Early-life environmental exposure determinants of child behavior in Europe: A longitudinal, population-based study

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A R T I C L E   I N F O

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A B S T R A C T

Background: Environmental exposures in early life influence the development of behavioral outcomes in children, but research has not considered multiple exposures. We therefore aimed to investigate the impact of a broad spectrum of pre- and postnatal environmental exposures on child behavior.

Methods and findings: We used data from the HELIX (Human Early Life Exposome) project, which was based on six longitudinal population-based birth cohorts in Europe. At 6–11 years, children underwent a follow-up to characterize their exposures and assess behavioral problems. We measured 88 prenatal and 123 childhood environmental factors, including outdoor, indoor, chemical, lifestyle and social exposures. Parent-reported behavioral problems included (1) internalizing, (2) externalizing scores, using the child behavior checklist (CBCL), and (3) the Conner’s Attention Deficit Hyperactivity Disorder (ADHD) index, all outcomes being discrete raw counts. We applied LASSO penalized negative binomial regression models to identify which exposures were associated with the outcomes, while adjusting for co-exposures. In the 1287 children (mean age 8.0 years), 7.3% had a neuropsychiatric medical diagnosis according to parent’s reports. During pregnancy, smoking and car traffic showing the strongest associations (e.g. smoking with ADHD index, aMR:1.31 [1.09; 1.59]) among the 13 exposures selected by LASSO, for at least one of the outcomes. During childhood, longer sleep duration, healthy diet and higher family social capital were associated with reduced scores whereas higher exposure to lead,
copper, indoor air pollution, unhealthy diet were associated with increased scores. Unexpected decreases in behavioral scores were found with polychlorinated biphenyls (PCBs) and organophosphate (OP) pesticides.

Conclusions: Our systematic exposome approach identified several environmental contaminants and healthy lifestyle habits that may influence behavioral problems in children. Modifying environmental exposures early in life may limit lifetime mental health risk.

1. Introduction

Childhood is a critical stage for the mental health and well-being of individuals with half of all adult neuropsychiatric problems starting by 14 years of age (World health organization, 2019). Yet there are large gaps in what we know about the causes of behaviour problems. Common genetic variants across the genome account for only about 5–25% of behavioral disorder risk in the general population (Uher and Zwicker, 2017) with small effects from thousands of genetic variants that are contingent on complex environmental interaction. Research into the environmental factors for behavioral disorders has found a complex picture of multiple social and physical exposures occurring at different stages of life, in particular during sensitive prenatal and early childhood periods when brain development accelerates (van Os et al., 2010; Rapport et al., 2012). These risk factors include unfavourable urban environment (Vassos et al., 2016), immigration and ethnic minority (Bourque, 2011); obstetric and pregnancy complications (Cannon et al., 2002), and exposure to industrial chemicals such as lead and mercury (Grandjean and Landrigan, 2014) but also protective factors such as stimulating home environments (Bradley et al., 1989) and healthy diets (Cohen et al., 2016) that may benefit brain development and child behavior. However, these factors have often been studied independently and cross-sectionally while the aetiology of behavioral disorders is known to consist of networks involving many elements. These elements are complex to measure since they belong to different domains such as psycho-social, physical and chemical environments, and their effect is often dependent on their timing (e.g. pregnancy vs. adulthood), duration, and to which extent they are repeated exposures over time (Guloksuz et al., 2018).

There is a large body of epidemiological literature on the effects of early-life exposures on behavioural problems in children, but few studies have investigated several families of exposure, or applied an exposome approach—i.e., one encompassing all environmental exposures from conception onwards. Populations are simultaneously exposed to a wide range of environmental factors, some of which are suspected to affect child behaviour, especially during early life. Even in its partial forms, the exposome provides a useful framework to systematically evaluate many associations (Wild, 2005), and may be used to avoid problems of selective reporting, publication bias of only positive results, and, to some extent, confounding by co-exposures, ingrained in the typical one-by-one reporting of associations. Consequently, the exposome may help both in discovery and in setting priorities for prevention.

Recent examples of exposome, or multiple exposure, approaches, in child behavior research, particularly focused on pregnancy exposure to specific exposure families such as indoor air pollutants (Gonzalez-Casanova et al., 2018), persistent organic pollutants (“POPs”, e.g. polychlorinated biphenyls (PCBs), organochlorine (OC) pesticides, polybrominated diphenyl ethers (PBDEs), and perfluoralkyl substances (PFAS) (Lenters et al., 2019) have applied a wider-exposure selection based on available cohort questionnaire-data (Steer et al., 2015). None of the previous studies combined chemical and non-chemical environmental stressors. A recent scoping review on human exposome studies identified mental health to be understudied in exposome research despite its crucial role in the fields of public health, health care, and treatment (Haddad et al., 2019).

Therefore, this study aims to evaluate the association between a wide array of pregnancy and childhood environmental exposures related to chemical and non-chemical environmental stressors and link this to a range of child behavioral symptoms in the large European Human Early-Life Exposome (HELIX) cohort (Vrijheid et al., 2014; Maître et al., 2018).

2. Method

2.1. Study population

The study population is based on the HELIX (Human Early-Life Exposome) project, which pooled data on existing six population-based birth cohorts (BiB in United Kingdom, EDEN in France, KANC in Lithuania, INMA in Spain, MoBa in Norway, and RHEA in Greece) where large sets of previously collected longitudinal data from early pregnancy to childhood were available. Background information on the HELIX cohorts and the full HELIX protocols are described elsewhere (Haug et al., 2018; Maître et al., 2018; Robinson et al., 2018; Tamayo-Uria et al., 2019). The HELIX subcohort consists of 1301 mother-child-pairs that were followed-up at 6–11 years of age using common protocols for all the cohorts (between 12/2013 and 02/2016) including biological samples collection, face-to-face questionnaire, health examination and additional characterization for a wide range of exposures (Vrijheid et al., 2014). The study was approved by the individual national ethics committees for cohort recruitment, follow-up visits and secondary use of pre-existing data for HELIX. Written informed consent was obtained from all HELIX subcohort participants.

2.2. Child emotional and behavioral problems

Parents completed questionnaires related to child’s behavior, including the Conner rating scale’s (N = 1287) and child behavior checklist (CBCL, N = 1298), within a week before the follow-up visit at 6–11 years of age. The 99-item CBCL/6–18 version for school children was used to obtain standardized parent reports of children’s problem behaviours, translated and validated in each native language of the participating six cohort populations (Achenbach and Rescorla, 2001). The parents responded along a 3-point scale with the code of 0 if the item is not true of the child, 1 for sometimes true, and 2 for often true. The internalizing score includes the subscales of emotionally reactive and anxious/depressed symptoms, as well as somatic complaints and symptoms of being withdrawn. The externalizing score includes attention problems and aggressive behaviors. In addition, an ADHD index based on the short form of the Conners’ rating scales of 27 items provided information on inattention and hyperactivity symptoms (Conners, 1997). The internal consistency (Cronbach’s alpha) of each of the study scales was >0.80. All outcomes were analyzed as raw count scores.

2.3. Characterisation of the exposome

A wide range of exposures were evaluated during pregnancy and childhood (6–11 years) as presented in Table 1. This set of exposures was selected at the start of the HELIX project, because they were of concern for more than one of the health outcomes under study and because population exposure was widespread. We assessed five domains of the exposome: the outdoor, indoor, chemical, lifestyle and social domains, representing a total of 88 pregnancy exposures and 123 childhood exposures (Haug et al., 2018; Tamayo-Uria et al., 2019). An extended description on all exposure assessments and methods are provided in Annex 1, and questions are presented elsewhere in previous publications regarding chemical assessment (Haug et al., 2018), questionnaires and general description on all exposure assessments and methods are provided in Annex 1.
Abbreviations used: NO\textsubscript{2}: nitrogen dioxide; PM\textsubscript{2.5}: particulate matter with an aerodynamic diameter of <2.5 \textmu m; PM\textsubscript{10}: particulate matter with an aerodynamic diameter of <10 \textmu m; PM\textsubscript{2.5,abs}: absorbance of PM\textsubscript{2.5} filters; TEX: toluene, ethylbenzene, xylene; DDE: 4,4′-dichlorodiphenyl dichloroethylene; DDT: 4,4′-dichlorodiphenyltrichloroethane; HCB: hexachlorobenzene; PCB: polychlorinated biphenyl – 118, 138, 153, 170, 180; PBDE 47, 2,2,4′,4,5,5′-hexa-bromodiphenyl ether; PBDE 153: 2,2′,4,4′,5,5′-hexa-bromodiphenyl ether; PFOA: perfluorooctanoic acid; PFOS: perfluorooctane sulfonate; As: arsenic; Cd: cadmium; Co: cobalt; Cs: cesium; Cu: copper; Hg: mercury; Mn: manganese; Mo: molybdenum; Pb: lead; Ti: thallium; MEP: monoethyl phthalate; MiBP: mono-iso-butyl phthalate; MnBP: mono-n-butyl phthalate; MBzP: mono benzyl phthalate; MEHP: mono-2-ethyl-hexyl phthalate; MEHHP: mono-2-ethyl-5-hydroxyethyl phthalate; MEDEHP: mono-2-ethyl-5-oxoethyl phthalate; MECCP: mono-2-ethyl-5-carboxyethyl phthalate; OHMNP: mono-4-methyl-7-hydroxyoctyl phthalate; OXOINP: mono-4-methyl-7-oxooctyl phthalate; MEPA: methyl paraben; ETPA: ethyl paraben; BPA: bisphenol A; PRPA: propyl paraben; BUPA: N-butyl paraben; OXBE: oxybenzone; TRCS: triclosan; DMP: dimethyl phosphate; DMTP: dimethyl thiophosphate; DMDTP: dimethyl dithiophosphate; DETP: diethyl phosphate; DETP: diethyl thiophosphate; THMs: trihalomethanes.

Exposure assessment methods for each variable are described in Annex 1.

Exclusion of food contamination due to very high correlation with facility density (r > 0.9).

During childhood, the urine sample analysed was a pool of equal amounts of two spot urine samples collected at bed time the day before and in the morning the day of the clinical examination.

Table 1
List of prenatal and childhood exposures assessed in this study.

| Exposure Group                  | List of variables                                      | Preg. | Child | Where(buffer) - Matrix | Sampling pregnancy period | Sampling childhood period | Method * |
|--------------------------------|--------------------------------------------------------|-------|-------|------------------------|---------------------------|---------------------------|---------|
| **External general environment**|                                                        |       |       |                        |                           |                           |         |
| Atmospheric pollutants         | NO\textsubscript{2}, PM\textsubscript{2.5}, PM\textsubscript{10}, PM\textsubscript{2.5,abs} | 4     | 4     | Home                   | Pregnancy                 | Annual average             | LUR model |
| UV                             | Ambient ultraviolet radiation (UVR) levels             | 0     | 1     | Home(300 m)            | Pregnancy                 | Month                     | Average of satellite images |
| Surrounding natural spaces     | Presence of a major green or blue space in a distance of 300 m, NDVI | 3     | 6     | Home/ School (green and blue 300 m; NDVI 100 m) | Pregnancy | Childhood | Spatial analysis of GIS layers |
| Meteorology                    | Temperature, humidity, pressure                        | 3     | 2     | Home                   | Pregnancy                 | Month                     | Weather stations |
| Built environment              | Population and building density, street connectivity, accessibility and facility richness⁎, walkability | 9     | 15    | Home/ School (300 m)   | Childhood                 |                           | Spatial analysis of GIS layers |
| Road traffic                   | Traffic density and inverse distances to nearest roads  | 3     | 5     | Home/ School (100 m)   | Pregnancy                 | LUR model                 |         |
| Road traffic noise             | 24 h and evening time road noise levels                 | 1     | 3     | Home/ School           | Pregnancy                 | Childhood                 | Local noise map + LUR model |
| **External individual environment** (contaminants) | | | | | | | |
| OCs                            | DDE, DDT, HCB, PCB (118, 138, 153, 170, 180)         | 8     | 9     | Serum/Plasma           | Trim 1                    | Childhood                 | GC-MS/MS |
| PBDE                           | PBDE(47, 153)                                         | 2     | 2     | Serum/Plasma           | Trim 1                    | Childhood                 | GC-MS/MS |
| PFAS                           | PFOA, PFNA, PFUNDA, PFHxS, PFOS, A\textsubscript{c},Cd,Co,Cu,Hg,Mn,Pb | 5     | 5     | Serum/Plasma/ Blood    | Trim 1                    | Childhood                 | LC-MS/SP-E |
| Metals and Elements            | Whole and cord blood                                  | 10    | 11    | Urine⁎                 | Trim 1 and 3              | Childhood                 | Q-ICP-MS/AAS |
| Phthalates                     | MEP, MiBP, MnBP, MiBP, MEHP, MEHHP, MEHP, MEHP, MEHP, MEHHP, MEHHP, MEHHP, MEHHP, OHMNP, OXOINP | 10    | 11    | Urine⁎                 | Trim 1 and 3              | Childhood                 | HPLC-MS |
| Phenols                        | MEPA, ETPA, BPA, PRPA, BUPA, OXBE, TRCS               | 7     | 7     | Urine⁎                 | Pregnancy                 | Childhood                 | HPLC-MS |
| OPs                            | DMP, DMTF, DMTF, DEP, DETP, DEDTP                      | 4     | 5     | Urine⁎                 | Pregnancy                 | Childhood                 | UPLC-MS |
| Tobacco smoking                | Cotinine, questionnaire on active and passive         | 4     | 3     | Urine⁎, questionnaire  | Pregnancy                 | Childhood                 | Immulite® 2000 Nicotine Multivariate + Spatial model |
| Water DBPs                     | THM, Chloroform, Brominated THMs                      | 3     | 0     | Home                   | Pregnancy                 | –                         | Passive samples + Multivariate model |
| Indoor AP                      | NO\textsubscript{2}, TEX, Benzene, PM\textsubscript{2.5}, PM\textsubscript{2.5,abs} | 0     | 5     | Home                   | –                         | 1 week × 2 24 h × 2 |          |
| **Other**                      | Diet, physical activity                               | 12    | 26    | Questionnaires          | Pregnancy                 | Childhood                 | –          |
| Social-economic capital        | Family affluence, social contact, house crowding      | 0     | 4     |                         |                           |                           | –          |

Total: 211

88

123

Abbreviations used: NO\textsubscript{2}: nitrogen dioxide; PM\textsubscript{2.5}: particulate matter with an aerodynamic diameter of <2.5 \textmu m; PM\textsubscript{10}: particulate matter with an aerodynamic diameter of <10 \textmu m; PM\textsubscript{2.5,abs}: absorbance of PM\textsubscript{2.5} filters; TEX: toluene, ethylbenzene, xylene; DDE: 4,4′-dichlorodiphenyl dichloroethylene; DDT: 4,4′-dichlorodiphenyltrichloroethane; HCB: hexachlorobenzene; PCB: polychlorinated biphenyl – 118, 138, 153, 170, 180; PBDE 47, 2,2,4′,4,5,5′-hexa-bromodiphenyl ether; PBDE 153: 2,2′,4,4′,5,5′-hexa-bromodiphenyl ether; PFOA: perfluorooctanoic acid; PFOS: perfluorooctane sulfonate; As: arsenic; Cd: cadmium; Co: cobalt; Cs: cesium; Cu: copper; Hg: mercury; Mn: manganese; Mo: molybdenum; Pb: lead; Ti: thallium; MEP: monoethyl phthalate; MiBP: mono-iso-butyl phthalate; MnBP: mono-n-butyl phthalate; MBzP: mono benzyl phthalate; MEHP: mono-2-ethyl-hexyl phthalate; MEHHP: mono-2-ethyl-5-hydroxyethyl phthalate; MEDEHP: mono-2-ethyl-5-oxoethyl phthalate; MECCP: mono-2-ethyl-5-carboxyethyl phthalate; OHMNP: mono-4-methyl-7-hydroxyoctyl phthalate; OXOINP: mono-4-methyl-7-oxooctyl phthalate; MEPA: methyl paraben; ETPA: ethyl paraben; BPA: bisphenol A; PRPA: propyl paraben; BUPA: N-butyl paraben; OXBE: oxybenzone; TRCS: triclosan; DMP: dimethyl phosphate; DMTP: dimethyl thiophosphate; DMDTP: dimethyl dithiophosphate; DETP: diethyl phosphate; DETP: diethyl thiophosphate; THMs: trihalomethanes.

⁎ Excluded from childhood exposure due to very high correlation with facility density (r > 0.9).

Exposure assessment methods for each variable are described in Annex 1.
biological samples from the mother during pregnancy and the child. The chemical list was based on concern for child health and included organochlorine compounds (PCBs and OC pesticides), PBDEs, PFAS, metals and essential elements, phthalate metabolites, phenols (triclosan, oxybenzone, parabens and bisphenol A), organophosphate (OP) metabolites, and cotinine. When appropriate, the concentrations were adjusted for lipids or creatinine. Lifestyle factors were collected by questionnaire and included smoking habits, diet, physical activity, allergens, and sleep. During childhood, questionnaire information was collected on socio-economic capital of the family based on the Family Affluence Score (FAS, Boyce et al., 2006) and through summary variables for social participation, social contact and house crowding.

2.4. Statistical analysis

Exposure variables were transformed to achieve normality; when normality could not be achieved with a transformation, the variable was categorized, and then standardized by the inter-quartile range (IQR) (Hernandez-Ferrer et al., 2019). Missing exposure and covariate data were imputed using chained equations (White et al., 2011) with the mice package in R (van Buuren and Groothuis-Oudshoorn, 2011). Each imputation models included between 5 and 25 variables selected based on their correlation to the variable to impute, their proportion of missing and the main health outcomes. (20 imputed datasets were created, see more details in Annex 2). Missing values, ranged from no missing values for some child phthalate metabolites to 65% for fast-food intake during pregnancy (Table A2.1 in Tamayo-Uria et al., 2019). The mean percentage of missing values per exposure was 12% (first quartile 0.9% and third quartile 16.8%). None of the participants had complete data on all exposures. Yet, for 98% of individuals <30% of exposure variables had missing values. Results of the complete case analysis are also available in Tables A4.2 and A4.3 with the number of complete cases for each exposure before imputation. In all subsequent regression models, Rubin’s rules were used to combine the results from the 20 imputed datasets, which allows the quantification of the uncertainty in results associated with imputation in the final standard errors, confidence intervals and p-values (White et al., 2011).

All analyses were based on negative binomial regression with adjustment for cohort, maternal age, child age, child gender, season of conception and maternal education. To explore separately the associations between the prenatal and childhood exposures with externalizing, internalizing behavior scores and ADHD index we applied two methods. Our primary analysis relied on a maximum likelihood function plus a penalty, the least absolute shrinkage and selection operator (LASSO) to account for the complex correlation pattern of the exposome and to mitigate potential estimation problems due to collinearity (multi-exposure models, see Annex 2 for more details). We determined the overall penalty parameter, lambda, by maximizing the prediction log-likelihood using 10-fold cross-validation and providing the maximum mean cross-validated log-likelihood values. This type of variable selection was shown to be more robust than the traditional stepwise variable selection and also well adapted to count data with overdispersion as observed with CBCL and ADHD index data (Wang et al., 2016). For descriptive and comparative purposes, we also estimated associations by modeling one exposure at the time (single-exposure model, also called exposome-wide association study (ExWAS) analysis). To account for multiple comparisons in the single exposure models, a family-wise error rate was calculated to provide an indicative p-value threshold (5% divided by the effective number of tests, based on Li et al. (Li et al., 2012)).

The results are presented as adjusted mean ratios (aMR) with corresponding 95% confidence intervals (CI). In all of these models, the exposure estimate was reported for one interquartile range (IQR) increase in exposure levels—for example, a ratio of mean score of 1.25 for an IQR increase in the exposure meant that those with an exposure at the 75th percentile had a 25% higher mean score, compared with those with an exposure at the 25th percentile.

2.5. Sensitivity analyses

Three sets of sensitivity analyses were performed to address the robustness of our results for the multi-exposure models: 1) further adjustment for child standardized body mass index (BMI), breastfeeding duration, prenatal smoking (only postnatal models) and maternal perceived stress (Cohen et al., 1988) (although some of these factors could act as mediators in the association between the exposure and child behavior); 2) evaluating the impact of outliers in exposure values and the linearity of the association between exposure and outcome were checked using generalized additive models (GAMs); 3) stratifying the models by cohort and testing for heterogeneity.

3. Results

3.1. Study population

At the time of delivery, pregnant women were aged 31 years on average, mostly highly educated (52%) and were a native of the cohort country (84%) (Table 2). Children were 45% girls and 55% boys. At the time of behavior assessment, children were on average 8 years old (range: 6.5 to 11 years). 3.9% regularly visited the psychologist and 7.3% had a neuropsychiatric diagnosis by the time of visit (according to parent’s reports, besides the CBCL screening). Boxplots with exposure level distribution (untransformed) by cohort are available online (Tamayo-Uria et al., 2019 Fig. A2.1 and Fig. A2.2).

3.2. Are pregnancy exposures associated with children behavioral symptoms?

Out of the 88 pregnancy exposures, few were associated with behavioral symptoms in the main models adjusting for LASSO selected co-exposures (Table 3) or in the single exposure models as illustrated in Fig. 1 (eTable 3.4). In our main multiple exposure models, pregnancy tobacco exposure was associated with increased externalizing symptoms (Table 3); compared to non-smoking, passive smoking was associated with a 25% increase (adjusted mean ration, aMR = 1.25[1.09; 1.44]) and active smoking with a 31% increase (aMR = 1.31[1.09; 1.59]) in externalizing symptoms. Results were similar for internalizing symptoms. Residential car traffic density on nearest road was associated with increased externalizing symptoms (aMR = 1.07[1.00–1.14]) and ADHD index (aMR = 1.10[1.03; 1.17]). The presence of green space near the home address (within 300 m) was associated with externalizing symptoms only (aMR = 1.15[1.01–1.31]; Table 3) and not significant in the single-exposure model (aMR = 1.12[0.98; 1.28]).

Other exposures that were selected by LASSO, but with confidence intervals including 1 (Table 3), were meat intake (increased internalizing and externalizing symptoms), bisphenol A and DMP exposures, an OP pesticide metabolite (increased externalizing symptoms). Lower ADHD scores were observed in children whose mothers had low fish intake during pregnancy (<2 times a week) compared to 2–4 times a week.

3.3. Are childhood exposures associated with children behavioral symptoms?

Out of 123 childhood exposures, 11 exposures were selected in multi-pollutant penalized models for internalizing symptoms, 9 for externalizing symptoms, and 17 for the ADHD index (Table 4). In the single exposure models, as illustrated in Fig. 1 (eTable 3.4), only few exposures passed multiple correction testing. Exposures that were selected by LASSO and with a confidence interval not including 1 (Table 3) were: indoor air PM2.5 and high ready-made food intake, associated with increased externalizing symptoms; second-hand smoking, intake of bakery products, and copper exposure, associated with increased internalizing symptoms; and second-hand smoking, ready-made food,
sweets, caffeinated drinks and lead exposure, associated with an increased ADHD index. A few chemical exposures were also associated with decreased symptoms (Table 4): the sum of PCBs was associated with all outcomes (e.g. for internalizing 0.84 [0.77–0.93]) and DMTP with ADHD index (0.92 [0.86–1.00]). Among exposures associated with reduced behavioral symptoms, sleep duration was associated with all the outcomes. Children sleeping one hour more per night (IQR: 0.93 h) had 11% lower internalizing symptoms score. Family social participation to organizations was associated with reduced internalizing symptoms, contact with family and friends (daily (almost) vs. less than once a week) and house crowding, i.e. number of persons living in the household, were associated with lower ADHD mean scores. The healthy diet score (KIDMED), was associated with decreased externalizing scores (4).

### 3.4. Sensitivity analyses

We observed similar findings after adjusting childhood models for breastfeeding duration, child’s BMI, perceived stress score of the mother at the time of child behavioral rating and pregnancy tobacco exposure (only tested for childhood exposome models) as presented in eTable 4.1. Adjustment for perceived maternal stress slightly attenuated the association between pregnancy tobacco smoking exposure with internalizing and externalizing symptoms (e.g. 1.31 [1.09; 1.59] vs. 1.19 [0.99;1.44] for active tobacco exposure with externalizing scale).

Most of the associations were consistent across cohorts, but some heterogeneity can be highlighted (i.e. $I^2 > 60$%, eTable 4.2): 1) the association between pregnancy copper was driven by the BiB and KANC cohorts; 2) the association between child caffeinated drink consumption and ADHD was driven by the EDEN cohort (eFig. 4.1).

Linearity of the associations was observed for all exposures except for pregnancy levels of bisphenol A, PFUnDA, PFOA with externalizing scale (non monotonic inverted U-shaped curves), for traffic density on nearest road with ADHD index (eFig. 4.3) and for child levels of PCB 180 with internalizing scale and ADHD (monotonic multiphasic curve, eFig. 4.2) and for the KIDMED score with externalizing scale (non monotonic U-shaped curve).

### 4. Discussion

Our study is the first to comprehensively and systematically evaluate a wide range of exposures associated with behavioral problems, covering social, physical and chemical exposures occurring at different stages of development. Our results confirm the important harmful role of maternal smoking during pregnancy in multiple behavior symptoms in children, but also highlight the potential protective role of a healthy family lifestyle during childhood (diet, sleep, regular social contact). We report new evidence about the association between car traffic density, indoor air pollution and copper with increased behavioral problems in children.

Maternal tobacco smoke exposure during pregnancy was the most important prenatal exposure related to behavioral and emotional problems in children. The adverse effects of active maternal smoking during pregnancy are well established, including adverse birth outcomes and increased respiratory illness in children (Kramer, 1987; Burke et al., 2012). However, reports on adverse behavioral outcomes are inconsistent (Rice et al., 2018). Previous studies demonstrated that maternal smoking exposure is closely linked to other co-exposures such as parental psychopathology symptoms, socio-economic factors, father smoking habits and the home environment (quality of attachment, support, and stimulation that a child is exposed to at home) which may account for a large part of the effect of maternal smoking during pregnancy on child behavior (Roza et al., 2009; Hopson et al., 2016). In our study, we found that maternal perceived stress at the time of rating explained some of the association between pregnancy tobacco exposure and child behavior, and we cannot entirely rule out a role of other, unmeasured confounders.

We also found that increased residential car traffic density on the nearest road during pregnancy, was associated with higher externalizing and ADHD scores. This exposure might represent an umbrella of other factors based on its correlation with other exposures in our dataset such as outdoor air pollutants (PM absorbance: r(spearman) = 0.4) and traffic

### Table 2

Description of the Study Population.

| Parental characteristics | N (%) or median (quartile range) |
|--------------------------|--------------------------------|
| **Cohort of inclusion**  |                                |
| BiB, United Kingdom      | 205 (15.9)                     |
| EDEN, France             | 197 (15.3)                     |
| INMA, Spain              | 218 (17)                       |
| KANC, Lithuania          | 202 (15.7)                     |
| MoBa, Norway             | 272 (21.2)                     |
| Rhea, Greece             | 192 (14.9)                     |
| **Family native from the country of the cohort** | |
| Both native parents      | 1056 (84)                      |
| 1 native parent          | 62 (4.9)                       |
| No native parent         | 139 (11.1)                     |
| Maternal age at inclusion, yrs | 31 (27.2–34)                   |
| **Maternal education**   |                                |
| Primary                  | 171 (13.8)                     |
| Secondary                | 429 (34.5)                     |
| Higher                   | 643 (51.7)                     |
| **Paternal education**   |                                |
| Primary                  | 207 (17.3)                     |
| Secondary                | 465 (38.8)                     |
| Higher                   | 526 (43.9)                     |
| **Marital status**       |                                |
| Cohabitant               | 183 (14.2%)                    |
| Divorced/separated       | 81 (6.3%)                      |
| Married                  | 958 (74.5%)                    |
| Single                   | 54 (4.2%)                      |
| Other                    | 8 (0.6%)                       |
| **Trimester of conception** |                             |
| Jan-March                | 409 (32.1)                     |
| April-June               | 257 (20.2)                     |
| July-Sept                | 275 (21.6)                     |
| Oct-Dec                  | 332 (26.1)                     |
| **Who completed the behavioural questionnaires?** | |
| Father                   | 63 (4.9%)                      |
| Mother                   | 1215 (94.5%)                   |

| **Child Characteristics** |                                |
| Child’s ethnicity        |                                |
| Caucasian                | 1155 (89.9)                    |
| Non-Caucasian            | 130 (10.1)                     |
| Child’s BMI category*    |                                |
| Thinness                 | 100 (7.8)                      |
| Normal wt                | 929 (72.7)                     |
| Overweight               | 180 (14.1)                     |
| Obese                    | 69 (5.4)                       |
| Sex                      |                                |
| Female                   | 580 (45.1)                     |
| Male                     | 706 (54.9)                     |
| Age (years)              | 8.05 (6.4–8.9)                 |
| Visit psychologist regularly |                            |
| No                       | 1235 (96.0%)                   |
| Yes                      | 50 (3.9%)                      |
| Any previous neuropsychological diagnosis | |
| No                       | 1191 (92.6%)                   |
| Yes                      | 94 (7.3%)                      |
| Internalizing scales     | 5 (2–9)                        |
| Externalizing scales     | 5 (2–10)                       |
| ADHD Index (Conners)     | 6 (2–11)                       |

*BIB = Born in Bradford; EDEN = Etude des Déterminants pré et postnatals du développement et de la santé de l’Enfant; INMA = INfancia y Medio Ambiente; KANC = Kaunas Cohort; MoBa = Norwegian Mother, Father and Child Cohort Study; Rhea = Mother-Child Cohort in Crete.

According to the WHO growth reference for children age between 5 and 19 years. World Health Organization. Growth reference 5-19 years. Available at: http://www.who.int/growthref/who2007_bmi_for_age/en/. Accessed August 2, 2019.
The presence of green space (300 m to residence) during pregnancy was associated with child neurobehavioral disorders. This result is not in line with previous studies which tend to find that greater access to or quantity of neighborhood nature or public open space were associated with increased externalizing scale. Finally, the associations between pregnancy exposures and change in behavioral problems, showing exposures selected into the multiple-exposure models (out of 88 candidate exposures, N = 1287).

Table 3

| Exposure (IQR or reference category) | Exposure family | Internalizing scale | Externalizing scale | ADHD index |
|-------------------------------------|-----------------|---------------------|---------------------|------------|
|                                     |                 | aMR [95%CI]         | p-value             | aMR [95%CI] | p-value |
| Traffic density on nearest road     | Road traffic    | 1.07 [1.00; 1.14]   | 0.050               | 1.10 [1.03; 1.17] | 0.004 |
| (3598 vehicles/day)                 |                 |                     |                     |            | |
| Green spaces (300 m)                | Natural spaces  | 1.15 [1.01; 1.31]   | 0.034               |            | |
| Meat intake                         | Lifestyle/Diet  | 0.96 [0.84; 1.10]   | 0.578               | 0.62 [0.87; 1.20] | 0.808 |
| Medium (vs. low)                    |                 |                     |                     |            | |
| High (vs. low)                      |                 | 1.11 [0.97; 1.27]   | 0.131               | 1.16 [0.99; 1.36] | 0.060 |
| Fish and seafood                    | Lifestyle/Diet  |                     |                     |            | |
| Medium (vs. low)                    |                 | 0.96 [0.84; 1.10]   | 0.578               | 0.62 [0.87; 1.20] | 0.808 |
| High (vs. low)                      |                 | 1.11 [0.97; 1.27]   | 0.131               | 1.16 [0.99; 1.36] | 0.060 |
| Fastfood intake                     | Lifestyle/Diet  |                     |                     |            | |
| Moderate physical activity, t3      | Lifestyle/Physical activity | 0.91 [0.78; 1.07] | 0.253             | 1.09 [0.91; 1.29] | 0.347 |
| Very often (vs. none or sometimes)  | Tobacco smoking | 1.10 [0.94; 1.29]   | 0.244               | 0.93 [0.78; 1.11] | 0.406 |
| Smoking in pregnancy                |                 |                     |                     |            | |
| Second-hand smoking (vs. none)      | Tobacco smoking | 1.28 [1.12; 1.45]   | <0.001              | 1.25 [1.09; 1.44] | 0.002 |
| Active smoking (vs. none)           |                 | 1.20 [1.03; 1.39]   | 0.020               | 1.31 [1.09; 1.59] | 0.005 |
| Cotinine (µg/L)                     | Tobacco smoking | 1.05 [0.96; 1.15]   | 0.262               |            | |
| 18.4-50 (vs. < 18.4)                |                 |                     |                     |            | |
| 50 (vs. > 18.4)                     |                 |                     |                     |            | |
| Bisphenol-A (4.96 µg/L)             | Phenol          | 1.06 [0.98; 1.15]   | 0.152               |            | |
| Caesium (0.97 µg/L)                 | Metal           | 0.94 [0.83; 1.06]   | 0.281               |            | |
| Dimethyl phosphate (11.37 µg/g)     | Organophosphate pesticides | 1.07 [1.00; 1.15] | 0.067             |            | |
| Perfluorooctanoate (1.96 µg/L)      | Perfluoralkyl substances (PFAS) | 0.95 [0.86; 1.05] | 0.301           |            | |
| Perfluoroundecanoate (0.18 µg/L)    | Perfluoralkyl substances (PFAS) | 0.93 [0.84; 1.03] | 0.162             |            | |

The selection of exposures was determined by the least absolute shrinkage and selection operator (LASSO) algorithm applying log link function. The change in behavioral problems was modelled with negative binomial regression which model the ratio of the mean score for one unit change in the exposure (the coefficients obtained needed to be exponentiated). The results are presented as adjusted mean ratios (aMR) with corresponding 95% confidence intervals (CIs). In all of these models, the exposure estimate was reported for one interquartile range (IQR) increase in exposure levels—for example, a ratio of mean score of 1.25 for an IQR increase in the exposure mean that those with an exposure at the 75th percentile had a 25% higher mean score, compared with those with an exposure at the 25th percentile. These coefficients are adjusted for the other exposures and for cohort, maternal age, maternal education level, maternal pre-pregnancy body mass index, parity, parental country of birth, child age, child sex, and child height. Abbreviations: CI = confidence interval; IQR, interquartile range; cat., categories.

Related noise level (r = 0.17). Previous studies have demonstrated that traffic-related pollution was associated with child neurobehavioral disorders such as inattention and higher prevalence of ADHD and Autism Spectrum Disorder (Suades-González et al., 2015). Traffic-related airborne particulate matter (PM) is a complex and heterogeneous mixture that includes residues from fossil fuel combustion, organic chemicals, trace metals, nitrate, and sulphate. During gestation, PM adverse effects may be provoked in the fetus through various mechanisms including both direct (e.g., PM placenta translocation) and/or indirect (e.g., maternal systemic and intrauterine inflammation) and/or direct (e.g., PM placenta translocation) manners, with recent evidence for translocation of black-carbon PM in human placenta (Bové et al., 2019). Therefore, an etiological role for traffic-related pollution during pregnancy in child behavioural disorders is biological plausible, although exact mechanisms in humans remains elusive. Finally, the presence of green space (300 m to residence) during pregnancy was associated with increased externalizing scale. This result is not in line with current studies which tend to find that greater access to or quantity of neighborhood nature or public open space were associated with better mental health (Alderton et al., 2019). In our study, there may be some residual negative confounding effect in relation to different geo-localization patterns according to socio-economic position amongst the European countries, for example, MoBa sub-cohort sample showed higher greenness degree in lower social classes, an opposite pattern to other HELIX sub-cohorts (Robinson et al., 2018).

Postnatal tobacco exposure and car traffic density, were not as strongly associated with child behavior as pregnancy measurements, although the probability to be exposed at both periods for a child was high. Our results may indicate that the pregnancy period is most sensitive to the harmful effects of these exposures which is in line with the very rapid development of the nervous system during this time window but also the different ways individuals are exposed, for example in utero and/or passively to tobacco smoke (Peterson et al., 2015; Allen et al., 2017). In line with our results, childhood second-hand smoke (SHS) was not related to blood methylation in HELIX children (biomarkers of past exposure), indicating much weaker effects of recent SHS with respect to active maternal smoking in pregnancy (Vives-Usano et al., 2020).

Among childhood associations, we observed that healthy dietary habits (Mediterranean diet adherence score) and longer hours of sleep had protective effects on internalizing problems. Similarly, an unhealthy diet (readymade food, sweets, and caffeinated drinks) was associated with higher risk of ADHD symptoms. These results and previous studies support the hypothesis that an inadequate micronutrient intake, which may result from a Western dietary pattern, could cause suboptimal brain function in children (Howard et al., 2011). Impulsivity traits in children with ADHD may also lead to poor dietary choices and emotional eating. As previously found, young adults who displayed greater impulsivity were found to be more likely to choose snack food when hungry, and hence consume more calories, than were less impulsive participants (Nederkoorn et al., 2009). However, diet modification, e.g. reducing carbohydrate intake and as a result obesity, has been demonstrated to improve symptoms of ADHD patients (Pelsser et al., 2017). Furthermore, one of the strongest association with ADHD was in relation to the social
Fig. 1. Exposome-wide associations (ExWAS) with child internalizing and externalizing scales (single-exposure models). On the left panel are the results as dot plots for the pregnancy exposome and the right panel the cross-sectional childhood exposome. The direction of the association is indicated by the color of the text label and the dot contour (blue, negative and red, positive associations). The horizontal red line across the plots represents the multiple testing threshold correction (based on effective number of tests). Any exposures above this line are considered significant. All the exposures with a p-value below 0.05 are labeled. The 0.05 p-value threshold is also represented by a grey line across the plots. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
### Table 4

Associations between childhood exposures and change in behavioral problems, showing exposures selected into the multiple-exposure models* (out of 123 candidate exposures, N = 1287).

| Childhood exposure (IQR or reference category) | Exposure family | Internalizing scale | Externalizing scale | ADHD index |
|-----------------------------------------------|-----------------|---------------------|--------------------|------------|
|                                               |                 | aMR [95%CI]         | p-value            | aMR [95%CI] | p-value |
| Road traffic load (100 m)                     | Road traffic    |                     |                    | 1.08 [0.98; 1.18] | 0.123 |
| (1138814 vehicles/day m)                      | Indoor air      | 1.06 [0.99; 1.13]   | 0.121              | 1.09 [1.03; 1.16] | 0.005 |
| Indoor benzene (0.99 µg/m³)                   | Indoor air      | 0.98 [0.94; 1.02]   | 0.300              | 1.05 [0.94; 1.18] | 0.395 |
| Indoor PM2.5 (6.5 µg/m³)                      | Tobacco smoking | 1.13 [1.01; 1.26]   | 0.034              | 1.08 [0.95; 1.23] | 0.217 |
| Second-hand smoking                           | Tobacco smoking | 1.09 [1.03; 1.26]   | 0.297              |            |        |
| Cotinine                                      | Tobacco smoking |                     |                    |            |        |
| Total hours of sleep                          | Lifestyle       | 0.89 [0.83; 0.96]   | 0.001              | 0.91 [0.85; 0.99] | 0.027 |
| Bakery products                               | Lifestyle/Diet  | 1.00 [0.88; 1.13]   | 0.997              | 1.03 [0.90; 1.18] | 0.649 |
| Medium (vs. low)                              | Lifestyle/Diet  | 1.18 [1.03; 1.36]   | 0.017              | 1.15 [0.98; 1.34] | 0.081 |
| High (vs. low)                                | Lifestyle/Diet  |                     |                    |            |        |
| Dairy products                                | Lifestyle/Diet  | 0.98 [0.87; 1.12]   | 0.060              | 0.98 [0.87; 1.12] | 0.060 |
| Meat intake                                   | Lifestyle/Diet  | 0.99 [0.87; 1.11]   | 0.030              | 0.98 [0.87; 1.11] | 0.030 |
| High (vs. low)                                | Lifestyle/Diet  | 1.11 [0.99; 1.25]   | 0.067              |            |        |
| Ready made food                               | Lifestyle/Diet  |                     |                    |            |        |
| Medium (vs. low)                              | Lifestyle/Diet  | 0.89 [0.79; 1.01]   | 0.060              | 0.90 [0.79; 1.01] | 0.060 |
| High (vs. low)                                | Lifestyle/Diet  | 0.98 [0.87; 1.12]   | 0.006              | 0.98 [0.87; 1.12] | 0.006 |
| Fastfood                                      | Lifestyle/Diet  |                     |                    |            |        |
| Medium (vs. low)                              | Lifestyle/Diet  | 0.99 [0.87; 1.11]   | 0.018              | 0.99 [0.87; 1.11] | 0.018 |
| High (vs. low)                                | Lifestyle/Diet  | 1.11 [0.99; 1.25]   | 0.067              |            |        |
| Fish and seafood                              | Lifestyle/Diet  |                     |                    |            |        |
| Medium (vs. low)                              | Lifestyle/Diet  | 1.00 [0.86; 1.16]   | 0.974              | 1.00 [0.86; 1.16] | 0.974 |
| High (vs. low)                                | Lifestyle/Diet  |                     |                    |            |        |
| Healthy diet (KIDMED score)                   | Lifestyle/Diet  | 0.93 [0.87; 0.99]   | 0.034              | 0.93 [0.87; 0.99] | 0.034 |
| Lead (5.91 µg/L)                              | Metal           | 1.06 [1.00; 1.13]   | 0.061              | 1.06 [1.00; 1.13] | 0.061 |
| Copper (350 µg/L)                              | Metal           | 1.10 [1.03; 1.17]   | 0.002              | 1.10 [1.03; 1.17] | 0.002 |
| Dimethylphosphite (4.89 µg/g)                 | OP pesticides   | 0.91 [0.85; 0.97]   | 0.000              | 0.92 [0.86; 0.98] | 0.013 |
| Sum of polychlorinated biphenyls (27.6 ng/g lipids) | OC compounds | 0.89 [0.82; 0.98]   | 0.016              | 0.93 [0.84; 1.02] | 0.131 |
| Dichlorodiphenyldichloroethylene, DDE (34.0 ng/g lipids) | OC compounds | 0.93 [0.84; 1.02]   | 0.131              | 0.93 [0.84; 1.05] | 0.259 |
| Polybrominated diphenyl ether 47 (0.21 µg/g) lipids | OC compounds | 1.04 [1.00; 1.09]   | 0.050              |            |        |
| Monoethyl phthalate (59.84 µg/g)              | Phtalates       |                      |                    |            |        |
| Social participation                           | Social capital  | 0.98 [0.88; 1.09]   | 0.775              | 0.98 [0.88; 1.09] | 0.775 |
| Social participation                           | Social capital  | 0.83 [0.71; 0.97]   | 0.016              |            |        |
| Contact with family and friends               | Social capital  |                      |                    |            |        |
| Once a week (vs. less than once a week)       |                 | 0.79 [0.63; 1.00]   | 0.051              | 0.69 [0.55; 0.87] | 0.002 |
| Almost daily (vs. less than once a week)      |                 | 0.69 [0.55; 0.87]   | 0.002              |            |        |
| House crowding                                |                 | 0.89 [0.85; 0.94]   | <0.001             |            |        |

Abbreviations: CI = confidence interval; IQR, interquartile range; cat., categories.

* The selection of exposures was determined by the least absolute shrinkage and selection operator (LASSO) algorithm applying log link function.

** The change in behavioral problems was modelled with negative binomial regression which model the ratio of the mean score for one unit change in the exposure (the coefficients obtained needed to be exponentiated). The results are presented as adjusted mean ratios (aMR) with corresponding 95% confidence intervals (CIs). In all of these models, the exposure estimate was reported for one interquartile range (IQR) increase in exposure levels—for example, a ratio of mean score of 1.25 for an IQR increase in the exposure meant that those with an exposure at the 75th percentile had a 25% higher mean score, compared with those with an exposure at the 25th percentile. The results are presented as adjusted mean ratios (aMR) with corresponding 95% confidence intervals (CIs).
capital of the parent (mostly of the mother), indeed children whose parent had contact with family or friends less than once a week had a 31% greater probability to have children with ADHD symptoms than average. This is an interesting novel finding in the literature among healthy children, in which the ‘psychosocial exposome’ is important in order to improve child behavior. When parents have strong ties with other families or friends, parents may receive helpful social support and feel less isolated and stressed. In addition, other adults provide additional social control and collective socialization, resulting in children less likely to develop behavioral problems (Bussing et al., 2015).

Another possible explanation is that parents’ ratings of their child’s social skills is often positively associated with ratings of their own social skills (Mikami et al., 2010) and ADHD is generally considered a highly hereditable trait (~70–80%) (Paranoa and Larsson, 2019).

We also confirmed established associations such as with lead. Since the removal of lead from fuel, a sharp decrease in blood lead levels worldwide has been observed (Landrigan, 2002), however, lead effects may occur at very low dose levels (Needleman and Bellinger, 1991) and potential new sources of exposures need to be considered, such as lead recycling activities (World Health Organization (WHO), 2017: The toxic truth, 2021). In a previous exposome study, lead was the only exposure adversely influencing children’s language and cognition among many other metal and lifestyle exposures, and so consistently across three European countries (Calamandrei et al., 2020). Passive smoking and home indoor air pollution (PM2.5) exposures were also associated with increased internalizing and externalizing scales, respectively. These exposures are known as potential hazardous agents affecting neurodevelopment and cognitive function through enhancing pro-inflammatory reactions in the brain (Morales et al., 2009). Finally, child copper (Cu) exposure was associated with higher internalizing symptoms. Cu is an essential trace element required for numerous biological processes, including the proper development and functioning of the central nervous system (De Bie et al., 2007). Among the body organs, the brain is the most copper-rich. In line with the numerous Cu biological functions, we found in HELIX that child Cu was associated with multiple child health outcomes (Agier et al., 2019; Vrijheid et al., 2020; Cadiou et al., 2020; Warembourg et al., 2019) and with molecular omics signatures associated with immune response, lipid storage and sequencing of metal ions (Maire et al. in preparation). Cu (and zinc) has been a candidate biomarker in mood disorders because of its fundamental role in the oxidative and nitrosative stress and consequent activation of the inflammatory response and acute phase proteins such as ceruloplasmin (a key protein involved in the storage of copper) (Styczyn, 2016).

Some childhood exposures, such as PCBs and OP pesticides, were unexpectedly associated with better child behavior. These results highlight a more complex exposure system than expected. Child DMTP was associated with decreased externalizing scale and ADHD score in the childhood exposome but pregnancy DMP was associated with increased externalizing scale (1.07 [1.00; 1.15]). Both DMTP and DMP are OP dialkylphosphate (DAP) metabolites that arise from exposure to OP pesticides, a class of insecticides widely used throughout the world, still thought in agriculture for insect control on food crops and mainly found in the general population through ingestion of residues on food products. Intake of fruits and vegetables is a main determinant of OP urinary metabolites in children and pregnant women, as demonstrated in the HELIX cohorts (Papadopoulou et al., 2019). This suggests that the observed association between cross-sectional child DMTP level and ADHD is likely to arise from a higher fresh fruit and vegetable intake which is beneficial for child mental health (Cohen et al., 2016), but also likely to represent a general healthier lifestyle (less ReadyMade meals and fastfood) and more generally a better home environment. Additionally, the cross-sectional nature of the measurements might further explain this association because child behaviour can influence child eating habits (e.g. ADHD and sweet intake). However, we also observed a trend for an association between pregnancy DMP and child externalizing scale. This association is in line with previous studies indicating the detrimental role of OP pesticides on brain development, when the exposure occurs in utero (Philippat et al., 2018). Another unexpected association was between PCB 170 and 180 during childhood and a decrease in all behavioural outcomes, suggesting a protective effect on behaviour. PCBs, among other persistent organic pollutants (POPs), are considered developmental neurotoxicants, mainly based on studies looking at the association between in utero exposure and neuropsychological development (Grandjean et al., 2014). Results with postnatal exposures (in 0–2 years olds exposed through breast milk) have been inconsistent (Lenters et al., 2019; Forns et al., 2016) and no studies were found on school-age PCB exposure for comparison. In our data, when stratifying the population by child BMI categories, the protective effect of PCB 180 with externalizing scale was mainly seen among obese children (0.92 [0.82;1.03]), but opposite in overweight and normal weight children (1.11 [0.88;1.41]). PCBs are highly lipophilic (they store in fat tissue), and the amount of body fat and growth trajectories in children are expected to affect the toxicokinetics of POPs (Vrijheid et al., 2020). Therefore, residual confounding in PCB-behaviour associations may be present and these results should be interpreted with caution.

4.1. Strengths and limitations

This study used standardized outcome and exposure measurement methods across six different countries with a wide range of exposure measurement techniques such as geographic information systems and satellite data for collecting outdoor exposure (road traffic, meteorological factors, natural spaces) and sensors (indoor air pollution). These exposure assessments were completed by more traditional methods such as standardized questionnaires for lifestyle and social capital variables and targeted biomonitoring for selected known (e.g. heavy metals and OP pesticides) and less known neurotoxic chemicals (e.g. phthalates and parabens). In addition, this prospective birth cohort design allowed the collection of objective prenatal exposure measures free of recall and response biases.

Limitations of this study include exposure misclassification, which is expected to be differential across exposures (e.g., stronger for the least persistent chemicals and for exposures based on modelling of outdoor levels). The cross-sectional aspect of the childhood exposures particularly limited our interpretation of results for lifestyle (sleep, diet) because they may suffer from reverse causation bias, whereby children’s health symptoms would influence sleep duration and dietary choices, instead of vice versa. For other cross-sectionally assessed exposures, the temporality of exposure and relationship with the outcome is more complex. For exposure misclassification, we used multiple exposure methods across six different countries with a wide range of exposure measurement techniques such as geographic information systems and satellite data for collecting outdoor exposure (land use, meteorological factors, natural spaces) and sensors (indoor air pollution). These exposure assessments were completed by more traditional methods such as standardized questionnaires for lifestyle and social capital variables and targeted biomonitoring for selected known (e.g. heavy metals and OP pesticides) and less known neurotoxic chemicals (e.g. phthalates and parabens). In addition, this prospective birth cohort design allowed the collection of objective prenatal exposure measures free of recall and response biases. Other cross-sectionally assessed exposures, the temporality of exposure and relationship with the outcome is more complex.
selection method which aims to optimally predict the outcome – has the advantage that it mutually adjusts for the effect of other exposures, but will tend to select only one predictor in the case that a true predictor is highly correlated with other exposures (e.g., within an exposure family) (Agier et al., 2016). Overall the estimates obtained in the multiple exposure models were similar to the ones obtained in the single-exposure models (i.e., ExWAS). However, changes in estimates (>5%) were observed in the multiple-exposure model for the following pregnancy exposures: active smoking (aMR single- vs. multi-exposure models = 1.21 vs. 1.31, externalizing scale) and fish consumption low vs medium intake (0.98 vs. 1.15), and for the following childhood exposures: bakery products (0.88 vs. 0.93), readymade food (1.18 vs. 1.25), childhood caffeinated drinks (1.21 vs. 1.14), fastfood (1.26 vs. 1.11), DDE (0.89 vs. 0.95) and PCB 180 (0.85 vs. 0.91) with ADHD index, possibly due to residual confounding in the single-exposure models. Correlations between exposures are known to present a challenge for exposome research. Statistical methods similar to the penalized models we used have been shown to provide a good compromise between minimizing false positives and maximizing the selection of truly associated exposures in comparison with other approaches, although in a context of many correlated exposures all methods have a limited ability to efficiently differentiate true predictors from correlated covariates (Agier et al., 2016).

5. Conclusion

In this first study of many environmental risk factors, child behavior problems were influenced by an array of early life environmental exposures in particular prenatal tobacco smoke and car traffic density. During childhood, a healthy lifestyle including healthy dietary habits, longer hours of sleep and a rich social capital of the family were protective, whereas indoor air pollution, copper and blood lead levels were associated with increased symptoms. Promoting early on healthy family habits at individual level and regulating air quality, lead exposure at population level could help improve people’s exposome and prevent the future development of mental health disorders.

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.2021.106523.

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