ENTRY AGE INTO DAY CARE AND LATER DEVELOPMENT OF ALLERGIC DISORDERS – RESULTS FROM THE CITY OF LEIPZIG COHORT OF THE LISA STUDY

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SUMMARY

The situation in early childhood is supposed to be a risk factor for later development of allergic diseases. The birth cohort from the LISA (Lifestyle – Immune System – Allergy) study gave us the opportunity to investigate the relationship between early childhood situation and the development of allergic diseases.

This paper describes our findings regarding to the relationship between entry age into day care and obstructive bronchitis as well as allergic rhinitis and atopic eczema.

Study was designed as a longitudinal birth cohort study. Children were examined by a physician at birth, ½ year, 1 year, 1 ½ year, 2 years, 3 years and 4 years. Further information was collected using a structured questionnaire which was answered by the parents.

Outcomes under investigation were atopic eczema, allergic rhinitis, wheezing (with and without cold), obstructive bronchitis and asthma (after 2nd year of life).

Logistic regression analysis adjusted for infectious diseases of the mother during pregnancy, vaccination of the mother during pregnancy, antibiotics, prenatal smoking of the mother or other persons in the apartment and vaccination state showed an significant promoting effect of entry age into day care against obstructive bronchitis over the first three years of life (OR ½ year: 8.55; 95%CI: 2.93…24.97; OR 1st year: 4.96; 95% CI: 1.73…14.24; OR 2nd year: 3.06 95% CI: 1…9.37). A further significant effect was found for crowding against asthma in the fourth year of life (OR 25.7; 95%CI: 1.65…400.17).

No significant effects were found between the other periods under investigation (1st year until 4th year of life).

On the base of our findings we recommend an entry age into day care of more than six months to prevent effects shown.

Key words: atopic eczema, allergic rhinitis, wheezing, asthma, entry age into day nursery, crowding, logistic regression model

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INTRODUCTION

Allergic diseases are common and appear to be increasing in prevalence over the past decades (1–4). Some authors report an increase in severity of these diseases (5, 6), whereas others report no further increase (7, 8).

Since Strachan (9) introduced the hygiene hypothesis in 1989, there were many attempts to verify this hypothesis. Illi et. al. (10) found a protective effect of running nose and other infectious diseases (excluding lower respiratory tract infections) against wheezing, asthma and bronchial hyperreactivity . Klimmert et. al. (11) showed a dependency between first year variables and school-age asthma support. Other publications investigated the relationship between entry age into day care, family size and later development of allergic diseases under the topic “hygiene hypothesis” (12–15). They concluded, that earlier exposure to allergens protects against atopy.

These reports guided us to the question: Can we reproduce similar effects using the data from the children in our LISA study?

The overall objective of the LISA (Lifestyle – Immune System – Allergy) study is to find reasons for atopic diseases. Atopy hereby refers to asthma, eczema, hay fever and other allergic disorders, which tends to accumulate within families (16).

In this paper we looked for a relationship between atopic situations (outcome variable) and the situation (independent variables) in early childhood. Independent variables under investigation were entry age into day care and crowding (person density in flats).

MATERIALS AND METHODS

Study Design

Investigations behind this paper are based on a subgroup (whose members were born and lived in the city of Leipzig,
Germany) of the LISA study. It consists of 976 newborn infants (494 girls and 482 boys) from the same year of birth (1998). They were recruited by the LISA team in the main hospitals with obstetrics department in the German city of Leipzig. Local ethics committees guided the study.

Only healthy, fully developed children with German origin were included into study cohort. The following exclusion criteria were applied (17):
- birth prior to 37th week of pregnancy,
- birth weight less than 2500 g,
- postnatal infection,
- dysplasia,
- mother with chronic disease or defect of the immune system,
- mother with persistent medication.

The cohort children were followed up at the age of 1/2, 1, 2, 3, 4, and 6 years within 3 months of the child’s birthday. The participation rate at the second birthday of the children was 86 %. All parents gave their consent before entering the study. Only 521 children had a complete longitudinal record. For all other children our data set missed one or more periods. Therefore the number of children differs in each period.

A questionnaire was sent to the parents together with an invitation for physical examination. The questionnaire consisted of the main topics “allergic heredity”, “residential characteristics”, “pregnancy”, “feeding”, “smoking/ETS”, “allergic symptoms (respiratory, nasal, skin, physician diagnosis)”, “pets”, “socioeconomic family situation”.

The physical examination of children took place close to their birthday in the respective year. This was done to exclude the effects of seasonal variations. During this examination the variables called “physicians diagnosis” were collected (17).

Definition of Variables

This section describes, how variables under investigation were defined. The description distinguishes between dependent variables (outcomes), independent variables and confounders.

Dependent variables

Atopic eczema:

According to the diagnostic criteria published by Hanifin and Rajka (18), a case was taken for as atopic eczema in the respective period, if there was skin rash with itch over more than 15 days at least on one of the following areas of the body:
- face, neck or nape;
- outer side of leg or arm;
- inner side of elbow or popliteus;
- hands or feet;
- at the trunk, not at the buttocks or
- other parts of the human body.

Allergic rhinitis:

Allergic rhinitis was supposed to be true if a child had discomfort of runny nose in conjunction with tearing eyes in the respective period. It was defined from the period 1 1/2 years onwards.

Wheezing (with/without cold):

If a child had wheezing over more than three days, we counted it as wheezing case in the respective period. We subdivided wheezing into wheezing with cold and wheezing without cold if there was/wasn’t cold reported in conjunction with wheezing.

Gastrointestinal infection:

We used helminth infection as an indicator for gastrointestinal infection.

Respiratory infection:

Respiratory infection was considered in the respective period if there was pneumonia, pseudo-croup, pertussis or sinusitis present.

Obstructive bronchitis:

In contrast to wheezing, obstructive bronchitis was relied upon physicians diagnosis, if a physician gave the ICD-10-code (19) J45.9.

Asthma:

Because of the controversial definition of asthma in different sources (16, 20-25), we distinguished between obstructive bronchitis, which embraces the widely accepted clinical effect: airway obstruction (26) and asthma.

For the widely used atopic conditions urticaria and food-incompatibility we did not have the required variables for a consistent definition over all periods. Therefore these outcomes were not included in our longitudinal analysis.

IgE:

IgE was measured using a Pharmacia ImmunoCAP-FEIA-System. IgE-values below the lower detection threshold of 0.35 kU/l were set to a value of 0.175 kU/l (27). Values above the upper detection threshold of 5,000 kU/l were set to 5,000 kU/l. We found a 90% percentile of 121 kU/l and a median of 19.3 kU/l at an age of two years in our cohort. Liappis et. al. found a reference value (90% percentile) for two years old children of up to 16 kU/l with a median of 6.62 kU/l (28). Because of the fact that Liappis et al. selected only children with absence of atopic heredity the values are not directly comparable. For better comparison we also did a selection for children with no atopic heredity and found 18.8 kU/l and a 90% percentile of 111 kU/l. This difference is possibly influenced by a further selection criterion in the article of Liappis et al. which excluded children with increased IgE-level (which was not defined in detail) from their analysis.

Specific IgE:

We used specific IgE measurements as a further reference for allergic situation in our cohort. If there were the RAST Classes for HX2 or E1 or MX1 or FX5 or RX1 above zero then we concluded that the child was sensitized against a specific allergen and therefore we counted it in the respective variable.

Independent variables

We investigated entry age into day care and crowding as independent variables.

Entry age into day care:

We investigated the effect of entry age into day care on atopic situations because of the findings of Krämer et al.(14). They concluded that age of entry into day care affects the development of bronchitis.

Entry age into day care was subdivided into six classes: less than 6 months, 6 to 12 months, 12 to 18 months, 18 to 24 months, 24 to 36 months and more than 36 months.

Crowding:

Unlike Krämer et al., who defined crowding as more than three people living together in an apartment, we defined crowding as a binary variable derived from the family size.

Because of absence of a variable with information about the number of children in family, crowding was constructed using the
variables “number of persons in household” and “marital status” respectively “living together with partner” in the following way: If “single mother/father” was true then crowding was counted if more than 5 persons live together in one family (single mother/father + 4 children). In the case that both parents are living together in the family, we counted crowding with 6 and more persons in the family (2 parents + 4 children).

Confounders
The following variables were taken into consideration as possible confounders:
- “allergic heredity” (29),
- “infectious diseases of the mother during pregnancy”,
- “vaccination of the mother during pregnancy”,
- “ingestion of antibiotics”,
- “prenatal smoking of the mother or other persons in the apartment”,
- “renovation of the apartment” and
- “vaccination state of the child”.

All models in a longitudinal series were calculated with the whole set of confounders for comparability.

Analysis
Before analysis took place, pre-processing was carried out. This includes selecting, concatenating and recoding of original variable values into a uniform coding scheme for this analysis. This pre-processing was necessary because analysis was performed using data from different data sources (periods) with different coding schemes.

Each original variable was described with its data type, scale level and coding scheme. Based upon this meta data we are able to analyze the specification of each variable for concatenation and recode them to a common coding scheme. Furthermore constraints such as missing value codes or values outside specification have to be considered.

Odds ratios (OR) with 95% confidence intervals were calculated to assess the risk estimates for allergic diseases. We performed stepwise logistic regression for each period in our longitudinal analysis with adjustment for the variables described in “Confounders”. All results in this paper are based on a data set consisting of 976 children’s data from birth to 4th year of life. In the case of missing data we deleted the entire case for the analysis in the respective period. If the significance level of the model (p) was 0.05 or less, then the model was judged to be significant.

All models were calculated with the cases which did not fulfil the criterion for the outcome variable as reference group. Therefore the results for a specified situation (outcome=true) have to be interpreted as referring to the cases which did not have this outcome (outcome=false).

Statistical analysis was performed using Statistica release 6.1 from Statsoft Inc (Tulsa, OK, USA) (30).

We performed a longitudinal analysis for each outcome variable using each independent variable and using the described confounders. Longitudinal analysis was done using logistic regression models (31).

RESULTS

Descriptive Statistics
Table 1 shows the frequencies of outcome variables subdivided into overall frequency, frequency reported by parents and frequency reported by physician.

We observed a relatively constant behaviour of wheezing, a slight decrease in allergic rhinitis and in atopic eczema over the first two years of life. The prevalence of atopic eczema, obstructive bronchitis and asthma is increasing in the fourth year of investigation.

Respiratory infections were very common during the observed period, whereas gastrointestinal infections were very rare.

We observed a large number of children that first entered day care before their 2nd year of life (42.1%). Unfortunately we do not have exact information about the distribution of entry age into day care in the population. But we know that there is a difference between the eastern and western part of Germany. Early entry into day care is very common in the city of Leipzig as well as in the entire area of former East Germany. This may have historical reasons because in former East Germany working women were common practice. Therefore many day care centres in eastern part of Germany are able to assist babies and children with diapers.

To control unknown bias we analyzed known risk factors over all periods. Atopic heredity was counted if both parents had atopic symptoms in the past. With the variable smoking we controlled the bias of smoking during pregnancy (32). The variable “educational level of parents” was calculated for controlling the effect of different educational levels within the families (33). It refered to the achieved level of education in conjunction with the achieved level of professional training for both parents using the scheme in Table 2 (34).

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Age (years)</th>
<th>0.5</th>
<th>1</th>
<th>1.5</th>
<th>2</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n=849</td>
<td>n=820</td>
<td>n=816</td>
<td>n=794</td>
<td>n=670</td>
<td></td>
</tr>
<tr>
<td>Atopic eczema</td>
<td>84 (9.9%)</td>
<td>104 (12.7)</td>
<td>114 (14)</td>
<td>144 (18.1)</td>
<td>93 (13.9)</td>
<td></td>
</tr>
<tr>
<td>Allergic rhinitis</td>
<td>N/A*</td>
<td>N/A</td>
<td>22 (2.7)</td>
<td>19 (2.4)</td>
<td>15 (2.2)</td>
<td></td>
</tr>
<tr>
<td>Wheezing without cold</td>
<td>15 (1.77)</td>
<td>7 (0.8)</td>
<td>1 (0.1)</td>
<td>10 (1.3)</td>
<td>11 (1.6)</td>
<td></td>
</tr>
<tr>
<td>Wheezing with cold</td>
<td>60 (7.1)</td>
<td>85 (10.4)</td>
<td>90 (11)</td>
<td>85 (10.7)</td>
<td>38 (5.7)</td>
<td></td>
</tr>
<tr>
<td>Gastrointestinal infection</td>
<td>N/A</td>
<td>0 (0)</td>
<td>4 (0.4)</td>
<td>3 (0.3)</td>
<td>16 (1.6)</td>
<td></td>
</tr>
<tr>
<td>Respiratory infection</td>
<td>N/A</td>
<td>573 (69.9)</td>
<td>648 (79.4)</td>
<td>640 (80.6)</td>
<td>561 (57.5)</td>
<td></td>
</tr>
<tr>
<td>Obstructive bronchitis</td>
<td>N/A</td>
<td>47 (5.8)</td>
<td>41 (5)</td>
<td>50 (6.3)</td>
<td>67 (10)</td>
<td></td>
</tr>
<tr>
<td>Asthma</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>1 (0.1)</td>
<td>7 (1)</td>
<td></td>
</tr>
</tbody>
</table>

*N/A: not available; These values represent, that we did not have any child with this diagnosis.
All values of 5 and above for at least one parent were counted as a high educational level within the family (confounder value=0). All other values were counted as low educational level (confounder value=1, which means additional risk).

Prevalences for confounding variables are shown in Table 3. We also computed a Mann-Whitney U-test for all periods, which had shown no significant differences between the risks in the respective periods.

We screened all periods which were covered by data from birth to 4th year of life for significant models.

Infections

We used obstructive bronchitis as sign for infection in the lower respiratory tract. The adjusted OR for obstructive bronchitis had shown a significant decrease over the following periods for children with entry age into day care 6 months or less (Table 4).

We attribute these effects to the fact that the immune system of these young children is not able to defend against the enormous invasion of pathogens. Older children do not show this effect.

Stratification by sex discovered a difference between both sexes: boys showed the described relationship whereas in girls we did not come across models with this behaviour.

**Allergic Disorders**

Entry age into day care was used for describing the contact with other children and the associated influence onto the immune system.

We found significant effects for obstructive bronchitis and allergic rhinitis in relationship to the entry age into day nursery. The risk for obstructive bronchitis was significantly increased

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**Table 2. Conversion scheme for the variable “educational level”**

<table>
<thead>
<tr>
<th>Level of professional training</th>
<th>without graduation</th>
<th>secondary school</th>
<th>O-level</th>
<th>advanced technical college entrance qualification</th>
<th>university-entrance diploma / A-level</th>
</tr>
</thead>
<tbody>
<tr>
<td>without graduation</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>vocational school</td>
<td>3</td>
<td>3</td>
<td>4</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>technical school</td>
<td>4</td>
<td>4</td>
<td>5</td>
<td>6</td>
<td>6</td>
</tr>
<tr>
<td>university of applied sciences</td>
<td>N/A</td>
<td>7</td>
<td>7</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>university</td>
<td>N/A</td>
<td>8</td>
<td>8</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>other</td>
<td>3</td>
<td>3</td>
<td>4</td>
<td>5</td>
<td>7</td>
</tr>
</tbody>
</table>

**Table 3. Period-prevalences of risks (values are absolute numbers; percentages in parentheses) in the LISA study Leipzig, Germany 1998-2002**

<table>
<thead>
<tr>
<th>Risk Factor</th>
<th>Age (years)</th>
<th>0.5 n=849</th>
<th>1 n=820</th>
<th>1.5 n=816</th>
<th>2 n=794</th>
<th>4 n=670</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atopic heredity</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Smoking</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Educational level of parents</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Renovation of apartment</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Entry age</td>
<td>Frequency</td>
<td>47</td>
<td>47</td>
<td>41</td>
<td>50</td>
<td>67</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(2.93..24.97)</td>
<td>(1.73..14.24)</td>
<td>(1.73..14.24)</td>
<td>(1.73..14.24)</td>
<td>(1.73..14.24)</td>
</tr>
<tr>
<td>6 months</td>
<td>44</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>12 months</td>
<td>84</td>
<td>N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>18 months</td>
<td>283</td>
<td>N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>24 months</td>
<td>4</td>
<td>N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>36 months</td>
<td>178</td>
<td>N/A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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**Table 4. Odds ratios (95% confidence intervals in parentheses) with significance levels of the independent variable (p) of obstructive bronchitis depending on entry age into day care (cases which did not fulfil the criterion for the outcome variable served as reference group for OR) in the LISA study Leipzig, Germany 1998-2002; models with statistically significant result are marked by italic font**

<table>
<thead>
<tr>
<th>Entry age</th>
<th>Frequency</th>
<th>Outcome obstructive bronchitis</th>
<th>Outcome obstructive bronchitis</th>
<th>Outcome obstructive bronchitis</th>
<th>Outcome obstructive bronchitis</th>
<th>Outcome obstructive bronchitis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>47</td>
<td>47</td>
<td>41</td>
<td>50</td>
<td>67</td>
<td></td>
</tr>
<tr>
<td>6 months</td>
<td>44</td>
<td>8.55</td>
<td>4.96</td>
<td>2.55</td>
<td>3.06</td>
<td>0.78</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(2.93..24.97)</td>
<td>(1.73..14.24)</td>
<td>(1.73..14.24)</td>
<td>(1.73..14.24)</td>
<td>(1.73..14.24)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>p&lt;0.01</td>
<td>p&lt;0.01</td>
<td>p&lt;0.01</td>
<td>p&lt;0.01</td>
<td>p&lt;0.01</td>
</tr>
<tr>
<td>12 months</td>
<td>84</td>
<td>1.09</td>
<td>1.43</td>
<td>1.43</td>
<td>1.43</td>
<td>1.43</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(0.52..3.72)</td>
<td>(0.46..4.49)</td>
<td>(0.46..4.49)</td>
<td>(0.46..4.49)</td>
<td>(0.46..4.49)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>p=0.89</td>
<td>p=0.33</td>
<td>p=0.33</td>
<td>p=0.33</td>
<td>p=0.33</td>
</tr>
<tr>
<td>18 months</td>
<td>283</td>
<td>1.07</td>
<td>0.75</td>
<td>1.26</td>
<td>1.26</td>
<td>1.26</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(0.37..3.02)</td>
<td>(0.35..1.61)</td>
<td>(0.35..1.61)</td>
<td>(0.35..1.61)</td>
<td>(0.35..1.61)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>p=0.74</td>
<td>p=0.46</td>
<td>p=0.48</td>
<td>p=0.48</td>
<td>p=0.48</td>
</tr>
<tr>
<td>24 months</td>
<td>4</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>36 months</td>
<td>178</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(0.54..2.07)</td>
<td>(0.54..2.07)</td>
<td>(0.54..2.07)</td>
<td>(0.54..2.07)</td>
<td>(0.54..2.07)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>p&lt;0.01</td>
<td>p&lt;0.01</td>
<td>p&lt;0.01</td>
<td>p&lt;0.01</td>
<td>p&lt;0.01</td>
</tr>
</tbody>
</table>
with higher absolute values, whereas in girls we did not find the described effects.

To prevent any misleading and to have real measures instead of answers to questions, we observed the behaviour of specific IgE in the 2nd year of life. Specific IgE level was constructed from the RAST classes of HX2 (house dust mite), E1 (cat dander), MX1 (mould), FX5 (food allergens) and RX1 (pollen). If one of these specific IgE had a RAST class more than zero the variable specific “IgE level raised” was supposed to be true. We found a significant model for entry age into day care between 12th and 18th month which was associated with 1.93-fold (95% CI: 0.99...3.77) risk for raised level of specific IgE in the 2nd year of life (see Table 7).

**Crowding**

We investigated all periods from ½ year of life until 4th year of life for a significant effect between atopic situations and crowding (number of children in an apartment). Crowding, which was found in 29 cases, showed a significant effect on the outcome asthma in the 4th year of life: OR was 25.7 (p=0.02) with an 95% CI from 1.65 ... 400.17 (adjusted for infectious diseases during pregnancy, smoking during pregnancy, ingestion of antibiotics during the child, vaccination state of the child, educational index of the parents, atopic heredity and renovation during pregnancy). The large confidence interval results from a relatively low number of asthma cases (only 7) in the 4th year of life. Because of the low case numbers and the large confidence interval, these findings are not suitable for recommendations.

**DISCUSSION AND RESULTS**

Entry age into day care was the exposure variable which yielded most significant models. Bias in this variable is unlikely because the information was collected directly in the questionnaire. This variable was collected in every relevant period. The results of this study indicate that early entry into day care (up to 6th month of life) is positively associated with an increased risk of obstructive bronchitis in later childhood. For an entry

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**Table 5. Odds ratios (95% confidence intervals in parentheses) with significance levels of the independent variable (p) of allergic rhinitis depending on entry age into day care (cases which did not fulfil the criterion for the outcome variable served as reference group for OR) in the LISA study Leipzig, Germany 1998-2002; models with statistically significant result are marked by italic font**

<table>
<thead>
<tr>
<th>Entry age</th>
<th>Frequency</th>
<th>Outcome allergic rhinitis 1.5 years</th>
<th>Outcome allergic rhinitis 2 years</th>
<th>Outcome allergic rhinitis 4 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>6 months</td>
<td>44</td>
<td>N/A</td>
<td>0.95 (0.11...8.14)</td>
<td>p=0.96</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p(M)=0.58</td>
<td>N/A</td>
</tr>
<tr>
<td>12 months</td>
<td>84</td>
<td>N/A</td>
<td>4.73 (1.28...17.5)</td>
<td>2.07 (0.4...10.64)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p=0.02</td>
<td>p=0.38</td>
</tr>
<tr>
<td>18 months</td>
<td>283</td>
<td>N/A</td>
<td>1.6 (0.5...5.12)</td>
<td>1.14 (0.34...3.05)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p=0.43</td>
<td>p=0.83</td>
</tr>
<tr>
<td>24 months</td>
<td>4</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>36 months</td>
<td>178</td>
<td>N/A</td>
<td>0.94 (0.26...3.39)</td>
<td>p=0.92</td>
</tr>
</tbody>
</table>

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**Table 6. Odds ratios (95% confidence intervals in parentheses) with significance levels of the independent variable (p) of atopic eczema depending on entry age into day care cases which did not fulfil the criterion for the outcome variable served as reference group for OR) in the LISA study Leipzig, Germany 1998-2002**

<table>
<thead>
<tr>
<th>Entry age</th>
<th>Frequency</th>
<th>Outcome atopic eczema ½ year</th>
<th>Outcome atopic eczema 1 year</th>
<th>Outcome atopic eczema 1.5 years</th>
<th>Outcome atopic eczema 2 years</th>
<th>Outcome atopic eczema 4 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>6 months</td>
<td>44</td>
<td>0.77 (0.21...2.82)</td>
<td>0.88 (0.28...2.72)</td>
<td>1.15 (0.4...3.28)</td>
<td>p=0.79</td>
<td>1.16 (0.44...3.09) p=0.76</td>
</tr>
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<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>12 months</td>
<td>84</td>
<td>N/A</td>
<td>1.04 (0.43...2.5)</td>
<td>1.15 (0.47...2.78)</td>
<td>p=0.93</td>
<td>1.36 (0.63...2.97) p=0.43</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>18 months</td>
<td>283</td>
<td>N/A</td>
<td>N/A</td>
<td>1.51 (0.87...2.64)</td>
<td>p=0.15</td>
<td>0.98 (0.58...1.63) p=0.93</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>24 months</td>
<td>4</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>36 months</td>
<td>178</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>0.91 (0.53...1.56) p=0.72</td>
</tr>
</tbody>
</table>

---

in the 1st and 2nd year of life if the children started their daily stay in day care until 6th month of life (see Table 4). The risk for allergic rhinitis was significant increased from in the 2nd year of life if the children started their daily stay in day care until 12th month of life (Table 5).

The pattern of the relationship between atopic eczema (Table 6) and entry age into day care is nonlinear. In the first year of life we see a protective effect of day care. For later periods we see an increase of the OR in this relationship with its top in the 2nd year of life. This leads to the assumption, that there is a critical timeframe around the 1st birthday.

Stratification by sex discovered a difference between boys and girls. Boys showed behaviour similar to that of the whole cohort...
Van Schayck et al. (36) analyzed three randomized clinical trials (37–39) and concluded that these trials have shown a relationship between allergen exposure reduction and a reduction in the development of allergy. This stands in contrast to the hygiene hypothesis, just like our findings concerning the relationship between crowding (more than 4 children in a family) on the development of asthma in the 4th year of life.

Svanes et al. (40) found a promoting effect of day care before age of 5 years against asthma in juvenile age. This result is not directly comparable with ours, because the observed range of age is different. In the context with the other findings of Svanes et al. that childhood exposure to other children in day care has a preventive effect against hay fever one can speculate, that there are possibly genetic factors which influence the development of allergic disorders. We suppose that these factors could affect the results of different studies in various ways so that conclusions of the authors of these papers differ.

This positive related effect of crowding on the development of atopic disorders (i.e. asthma) agrees with the results found by other studies regarding crowding and atopic situations (41–44). The findings of Golding et al. (45), von Mutius et al. (46) and (47) are in contradiction: they found a protective effect of crowding for asthma.

The main difference between the studies which found a protective effect and those who found a promoting effect is the geographical region and related with this the economic development status of the country. The cohorts of Golding et al. (UK), Mutius et al. (Germany) and Brabæk (Baltic region) live in Europe, whereas the probationers of Victora (Brazil), Cerqueiro (Argentina), Ballard (Kenya), Cardoso (Brazil) live in South America and Africa.

Obviously, the region and the status of economic development of a country plays a role in whether crowding has a promoting or protecting effect against atopic situations. This implicates that there must be hidden factors behind crowding.

Because of our relatively low number of cases we did not provide clear evidence in this area. Further investigations of this issue may be interesting. On the basis of our findings we recommend an entry age into day care to be more than six months to prevent effects shown.

ACKNOWLEDGEMENT

We thank all families for their participation in the LISA Study. The study was supported by grants of the Federal Ministry for Education, Science, Research and Technology (no. 01 EG 9705/2 and 01 EG 9732).

APPENDIX

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Received September 1, 2005
Received in revised form and accepted January 30, 2006