 0.3 mg, Bid).

Results Total 304 infants (58.1%) had raised levels of inflammatory cells, eosinophilic 30.6%, neutrophilic 65.2%, mixed granulocytic 4.2%. Variabilities in sputum inflammatory phenotype were observed in both the severe and the mild to moderate wheezing groups. Changes in phenotype were not related to inhaled ICS or oral montelukast sodium, nor were it reflected in a change in tidal pulmonary function. About 27.3% infants fulfilled the criteria for eosinophilia and there were no differences in severity even atopy between non-eosinophilic and eosinophilic wheezing.

Conclusions Raised levels of inflammatory cells were frequently found in infants with recurrent wheezing. Sputum inflammatory