ORIGINAL ARTICLE

Epidemiology and Genetics

A strategy for high-dimensional multivariable analysis classifies childhood asthma phenotypes from genetic, immunological, and environmental factors

Norbert Krautenbacher^{1,2} | Nicolai Flach^{1,2} | Andreas Böck³ | Kristina Laubhahn^{3,4} | Michael Laimighofer^{1,2} | Fabian J. Theis^{1,2} | Donna P. Ankerst^{2,5} | Christiane Fuchs^{1,2,6} | Bianca Schaub^{3,4}

¹Institute of Computational Biology, Helmholtz Zentrum München, German Research Center for Environmental Health GmbH, Neuherberg, Germany

²Technische Universität München, Center for Mathematics, Chair of Mathematical Modeling of Biological Systems, Garching, Germany

³Department of Pulmonary and Allergy, Dr. von Hauner Children's Hospital, LMU, Munich, Germany

⁴Member of German Lung Centre (DZL), CPC, Munich, Germany

⁵University of Texas Health Science Center at San Antonio, San Antonio, Texas

⁶Faculty of Business Administration and Economics, Bielefeld University, Bielefeld, Germany

Correspondence

Christiane Fuchs, Institute of Computational Biology, Helmholtz Zentrum München, German Research Center for Environmental Health GmbH, Neuherberg, Germany. Email: christiane.fuchs@helmholtzmuenchen.de and Bianca Schaub, Department of Pulmonary

and Allergy, Dr. von Hauner Children's Hospital, LMU, Munich, Germany. Email: bianca.schaub@med.uni-muenchen.de

Funding information

Bundesministerium für Bildung und Forschung, Grant/Award Number: 01DH17024 (CF); Deutsche Forschungsgemeinschaft, Grant/Award Number: SFB 1243 (CF, FJT), SFB-TR22 (BS), SCHA-997/8-1 (BS); Else-Kröner-Fresenius Foundation (BS, AB)

Abstract

Revised: 22 December 2018

Background: Associations between childhood asthma phenotypes and genetic, immunological, and environmental factors have been previously established. Yet, strategies to integrate high-dimensional risk factors from multiple distinct data sets, and thereby increase the statistical power of analyses, have been hampered by a preponderance of missing data and lack of methods to accommodate them.

Methods: We assembled questionnaire, diagnostic, genotype, microarray, RT-qPCR, flow cytometry, and cytokine data (referred to as data modalities) to use as input factors for a classifier that could distinguish healthy children, mild-to-moderate allergic asthmatics, and nonallergic asthmatics. Based on data from 260 German children aged 4-14 from our university outpatient clinic, we built a novel multilevel prediction approach for asthma outcome which could deal with a present complex missing data structure.

Results: The optimal learning method was boosting based on all data sets, achieving an area underneath the receiver operating characteristic curve (AUC) for three classes of phenotypes of 0.81 (95%-confidence interval (CI): 0.65-0.94) using leave-one-out cross-validation. Besides improving the AUC, our integrative multilevel learning approach led to tighter CIs than using smaller complete predictor data sets (AUC = 0.82 [0.66-0.94] for boosting). The most important variables for classifying childhood asthma phenotypes comprised novel identified genes, namely PKN2 (protein kinase N2), PTK2 (protein tyrosine kinase 2), and ALPP (alkaline phosphatase, placental).

Conclusion: Our combination of several data modalities using a novel strategy improved classification of childhood asthma phenotypes but requires validation in external populations. The generic approach is applicable to other multilevel databased risk prediction settings, which typically suffer from incomplete data.

KEYWORDS

childhood asthma, complex study design, immunology, machine learning, risk prediction

This is an open access article under the terms of the Creative Commons Attribution-NonCommercial License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited and is not used for commercial purposes. © 2019 The Authors. *Allergy* Published by John Wiley & Sons Ltd.



GRAPHICAL ABSTRACT

Statistical learning on immunological, genetic, and environmental data classifies asthma well. Risk estimation is most precise when incorporating all given data with the novel multi-modality strategy (area under the receiver operating characteristics curve = 0.81). Best predictors are three target genes of microarray data, comprising novel identified genes protein kinase N2, protein tyrosine kinase 2, and alkaline phosphatase, placental. These show the highest importance for childhood asthma classification. ALPP-alkaline phosphatase, placental; AUC-area under the receiver-operator-characteristics curve; CLARA-clinical asthma research association; PKN2-protein kinase N2; PTK2-protein tyrosine kinase 2; SNP-single nucleotide polymorphism.

1 | INTRODUCTION

Asthma, a complex chronic pulmonary disorder, is the most common airway inflammatory disease in children worldwide, with increasing prevalence. It is characterized by bronchial hyperresponsiveness and reversible airway obstruction, causing recurrent episodes of wheezing, cough, shortness of breath, and chest tightness.^{1,2} Several subphenotypes of childhood asthma were suggested in various epidemiological studies.^{3,4} However, clinical practice and also molecular studies still divide children into two main phenotypes, namely allergic and nonallergic asthma.^{5,6} Attempts were made to disentangle distinct underlying pathophysiological mechanisms, but were hampered by the complex nature of the disease.⁶⁻⁹ While singular targets were identified, one could not consistently pinpoint a reliable pattern of relevant pathways critical for asthma phenotype differentiation and in the long-term potentially patient-tailored treatment of the disease. However, this is important as to date, several children with asthma are not well controlled, potentially due to uniform, non-patient-specific therapies with mainly steroids.

Omics data, such as genomics and transcriptomics, have become increasingly available in human cohorts and thus more critical for understanding the pathogenesis of childhood asthma.¹⁰ Inherent high dimensionality, incomplete data, and multiple platforms make the analysis of prediction models complex. Reliable analysis strategies for multi-omics data from multiple platforms in large cross-sectional studies are urgently needed to predict the risk of this multifaceted disease. Tools for integration of multiple omics data sets exist in literature¹¹ but are often restricted to analyzing correlation structures rather than building multi-variable prediction models. Methods have been proposed to do so, that is, using several modalities for prediction.¹² Acharjee et al¹³ use the machine learning method random forest and preselect significant variables. Zhao et al¹⁴ analyze each modality separately and merge the single components. Boulesteix et al¹⁵ incorporate each modality via penalized regression estimating weights for each modality. However, successful solutions are not yet available for cases where different modalities are assessed for different individuals. Strategies to build and validate multivariable prediction models incorporating all individuals and all variables simultaneously are needed for classifying asthma in children.

1365

In this study, we propose a novel approach to optimize prediction of childhood asthma phenotypes when different modalities are used as input factors. Prediction in the context of this paper refers to describing and distinguishing childhood asthma phenotypes in terms of classifying them into the corresponding clinical phenotype category rather than predicting the development of asthma. Our data include questionnaire, clinical diagnostic, genotype, expression microarray, quantitative real-time RT-PCR (RT-qPCR), flow cytometry, and cytokine secretion data. Combining multilevel data types by a reliable analysis strategy for large human cohorts will contribute to detailed understanding of childhood asthma, potentially relevant for novel therapeutic strategies. The strategy can also be translated into numerous other complex diseases.

2 | METHODS

2.1 | Study population

Children between 4 and 15 years from southern Germany were recruited in the University Children's Hospital Munich from the CLARA/

VILEY-Allergy CONTRACTORS OF ALLERY

CLAUS (Clinical Asthma Research Association) study⁶ in three clinical groups, namely healthy children (HC), mild-to-moderate allergic asthmatics (AA), and nonallergic asthmatics (NA). Parents completed a detailed questionnaire assessing health data on allergy, asthma, and socioeco-nomic factors. Asthmatic patients were diagnosed according to GINA guidelines.¹⁶ Inclusion criteria for asthmatics were classical asthma symptoms, including at least three episodes of wheeze and/or a doctor's diagnosis and/or history of asthma medication in the past and lung function indicating significant reversible airflow obstruction according to American Thoracic Society (ATS)/European Respiratory Society (ERS) guidelines.¹⁷ Allergy was defined based on a positive specific IgE level in accordance with clinical symptoms. Blood specimen was collected during the children's recruitment and processed identically.

2.2 | PBMC isolation, RNA and DNA extraction

Peripheral blood mononuclear cells (PBMCs) were isolated within 24 hours after blood withdrawal, cultured in X-Vivo (48 hours) unstimulated (U), stimulated with plate-bound anti-CD3 (3 μ g/mL) plus soluble anti-CD28 (1 μ g/mL), lipid A (LpA, 0.1 μ g/mL), or peptidoglycan (PGN, 1 mg/mL, OR) at 37°C. Cell pellets were used for RNA isolation utilizing the RNeasy Mini Kit (Qiagen, Hilden, Germany), and supernatants were frozen at -80° C. Genomic DNA was extracted from whole blood (Flexigene DNA-Kit, Qiagen).

2.3 | Modalities

We investigated seven data modalities: questionnaire, diagnostic, genotype, microarray, RT-qPCR, flow cytometry, and cytokine data. Diagnostics included weight, height, blood count, immunoglobulins, CrP and IL-6 as well as FeNo.

2.4 Genotyping

Extracted DNA was genotyped for 101 loci using matrix-assisted laser desorption/ionization time-of-flight-mass spectrometry (Sequenom, Inc., San Diego, CA). Deviations from Hardy-Weinberg equilibrium were assessed for quality control of genotyping procedures. Loci were selected based on known biological relevance and genome-wide association study results.¹⁸

2.5 | Microarrays

RNA of PBMC from a subgroup (14AA/8NA/14HC), comparable to the whole population, was analyzed by Affymetrix-GeneChip[®] Human-Gene 1.0 ST-arrays. Quality of scanned arrays was checked by MvA, density, RNA degradation plots, using R and Bioconductor.^{19,20} Robust multichip averages were used for background correction, normalization, and control of technical variation.

2.6 | RT-qPCR, flow cytometry, and cytokines

Isolated RNA was processed (1 μg) with reverse transcriptase (Qiagen). Gene-specific PCR products were measured by CFX96

TouchTM Real-Time PCR Detection System (Bio-Rad, Munich, Germany) for 40 cycles. Subpopulations of 2.5×10^6 PBMC were counted on a FACSCanto II flow cytometer (Becton Dickinson). Cytokine levels were determined in supernatants of cultured PBMCs with Human Cytokine Multiplex Assay Kit (Bio-Rad) using LUMINEX.

2.7 Computational and statistical analysis

The statistical analyses were performed with R software.¹⁹ Details of this section are provided in the article's Supplement. The complex sparse data structure required strategies for handling missing values. Variables containing more than 25% missing values within one modality data set were removed. Remaining missing values were handled via multiple imputation²¹ (without using any information on the outcome variable) since we assumed missingness at random. This yielded a basic structure of the full data set (Figure 1). We could rule out the possibility that this remaining complex missing data suffered from sample selection bias.⁶ The intersection data set containing complete observations from all modalities embraced 33 children.

For classification of the three categorical outcome variable with categories AA, NA, and HC, we utilized four state-of-the-art classification algorithms suitable for high-dimensional predictors: the least absolute shrinkage and selection operator (LASSO) and elastic net,²² both representing penalized regression methods (in our case multi-class logistic regression); and random forest²³ and (stochastic gradient) boosting,²² both machine learning ensemble methods based on decision trees.

The area under the receiver operating characteristics curve (AUC) was used as metric for comparing prediction accuracy.²⁴ As we compared three outcome categories instead of the standard number two, we obtained an overall AUC by calculating a weighted average over the three one-category-vs-all-categories combinations.^{24,25} Prediction models were validated via leave-one-out cross-validation (Figure S1).

For the complex data structure, we utilized two standard modeling strategies and combined them to a novel one. We compared the resulting three approaches to the four mentioned statistical learning approaches (in short: LASSO, elastic net, random forest, boosting). In Strategy A, each modality was analyzed independently, so that all observations were used but training and validation were possible only modality-wise. Strategy B is a complete case model, that is, we used only complete observations where all seven modalities were measured. Here, all modalities were analyzed at once, but only the completely measured cases were left for analysis. The newly developed Strategy C combined the former two: Classifiers were trained on each modality separately in a first step on a training data set. Applying an inner validation, each modality obtained an optimized weight. The weighted classifiers were combined to a single prediction model, which was evaluated on the complete observations. The three strategies are illustrated in Figure 2.

3 | RESULTS

Two hundred and sixty individuals of the CLARA/CLAUS population with well-defined phenotypes (AA/NA/HC) in total were available for

FIGURE 1 Structure of the given data after imputation within each modality. The blue-colored areas depict the given data values (all white areas correspond to missing data). The given data consist of seven groups of variables of the same type (modalities). There are only few subjects containing data for all modalities. The given gene expression by microarray data is the restricting component regarding complete cases and contains the most variables (reduced in figure for illustration reasons)



Variables/modalities



Variables/Modalities

FIGURE 2 Schematic illustration of data partitions taken into account for prediction modeling at a time. A, All observations per modality were included, but training and validation were done separately for each block. B, Only complete observations were used, and classifiers were trained on all modalities at once. C, All modalities and all observations were incorporated in a single prediction model and validated on complete observations

the present analyses. AA cases (47%), NA cases (11%), and HC (43%) in the data differed with respect to variables from seven data modalities (Table S4). Full information on all variables was available for 33 children. The most complete modality was the questionnaire with all 260 individuals being measured. The smallest modality data set regarding the number of measured individuals was the microarray data set with 36 observations. The remaining modality data sets, cytokines, flow cytometry, diagnostics, questionnaire, and RT-qPCR contained 148, 172, 162, 248, and 107 observations, respectively (Table S4).

3.1 | Prediction modeling

For preventing from severe overoptimistic bias regarding performance of a best model, we report results for all models^{26,27}: Strategy A performed prediction on single modalities separately (Figure 2A). On a stand-alone basis, there was no discriminatory power shown for any classifier on flow cytometry (AUC for best classifier boosting 0.54 [0.45-0.64]) and RT-qPCR (AUC for LASSO 0.47 [0.36-0.59], Figure 3A). Here, all Cls crossed the AUC = 0.5 line, indicating that the prediction models did not do better than random guessing. There were moderate performances (mean AUC less than 0.7) for cytokines (boosting 0.60 [0.51-0.70]), SNPs (random forest 0.66 [0.57-0.75]), and diagnostics (LASSO 0.69 [0.61-0.75]). Mean AUCs higher than 0.7 were yielded by modalities environment with an AUC for boosting of 0.75 [0.69-0.82] and microarray with an AUC of 0.74 and a comparatively large confidence interval [0.54-0.90] (Figure 3A).

Strategy B considered only observations with values of all modalities given (Figure 2B) and achieved a higher AUC than Strategy A



FIGURE 3 Comparison of prediction for different modalities for different statistical methods and strategies. A, Performance of prediction models on each modality analyzed separately (Strategy A). B, Performance for complete case model (Strategy B). C, Performance of combination strategy (Strategy C)

for LASSO (0.77 [0.60-91]) and boosting (0.81 [0.65-0.94], Figure 3B), again with large confidence intervals.

Strategy C combined A and B. Here, as in B, boosting outperformed the other classifiers clearly with an AUC of 0.82 [0.66-0.94] (Figure 3C). Performance did not significantly increase from Strategy B to C. However, the classifiers' variance for C decreased slightly as shown by the narrower confidence intervals (Table S1).

3.2 | Variable importance

Strategy B presents a reasonable trade-off between convenient interpretability and good prediction performance. Hence, we investigated its best prediction model with respect to its most important predictor variables. For meaningful interpretation, we considered annotated genes only for the microarray modality set here.

Figure 4 shows the performance of the refitted modified model, that is, Strategy B with annotated genes only. Boosting, which originally performed best (AUC = 0.81 [0.65-0.94]), predicted slightly worse in the modified version (AUC = 0.77 [0.58-0.93]). Here, LASSO performed similarly to boosting (AUC = 0.77 [0.60-0.91]). Therefore, we analyzed the most important variables of both classifiers. As we based our investigations on variable importance on the two prediction models, we looked in detail at the sensitivities and specificities in terms of ROC curves for these two models (Figure 5); even though the overall AUC was equal in both prediction models, their values differed regarding their one-vs-all comparisons. Generally, the predictive quality was higher for discriminating healthy controls from both kind of asthmatics (Figure 5A) and for discriminating allergic asthmatics



FIGURE 4 Performance of prediction models on the 33 complete cases (Strategy B). The procedure was run twice—once the modified model including genes which only contained annotated genes (left), once the original model including nonannotated genes in addition (right). The AUCs are calculated as the average over the 5 imputations; the error bars show 95% bootstrap confidence intervals

from healthy controls and nonallergic asthmatics (Figure 5B) than for discriminating nonallergic asthmatics from healthy controls and allergic asthmatics (Figure 5C) (for boosting: AUC = 0.79 for HC vs all, AUC = 0.78 for AA vs all, AUC = 0.72 for NA vs all).

Over all imputations, LASSO selected 22 non-highly correlated variables, which were exclusively genes from the microarray modality (Table S2, Figure 6B). In contrast, boosting used all variables by preferring and ranking them according to their importance without excluding correlated variables. Here, we took those 50 variables into consideration which were ranked highest (Figure S2). The selection contained variables from modalities microarray, cytokines,



FIGURE 5 Sensitivities and specificities in terms of ROC curves for the two best-performing prediction models, LASSO and boosting, on the 33 complete cases (Strategy B), when all variables were used but nonannotated genes were excluded. ROC curves were calculated separately (aggregated over all 5 imputations) as (A) Healthy controls (HC) vs all others, (B) Allergic asthmatics (AA) vs all others and (C) Nonallergic asthmatics (NA) vs all others. The overall AUC of 0.77 for both prediction models is a weighted average over the three single AUC comparisons. The weights correspond to the proportions of HC (0.36), AA (0.39), and NA (0.24), respectively

diagnostics, environment, and RT-qPCR (Table S3, Figure 6A). The two lists overlapped in three variables, illustrated by Figure 6C and Tables S2 and S3, all of them were genes from the microarray modality: PKN2, PTK2, and ALPP. Thus, we considered these as model-independent most important variables for prediction of childhood asthma.

A wider overlap could be determined with more relaxed assumptions (s. details in Table S5 and Figure S3), that is, when variables in the two sets were considered as corresponding to each other when their correlation coefficient exceeded a predefined threshold (Figure S2). Besides breastfeeding, other characteristics considered as potential confounders in a standard analysis (such as age and sex) did not show high variable importance.

DISCUSSION 4

This study contains a novel proposal for prediction analyses of childhood asthma using cytokine, genotype, flow cytometry, diagnostic, questionnaire, RT-PCR, and microarray data simultaneously. Many studies on childhood asthma currently analyze phenotypes based on assessment of singular measurements only.²⁸

Combining several data types has optimized prediction of childhood asthma phenotypes in the CLARA study. The most important variables for prediction of childhood asthma phenotypes comprised novel identified genes, namely PKN2 (protein kinase N2), PTK2 (protein tyrosine kinase 2), and ALPP (alkaline phosphatase placental).

The need for a new strategy arose from the complex data design with seven groups of variables (modalities) of various dimensions on the one hand, and the comparably rare number of complete cases, where observations were given for all modalities, on the other side. The novel strategy incorporated all individuals and all variables simultaneously. The employed classifiers (LASSO, elastic net, random forest, boosting) were capable of handling biomedical data difficulties such as highly correlated and large numbers of variables, possibly exceeding the number of observations, and of filtering important variables from big amounts of noisy variables, which is especially important for the huge amount of predictor variables and the additional heterogeneity in the variables.

4.1 Prediction by seven modalities—best prediction obtained by boosting

The single-modality approach (Strategy A) showed differences in prediction quality for the various modalities and four classifiers. Prediction was unambiguously successful for environment and microarray, partly successful for cytokines, genetics, and diagnostics, and unsuccessful for flow cytometry and RT-qPCR. This is crucial as several studies are analyzed based on singular modalities.

The complete case approach (Strategy B) proved that combining all variables of all modalities to one model is more predictive than using only single modalities.

Both strategies were trade-offs between using all observations per modality and using all modalities simultaneously. Combining both aspects led to the novel combined approach (Strategy C), using the complete data for the training process (Figure 2C) by training a classifier and optimizing a weight via internal model validation for each modality separately in a first step and aggregating all established components in a second step (Figure S1). This strategy tended to decrease the variability of asthma prediction on independent data (Table S1). Thus, including not only all data modalities but also all observations per modality, as Strategy C does, may offer the chance to improve precision in risk estimates for asthma rather than it is possible by using, for example, only clinical or only diagnostic measures, or otherwise using all possible modalities but taking only those observations into account where all values for all these modalities are measured. Even though the decrease in the variability in terms of smaller confidence intervals was small in our data, in further



FIGURE 6 Variable importance for best models on complete observations. Genes are denoted by their names with the type of stimulation in parentheses. A, Boosting variable importance: Variables ranked under the top 50 by boosting in the complete case model averaged over all five imputations. B, LASSO-selected variables: Variables selected by LASSO in the complete case model over all five imputations. C, Venn diagram/pie charts for sets of variables ranked highest by boosting (50 variables) and of variables selected by LASSO (19 variables). Three variables (genes) were selected in both prediction models

applications, the strategy will generally guarantee at least as good precision as Strategy B, as more information in the data is used. The strategy is especially advantageous when the number of complete cases is substantially smaller than the number of overall individuals in the study. It may even be the only solution when this number is too small for Strategy B.

Boosting showed best performance for both Strategies B and C (Figure 3B/C). This method is convenient for clinical data sets where a multitude of immune-related measurements are available, but missing or small numbers of subjects pose a problem for common analysis strategies.

4.2 | Contributional influences—gene expression is most predictive

Prediction on complete cases using annotated genes only was comparable to the original model using also nonannotated genes and yielded high interpretability regarding the most important variables for prediction. We thus repeated prediction by Strategy B on the adjusted selection of genes. Evaluation by two conceptually different methods, the variable selection via LASSO and the relative influence determined by decision trees in the framework of boosting, yielded three model-independent most important variables for prediction: the genes *PTK2*, *PKN2*, and *ALPP*.

PTK2, a member of focal adhesion kinase (FAK), encodes a cytoplasmic protein tyrosine kinase that localizes to focal adhesions and contributes to integrin-mediated cell processes related to cell survival. The activation of this gene regulates a wide variety of cellular responses and is assumed to be important in the early step of cell growth and intracellular signal transduction pathways.²⁹ Although tyrosine kinases play an important role in several pulmonary mechanisms like in airway hyperresponsiveness and airway remodeling, no correlation between *PTK2* gene and asthma has been described so far.³⁰ *PKN2*, also called protein kinase C-related kinase 2 (*PRK2*), is a Rho target protein which regulates the apical junction formation in human bronchial epithelium. It has been shown critical for human

cancer and would represent a novel gene pathway potentially relevant for childhood asthma.³¹ *ALPP* is a gene which encodes the placental alkaline phosphatase that catalyzes the hydrolysis of phosphoric acid monoesters and was previously identified to be potentially involved in recurrent spontaneous abortion.³² We acknowledge that the identification of the novel genes *PKN2*, *PTK2*, and *ALPP* is based on a limited number of children and requires confirmation in future cohorts. Although these three genes have not been associated with childhood asthma yet, the findings in this study could be a first hint for future investigations.

Further model-specific variables contributing to prediction were obtained (Tables S2, S3). Contrary to the LASSO model which only labeled genes as most important, boosting found variables also from other modalities. One of them is the number of months of breastfeeding. This may have an influence on asthma, however can be a case of translucent correlation since mothers with family history may be biased in their decisions for breastfeeding. Besides this, selected cytokines such as IL-1 β and IL-5, diagnostics variables, and RT-qPCR variables such as *IRF8* have been identified as important by boosting (Figure 6A).

In our results, no genotype variables (SNPs) turned out to be important for prediction. This is not surprising as in our and in previous analyses SNPs on a stand-alone basis did not exceed AUC values of around 0.60.¹⁸ The low predictive effects of SNPs may be covered by effects from other modalities in our analysis.

4.3 | Prediction techniques—using well-established algorithms and all data information

We have used four of the most powerful instruments for prediction in terms of classification from regularization regression methodology to machine learning. In practice, classical approaches as (multivariable) nonpenalized logistic regression can bias parameter estimates and make models instable when variables are highly correlated. Furthermore, there is no maximum-likelihood estimator when the number of variables exceeds the number of observations. Particularly the microarray data set represents both difficulties. Penalized regression, such as LASSO and elastic net, solves these problems: Variable selection generally ensures stability and prevents from overfitting.

Conceptually different but equally sufficient prediction methods are ensembles of decision trees, commonly random forest and boosting, as used here. Both belong to the most popular methods in machine learning and are now used in immune-related analysis. They can handle highly correlated variables and high-dimensional data as well and incorporate interactions between contributing variables. The ensembling principle combines many decision trees at once and thus makes the two methods highly robust.

Applying efficient classification algorithms in combination with running and comparing three modeling strategies complements the methodology of predicting childhood asthma: Multi-omics approaches for childhood asthma have been proposed³³ but rather for finding associations than for building multivariable prediction models. Predicting on each modality separately revealed first answers on the predictive power of each modality. However, this ignored the multivariate -WILEY

structure between the modalities and could hence cause an information loss. The obvious solution to only use complete observations with respect to all modalities, again, came at the cost of a lack of information due to a smaller number of observations. Prediction seemed complete and fully efficient only if all variables and all observations were included in the analysis. Our novel approach, combining weighted prediction scores obtained from the full information of each modality, fulfilled this requirement.

The rigorous use of cross-validation performance to select optimal models brings some limitations, though. Single variables found to be relevant for prediction have no *P*-values attached. Although there are concepts to derive them empirically, their validity is doubtful in the context of statistical modeling with intense variable selection. The different prevalence of phenotypes affected the ability of the model to discriminate between HC, AA, and NA. The smallest group, NA, could not be identified satisfactorily in the presented three-class prediction model, and further efforts are needed to improve this behavior. As another consequence of small sample sizes, we focused on the assessment of main clinical phentoypes and suggest in-depth analysis of additional subgroups such as distinct wheeze and asthma phenotypes in larger studies.

For predicting asthma from seven modalities from genetics, immunology, and environment, we applied robust classification algorithms in concordance with strategies for fully exploiting all information of the data. Penalized regression methods complemented with machine learning approaches have not been used in this context so far and should be considered as efficient prediction methods for this kind of application and beyond. Prediction analysis on incomplete data with respect to different modalities is feasible with certain strategies. We developed a novel strategy combining all information from the data leading to smaller prediction variability. However, the sufficient performance of the complete case prediction model suggests focusing future data collection on enriching complete observations rather than enlarging the number of (at least partially) investigated individuals in total. This is important and requires a strict and thorough recruiting protocol, which is particularly difficult in children and if multicenter studies are envisioned.

Microarray data in terms of three target genes responsible for integrin-mediated cell processes, regulation of apical junction formation in human bronchial epithelium, and placental alkaline phosphatase are predictive for asthma independently of the model approach, even though model-specific results show contributions from other modalities, such as breastfeeding months, IL1-beta and IL-5 cytokine and *IRF-8* gene expression.

For the future, we suggest to implement our novel analysis strategy to more comprehensively understand and analyze complex human immune regulation with respect to childhood asthma phenotypes. The method is also applicable for other cohort studies aiming to assess multi-omics data sets in medium or large cohort studies. Further, when more data like in the given study can be made available, there is high potential for building and improving current risk tools for childhood asthma which can be optimized by distinguishing for pairs of outcome categories as in Ref. 34.

In conclusion, with our approach of combining seven data modalities (cytokines secretion, candidate SNPs, flow cytometry, clinical diagnostics, questionnaires, RT–qPCR gene expression, and microarray gene expression) using a novel strategy, it was possible to improve the classification of childhood asthma phenotypes in contrast to using only single aspects of the data. A rigorous cross-validation scheme was implemented to assess the performance. Of note, a validation in external populations is important. This generic approach is applicable to other risk prediction or classification settings with incomplete data sets, typically arising in circumstances where collection of specimen depends on clinical feasibility and availability of advanced laboratory techniques. The outlined strategy of this manuscript offers the chance to overcome these challenges and provides a quantitative method making use of the entire information at hand.

ACKNOWLEDGMENTS

Our research was supported by the German Research Foundation within SFB 1243, Subproject A17 (CF, FJT), and the SFB TR22 (BS) and Grant Number SCHA-997/8-1; by the Else-Kröner-Fresenius Foundation (BS, AB); by the DZL (German Lung Center, BS, KL) and by the Federal Ministry of Education and Research, Grant Number 01DH17024 (CF).

CONFLICTS OF INTEREST

The authors declare that they have no conflicts of interest.

AUTHOR CONTRIBUTIONS

BS, AB, and KL designed and implemented the CLARA study. NK and NF performed the statistical analyses.CF supervised the statistical analyses. AB, ML, FJT, and DA advised with respect to statistical questions. NK, AB, CF, and BS drafted the manuscript for important intellectual content.

ORCID

Andreas Böck D https://orcid.org/0000-0002-4511-7769 Christiane Fuchs D https://orcid.org/0000-0003-3565-8315 Bianca Schaub D https://orcid.org/0000-0003-1652-8873

REFERENCES

- Asher MI, Montefort S, Björkstén B, et al. Worldwide time trends in the prevalence of symptoms of asthma, allergic rhinoconjunctivitis, and eczema in childhood: ISAAC Phases One and Three repeat multicountry cross-sectional surveys. *Lancet*. 2006;368(9537):733-743.
- Ober C, Yao TC. The genetics of asthma and allergic disease: a 21st century perspective. *Immunol Rev.* 2011;242(1):10-30.
- Depner M, Fuchs O, Genuneit J, et al. Clinical and epidemiologic phenotypes of childhood asthma. Am J Respir Crit Care Med. 2014;189(2):129-138.

- Haldar P, Pavord ID, Shaw DE, et al. Cluster analysis and clinical asthma phenotypes. Am J Respir Crit Care Med. 2008;178(3):218-224.
- Romanet-Manent S, Charpin D, Magnan A, Lanteaume A, Vervloet D; EGEA Cooperative Group. Allergic vs nonallergic asthma: what makes the difference? *Allergy*. 2002;57(7):607-613.
- Raedler D, Ballenberger N, Klucker E, et al. Identification of novel immune phenotypes for allergic and nonallergic childhood asthma. J Allergy Clin Immunol. 2015;135(1):81-91.
- Landgraf-Rauf K, Anselm B, Schaub B. The puzzle of immune phenotypes of childhood asthma. *Mol Cell Pediatr.* 2016;3(1):27.
- Anderson GP. Endotyping asthma: new insights into key pathogenic mechanisms in a complex, heterogeneous disease. *Lancet*. 2008;372 (9643):1107-1119.
- Busse WW, Holgate S, Kerwin E, et al. Randomized, double-blind, placebo-controlled study of brodalumab, a human Anti–IL-17 receptor monoclonal antibody, in moderate to severe asthma. *Am J Respir Crit Care Med.* 2013;188(11):1294-1302.
- 10. Vercelli D. Gene-environment interactions in asthma and allergy: the end of the beginning? *Curr Opin Allergy Clin Immunol.* 2010;10 (2):145-148.
- Hieke S, Benner A, Schlenl RF, Schumacher M, Bullinger L, Binder H. Integrating multiple molecular sources into a clinical risk pre- diction signature by extracting complementary information. *BMC Bioinformatics*. 2016;17(1):327.
- Vazquez AI, Veturi Y, Behring M, et al. Increased proportion of variance explained and prediction accuracy of survival of breast cancer patients with use of whole-genome multiomic profiles. *Genetics*. 2016;203(3):1425-1438.
- Acharjee A, Kloosterman B, Visser RG, Maliepaard C. Integration of multi-omics data for prediction of phenotypic traits using random forest. *BMC Bioinformatics*. 2016;17(5):180.
- Zhao Q, Shi X, Xie Y, Huang J, Shia B, Ma S. Combining multidimensional genomic measurements for predicting cancer prognosis: observations from TCGA. *Brief Bioinform*. 2015;16(2):291-303.
- Boulesteix AL, De Bin R, Jiang X, Fuchs M. IPF-LASSO: integrative L1-penalized regression with penalty factors for prediction based on multi-omics data. *Comput Math Methods Med.* 2017;2017:7691937.
- 16. Global Initiative for Asthma. Global Strategy for Asthma Management and Prevention. 2017. cited 2017.
- Beydon N, Davis SD, Lombardi E, et al. An Official American Thoracic Society/European Respiratory Society Statement: pulmonary function testing in preschool children. Am J Respir Crit Care Med. 2007;175(12):1304-1345.
- Moffatt MF, Gut IG, Demenais F, et al. A large-scale, consortiumbased genomewide association study of asthma. N Engl J Med. 2010;363(13):1211-1221.
- 19. R Core Team. R: A Language and Environment for Statistical Computing. Vienna, Austria: R Foundation for Statistical Computing; 2016.
- Huber W, Carey VJ, Gentleman R, et al. Orchestrating high-throughput genomic analysis with Bioconductor. *Nat Methods*. 2015;12 (2):115-121.
- 21. van Buuren S, Groothuis-Oudshoorn K. mice: multivariate imputation by chained equations in R. J Stat Softw. 2011;45(1):1-67.
- Hastie T, Tibshirani R, Friedman J. The Elements of Statistical Learning. Springer Series in Statistics. New York, NY: Springer New York Inc; 2001.
- 23. Breiman L. Random forests. Mach Learn. 2001;45(1):5-32.
- Fawcett T. An introduction to ROC analysis. Pattern Recogn Lett. 2006;27(8):861-874.
- Provost F, Domingos P. Well-trained PETs: Improving prob- ability estimation trees. CeDER Working Paper #IS-00-04, Stern School of Business, New York University, New York, NY, 10012, 2001.
- Jelizarow M, Guillemot V, Tenenhaus A, Strimmer K, Boulesteix AL. Over-optimism in bioinformatics: an illustration. *Bioinformatics*. 2010;26(16):1990-1998.

- Dupuy A, Simon RM. Critical review of published microarray studies for cancer outcome and guidelines on statistical analysis and reporting. J Natl Cancer Inst. 2007;99(2):147-157.
- Brown KR, Krouse RZ, Calatroni A, et al. Endotypes of difficult-tocontrol asthma in inner-city African American children. *PLoS One*. 2017;12(7):e0180778.
- 29. Parsons JT, Parsons SJ. Src family protein tyrosine kinases: cooperating with growth factor and adhesion signaling pathways. *Curr Opin Cell Biol.* 1997;9(2):187-192.
- 30. Guntur VP, Reinero CR. The potential use of tyrosine kinase inhibitors in severe asthma. *Curr Opin Allergy Clin Immunol.* 2012;12(1):68-75.
- Wallace SW, Magalhaes A, Hall A. The Rho target PRK2 regulates apical junction formation in human bronchial epithelial cells. *Mol Cell Biol.* 2011;31(1):81-91.
- Vatin M, Bouvier S, Bellazi L, et al. Polymorphisms of human placental alkaline phosphatase are associated with in vitro fertilization success and recurrent pregnancy loss. *Am J Pathol.* 2014;184(2):362-368.
- Forno E, Wang T, Yan Q, et al. A multiomics approach to identify genes associated with childhood asthma risk and morbidity. *Am J Respir Cell Mol Biol.* 2017;57(4):439-447.

 Ankerst DP, Hoefler J, Bock S, et al. Prostate cancer prevention trial risk calculator 2.0 for the prediction of low- vs high-grade prostate cancer. Urology. 2014;83(6):1362-1368.

SUPPORTING INFORMATION

Allera

Additional supporting information may be found online in the Supporting Information section at the end of the article.

How to cite this article: Krautenbacher N, Flach N, Böck A, et al. A strategy for high-dimensional multivariable analysis classifies childhood asthma phenotypes from genetic, immunological, and environmental factors. *Allergy*. 2019;74: 1364–1373. https://doi.org/10.1111/all.13745