Predicting asthma in preschool children at high risk presenting in primary care: development of a clinical asthma prediction score

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Abstract
Background: A setting-specific asthma prediction score for preschool children with wheezing and/or dyspnoea presenting in primary healthcare is needed since existing indices are mainly based on general populations.

Aims: To find an optimally informative yet practical set of predictors for the prediction of asthma in preschool children at high risk who present in primary healthcare.

Methods: A total of 771 Dutch preschool children at high risk of asthma were followed prospectively until the age of six years. Data on asthma symptoms and environmental conditions were obtained using validated questionnaires and specific IgE was measured. At the age of six years the presence of asthma was assessed based on asthma symptoms, medication, and bronchial hyper-responsiveness. A clinical asthma prediction score (CAPS) was developed using bootstrapped multivariable regression methods.

Results: In all, 438 children (56.8%) completed the study; the asthma prevalence at six years was 42.7%. Five parameters optimally predicted asthma: age, family history of asthma or allergy, wheezing-induced sleep disturbances, wheezing in the absence of common colds, and specific IgE. CAPS scores range from 0 to 11 points; scores <3 signified a negative predictive value of 78.4% while scores of >7 signified a positive predictive value of 74.3%.

Conclusions: We have developed an easy-to-use CAPS for preschool children with symptoms suggesting asthma who present in primary healthcare. After suitable validation, the CAPS may assist in guiding shared decision-making to tailor the need for medical or non-medical interventions. External validation of the CAPS is needed.

Keywords preschool children, asthma, prediction, wheeze, clinical studies, sensitivity and specificity, diagnosis

Introduction
Childhood asthma usually starts during the preschool years. Unfortunately, it is currently not possible to test for or alter the natural history of asthma accurately in young children, so asthma during the preschool years is frequently underdiagnosed. About 10 asthma risk prediction indices for young children have been developed, although only one was developed in a primary care setting where most of these children present with respiratory symptoms for the first time and where most are treated. Risk stratification using prediction indices may be useful for clinical and research purposes. For example, risk stratification supports the tailoring of treatment strategies such that benefits and harms may be properly weighed. It may therefore improve the follow-up of...
Children at risk by decreasing the disease burden and improving management, resulting in fewer respiratory symptoms, exacerbations, and emergency medical visits.\(^5\)\(^6\)

Prediction indices are valuable because the information itself may contribute to patients’ quality of life.\(^4\) Patients’ emotional response to medical testing is a natural effect. The information involved in a high predicted probability of a young child having asthma at six years of age may enable them (or their parent/guardians) to make sense of the symptoms or to adjust their lifestyle and living conditions in light of the results, while a low predicted probability may alleviate their concerns.\(^6\)\(^7\)

In addition, various research groups are investigating the underlying pathogenic mechanisms and preventive strategies for asthma,\(^4\) and as soon as effective therapy that is either costly and/or has adverse effects emerges, risk stratification will be useful.\(^3\)

Existing predictive indices have an uncertain value for the primary healthcare setting because the majority of the cohort studies were based on parental questionnaires in a general population,\(^6\)\(^8\)\(^9\) so they are of modest clinical value for preschool children.\(^10\) For example, the most well-known indices – the loose and stringent Asthma Predictive Indices (API) – were developed in a general population and are based on questionnaires.\(^9\) API predicts persistent wheezing in children who ever wheezed and does not predict asthma in children with recurrent wheeze presenting to a physician.\(^11\)

In 2004 we started a prospective cohort of young children at high risk of developing asthma. The objective was to construct a clinical index for predicting asthma in preschool children at risk of asthma presenting in primary care. Here we describe the development and performance of a novel clinical asthma prediction score (CAPS).

**Methods**

**Design and subjects**

The subjects were children aged one to five years in the Netherlands who were participating in the ARCADE (AiRway Complaints and Asthma Development) prospective cohort study.\(^4\) They had presented at one of the pertinent 14 primary care clinics in the previous 12 months with complaints of recurrent coughing (>2 visits), wheezing (>1 visit), and/or shortness of breath (>1 visit). The study was approved by the Netherlands Central Committee on Research Involving Human Subjects (CCMO/P04.0098C) and the details of the study have been published elsewhere.\(^4\) Written informed consent was obtained from the parents before all measurements.

**Data collection**

**Questionnaires**

For this study we derived demographic data and data with respect to the child’s clinical history from questionnaires obtained by the Core Questionnaire of the International Study of Asthma and Allergy in Children (ISAAC), which was administered at baseline and at the age of six years.

**Allergy test**

Total immunoglobulin E (IgE) and specific IgE directed against cat, dog, and house dust mite were determined by radioallergosorbent test (RAST) in all the children at baseline. IgE positivity to cat, dog, and house dust mite was defined as >0.35kU/L.\(^4\)

**Spirometry and bronchial hyper-responsiveness**

At six years of age, spirometry and bronchial hyper-responsiveness (BHR) assessments were obtained in children with wheezing, shortness of breath, recurrent coughing, or use of asthma medication (\(\beta_2\)-agonists, inhaled corticosteroids) during the previous 12 months. BHR to increasing doses of methacholine was measured according to Cockcroft et al. (online supplement reference 3).

**Outcome: asthma diagnosis at age six**

The diagnosis of asthma at six years of age was defined as: having persistent symptoms of asthma and/or using asthma medication in the last year in combination with BHR (defined as the methacholine dose (>8.0mg/ml) causing a 20% reduction in forced expiratory volume in one second (PC\(_{20}\)-FEV\(_1\))) or reversibility (>10% increase in FEV\(_1\) after short-acting \(\beta_2\)-agonists).\(^4\) Persistent symptoms were defined as self-reported complaints during the previous 12 months of wheezing and/or shortness of breath and/or recurrent coughing.

Children who had not experienced asthma-related symptoms in the previous year or who had not used asthma medication in the previous year were not invited for lung function measurement. They were considered as not having asthma.

**Probability of asthma at the age of six**

Depending on the direction and magnitude of the selected predictors as measured by regression coefficients, we developed the Clinical Asthma Prediction Score (CAPS) as a points system which is easy to use, without requiring a computer or calculator.\(^16\) To use the CAPS in clinical practice, thresholds are needed to either rule in or rule out asthma at the age of six years. Therefore we set two thresholds – a lower limit at 30% below which we assumed that many physicians may well choose a ‘wait and see’ policy, and an upper limit at 60% above which it was assumed that physicians may pursue a more active management such as a trial of asthma treatment. These thresholds imply that children with a predicted risk of asthma risk between 30% and 60% will be assigned to the area of clinical indecision in which a ‘watchful waiting’ policy may be defensible.

**Statistical analysis**

Details of the statistical analysis are described in Appendix 1, available online at www.thepcrj.org. Briefly, we analysed whether dropout was selective and missing predictor data were imputed using multiple imputation.\(^17\) Twelve candidate predictors of all available data were preselected based on clinical expert opinion after removing five due to collinearity (Table 1). Bootstrapped (1000x) backward selection was used to select the final important predictors whose regression coefficients were shrunk to facilitate better performance when the model is applied to other data or future patients.\(^18\) The predictive performance of the prediction index was assessed by its calibration and discrimination abilities.

Furthermore, the additional value of specific IgE was calculated by the additional area under the ROC curve (AUC\(_{\text{adj}}\)) and the net reclassification improvement (NRI).\(^19\) All statistical analyses were carried out in StataSE Version 10.1 (Stata Corp, College Station, TX, USA).
Table 1. Candidate predictors of asthma diagnosed at 6 years of age in preschool children

| Candidate variables | Asthma diagnosis at age 6 years |
|---------------------|--------------------------------|
|                     | No (n=251) | Yes (n=187) | All (n=438) |
| Mean (SD) age at baseline, years | 2.71 (1.23) | 3.39 (1.26) | 3.00 (1.29) |
| 1–2                 | 98 (39)    | 34 (18)     | 132 (30)    |
| 2–3                 | 63 (25)    | 35 (19)     | 98 (22)     |
| 3–4                 | 37 (15)    | 47 (25)     | 84 (19)     |
| 4–5                 | 36 (14)    | 54 (29)     | 90 (21)     |
| ≥5                  | 17 (7)     | 17 (9)      | 34 (8)      |
| Perinatal factors   |            |             |             |
| Female              | 110 (44)   | 79 (42)     | 189 (43)    |
| Breastfeeding >3 months (2 missing) | 107 (43) | 86 (46) | 193 (44) |
| Reported symptoms in past year |            |             |             |
| Nightly persistent mucous cough or phlegm apart from a cold* (5 missing): yes | 78 (31) | 85 (46) | 163 (38) |
| Dry cough at night, apart from a cold* (6 missing): yes | 112 (45) | 109 (59) | 221 (51) |
| Coughing episode ≥ 1 month** (4 missing): yes | 31 (12) | 41 (22) | 72 (17) |
| Wheezing*:yes      | 133 (53)   | 126 (67)    | 259 (59)    |
| Frequency of wheezing attacks*: (1 missing) |            |             |             |
| None                | 117 (47)   | 61 (33)     | 178 (41)    |
| 1–3 times           | 103 (41)   | 78 (42)     | 181 (41)    |
| >4 times            | 30 (12)    | 48 (25)     | 78 (18)     |
| Wheezing-induced sleep disturbances*: (3 missing): yes | 61 (25) | 82 (44) | 143 (33) |
| Wheezing during or in absence of cold*: (3 missing): |            |             |             |
| No                  | 121 (49)   | 62 (33)     | 182 (42)    |
| Yes, only with cold | 100 (40)   | 80 (43)     | 180 (41)    |
| Yes, with and without cold | 28 (11) | 44 (24) | 72 (17) |
| Itch/eczema*: (3 missing) | 78 (31) | 74 (40) | 152 (35) |
| Allergy child       |            |             |             |
| Allergy (ever)** (2 missing) (doctor's diagnosis) | 31 (12) | 45 (24) | 76 (17) |
| Allergy child – IgE dichotomous |            |             |             |
| Specific IgE against cats, dogs, and/or house dust mite* (96 missing) | 23 (12) | 65 (41) | 88 (26) |
| Environmental factors |            |             |             |
| Parental smoking in child's presence* (3 missing) | 33 (13) | 30 (16) | 63 (14) |
| Child has older sibling | 132 (53) | 96 (51) | 228 (52) |
| Family history      |            |             |             |
| Asthma, parents, and/or siblings* (3 missing) | 76 (31) | 80 (43) | 156 (36) |
| Allergy, parents, and/or siblings* (3 missing) | 98 (28) | 93 (33) | 191 (31) |

*Variables not used in bootstrapped backward selection. Numbers are n (percentages in parentheses) unless stated otherwise. IgE = immunoglobulin E.
1. In the last 12 months, has your child usually seemed congested in the chest or has she/he coughed up phlegm (mucus) during the night when she/he did not have a cold?
2. In the last 12 months, has your child had a dry cough at night, apart from a cough associated with a cold or chest infection?
3. In the last 12 months, has your child coughed for more than 5 days a week over a period longer than 1 month?
4. Has your child had attacks of wheezing in the last 12 months?
5. How many attacks of wheezing has your child had in the last 12 months?
6. In the last 12 months, has your child's sleep been disturbed due to wheezing?
7. Combination of questions (1) In the last 12 months, has your child's chest sounded wheezy associated with a cold? and (2) In the last 12 months, has your child's chest sounded wheezy not associated with a cold?
8. Has your child had an itchy rash which was coming and going for at least six months in the last 12 months?
9. Has a doctor ever diagnosed one or more allergies?
10. Specific IgE positivity to house dust mite, cat, and/or dog dander was defined as ≥0.35kU/L.
11. Does at least one of the parents smoke in the parental house?
12. Does at least one of the parents and/or siblings have asthma? Variable was collected by questionnaire. In the analyses it was combined with ‘Allergy, parents, and/or siblings’.
13. Does at least one of the parents and/or siblings have allergies to pets, pollen, and/or house dust mites? Variable was collected by questionnaire. In the analyses it was combined with ‘Allergy, parents, and/or siblings’.
**Results**

Of the 771 children, 438 (56.8%) had complete information at baseline and an asthma diagnosis at the endpoint (Figure 1). Of these, 187 (42.7%) had asthma at the age of six. Table 1 shows the univariable frequencies of the preselected candidate predictors. The mean (SD) age at baseline was 3.00 (1.29) years and 189 (43.1%) were female. In 156 children (35.6%), asthma was present in either the parents or siblings. Missingness analyses showed no strong indications for selective drop out (Appendix 1 available online at www.pcrj.org).

**Multivariable analysis**

Five predictors were included (Table 2): age, family history of asthma or allergy, wheezing-induced sleep disturbances in the previous 12 months, wheezing in the absence of a cold in the previous 12 months, and specific IgE test. The discrimination of the model expressed as area under the ROC curve was 0.73 (SE=0.024). Calibration was satisfactory (Table 2).

**IgE and no IgE in the model**

The model performance was calculated for the model including age, family history, wheezing-induced sleep disturbances, and wheezing in the absence of a cold but excluding IgE. The discrimination of the model expressed as area under the ROC curve was 0.71 (SE=0.024). Calibration was good (Table 2). Adding specific IgE to the model increased the AUC by 2.8 percentage points (AUCdiff= 0.028, 95% CI 0.0047 to 0.051, SE=0.012). Net reclassification analysis showed that 11.23% (p=0.09) of the children were correctly reclassified when IgE was added to the model based on history items only (Table 3).†

**Probability of asthma at the age of six using CAPS**

A points system ranging from 0 to 11 was developed. Thresholds to

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*Figure 1. Flow diagram*
The clinical asthma prediction score (CAPS) was developed in a prospective cohort study to facilitate the risk stratification of preschool children who attend primary healthcare with wheezing, dyspnoea, or recurrent coughing. The predictive ability of the CAPS may assist physicians to distinguish between a relatively low risk (14%) and a very high risk (90%) of developing asthma. The CAPS uses easily accessible information on age, family history of asthma or allergy, wheezing-induced sleep disturbances, and wheezing in the absence of cold.

### Table 2. Predictors for asthma at age of six years and prediction model performance measures

| Predictors                                      | OR*       | p Value | 95% CI            |
|-------------------------------------------------|-----------|---------|-------------------|
| Age (year): linear                              | 1.47      | 0.000   | 1.24 to 1.75      |
| Specific IgE positive: yes or no                | 2.45      | 0.003   | 1.38 to 4.33      |
| Asthma or allergy, parents and/or siblings      | 1.54      | 0.060   | 0.98 to 2.41      |
| Wheezing-induced sleep disturbances             | 2.08      | 0.002   | 1.31 to 3.29      |
| Wheezing, in absence of cold                    | 2.22      | 0.008   | 1.23 to 3.99      |

**Discrimination, calibration, reclassification**

AUC (SE)  | HL (p) | NRI (SE)**
---|---|---
Extended model (including all predictors) | 0.73 (0.024) | 4.51 (>0.75) |
Simple model (excluding IgE) | 0.71 (0.024) | 7.15 (>0.61) |
Comparisons between the models AUCdiff | 0.028 (0.012)# | 11.23 (0.05)#

*OR = odds ratio before fixed shrinkage (shrinkage factor k=0.949), NRI=net reclassification improvement. **Thresholds are 30% and 60%. # p=0.019; ##p=0.09.

### Table 3. Reclassification tables for model with or without IgE in children with and without asthma

| Risk factor | Categories | CAPS | OR*       | p Value | 95% CI            |
|-------------|------------|------|-----------|---------|-------------------|
| Age         | 1-2        | 0    |           |         |                   |
|             | 2-3        | 1    |           |         |                   |
|             | 3-4        | 2    |           |         |                   |
|             | 4-5        | 3    |           |         |                   |
|             | >5         | 4    |           |         |                   |
| sIgE        | Negative   | 0    |           |         |                   |
|             | Positive    | 2    |           |         |                   |
| Asthma or allergy, parents and/or siblings    | Negative   | 0    |           |         |                   |
|             | Positive    | 1    |           |         |                   |
| Wheezing-induced sleep disturbances           | No         | 0    |           |         |                   |
|             | Yes         | 2    |           |         |                   |
| Wheezing in absence of cold                   | No         | 0    |           |         |                   |
|             | Yes         | 2    |           |         |                   |
| Score (CAPS) |            | 0-11 |           |         |                   |

95% CI=95% confidence interval.

### Table 4. Risk estimates associated with Clinical Asthma Prediction Score (CAPS) points

| Risk factor                                      | Categories | CAPS | OR*       | p Value | 95% CI            |
|--------------------------------------------------|------------|------|-----------|---------|-------------------|
| Age                                              | 1-2        | 0    |           |         |                   |
|                                                  | 2-3        | 1    |           |         |                   |
|                                                  | 3-4        | 2    |           |         |                   |
|                                                  | 4-5        | 3    |           |         |                   |
|                                                  | >5         | 4    |           |         |                   |
| sIgE                                             | Negative   | 0    |           |         |                   |
|                                                  | Positive    | 2    |           |         |                   |
| Asthma or allergy, parents and/or siblings       | Negative   | 0    |           |         |                   |
|                                                  | Positive    | 1    |           |         |                   |
| Wheezing-induced sleep disturbances              | No         | 0    |           |         |                   |
|                                                  | Yes         | 2    |           |         |                   |
| Wheezing in absence of cold                      | No         | 0    |           |         |                   |
|                                                  | Yes         | 2    |           |         |                   |

95% CI=95% confidence interval.
Clinical asthma prediction score

The use of the CAPS may enable physicians to reduce the number of children unnecessarily labelled with asthma (low CAPS). For this group, the prior probability went from 42.7% to 21.6%; this helps to reduce unnecessary testing, treatment, and uncertainty. On the other hand, CAPS may allow physicians to identify preschool children who may need proper asthma management (high CAPS). For this group, the prior probability went from 42.7% to 74.3%; a high CAPS may therefore reduce the burden of asthma in these children through better symptom control and fewer exacerbations and hospital admissions. In both situations there will be a reduction in costs. Children with an intermediate CAPS (3>7) retain a probability similar to the probability before applying the CAPS. In this group a watchful waiting policy seems justified, in line with the British guideline, which subsequently suggests a watchful waiting approach with review in the majority of cases with intermediate risk predictions.

Strengths and limitations of this study

Our study has the following strengths. First, we selected children from a relevant population – namely, those children in whom an attending primary care physician had considered asthma. Second, all predictors collected are easily accessible in primary care. Third, we used a stringent definition of asthma to ensure a reliable reference standard diagnosis for research purposes. Fourth, we followed the relevant STARD checklist criteria – a strict protocol with well-defined inclusion and exclusion criteria for the setting and population and a generally accepted reference standard. Fifth, shrinkage of the magnitude of the regression coefficients was used to counteract over-optimism of our model, thus enhancing its validity outside our data set. Sixth, we used NRI to measure the additional value of specific IgE to the model. NRI is a marker to express the model performance and to compare models, focusing on the correct reclassification of patients with and without asthma; it appears to be a clinically more sensible approach than the differences between AUCs.

We see the following limitations to our work. First, our study is a model development study without external validation. We did, however, use bootstrapped variable selection and shrinkage of regression coefficients to temper an over-optimistic performance of CAPS in external populations. Second, we have a high dropout rate. Missingness analyses showed that dropout was higher in children of parents who smoked (OR=1.88) and those with persistent (>1 month) coughing (OR=1.62). Parental smoking may be explained as a psychological cause for dropout; parents who smoke may feel guilty or embarrassed that smoking might have a negative influence on their child's physical condition. We assumed that persistent coughing as a cause of dropout was a coincidental finding. The high dropout rate may also be explained by the fact that, due to an extension of follow-up, participants had to explicitly renew their informed consent. Third, although some combinations of predictor values yield low (<30%) or high (>60%) asthma probabilities, almost 58% of the children in our study population had a predicted probability between 30% and 60%, close to the overall prevalence of 42%. For such children, CAPS results are not likely to change clinical management. Finally, the choice of different thresholds may lead to different results. As far as we are aware, there is no evidence-based advice on thresholds for this type of decision-making in the asthma literature. We decided to use thresholds as a clinically intuitive alternative to more formal methods of reclassification and as an attractive alternative for classical methods of predictive performance such as the area under the ROC curve. We chose a lower limit of 30% to rule out asthma and an upper limit of 60% to rule it in, which implies an area of indecision between the probabilities of 30% and 60%. We think that these limits – or limits close to these – may be relevant in daily primary care practice. However, evidence-based thresholds would require formal benefit-harm or cost-utility analyses that incorporate treatment effects, potential harms (and costs), and children's (and their parents') valuation of all different health outcomes. Although utility measurement in children is not simple, instruments to measure quality of life in children do exist.

Interpretation of findings in relation to previously published work

The API represents innovative research and is the most investigated index. This index was developed in the Tucson general population cohort and has also been tested in other populations (e.g. in the Leicester Respiratory Cohort, n=1,954). Unfortunately, one of the predictors – a blood sample for eosinophilia at one year of age – was omitted from the Leicester cohort.

Several predictive indices for preschool children have been published in the past years. However, the populations in which these indices were developed are atypical of primary care, which may limit their application in the relevant spectrum of high-risk children in primary care. Nevertheless, most indices were prospectively developed and a number of selected predictors correspond reasonably well with ours: family history of atopy or asthma, persistent airway complaints (mainly wheezing), eczema,
Allergic rhinitis, and allergic blood testing. An important difference between the CAPS and other indices is the endpoint: only Eysink et al. developed their index with a comparable outcome – namely, a strict definition of the diagnosis of asthma.21 The major difference between the CAPS and the prediction index of Eysink et al. is that we used clinically more relevant inclusion criteria, which is reflected in the higher overall prevalence of asthma. The other published indices used varying ‘asthma’ definitions, mainly persistent wheezing at a specific age (varying from six to 13 years),9-12,28,31 or asthma defined by asthma symptoms or a physicians’ diagnosis of asthma at age six to 10 years.9,27,30,31 Using weak and varying definitions may cause variation in prevalences and prediction indices, hampering generalization of findings.23

Most research groups used different populations to develop their indices and did not focus on the target population (i.e. preschool children presenting with respiratory complaints to their primary care physician). Eysink et al. included children aged one to four years visiting their primary care physician for persistent coughing of at least five days, and followed them to develop a predictive index for persistent asthma.21 Vial Dupuy et al. retrospectively selected wheezing infants from the medical files of an outpatient department of a paediatric pulmonology and allergy centre. The present study resembles the earlier Dutch study in the primary care setting except that the previous study was retrospective and the children were younger.21 Frank et al. selected their patients by postal survey on parent-reported wheeze, although this matched the clinicians’ findings in fewer than 50% of cases.12,26 In the seven remaining studies, the participants were selected from general population birth cohorts,9,10,12,27,28,30,31 which makes these indices of doubtful applicability to most clinical settings unless major adaptations are made.

The predictive abilities of existing indices are, in general, not high. For example, Castro-Rodriguez et al. claim that the API is a ‘very useful tool’ to predict the long-term outcome of preschool wheeze ‘in every healthcare setting worldwide’. However, it was calculated that the performance of the API in more high-risk populations would be poorer.25 A negative API result in the Tucson cohort (‘asthma’ prevalence at age six of 22.2%) yielded a 10% probability of asthma (NPV=0.90). Overall, earlier predictive indices had relatively high NPVs but disappointing PPVs, which makes these indices of doubtful interest in relation to this article.

Implications for future research, policy and practice
The implementation of a simple risk score in daily care is attractive to both physicians and parents. It allows physicians to provide tailored medical care and follow-up in children at high risk of developing asthma, and to apply a wait-and-see policy in those with a low or intermediate risk of asthma. The increased likelihood of predicting which children will develop asthma will help physicians to explain treatment and prognosis to parents, reducing uncertainties and improving adherence in their children.

Our study was a model development study, extended with internal validation using bootstrapped variable selection and shrinkage of regression coefficients, with the aim of developing a multivariable prediction model. Before the model can be confidently applied in daily clinical practice, external validation is needed.10 Also, further research should focus on new markers or tests to decrease the number of patients remaining inside the area of indecision (i.e. to increase discrimination).

Risk stratification serves three distinct functions in controlled trials: (1) assembling the appropriate spectrum of patients and allowing more efficient subgroup analysis;4 and better assessment of baseline comparability; (2) it may also distinguish persistent and transient complaints early on and therefore increase the accuracy of the RCT results;1 and (3) the CAPS may be used to re-analyse early intervention trials to assess treatment response variability as a function of the CAPS score.1 Such trials should, of course, allow the calculation of a CAPS score.

Conclusions
We have developed an easy-to-use CAPS for preschool children at high risk of asthma in a primary healthcare setting. After suitable validation, the CAPS may assist in guiding shared decision-making to tailor the need for medical or non-medical interventions.

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Contributorship LvdM was the lead investigator, participated in the design, coordinated the follow-up of the study, carried out all the statistical analyses, and drafted the manuscript. KEVV participated in the design, coordinated the follow-up of the study, and revised the manuscript critically. JM and WMCvA participated in preparing the data for statistical analysis and revised the manuscript critically. GtR participated in the design and coordination of the study, supervised it, participated in the statistical analysis, helped to draft the manuscript, and revised the manuscript critically. PIEB participated in the design of the study, supervised it, and revised the manuscript critically. All authors read and approved the final manuscript.

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Clinical asthma prediction score

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Methods

Design
The subjects were one- to five-year-old children in an urban agglomeration in the western part of the Netherlands who were participating in the ARCADE (AirWay Complaints and Asthma Development) prospective cohort study,1 and had presented at one of the pertinent 14 primary care clinics in the previous 12 months with complaints of recurrent coughing (>2 visits), wheezing(>1 visit), and/or shortness of breath (>1 visit).

The study was approved by the Netherlands Central Committee on Research Involving Human Subjects (CCMO/P04.0098C).

Subjects
Eligible for participation were children aged one to five years who visited their primary care physician (in the Netherlands always a primary care physician) with the respiratory complaints of wheezing and/or dyspnoea in the previous 12 months or following 6 months after the start of ARCADE. Children who visited their primary care physician at least twice with the symptom of coughing were also eligible. Exclusion criteria were a confirmed diagnosis of asthma at inclusion (based on spirometry and challenge test), known temporary stay in the region, or parents unable to read or understand Dutch/English. The primary care physician sent postal information about the study to the parents (or caregiver, from now on called ‘parent’) of eligible children, including a reply card, which they could return if they considered participation. After returning the card, further detailed written information and an informed consent form was sent.

Data collection
Questionnaires
At baseline (T0) and at age 6 years (Tend) the parents received a questionnaire on (changes in) housing conditions, family history of allergy, asthma and eczema, presence of pets, breastfeeding, and asthma-related symptoms. Also, information about wheezing, rhinitis, eczema, cough, and phlegm was obtained by the Core Questionnaire of the International Study of Asthma and Allergy in Children (ISAAC).2

Allergy test
Total immunoglobulin E and specific IgE directed against cat (C), dog (D), and house dust mite (HDM) were determined by radioallergosorbent test (RAST) in all the children at baseline.3 IgE positivity to house dust mite, cat and/or dog dander was defined as >0.35kU/L.1 The blood samples were investigated by independent chemical analysts.

Spirometry and bronchial hyper-responsiveness
At six years of age spirometry and bronchial hyper-responsiveness (BHR) assessments (using Pulmoassist 2 spirometer; Jaeger, Würzburg, Germany) was obtained in children with symptoms (wheezeing, shortness of breath or recurrent coughing) or use of asthma medication (β2-agonists, inhaled corticosteroids (ICS)) during the prior 12 months. BHR to increasing doses of methacholine was measured according to the modified method of Cockcroft et al. (using DeVilbiss 646 nebuliser; DeVilbiss, Somerset, MA, USA, with an output of 0.13mL/min).4 Tests were conducted by independent lung function assistants. The parents of children using asthma medication were asked to withhold all bronchodilators for 48 hours prior to the test. In the case of shortness of breath, children received ongoing short-acting β2-agonists up to 8 hours before the test.1

Table A1. Dropout analyses

| Candidate variables | OR   | p Value | 95% CI        |
|---------------------|------|---------|---------------|
| **Inclusion criteria** |      |         |               |
| Coughing            | 0.777| 0.268   | 0.498 to 1.214|
| Wheezing            | 0.721| 0.054   | 0.518 to 1.005|
| Shortness of breath | 1.056| 0.775   | 0.728 to 1.531|
| **Characteristics**  |      |         |               |
| Age                 | 0.62 | <0.001* | 0.53 to 0.73  |
| Female              | 1.06 | 0.747   | 0.74 to 1.51  |
| Breastfeeding >3 months | 1.02 | 0.926   | 0.71 to 1.46  |
| Parental smoking    | 1.89 | 0.001** | 1.32 to 2.71  |
| Child has sibling(s)| 1.01 | 0.977   | 0.7 to 1.45   |
| Asthma, parents, and/or sibling(s) | 0.915 | 0.660 | 0.62 to 1.36 |
| Allergy, parents, and/or sibling(s) | 0.7 | 0.065 | 0.48 to 1.02 |
| **Reported symptoms in past year** | | | |
| Nightly persistent mucous cough or phlegm apart from a cold | 1.0 | 0.995 | 0.67 to 1.48 |
| Dry cough at night, apart from a cold | 0.99 | 0.950 | 0.67 to 1.46 |
| Coughing episode >1 month | 1.62 | 0.038** | 1.03 to 2.55 |
| Wheezing            | 0.47 | 0.048*  | 0.22 to 0.99  |
| Frequency of wheezing attacks | 0.89 | 0.506 | 0.63 to 1.26 |
| Wheezing-induced sleep disturbances | 1.17 | 0.531 | 0.72 to 1.91 |
| Wheezing, in absence of cold | 1.39 | 0.211 | 0.83 to 2.32 |
| Wheezing, in absence of exercise | 0.79 | 0.300 | 0.50 to 1.24 |
| Shortness of breath-induced sleep disturbances | 1.15 | 0.502 | 0.76 to 1.74 |
| Complaints of rhinitis | 0.94 | 0.748 | 0.64 to 1.38 |
| Itch/eczema         | 0.95 | 0.771   | 0.66 to 1.36  |
| **Allergy child**   |      |         |               |
| Specific IgE against cats, dogs, and/or house dust mite | 1.036 | 0.931 | 0.466 to 2.301 |
| **Other**           |      |         |               |
| Drug use for airway complaints | 1.0 | 0.993 | 0.65 to 1.54 |
| Referral to hospital | 1.06 | 0.829 | 0.63 to 1.37 |

Multivariable logistic regression analyses were performed on 730 children for inclusion criteria, for all other variables on 597 children, except for specific IgE (n=388).

* Lower dropout rate, has no influence, because variables are selected by BBS in final model.1
** Higher dropout rate.
Outcome: asthma status at age six
The diagnosis of asthma (masked for predictor values) at six years of age was defined as having persistent symptoms of asthma and/or using asthma medication in the last year in combination with BHR (defined as the methacholine dose (<8.0mg/ml) causing a 20% reduction in FEV1 (PC20-FEV1)) or reversibility (>10% increase in FEV1 after short-acting β2-agonists). Persistent symptoms were defined as self-reported complaints during the previous 12 months of wheezing and/or shortness of breath and/or recurrent coughing.
Children who had not experienced the asthma-related symptoms in the previous year or who had not used asthma medication were not invited for lung function measurement. They were considered as not having asthma.

Probability of asthma at the age of six
Depending on the direction and power of the selected predictors measured by regression coefficients, we developed a clinical asthma prediction score (CAPS) as a points system that is easy to use and does not require a computer or calculator. To use the CAPS in clinical practice, thresholds are needed to either rule in or rule out asthma at the age of six years. We set two thresholds – a lower limit at 30% below which we assumed that many physicians may well choose a “wait and see” policy and an upper limit at 60%, assuming that physicians may pursue more active management such as a trial of asthma treatment. These thresholds imply that children with a predicted asthma risk of between 30% and 60% were assigned to the area of clinical indecision in which a “watchful waiting” policy may be defensible.

Analytical analysis
Analyses of dropouts
Multivariable logistic regression was used to analyse selective dropout (see Table A1). First we analysed the dropout on inclusion criteria (n=730), coughing, wheezing or shortness of breath. Second, we analysed for selective dropout using the other candidate variables (n=597).

Handling of missing data
Multiple imputations using all predicting covariates by iterative chained equations (ICE) to replace missing values were used. Ten imputed datasets were generated.

Development of the model
Candidate predictors were selected based on clinical expert opinion. In the case of highly correlated candidate predictors (if Spearman >0.7), only one was analysed. Further, predictors were selected using bootstrapped backward selection (BBS method) with a significance level of 0.157 (Akaike information criterion; AIC), by fractional polynomial multivariable modelling according to Royston and Sauerbrei. In each imputed dataset, backward variable selection was used in 1,000 bootstrapped samples; if the bootstrap inclusion fraction (BIF) of a covariate was more than 667/1,000 in each imputed dataset, it was selected as a predictor. After selecting the predictors, the coefficients of the predictors were multiplied by a fixed heuristic shrinkage factor.

Performance of asthma prediction model
The predictive performance of the asthma prediction model was assessed by calibration and discrimination. Calibration (Hosmer-Lemeshow) and discrimination (area under the ROC) were calculated on the test cohort.
Lemeshow test (11) describes the accuracy of the measurements between the observed probabilities and the predicted probabilities in independent datasets. Discrimination represents the model’s ability to discriminate between patients with and without asthma during our study. The most familiar measurement for discrimination is the area under the receiver operating characteristic (ROC) curve or concordance (c) statistic: a value of 0.5 represents chance and a value of 1 represents perfect discrimination.

Reclassification/NRI

The additional value of the most invasive predictor – namely, blood testing on specific IgE – was calculated using the area under the ROC curve (AUCdiff) and the NRI, an instrument that focuses on the reclassification of patients with and without asthma. The thresholds were <30% for a small probability of asthma at the age of six years and >60% for a very high probability of having asthma in later life. In the light of the upward movement for events and the downward movement for non-events, an index could be calculated.11 In addition, the results (including SE and 95% CI) were bootstrapped 1000 times in order to test the AUCdiff.11

Probability of asthma at age six by clinical asthma prediction score (CAPS)

To make complex statistical models useful to practitioners, Sullivan et al. developed a points system that is easy to use and does not require a computer or calculator5 (see Table A2). After organising risk factors into categories and determining reference values, referent risk factor profiles (WiREF, i=1,…,4) were selected for each risk factor. A constant (B) was defined, reflecting the increase in risk associated with a 1-year increase in age (B=1*(0.333)=0.333). Points associated with each of the categories of the risk factors were computed by:

\[ \text{Points}_{ij} = \frac{1}{1+\exp(-\sum_{k=1}^{n} \beta_i x_{ki})} \]

To estimate the risk of developing asthma based on the statistical model, the simplified CAPS could be used in daily practice.

All statistics were carried out in Stata/SE Version 10.1 (Stata Corp, College Station, TX, USA).

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Table A3: Positive predictive value (PPV) and negative predictive value (NPV) CAPS <3 and CAPS ≥7

| Category | <3 CAPS | 3 - 6 | ≥7 | Total |
|----------|---------|------|----|-------|
| No asthma | 87      | 145  | 19 | 251   |
| Asthma    | 24      | 108  | 55 | 187   |
| Total     | 111     | 253  | 74 | 438   |
| Risk of asthma | 21.6% | 42.7% | 74.3% | 42.6% |
| PPV       | 55/74=0.743 (95% CI 63.3 to 82.9) |
| NPV       | 87/111=0.784 (95% CI 96.8 to 85.0) |

Figure A1. Formulae to calculate the linear predictor and probability per CAPS category (0-11)

\[ \Sigma_{i=0}^{p} \beta_i X_i = -2.41144 + 0.367(\beta_{\text{age}}) + 0.849(\beta_{\text{IgE}}) + 0.409(\beta_{\text{family history}}) + 0.694(\beta_{\text{sleep}}) + 0.756(\beta_{\text{no cold}}) \]

\[ P = \frac{1}{1+\exp(-\Sigma_{i=0}^{p} \beta_i X_i)} \]