



Full length article



## Early-life exposome and health-related immune signatures in childhood

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### ABSTRACT

**Background:** Early-life environmental exposures are suspected to modify important immune processes related to child health. Yet, no study has investigated immunotoxicity in relation to the exposome and multiple health domains simultaneously.

**Methods:** Among 845 children (median age 8) from six European birth cohorts included in the Human Early-Life Exposome (HELIX) project, we identified immune signatures of a health score covering cardiometabolic, respiratory/allergic and neurodevelopmental health in children. Those signatures were identified from blood samples in three biological layers (white blood cell (WBC) composition, plasma proteins concentrations, DNA methylation of WBCs) using an advanced factorial analysis supervised on the child health score. Second, we estimated the association between the identified signatures and 91 pre- and postnatal environmental exposures.

**Results:** Three key immune signatures were associated with a better health score in children: a first protein signature characterizing a low inflammatory profile ( $R^2 = 17\%$ ), a second protein signature characterizing a low inflammatory profile with balanced antiviral Th response ( $R^2 = 2\%$ ), and a WBC signature characterizing an immuno-regulatory and naïve profile ( $R^2 = 2\%$ ). In childhood, less exposure to indoor air pollutants, proximity to blue spaces and public transport, healthy dietary habits and higher social capital were associated with the three immune signatures related to a better health score (regression p-values < 0.05). One signature was

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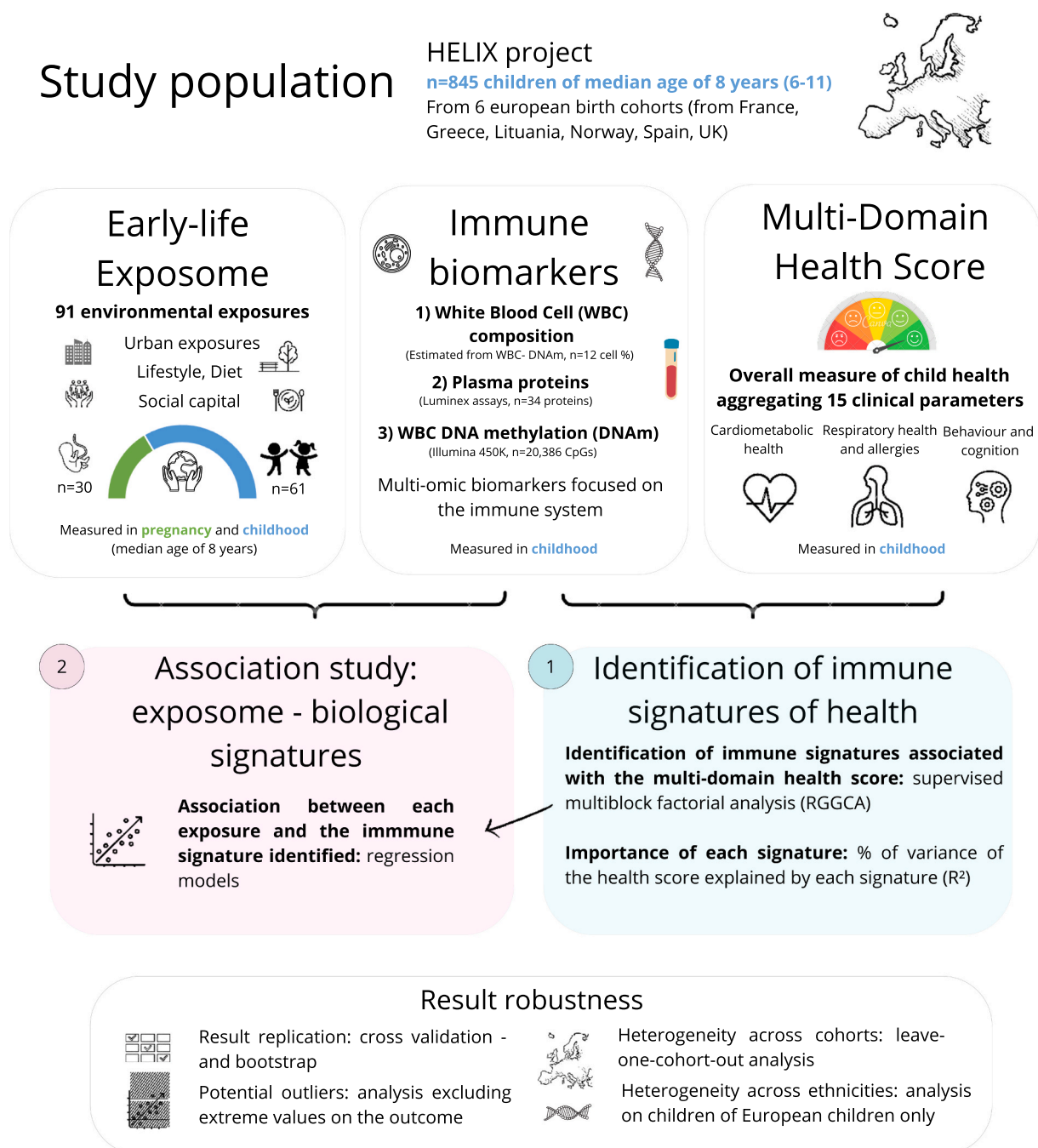
identified from DNA methylation, but was not significantly associated with the health score nor with the exposome.

**Conclusions:** These findings highlight the influence of early-life environmental exposures on key inflammatory processes associated with the cardiometabolic, respiratory and neurodevelopmental health of children.

### 1. Introduction

The exposome, encompassing all environmental exposures from conception onward, is a major determinant of human health across the lifespan (Wild, 2005; Silveira et al., 2007). The early-life environment, including exposures to chemicals, atmospheric pollutants, natural spaces, and lifestyle factors, is known to influence the risk of chronic diseases and cancers later in life (Wallace et al., 2020; Rojas-Rueda et al.,

2021; Wies et al., 2024; Dai et al., 2024). While studies have traditionally focused on one or few specific environmental exposures, holistic approaches are now integrating the entire exposome (Patel et al., 2010). Those novel approaches were made possible with the development of large exposome datasets spanning hundreds of pre- and post-natal environmental exposures (Vrijheid et al., 2014; Vrijheid et al., 2021; Vlaanderen et al., 2021; Ronkainen et al., 2022; van Kamp et al., 2022; Benjdir et al., 2021; Ronsmans et al., 2022; Pronk et al., 2022).



**Fig. 1. Project overview.** This figure describes the methodology used for this project.

**Table 1**  
Environmental exposures studied.

Type of exposure	Exposures	Number of exposures Pregnancy	Number of exposures Childhood
<b>OUTDOOR &amp; INDOOR EXPOSOME</b>			
Outdoor air pollution	NO <sub>2</sub> , PM <sub>10</sub> , PM <sub>2.5</sub> , PM <sub>2.5</sub> absorbance	4	4
Indoor air pollution	NO <sub>2</sub> , PM <sub>2.5</sub> , PM absorbance, Benzene, TEX	0	5
Meteorology	Temperature, humidity, pressure, UV-vit D	3	3
Surrounding natural spaces	NDVI, presence of a major greenspace and bluespace	3	3
Built environment	Population density, building density, street connectivity, accessibility, facility richness, walkability, land use index	9	9
Road traffic	Traffic load on all roads and nearest road, traffic density on nearest road, inverse distance to nearest road	4	4
Water DBPs	THMs, brominated THMs, chloroform	3	0
Total of indoor and outdoor exposures for each period		26	28
<b>LIFESTYLE</b>			
Tobacco	Active and passive smoking (pregnancy), child exposure to smoke (ETS) and parental smoking (childhood)	1	2
Diet	Food intakes (15 groups), food habits, supplements in folic acid, KIDMED score	3	20
Sleep	Average sleep	0	1
Physical activity	Moderate/vigorous activity, sedentary time	0	2
Allergens	Pet (all, dogs, cats)	0	3
Total of lifestyle exposures for each period		4	28
<b>SOCIO-ECONOMIC</b>			
Economic status	Family Affluence Score (FAS)	0	1
Social capital	Family contact, participation in organizations, family size, maternal stress	1	3
Total of socio-economic exposures for each period		1	4

Abbreviations: KIDMED: Mediterranean Diet Quality Index for children, NO<sub>2</sub>: Nitrogen dioxide, PM: Particulate matter, TEX: Toluene, Ethylbenzene, and Xylene.

Alongside its clinical impacts, the exposome can alter fundamental biological processes, particularly those related to the immune system (Johnson et al., 2021; Malesza et al., 2021). These immune-related effects are of particular concern given the central role of the immune function in chronic diseases such as diabetes, obesity and respiratory disorders (Furman et al., 2019). Recently, the Lancet Commission on Planetary Health has recognized immunotoxicity as one of the most worrying and least studied consequences of environmental pollutants (Fuller et al., 2022). Disruptions in immune function may contribute to shared pathways underlying a range of conditions across cardiometabolic, respiratory and mental health domains (Germolec et al., 2017; Piovani et al., 2019). Despite these insights, no exposome study has investigated immunotoxicity in relation to multiple health domains, with the aim of exploring the wide-ranging effects of environmental factors on health through modifications in immune profiles.

As immune processes span multiple biological layers (e.g. genetic, epigenetic, proteomics), the integration of multi-omics data could provide a more holistic characterization of the key mechanisms. While previous studies have explored the impact of the exposome on specific biological layers (Papadopoulou et al., 2021; Stratakis et al., 2020), few have used approaches integrating several biological layers (García-Serna et al., 2022; Maitre et al., 2022). To overcome this limitation, the use of advanced statistical methods, capable of simultaneously analyzing multiple biological and clinical data layers, are of high interest. For example, methods that reduce dimensionality (e.g. latent components, clusters) are particularly promising for identifying complex, yet interpretable, immune processes (Meng et al., 2016; Misra et al., 2021; Babin et al., 2024; Goodrich et al., 2024).

This study, leveraging the Human Early Life Exposome (HELIX) cohort, seeks to address these gaps, by assessing the associations between a large range of early-life environmental exposures and multi-omics immune profiles linked to several child health domains.

## 2. Methods

The methodological workflow adopted for this project is illustrated in Fig. 1 and detailed below.

### 2.1. Study population

This study is based on the HELIX project, including six population-based birth cohorts: Born in Bradford (BiB, UK, Wright et al. 2013), Étude des Déterminants pré et postnataux du développement et de la santé de l'Enfant (EDEN, France, Heude et al. 2016), Infancia y Medio Ambiente (INMA, Spain, Guxens et al. 2012), Kaunas Cohort (KANC, Lithuania, Grazuleviciene et al. 2011), The Norwegian Mother, Father and Child Cohort Study (MoBa, Norway, Magnus et al. 2016), and Mother-Child Cohort (RHEA, Greece, Chatzi et al. 2017). Initially, 32,000 mothers were recruited during pregnancy (2003–2009), from which 1,301 mother–child pairs were followed-up in 2014–2015 when the child was between 6 and 11 years, depending on the cohort (Vrijheid et al., 2014; Maitre et al., 2018). Standardized protocols were employed for biological sample collection, questionnaire administration, health examinations, and exposure characterization. The present study included 845 mother–child pairs with available data on the immune biomarkers and the outcome.

### 2.2. Exposome

As detailed in Table 1, a total of 13 families of exposures were assessed in each mother–child pair, with 31 prenatal and 60 childhood exposures. They include outdoor exposures (outdoor air pollution, built environment, road traffic, surrounding natural spaces, meteorological data, water disinfections by products), indoor air pollutants, lifestyle (tobacco exposure, diet, physical activity, sleep, pet owning) and socio-economic factors. Outdoor exposures were evaluated using geospatial

methods and remote sensing at home and school addresses (Robinson et al., 2018; Tamayo-Uria et al., 2019). Exposure to indoor air pollutants was assessed through predictive modeling based on a panel study of 150 mother–child pairs (more information in eMethods 1 and eTable 1). Lifestyle factors, exposure to water disinfection by products and socio-economic factors were collected via questionnaires. More details on exposure assessment can be found in eMethods 1.

### 2.3. Immune biomarkers

Three biological layers were analyzed from blood samples in childhood (6–11 years): **1) White Blood Cell (WBC)** composition estimated from DNA methylation measured in buffy coat, **2) plasma proteins (Prot)** concentrations and **3) DNA methylation (DNAm)** of white blood cells (genome-wide) in buffy coat. Table 2 details the biomarkers studied and the measurement methods used, with more details in eMethods 2. Briefly, plasma proteins were estimated by Luminex using the Cytokines 30-plex, the Apolipoprotein 5-plex, and the Adipokine 15-plex. DNAm of WBC was assessed in buffy coat using Illumina 450 K (genome-wide), from which WBC composition was derived using the Houseman method (Salas et al., 2022). For the study of DNAm, pre-processing included addressing technical issues such as batch effects and WBC composition through residualization analysis (Rao, 1969; Demissie and Cupples, 2011), filtering out CpGs with high technical variability (Bose et al., 2014) and those with no association with the exposome based on the study of Maitre (Maitre et al., 2022). A total of 20,386 CpG sites were selected from the 480,071 sites initially measured.

**Table 2**  
Immune biomarkers studied.

	1) White blood cell (WBC) composition	2) Plasma Proteins (Prot)	3) DNA methylation (DNAm)
<b>MEASUREMENT METHOD</b>	Estimated from buffy coat DNA methylation of WBCs using Houseman method (Salas et al., 2022)	Luminex: Cytokines 30-plex, Apolipoprotein 5-plex, and Adipokine 15-plex	DNA methylation of WBC in buffy coat using Illumina 450 K (genome-wide)
<b>LIST OF BIOMARKERS</b>	Cell type percentages (N = 12): Monocytes, neutrophils, eosinophils, basophils, memory B cell, naive B cell, memory CD4 + T cell, naive CD4 + T cell, memory CD8 + T cell, naive CD8 + T cell, T-regulatory cells, and natural killer cells	Cytokines (N = 23): IL1β, IL1RA, IL2, IL2RA, IL4, IL5, IL6, IL8, IL10, IL12, IL13, IL15, IL17, MCP1, Eotaxin, MIP, MIP1β, IFNγ, IFNα, TNF, IP10, MIG, G-CSF Other proteins (N = 12): FGF, EGF, HGF, ApoA1, ApoB, ApoE, adiponectin, CRP, leptin, C-peptide, serpine1/PAI-1, BAFF	N = 20,386 CpGs sites after filtering CpGs based on technical variability and association with the exposome

ApoA1: Apolipoprotein A1, ApoB: Apolipoprotein B, ApoE: Apolipoprotein E, BAFF: B-cell activating factor, CRP: C-reactive protein, EGF: Epidermal growth factor, Eotaxin: Eosinophil chemotactic protein, FGF: Fibroblast growth factor, G-CSF: Granulocyte colony-stimulating factor, HGF: Hepatocyte growth factor, IFN α: Interferon-alpha, IFN γ: Interferon-gamma, IL: Interleukin, IP10: Interferon gamma-induced protein 10, MCP1: Monocyte chemoattractant protein-1, MIG: Monokine induced by gamma interferon, MIP: Macrophage inflammatory protein, MIP1β: Macrophage inflammatory protein-1 beta, Serpine1/PAI-1: Plasminogen activator inhibitor-1, TNF alpha: Tumor necrosis factor-alpha.

### 2.4. Multi-domain health score

We integrated pre-clinical and clinical data through a multi-domain health score, thereafter called “health score”, developed in a previous HELIX study (Amine et al. 2023). This score is derived from fifteen health parameters, which were aggregated into three subscores (respiratory, cardiometabolic, and neurodevelopmental), each represented as z-scores (zero average, unit of 1 standard deviation, more details in eMethods 3). By design, a higher health score indicates better health status in children. This score is low-to-moderate for children who exhibit low-to-moderate health in cardiometabolic, respiratory/allergy and neurodevelopmental domains simultaneously, as well as for children who are severely affected in one health domain while being unaffected or poorly affected in the other two.

### 2.5. Covariates

Confounders adjusted for in the analysis included cohort, child age (continuous), sex, ethnicity (European vs non-European origin), maternal education (low, medium, high), breastfeeding duration (<11 weeks, 11–35 weeks, >35 weeks), pre-pregnancy BMI (kg/m<sup>2</sup>), maternal age (years), season of birth (winter, spring, summer, autumn), time since last meal (analyses on plasma proteins only), and hour of blood collection (analyses on plasma proteins only). Exposures, immune biomarkers and the health scores were pre-corrected on these covariates using a residualization approach (Rao, 1969; Demissie and Cupples, 2011).

### 2.6. Statistical analysis

Exposures and covariates were transformed to reach normality when needed (see eTable 2), and imputed using chained equations (White et al., 2011) (first imputed dataset used), as detailed in eMethods 4. Continuous exposures were centered and standardized by the inter-quartile range (see eTable 2).

In the first step, immune signatures predictive of the multi-domain health score were identified using a multi-block model (see more details in eMethods 5). Regularized Generalized Canonical Correlation Analysis (RGCCA) (RGCCA R package), an advanced multi-block supervised factorial analysis model, was applied to estimate such signatures, also called “latent components”, in each data block (WBC, proteins and DNAm) (Tenenhaus et al., 2014). The optimal number of signatures, and a sparsity parameter used for DNAm, were estimated based on cross-validation. Evaluation of the model was performed using cross validation: we reported the R<sup>2</sup> and p-value based on linear regression models to assess the ability of the immune signatures (individually or together) to predict the health score and its sub-scores (cardiometabolic, respiratory, and neurodevelopmental). Interpretation of the signatures was based on the contribution of each immune biomarker given by the loadings, i.e. the correlation between a biomarker and the signature, and 95 % intervals were estimated by bootstrap. DNAm results were further interpreted using functional enrichment analysis with the Kyoto Encyclopedia of Genes and Genomes (KEGG) database via ClusterProfiler v3.8.0 in R (Kanehisa and Goto, 2020).

In a second step, associations between environmental exposures and the identified immune signatures were assessed using univariate linear regression models.

We conducted further analyses to examine specific hypotheses and perform robustness checks. To further investigate the role of WBC correction on DNAm results, we conducted an analysis using DNAm levels uncorrected for WBC composition. To address sex-specific associations, the analysis was further stratified on sex. In addition, robustness checks were performed through sensitivity analyses, including leave-one-cohort-out, removal of the 4 % extreme values on the outcome and analysis restricted to children with European-origin. We

also compared the results to a more traditional “Meet in the middle” approach (Chadeau-Hyam et al., 2011), which aims to identify the immune biomarkers simultaneously associated with environmental exposures and the child health scores, based on regression models (see eMethods 5).

Analyses were done with R version 4.2.1, and codes are publicly available on GitLab (Amine, 2025).

### 3. Results

#### 3.1. Description of the population

The study population comprised 47 % girls, and children had a median age (Q25; Q75) of 8 years (6.5; 8.9) at follow-up (eTable 3). The median maternal age at childbirth was 31 years, with 51 % of mothers having a high degree of education. Description of the immune biomarkers (eFig. 1) and the multi-domain health score is available in Supplementary Data 1 (eFig. 2, eTable 3 and 4), and exposure levels were previously described (Robinson et al., 2018; Amine et al., 2023).

#### 3.2. Identification of immune signatures of the health score using a multi-block model

Based on cross validation, the multi-block model identified seven Latent Components (LC), representing seven health-related immune signatures: one for DNA methylation (DNAm), three for plasma proteins (Prot) and three for WBC composition, as shown in Fig. 2. The signatures were poorly correlated with each other (maximum  $\rho = 0.2$ , see eTable 5). Together, these immune signatures explained 19 % of the multi-domain health score (cross validated  $R^2$ ). By design, all signatures were associated with a better multi-domain health score.

The first protein signature (Prot-LC1) best predicted the multi-domain health score (cross-validated  $R^2 = 17$  %), followed equally by the second protein signature (Prot-LC2) and the first WBC signature (WBC-LC1) ( $R^2 = 2$  % each) (see Fig. 2). The other signatures (Prot-LC3, WBC-LC2, WBC-LC3, DNAm-LC1) were excluded because of their weak associations with the child health score (cross-validated  $p$ -value  $> 0.1$ ), but their description can be found in the Supplementary Data 1 (eFig. 3).

The main contributors to Prot-LC1 were IL1 $\beta$ , IL6 (pro-inflammatory cytokines) as well as leptin, Hepatocyte Growth Factor (HGF) and C-Reactive Protein (CRP), all with negative loadings. Therefore Prot-LC1 was labeled as a “Low inflammatory profile” (see Fig. 2 and Supplementary Data 2). The signature Prot-LC2 was characterized by high levels of IL1RA, IL2R (immune modulators), IL4, IL5, IL13 (Th2 response), IL17 (Th17), IL12, IL2, IFN $\gamma$  (Th1 response) and IFN $\alpha$  (antiviral), thus it was labeled as a “Low inflammatory and balanced antiviral Th response”. The term “balanced” was chosen because of the presence of several Th response, without specific polarity (e.g. Th1/Th2). Lastly, the signature WBC-LC1 was characterized by high proportions of T regulatory cells, CD4 naïve, CD8 naïve, and lower proportions of neutrophils, monocytes and eosinophils, therefore it was labeled as an “Immuno-regulatory and naïve profile”. As shown in Fig. 2 and detailed in eTable 6, Prot-LC1 and WBC-LC1 were significantly associated with all subscores (cardiometabolic, respiratory, neurodevelopmental). Additionally, Prot-LC1 and Prot-LC2 were more strongly associated with the cardiometabolic subscore than with the neurodevelopmental and respiratory subscores, as confirmed by a beta comparison test ( $p < 0.01$ , data not shown).

Association between the exposome and the health-related immune signatures.

As shown in Fig. 3, several environmental factors were associated with immune signatures related to a better health score (see also eTable 7).

Among outdoor and indoor exposures, childhood indoor exposure to particulate matters with a diameter less than or equal to 2.5  $\mu\text{m}$  (PM<sub>2.5</sub>), PM<sub>2.5</sub> absorbance and benzene was associated with a decrease in Prot-

LC1, with indoor PM<sub>2.5</sub> also associated with decreased WBC-LC1 and Prot-LC2. On the other hand, surrounding blue spaces (size greater than 10,000 m<sup>2</sup> in a 300 m buffer near residence) during pregnancy and childhood were associated with higher WBC-LC1 and Prot-LC1, respectively. Higher ambient temperature (the month preceding the visit) and better access to transports (bus lines in a 300 m buffer) during pregnancy were associated with higher WBC-LC1 and Prot-LC1, respectively.

Regarding lifestyle and diet, high intakes of vegetable, organic food, and breakfast cereals (vs low intake) in childhood was associated with higher Prot-LC1. Moderate intake of sweets (vs. low intake) and high intake of soda (vs. low intake) were associated with higher WBC-LC1. Moderate intake of bakery products was associated with higher Prot-LC2. The household presence of a dog in childhood was associated with lower Prot-LC2.

Finally, two social factors were associated with higher Prot-LC1, namely the child social participation in an organization (vs none) and the number of people living in the house.

#### 3.3. Supplementary analyses

Using DNA methylation (DNAm) values uncorrected for white blood cell (WBC) composition, the multiblock model identified a DNAm signature with a significantly higher  $R^2$  (2.7 %) compared to the main analysis (0.1 %) (see eTable 8). This signature was significantly associated with a higher health score (cross validated  $p$ -value  $< 0.05$ ). A total of 11,655 CpGs (approximately half of the CpGs considered) significantly contributed to this signature, with corresponding genes enriched in biological processes involved in neuroimmune interactions, PI3K-Akt signaling, hormone and insulin signaling and hematopoietic cell lineage (adjusted  $p$ -value  $< 0.05$ , see eTable 9). Notably, this signature showed a strong correlation with WBC-LC1 ( $\rho = 0.87$ , see eTable 10), and similar associations with environmental exposures than those observed with WBC-LC1 (see eTable 11).

The multi-block model revealed similar signatures for boys and girls, except for Prot-LC2, whose loadings displayed significant instability in females (see eFig. 4).

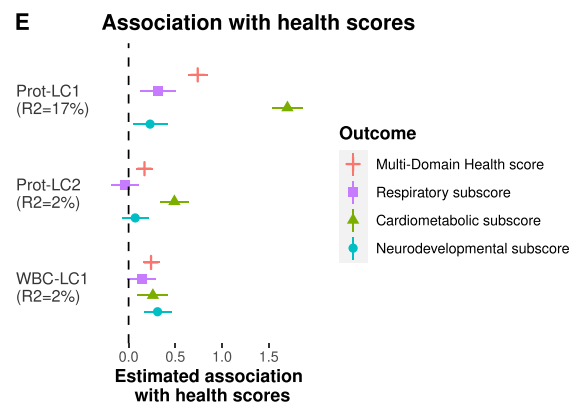
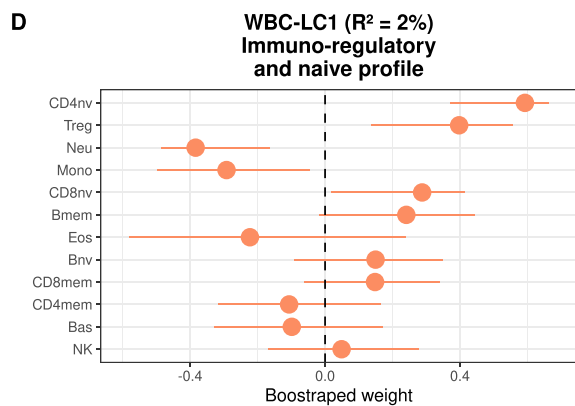
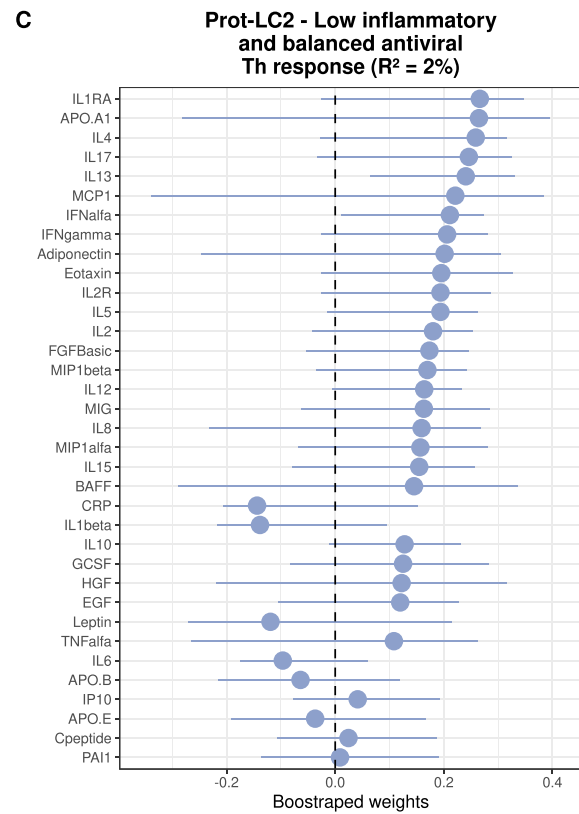
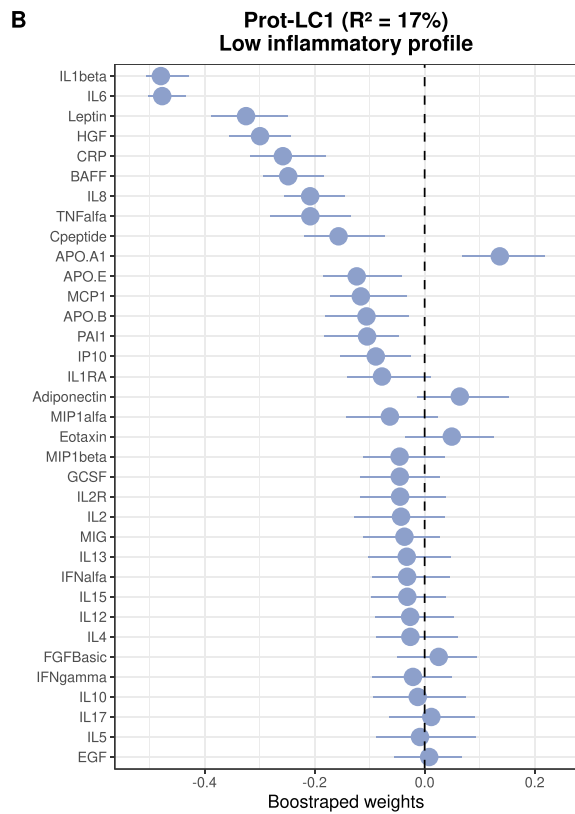
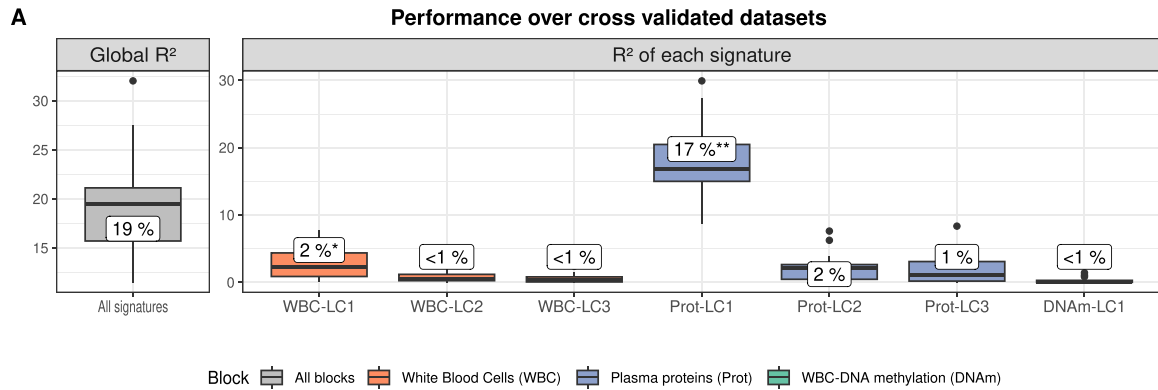
Results from the sensitivity analyses reinforced those from the main analysis (see eFig. 5–6 and Supplementary Data 2), with similar signatures identified. Similarly, the “Meet-in-the-Middle” analysis, based on univariate models, highlighted a similar set of biomarkers to the multi-block model (see the Supplementary Data 3).

## 4. Discussion

This study uniquely explores how environmental exposures influence key health-related immune signatures in children, identified using an advanced multi-block model (RGCCA). It simultaneously integrated exposome data, multi-omic biomarkers focused on the immune system, and a multi-domain health score covering the child cardiometabolic, respiratory and neurodevelopmental health. We identified three immune signatures associated with a better health score in children: Prot-LC1 (low inflammatory profile), WBC-LC1 (immuno-regulatory and naïve profile), and Prot-LC2 (low inflammation and balanced antiviral Th response). Among childhood exposures, better indoor air quality, proximity to blue spaces, healthy dietary intakes, and a strong social capital were associated with immune signatures related to a better health score. Although a DNA methylation signature was also identified, it showed no significant association with the health score or the exposome. These findings highlight the influence of several environmental factors on key inflammatory processes, which hold significant clinical relevance for cardiometabolic, respiratory and neurodevelopmental health in children.

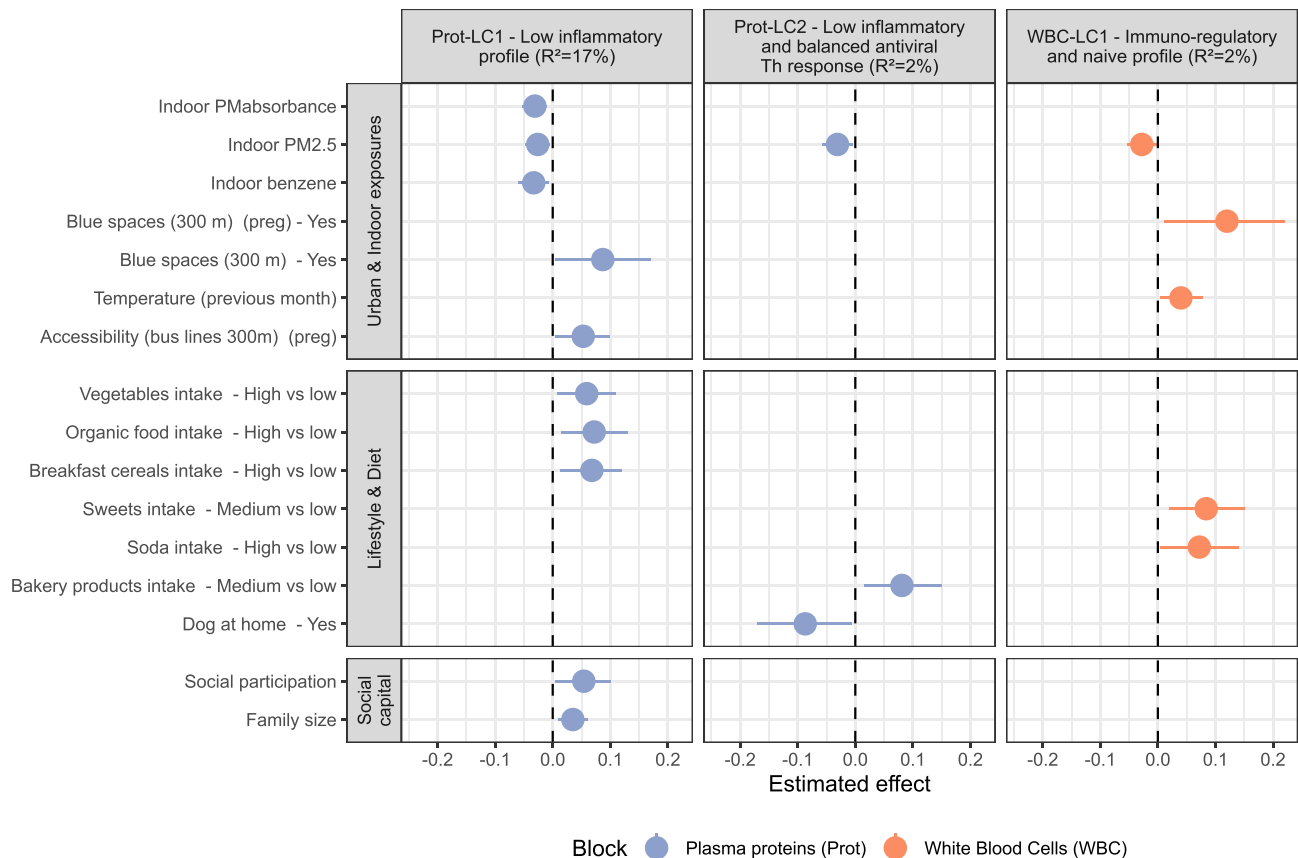
#### 4.1. Main findings

Using RGCCA, an advanced dimension reduction technique, we



(caption on next page)

**Fig. 2. Immune signatures estimated using a multi-block model.** This figure describes the immune signature (latent components) estimated using a multi-block model (Regularized Generalized Canonical Correlation Analysis). It includes the distribution of the performance indicators across the cross validated datasets (5-fold repeated 5 times) (panel A), as well as biomarker loadings for the main immune signatures which are Prot-LC1 (panel B), Prot-LC2 (panel C) and WBC-LC1 (panel D) and their association with the health scores (panel E). For panel A, Rsquared (R<sup>2</sup>) and p-values have been obtained by fitting a linear regression between the health score and the given signature (or all signature for “Global R<sup>2</sup>”), with \* or \*\* meaning that the p-value of association was below 0.05 or 0.01, respectively. For panels B to D, the point represents the biomarkers loading, i.e. the correlation between a given biomarker and a given signature, and 95% confidence intervals were computed using bootstrapping. Panel E represent the estimated coefficient of association and 95% confidence interval between Prot-LC1 and WBC-LC1 with the different health scores, obtained by linear regressions. Abbreviations: ApoA1: Apolipoprotein A1, ApoB: Apolipoprotein B, ApoE: Apolipoprotein E, BAFF: B-cell activating factor, Bas: Basophils, Bmem: B memory, Bnv: B naïve, CD4mem: CD4 memory, CD4nv: CD4 naïve, CD8mem: CD8 memory, CD8nv: CD8 naïve, CRP: C-reactive protein, EGF: Epidermal growth factor, Eos: Eosinophils, Eotaxin: Eosinophil chemotactic protein, FGF: Fibroblast growth factor, G-CSF: Granulocyte colony-stimulating factor, HGF: Hepatocyte growth factor, IFN $\alpha$ : Interferon-alpha, IFN gamma: Interferon-gamma, IL: Interleukin, IP10: Interferon gamma-induced protein 10, MCP1: Monocyte chemoattractant protein-1, MIG: Monokine induced by gamma interferon, MIP: Macrophage inflammatory protein, MIP1beta: Macrophage inflammatory protein-1 beta, Mono: Monocytes, Neu: Neutrophils, NK: Natural Killers, Prot: Proteins, Serpine1/PAI-1: Plasminogen activator inhibitor-1, TNF $\alpha$ : Tumor necrosis factor-alpha. Treg: T regulatory, WBC: white blood cell.



**Fig. 3. Significant associations between exposures and the main immune signatures.** Significant associations between exposures and the main immune signatures estimated using the multi-block model (RGCCA) in the study population (n = 845), with a p-value threshold of 5 %. Exposures labeled with “(preg)” refer to pregnancy-related exposures, while the others were measured during childhood (ages 6–11 years). The R<sup>2</sup> values represent the cross-validated R<sup>2</sup> from fitting a linear regression between the respective immune signature and the health score. Abbreviation: PM: particulate matter, Prot-LC1: first protein latent component, Prot-LC2: second protein latent component, WBC-LC1: first white blood cell latent component.

identified three key immune signatures—Prot-LC1, Prot-LC2, and WBC-LC1—associated with better health outcomes across three key health domains in children (cardiometabolic, respiratory, neurodevelopment). Prot-LC1, the primary signature (R<sup>2</sup> = 17 %), reflects a low inflammatory profile, characterized by reduced levels of pro-inflammatory cytokines (IL1 $\beta$ , IL6, TNF), CRP, and HGF, the latter suggesting a tissue repair response to inflammation-induced stress (Matsumoto and Nakamura, 1992; Molnarfi et al., 2015). The inclusion of leptin highlights the interplay between metabolism and inflammation, with metabolic disorders (e.g., obesity) driving cytokine production by fat cells (e.g., IL1 $\beta$ , IL6) (Febbraio, 2014; Hildebrandt et al., 2023). The secondary signatures, WBC-LC1 and Prot-LC2 (R<sup>2</sup> = 2 % each), further emphasize the key role of inflammatory regulation, as evidenced by the contribution of

regulatory T cell and myeloid cells (neutrophils, monocytes) in WBC-LC1 (Josefowicz et al., 2012; Noone et al., 2024) and anti-inflammatory cytokines (IL1RA, ILR2) in Prot-LC2. Interestingly, the identification of WBC-LC1 adds further evidence of the elevated presence of neutrophils in individuals with poor health, potentially representing either a cause or a consequence of existing conditions. This is in line with previous evidence showing associations between neutrophilic inflammation and respiratory-related diseases already in childhood, such as pneumonia (Lu et al., 2021), and multiple chronic diseases in adulthood (Herrero-Cervera et al., 2022). In addition, these signatures reflect the degree of exposure to previous viral infections, with elevated IFN $\alpha$  in Prot-LC2 and high naïve T cells in WBC-LC1, suggesting the presence and absence of infections, respectively. Except for Prot-LC2,

these immune signatures were associated with each sub-score (cardiometabolic, respiratory/allergic, and neurodevelopmental), highlighting the central role of the immune system, and inflammation in particular, across various health domains (Hildebrandt et al., 2023; Murdoch and Lloyd, 2010; Beurel et al., 2020).

The results of this study suggest that better indoor air quality, the absence of a dog, and higher ambient temperature, may significantly improve the health-related immune signatures identified. Better indoor air quality (PM<sub>2.5</sub>, PM<sub>2.5</sub> absorbance, benzene) was associated with favorable immune signatures at both protein and WBC levels, likely due to the PM effects on oxidative stress (Gangwar et al., 2020) and the link between oxidative stress and inflammation (Sies et al., 2017). Moreover, the absence of a dog was associated with better Prot-LC2, suggesting balanced Th response in absence of allergens, although the literature on this matter remains unclear (Chen et al., 2010). Lastly, higher ambient temperature in the month preceding the health visit was associated with better WBC-LC1, possibly reflecting decreased infections during warmer seasons (Moriyama et al., 2020; Xiong et al., 2023), as evidenced in WBC-LC1 by higher proportions of naive versus memory cells.

In addition, multiple types of child food intakes were associated with the immune signatures. A healthy diet, rich in vegetables, organic foods and including breakfast, was associated with higher Prot-LC1 (i.e. better health), reflecting their benefits for inflammation regulation and metabolic health (Rampersaud et al., 2005; Stanaway et al., 2022; Hosseini et al., 2018). On the other hand, moderate sweet and bakery product intakes and high soda consumption were associated with higher WBC-LC1 or Prot-LC2. While moderate sugar intake may have some benefits, the association with high soda intake, including artificially sweetened options, warrants further investigation (Basson et al., 2021; Kränkel and Rauch-Kroehnert, 2023).

Lastly, stronger social capital and proximity to natural spaces and public transports, were associated with immune signatures (Prot-LC1 and WBC-LC1) related to better health. Protective associations were found with the family size and the child participation in organizations, possibly reflecting the importance of a good social network for the child overall health. Possible underlying mechanisms involved enhanced antiviral immunity, potentially due to more pathogen exposure (Pressman et al., 2005; Leschak and Eisenberger, 2019), cortisol reduction (Pressman et al., 2009), and protection against social isolation and interpersonal stressors, which can exacerbate inflammation (Leschak and Eisenberger, 2019; Fuligni et al., 2009; Segerstrom and Miller, 2004; Miller et al., 2011; Uchino et al., 2018). Similarly, surrounding blue spaces and public transport near the residency appeared to be associated with healthier immune signatures, likely by promoting social interactions and encouraging physical activity (Gascon et al., 2017; Chastin et al., 2021).

While the exposome is suspected to influence epigenetic processes (Wu et al., 2023; Broséus et al., 2024), the signature from DNA methylation of WBCs showed no significant association with the health score (cross-validated  $p > 0.1$ ) or environmental exposures in the main analysis. One hypothesis is that correcting DNA methylation for WBC effects may have removed important variability. When WBC effects were retained in the supplementary analysis, the  $R^2$  of DNAm-LC1 was significantly improved (2.7 % vs. 0.1 % in the main analysis). The new DNAm and WBC signatures were highly correlated, reinforcing the hypothesis that the association between the new DNAm signature and the health score could be strongly confounded by WBC composition, a known determinant of both DNAm and health. This is further supported by the enrichment analysis, which suggested that CpGs contributing to the new DNAm signature were involved in biological processes related to cell type composition, such as the hematopoietic cell lineage (cell differentiation) (Dharampuriya et al., 2017) and PI3K-Akt signaling (regulation of Treg cells) (Liu et al., 2009). However, the DNAm signature still explained less variability than the protein signature, likely due to the lower intra-block correlation, which makes this layer less suited for the identification of latent components.

#### 4.2. Strength and weaknesses

This study has several strengths, including the analysis of numerous exposures, immune biomarkers, and health outcomes, providing a comprehensive view of immunotoxicity in relation to the exposome and multiple health domains. By integrating multi-omic data with the RGCCA method, an advanced multi-block dimension reduction technique, we were able to identify complex yet interpretable immune processes. We have made the codes used in this project publicly available, providing a valuable resource for future exposome-omics studies (Amine, 2025). Moreover, we ensured the strong clinical relevance of these immune processes by supervising the model on a novel health score that integrates clinical and pre-clinical parameters of cardiometabolic, respiratory and neurodevelopmental health in children.

However, we recognize that our study has some limitations. The multi-domain health score aims to study simultaneously three health areas; thus, it cannot identify immune signatures specific to an individual health domain. Additionally, WBC analysis relied on estimated cell proportions by using a validated model, rather than measured counts (Salas et al., 2022). Similarly, exposure to indoor air pollutant was assessed using prediction models from a subset of participants ( $n = 150$ ), thus may be subject to measurement errors suggesting cautious interpretation in the results observed, especially for pollutants with low  $R^2$  values in their underlying models (e.g. Benzene, with an  $R^2$  of 31 %, see eTable 3). Nevertheless, these errors are expected to be non-differential, thus potentially biasing the association results towards the null without generating spurious associations. Lastly, the cross-sectional study design limits causal inferences, with exposures (at the exception of those from pregnancy), immune biomarkers, and outcomes measured simultaneously in childhood. To avoid potential reverse causality due to this design, we preferred not to analyze chemical pollutants, which were measured in HELIX in the same biological samples as the immune biomarkers.

#### 5. Conclusion

This novel study investigated associations between the early-life exposome and health-related immune signatures in children. The findings suggest that better indoor air quality, proximity to natural spaces, healthy dietary habits, and strong social capital are associated with reduced and better-regulated inflammation, at the protein and white blood cell levels. Overall, these results underline the importance of these environmental factors in mitigating immunotoxicity related to multiple areas of health in children.

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### Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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### Appendix A. Supplementary material

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.envint.2025.109668>.

### Data availability

The authors do not have permission to share data.

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