

Genome-Wide DNA Methylation in Peripheral Blood and Long-Term Exposure to Source-Specific Transportation Noise and Air Pollution: The SAPALDIA Study

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BACKGROUND: Few epigenome-wide association studies (EWAS) on air pollutants exist, and none have been done on transportation noise exposures, which also contribute to environmental burden of disease.

OBJECTIVE: We performed mutually independent EWAS on transportation noise and air pollution exposures.

METHODS: We used data from two time points of the Swiss Cohort Study on Air Pollution and Lung and Heart Diseases in Adults (SAPALDIA) from 1,389 participants contributing 2,542 observations. We applied multiexposure linear mixed-effects regressions with participant-level random intercept to identify significant Cytosine-phosphate-Guanine (CpG) sites and differentially methylated regions (DMRs) in relation to 1-y average aircraft, railway, and road traffic day-evening-night noise (Lden); nitrogen dioxide (NO₂); and particulate matter (PM) with aerodynamic diameter < 2.5 μm (PM_{2.5}). We performed candidate (CpG-based; cross-systemic phenotypes, combined into “allostatic load”) and agnostic (DMR-based) pathway enrichment tests, and replicated previously reported air pollution EWAS signals.

RESULTS: We found no statistically significant CpGs at false discovery rate < 0.05. However, 14, 48, 183, 8, and 71 DMRs independently associated with aircraft, railway, and road traffic Lden; NO₂; and PM_{2.5}, respectively, with minimally overlapping signals. Transportation Lden and air pollutants tendentially associated with decreased and increased methylation, respectively. We observed significant enrichment of candidate DNA methylation related to C-reactive protein and body mass index (aircraft, road traffic Lden, and PM_{2.5}), renal function and “allostatic load” (all exposures). Agnostic functional networks related to cellular immunity, gene expression, cell growth/proliferation, cardiovascular, auditory, embryonic, and neurological systems development were enriched. We replicated increased methylation in cg08500171 (NO₂) and decreased methylation in cg17629796 (PM_{2.5}).

CONCLUSIONS: Mutually independent DNA methylation was associated with source-specific transportation noise and air pollution exposures, with distinct and shared enrichments for pathways related to inflammation, cellular development, and immune responses. These findings contribute in clarifying the pathways linking these exposures and age-related diseases but need further confirmation in the context of mediation analyses. <https://doi.org/10.1289/EHP6174>

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Introduction

Transportation-related noise (including road traffic, railway, and aircraft noise) and air pollution [including nitrogen dioxide (NO₂) and particulate matter (PM) with aerodynamic diameter < 2.5 μm (PM_{2.5})] both make the greatest contribution to the global environmental burden of disease (Fritschi et al. 2011; Hänninen et al. 2014; Vienneau et al. 2015). Both exposure groups have been linked to cross-systemic phenotypes, including respiratory (Adam et al. 2015; Eze et al. 2018; Hoek et al. 2013), cardiovascular (Beelen et al. 2014; Fiorito et al. 2018; Foraster et al. 2017; Heritier et al. 2019; Kempen et al. 2018), and metabolic diseases (Eze et al. 2015; Zare Sakhvidi et al. 2018); cancers (Andersen et al. 2018; Hegewald et al. 2017; Raaschou-Nielsen et al. 2013); and neurological disturbances (Clark and Paunovic 2018; Stafoggia et al. 2014; Zhang et al. 2018). The potential mechanisms linking air pollution and these phenotypes include inflammatory, immune, and oxidative stress responses following inhalation (Münzel et al. 2017a), whereas noise is thought to act through annoyance reactions, sleep disturbances, stress, and activation of the hypothalamic–pituitary–adrenal (HPA) axis and sympathetic nervous system, with subsequent

release of stress hormones and inflammatory molecules (Daiber et al. 2019).

DNA methylation alterations may mediate part of the physiological and biochemical changes leading from these traffic-related exposures to their associated preclinical and clinical phenotypes. Increasing evidence has linked short- and long-term air pollution exposure to various measures of DNA methylation in both children and adults (Abraham et al. 2018; de FC Lichtenfels et al. 2018; Gondalia et al. 2019; Gruzieva et al. 2017; Lee et al. 2019; Mostafavi et al. 2018; Panni et al. 2016; Plusquin et al. 2017, 2018; Sayols-Baixeras et al. 2019). Although findings from these studies were largely inconsistent with regard to single Cytosine-phosphate-Guanine (CpG) sites, the systematic review by Alfano et al. (Alfano et al. 2018) highlighted that air pollution was consistently linked to global hypomethylation in children (Breton et al. 2016; Cai et al. 2017; Janssen et al. 2015) and in adults (De Nys et al. 2018; De Prins et al. 2013) and accelerated epigenetic aging in adults (Nwanaji-Enwerem et al. 2016, 2017; Ward-Caviness et al. 2016). Overarching epigenomic effects of air pollution identified across these studies include inflammation, mitochondrial and DNA damage responses, and accelerated biological aging (Alfano et al. 2018).

Transportation noise, like air pollution, may also influence health outcomes via DNA methylation. First, they might share mechanistic pathways, because there is growing evidence for the inflammatory and oxidative downstream effects of noise exposure (Daiber et al. 2019; Münzel et al. 2017). Second, oxidative DNA damage, a correlate of altered DNA methylation and gene expression (Russo et al. 2016) was linked to occupational noise (Bagheri Hosseinabadi et al. 2019; Nawaz and Hasnain 2013) and traffic noise (Hemmingsen et al. 2015). Third, DNA methylation changes were reported for noise-related stressors such as vehicular traffic (Commodore et al. 2018) and insufficient sleep (Gaine et al. 2018). Methylation changes were also reported in circadian rhythm genes in conditions of acute sleep loss (Cedernaes et al. 2015) and night-shift work (Zhu et al. 2011). A study investigating brain tissue DNA methylation in relation to noise exposure in animal models reported gene-specific methylation changes, which were in turn associated with metabolic health (Guo et al. 2017).

No transportation noise epigenome-wide association study (EWAS) has been performed so far in population-based studies, and previous air pollution EWAS studies did not account for concurrent noise exposure (Eze and Probst-Hensch 2019). This partial scope of research is a limitation because both exposures share common sources and might be potential mutual confounders (Tetreault et al. 2013). As demonstrated by previous studies (Franklin and Fruin 2017; Heritier et al. 2019; Tetreault et al. 2013), the effect estimates of air pollution might be overestimated if noise level is unaccounted for, and vice versa. In addition, a parallel consideration of both exposure groups might elucidate directly shared and independent pathways of individual exposures and provide an improved understanding of mechanisms linking these exposures to disease.

In this paper, we aimed within the Swiss Cohort Study on Air Pollution and Lung and Heart Diseases in Adults (SAPALDIA) to

1. conduct a multiexposure EWAS (single CpGs and genomic regions) involving long-term exposure to source-specific transportation noise (aircraft, railway, and road traffic) and air pollution (NO₂ and PM_{2.5});
2. assess in a candidate approach, using *a priori*-curated CpGs, the pathway enrichment for “allostatic load”-related cardiometabolic, immunological, and renal phenotypes (Johnson et al. 2017; McCrory et al. 2019; Seeman et al. 2010), given that both traffic-related exposures are chronic stressors of several physiological systems;

3. assess in an agnostic approach, the functional pathway and network enrichment of EWAS-derived differentially methylated regions (DMRs); and
4. replicate previously reported CpGs associated with long-term exposure to NO₂ and PM_{2.5} in the LifeLines (de FC Lichtenfels et al. 2018), EPIC-ITALY and EPIC-NL (Plusquin et al. 2017), and Korean COPD (Lee et al. 2019) cohorts.

Methods

Study Population

The SAPALDIA cohort has been described elsewhere in detail (Ackermann-Lieblich et al. 2005; Martin et al. 1997). SAPALDIA is a population-based study that recruited 9,651 adults from 8 geographically diverse Swiss areas (Aarau, Basel, Davos, Geneva, Lugano, Montana, Payerne, and Wald) in 1991 (SAP1) to investigate the respiratory effects of air pollution exposure. The first (SAP2) and second (SAP3) follow-up surveys occurred in 2002 and 2010; included 8,047 and 6,088 participants, respectively; and additionally, assessed cardio-metabolic and quality of life phenotypes. At each survey, participants had physical examinations and health- and lifestyle-related interviews. Measures of transportation noise and air pollution exposures were modeled at participants' residences. At both SAP2 and SAP3, whole blood samples were collected, preprocessed, and stored in a biobank (−80°C) for biomarker assessments, including DNA methylation. In the context two SAPALDIA nested studies—Aging Lungs in European Cohorts study (ALEC; www.alecstudy.org) and the EXPOsOMICS study (Vineis et al. 2017)—SAP2 and SAP3 repeat blood samples from 987 representative participants (ALEC), and an independent 405 nonsmoking participants (for at least 1 y before SAP2; EXPOsOMICS), were applied toward DNA methylation assessment. The EXPOsOMICS sample was an asthma case-control sample of nonsmokers (204 cases and 201 controls), where the asthma cases had methylation assessed only in the SAP3 blood sample and the controls had methylation assessment in both the SAP2 and SAP3 blood samples. Following exclusions due to methylation data quality and covariate data availability, we included SAP2 data from 972 ALEC and 198 EXPOsOMICS (asthma controls) participants and SAP3 data from 970 ALEC and 402 EXPOsOMICS (204 asthma cases and 198 controls) participants.

The present study therefore includes 1,389 participants contributing 2,542 observations, with an average of 1.8 observations per participant. Figure 1 presents the details of participant selection and inclusion. The SAPALDIA study complies with the Declaration of Helsinki. All participants provided informed written consent before participating in any aspect of the SAPALDIA surveys and ethical approvals were obtained from the Swiss Academy of Medical Sciences and the Ethics committees of the participating cantons.

Assessment of DNA Methylation

DNA methylation was measured in the ALEC and EXPOsOMICS samples separately in different laboratories, but using consistent methodology, using Illumina Infinium 450K BeadChip and processed as previously described (Imboden et al. 2019; Jeong et al. 2019). In brief, paired blood samples from the same participant were randomized across the arrays and placed next to each other on the arrays. Dye-bias correction (Triche et al. 2013) and absolute methylation level (β -values, defined as the ratio of methylation intensity over total intensity, with offset of 100) were computed using the minfi R-package (R Development Core Team) (Aryee et al. 2014). Quality control criteria included call rate > 95%,

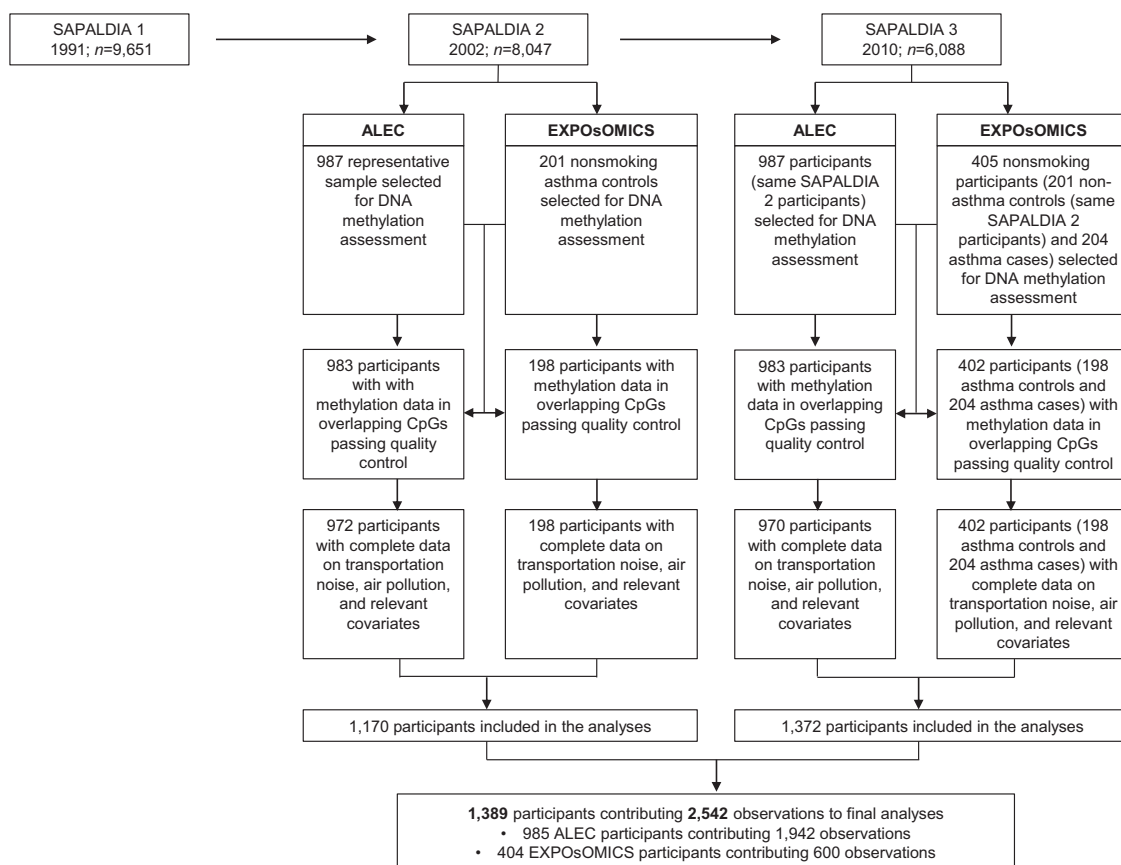


Figure 1. Selection of participants included in the present study. SAPALDIA: Swiss cohort study on air pollution and lung and heart diseases in adults. ALEC: Aging lungs in European cohorts. ALEC and EXPOsOMICS are European cohort consortia in which SAPALDIA participates. DNA methylation was measured using Illumina Infinium 450K BeadChip and processed in the same manner across the ALEC and EXPOsOMICS samples to derive the residuals of the beta values (corrected for technical bias) of overlapping 430,477 CpGs, which were subsequently applied to the present epigenome-wide association study.

detection p -value $< 10^{-16}$, sex consistency, autosomal chromosome location of probes, and restriction to probes identified in both ALEC and EXPOsOMICS samples. We applied beta mixture quantile normalization (BMIQ) of the β -values to correct for the Illumina probe design bias (Teschendorff et al. 2013). We further excluded 37,882 CpGs that are cross-reactive or target polymorphic (Chen et al. 2013). For technical bias (batch effect) correction, these β -values were then regressed on the first 30 principal components derived from a principal component analysis of the control probes incorporated on the methylation chip (Lehne et al. 2015). Figure S1 shows the distributions of the BMIQ-normalized β -values and technical bias-corrected residuals. In line with previous studies (Imboden et al. 2019; Jeong et al. 2019), we applied these residuals (in place of the β -values) in subsequent EWAS covering 430,477 CpG sites that overlapped between ALEC and EXPOsOMICS samples.

To minimize bias due to extreme values (EVs) of the residuals—the dependent variables in the present EWAS—while retaining the distribution of DNA methylation at the nonskewed CpG sites, we performed modified winsorization of the CpG sites containing EVs. We defined EVs as values lying beyond three times the interquartile range (IQR) below and above the first (Q1) and third quartiles (Q3), respectively (Tukey 1977). We replaced each EV with the corresponding threshold of detection, in a CpG-specific manner, that is:

If $EV < [Q1 - (3 \times IQR)]$, then EV is replaced with the value of $Q1 - (3 \times IQR)$ at the CpG site;

If $EV > [Q3 + (3 \times IQR)]$, then EV is replaced with the value of $Q3 + (3 \times IQR)$ at the CpG site.

Among the 430,477 included CpGs, 393,397 CpGs (91%) had at least one EV. The EVs comprised 0.04%–22% of observations across the “winsorized” CpG sites. We used these “winsorized” data as the primary outcome variables in subsequent analyses (including EWAS, enrichment, and replication), and only used the “nonwinsorized” data for sensitivity testing of the top CpG signals identified in the EWAS.

Assessment of Transportation Noise Exposure

As detailed elsewhere (Karipidis et al. 2014), annual average day-evening-night noise level [Lden, with respective 5 dB and 10 dB penalties for evening (19–23 h) and nighttime (23–07 h; Lnight)] were calculated for 2001 and 2011 (corresponding to SAP2 and SAP3, respectively), for aircraft, railway, and road traffic noise at the maximum-exposed façade of the residential floors of participants. This calculation was done in the framework of the Short and Long Term Effects of Transportation Noise Exposure (SiRENE) project (Röösli et al. 2017). Aircraft noise was modeled with the FLULA2 version 4 software (EMPA 2010) with input data covering four airports in Basel, Geneva, Zurich, and Payerne. Railway noise emission and propagation were modeled using the sonRAIL (Thron and Hecht 2010) and the SEMIBEL (FOEN 1990) models, respectively, whereas road traffic noise emission and propagation were separately modeled using sonROAD (Heutschi 2004) and StL-86 (FOEN 1987). Validation of noise calculations was done by comparing the calculated noise levels with measured levels from the field. Taking all measurements into account, a mean Lden difference of 1.6 ± 5 dB was observed (Schlatter et al. 2017). Lden

and Lnight values were respectively truncated at 30 dB and 20 dB for railway and aircraft noise, and 35 dB and 25 dB for road traffic noise. Calculated Lden and Lnight values below these limits were replaced by the respective truncation values. Participants with truncated values were assigned a truncation indicator (yes/no) for subsequent modeling, in line with previous analyses in the context of the SIRENE study (Eze et al. 2017; Foraster et al. 2017). Given the high correlations of source-specific Lden and Lnight, (Spearman's rank correlation, $r_{\text{aircraft}} = 0.76$, $r_{\text{railway}} = 0.97$, and $r_{\text{road traffic}} = 0.99$), and the generally lower Lnight levels (Table S1 and Figure S2); we focused on source-specific Lden as the noise parameters of interest in this EWAS.

Assessment of Air Pollution Exposure

Traffic-related air pollutants—NO₂ and PM_{2.5}—were also modeled at participant's residences at SAP2 and SAP3 as outdoor mean exposures. For SAP2, annual mean NO₂ was estimated (year 2003) using a hybrid model that regressed passive sampler measurements against dispersion model estimates, seasonal and climatic variables, as well as traffic and land use characteristics. Adjusted R² of the hybrid model was 0.8 (Liu et al. 2012). PM_{2.5} was derived from the Swiss PolluMap dispersion model (year 2000). Modeled values were validated against measured values with an R² of 0.9 (Heldstab et al. 2003). For SAP3, NO₂ and PM_{2.5} were modeled as average biennial exposures (2010/2011) using land use regression models. The best models were used in each case: the combined PM_{2.5} with an adjusted R² of 0.6, and the area-specific NO₂ models with adjusted R²'s of 0.5–0.9 across SAPALDIA study areas (Eeftens et al. 2016). We included both pollutants in our analyses, examining their associations in both single- and multiexposure models.

Assessment of Covariates

At the level of the individual, we considered groups of potential confounders measured at both SAP2 and SAP3, which might be associated with transportation noise, air pollution exposures, and DNA methylation. We considered sociodemographic factors including age (y), sex (male/female), and educational attainment (primary education/secondary education or apprenticeship/tertiary education). We considered lifestyle factors such as smoking status (never/former/current), smoking pack-years (calculated from cigarettes per day and duration of smoking), passive smoking (yes/no), alcohol consumption (beers, wines, liquors and spirits; ≤ 1 glass/d or > 1 glass/d), frequency of fruit intake (citrus or noncitrus fruits in any form; ≤ 3 d/wk or > 3 d/wk), vegetable intake (raw or cooked; ≤ 3 d/wk or > 3 d/wk), body mass index (BMI; kg/m²), and sufficient moderate-to-vigorous physical activity (≥ 150 min/wk of engagement in activities that makes one at least moderately sweat or breathless). To minimize the between-nested study differences, including residual batch effects and asthma status, we considered the contributing nested study (ALEC/EXPOSOMICS) and asthma status, defined as ever having a diagnosis of asthma. We also considered estimates of leukocyte composition for B cells, CD4T cells, CD8T cells, eosinophils, monocytes, natural killer cells, and neutrophils (derived from the DNA methylation data using the “estimatecellcounts” function in the minfi R package (Houseman et al. 2012; Reinius et al. 2012), to control the influence of cell proportions on methylation level (Adalsteinsson et al. 2012).

On the contextual scale, we considered some commonly indicated potential confounders of environmental health such as study area, a neighborhood index of socioeconomic position (SEP; %), and greenness index within a 1-km residential buffer. SEP was derived from a principal component analysis of household characteristics (education, occupation of household head, occupancy

and median rent of household) based on 2001 census data (Panczak et al. 2012), and assigned to residential geo-coordinates at SAP2 and SAP3. Greenness index was calculated for 2014 as normalized difference vegetation index based on surface reflectance and assigned to participants' geo-coordinates at SAP2 and SAP3 (Vienneau et al. 2017).

Statistical Analyses

EWAS. We described the characteristics of included participants by survey and by nested study, summarizing categorical variables as counts and proportions and summarizing continuous variables as medians and IQR. Using the combined sample, we performed EWAS by linear mixed-effects regressions on the “winsorized” technical-bias corrected residuals of 430,477 CpGs, applying random intercepts at the level of participants. We used the “lmer” function of the lme4 R package for the regressions (Bates et al. 2015). We performed single- as well as multiexposure EWAS for aircraft, railway, and road traffic Lden, NO₂ and PM_{2.5}, adjusting for age; sex; education; smoking status; pack-years; passive smoking; fruit, vegetable, and alcohol intake; study area; SEP; greenness index; survey; nested study; asthma; Lden truncation indicator; and Houseman estimates of leukocyte composition in the main model. We identified genome-wide significant CpGs at false discovery rate (FDR) and Bonferroni-corrected *p*-value thresholds of 0.05 and 1.16×10^{-7} , respectively.

We tested sensitivity of the top 10 CpGs to further adjustment for the potential mediators BMI and physical activity, which have been associated with transportation noise (An et al. 2018b; Foraster et al. 2016; Roswall et al. 2017) and air pollution exposures (An et al. 2018a, 2018c). We further stratified these CpGs by nested study to assess consistency in effect direction, limited these models to participants who reported regular nighttime opening of windows, and assessed the robustness of top signals in models using the “nonwinsorized” methylation data.

We tested for DMRs, using the “dmrca” function in the DMRcate R package (Peters et al. 2015) and the multiexposure EWAS-derived parameters for each CpG as input file. Thus, we performed the DMR analyses using the individual estimates of 430,477 CpGs from the main multiexposure model. We defined DMRs as significant if they contained at least two CpGs, within at least 1,000 base pairs, and had a minimum FDR *p*-value < 0.05 . Except when unannotated, all CpGs and DMRs are reported with gene annotation in parenthesis.

Pathway Enrichment

In a candidate pathway approach, we identified previously published EWAS signals for different physiological systems (captured by selected phenotypes) of potential relevance to transportation-related noise and air pollution effects. These systems (phenotypes) included *a*) immunological (C-reactive protein, CRP); *b*) metabolic [glycemia, insulin secretion/sensitivity, lipoprotein cholesterol, BMI, waist circumference (WC)/visceral adipose tissue mass, and metabolic syndrome]; *c*) renal (estimated glomerular filtration rate, eGFR); and *d*) cardiovascular and autonomic nervous systems [blood pressure and cardiac autonomic responses (CAR)]. For all phenotypes, we curated CpGs from the EWAS atlas (Li et al. 2019), at a *p*-value threshold of 1.10×10^{-5} , except for CRP where only the genome-wide significant signals at *p*-value of 1.15×10^{-7} were reported (Ligthart et al. 2016). CpGs that overlapped with those of the SAPALDIA study were finally curated for enrichment analyses (Table S1).

We defined a global “allostatic load” pathway as the entirety of unique CpGs assigned to at least one of the constituent pathways ($n = 1,626$). We applied the Weighted Kolmogorov-Smirnov

method (Charnpi and Ycart 2015) to test for enrichment of the candidate and “allostatic load” pathways using the absolute values of test statistics from multiexposure EWAS. These test statistics from CpGs mapped to the pathway were compared with the empirical null distribution derived by 10,000 permutation samples. If the Kolmogorov-Smirnov test statistic was larger than the 90th percentile of test statistics obtained from the permutation samples after permutation-based multiple testing correction (van der Laan et al. 2005), we declared enrichment for the pathway.

In an agnostic approach, we assessed the functional pathways of exposure-specific differential methylation, using the “Core Analysis” of Ingenuity pathway analysis (IPA; Ingenuity Systems) on genes annotated to significant DMRs. For each exposure, we identified the functional pathways that were significantly enriched at p -value < 0.05 , as well as top disease and functional networks related to these pathways.

Replication of Previously Reported Air Pollution–Associated CpGs

Using the SAPALDIA EWAS results, we looked up the single- and multiexposure EWAS signals for validation of 22 and 10 previously reported genome-wide significant CpGs for long-term exposure to NO₂ and PM_{2.5}, respectively. These include NO₂ signals from the LifeLines cohort [cg04908668, cg14938677, cg00344801, cg18379295, cg25769469, cg02234653, and cg08500171 (de FC Lichtenfels et al. 2018)], EPIC-ITALY [cg08120023, cg22856765, cg18164357, cg13918628, cg03870188, cg20939320, cg13420207, cg04914283, cg21156210, cg16205861, cg12790758, and cg18201392 (Plusquin et al. 2017)], and the Korean COPD cohort [cg05171937, cg06226567, and cg26583725 (Lee et al. 2019)]. PM_{2.5}-associated signals include in cg23890774 from EPIC-ITALY; and cg12575202, cg08630381, cg17629796, cg07084345, cg04319606, cg09568355, cg03513315, cg25489413, and cg00005622 from EPIC-NL (Plusquin et al. 2017). We identified CpGs as replicated in our study if they showed consistent effect direction at nominal p -value threshold of 0.05.

Results

The characteristics of the study sample at SAP2 and SAP3 are presented in Table 1. In general, participants tended to gain weight and smoke less with relatively lower pollutant exposures between surveys. ALEC and EXPOsOMICS participants mainly differed by design in their smoking habits and asthma status, but their environmental exposure profiles were comparable (Table S2). Figure S2 shows the overall distributions of the aircraft, railway, and road traffic Lden and NO₂ and PM_{2.5} exposures. The Spearman’s rank correlations (r) of the five exposures are shown in Table S1. Correlation of NO₂ and PM_{2.5} was 0.65. NO₂ had higher correlation with road traffic ($r=0.41$) than railway ($r=0.23$) and aircraft Lden ($r=0.10$), whereas PM_{2.5} had higher correlation with aircraft ($r=0.23$) than railway ($r=0.19$) and road traffic Lden ($r=0.19$). Correlation patterns were consistent across SAP2 and SAP3.

EWAS

In the single exposure models (Table S3), we observed one genome-wide significant (FDR = 0.040) PM_{2.5}-associated CpG [cg26704043 (FARS2)] and one borderline-significant (FDR = 0.077) railway Lden-associated CpG [cg25201280 (ATPBD4)]. There were no genome-wide significant signals in the multiexposure model (Table 2). Here, the PM_{2.5}-associated cg26704043 (FARS2) got considerably weaker (FDR = 0.180), whereas the railway Lden-associated cg25201280 (ATPBD4) remained borderline-significant (FDR = 0.075). The top 10 CpGs of the single-exposure

Table 1. Summary of the included SAPALDIA sample.

	SAPALDIA2 ^a	SAPALDIA3 ^a
Categorical variables [n (%)]	1,170 (100)	1,372 (100)
Women	623 (53)	737 (54)
Formal education (y)		
≤ 9	56 (5)	62 (4)
10–12	759 (65)	887 (65)
> 12	355 (30)	423 (31)
Smoking status		
Never	539 (46)	659 (48)
Former	355 (30)	496 (36)
Current	276 (24)	217 (16)
Passive smoke exposure	289 (25)	159 (12)
Alcohol intake > 1 glass/d	456 (39)	529 (38)
Fruit intake ≤ 3 d/wk	326 (28)	280 (20)
Vegetable intake ≤ 3 d/wk	86 (7)	100 (7)
Urban area	689 (59)	897 (60)
Prevalent asthma	161 (14)	398 (21)
MVPA < 150 min/wk	330 (28)	354 (26)
Regular nighttime opening of windows	971 (83)	1,131 (82)
Nested study		
ALEC	972 (83)	970 (71)
EXPOsOMICS	198 (17)	402 (29)
Continuous variables [median (IQR)]		
Age	50 (18)	58 (18)
Body mass index (kg/m ²)	24.9 (5)	25.8 (6)
Smoking pack-years	0.4 (15)	0 (14)
Neighborhood index of socioeconomic position (%)	64.6 (13)	64.8 (13)
Greenness index within 1-km buffer	0.61 (0.2)	0.62 (0.2)
Aircraft Lnight (dB)	20 (2)	20.1 (5)
Railway Lnight (dB)	22.9 (14)	20 (10)
Road traffic Lnight (dB)	44.9 (111)	45.1 (11)
Aircraft Lden (dB)	30 (9)	32.7 (8)
Railway Lden (dB)	30 (11)	30 (7)
Road traffic Lden (dB)	53.7 (11)	53.9 (11)
NO ₂ ($\mu\text{g}/\text{m}^3$)	20.2 (14)	16.7 (10)
PM _{2.5} ($\mu\text{g}/\text{m}^3$)	14.3 (5)	12.9 (2)

Note: ALEC and EXPOsOMICS are European cohort consortia in which SAPALDIA participates. ALEC, Aging Lungs in European Cohorts; MVPA, moderate to vigorous physical activity; PM_{2.5}, particulate matter with aerodynamic diameter < 2.5 μm ; NO₂, nitrogen dioxide; SAPALDIA, Swiss Cohort Study on Air Pollution and Lung and Heart Diseases in Adults.

^aPopulation included in the analysis was limited to participants with complete methylation, exposure, and covariate data.

models were not entirely consistent with those of the multiexposure models, with varying degrees of overlap across exposures. However, overlapping CpGs were directionally consistent (for all exposures), and the top CpG was positionally consistent (except for road traffic Lden) between the single- and multiexposure models (Tables 2 and S3).

In the multiexposure model, aircraft Lden (cg02286155), railway Lden [cg25201280 (ATPBD4)] and road traffic Lden cg09129334 (ARHGEF7) were associated with reduced methylation, whereas NO₂ [cg04337651 (ASBI)] and PM_{2.5} [cg26704043 (FARS2)] were associated with increased methylation, at their respective top CpG sites. Among the top 10 signals, we observed decreased methylation at 9 (90%), 8 (80%), and 8 (80%) CpGs in relation to aircraft, railway, and road traffic Lden, respectively, and increased methylation at 5 (50%) and 10 (100%) CpGs in relation to NO₂ and PM_{2.5}, respectively. These CpGs were generally robust to adjustment for BMI and physical activity (Table 2) and showed consistent effect direction when stratified by nested study (Table S4). Associations were also robust in the model limited to participants reporting regular nighttime window opening (Table S5). The “non-winsorized” model highlighted few CpGs where extreme values would potentially bias their estimates if unaddressed. These included directionally consistent but considerably weaker effects on aircraft Lden-associated cg11944797 (STK24) and cg10975000; road traffic Lden-associated cg08351004 (DLX2); NO₂-associated

Table 2. Top ten CpGs independently associated with source-specific transportation noise and air pollution in the SAPALDIA study, multiexposure models.

Exposure	CpG ID	CHR	Location	Gene	Feature	Model 1				Model 2			
						Beta	SE	<i>p</i> -Value	P (FDR)	Beta	SE	<i>p</i> -Value	
Aircraft Lden	cg02286155	5	176826262	NA	NA	-0.007	0.001	1.48 × 10 ⁻⁶	0.637	-0.007	0.001	2.13 × 10 ⁻⁶	
	cg15063530	2	17716941	NA	NA	-0.009	0.002	8.14 × 10 ⁻⁶	0.659	-0.009	0.002	8.14 × 10 ⁻⁶	
	cg16218477	7	1066167	<i>C7orf50</i>	Body	-0.004	0.001	8.53 × 10 ⁻⁶	0.659	-0.004	0.001	7.05 × 10 ⁻⁶	
	cg21602842	18	46291908	<i>KIAA0427</i>	Body	-0.005	0.001	9.84 × 10 ⁻⁶	0.659	-0.005	0.001	9.84 × 10 ⁻⁶	
	cg09042449	10	44064225	<i>ZNF239</i>	5'UTR	-0.002	0.0004	1.06 × 10 ⁻⁵	0.659	-0.002	0.0004	1.18 × 10 ⁻⁵	
	cg10975000 ^a	13	28371375	NA	NA	-0.002	0.0005	1.12 × 10 ⁻⁵	0.659	-0.002	0.0005	1.16 × 10 ⁻⁵	
	cg06220958	17	10452851	<i>MYH2</i>	5'UTR	0.011	0.002	1.24 × 10 ⁻⁵	0.659	0.010	0.002	1.43 × 10 ⁻⁵	
	cg25462190 ^a	5	177547067	<i>N4BP3</i>	Body	-0.007	0.002	1.32 × 10 ⁻⁵	0.659	-0.007	0.002	1.06 × 10 ⁻⁵	
	cg11944797	13	99135711	<i>STK24</i>	Body	-0.001	0.0003	1.38 × 10 ⁻⁵	0.659	-0.001	0.0003	1.70 × 10 ⁻⁵	
	cg04635504	11	2829241	<i>KCNQ1</i>	Body	-0.005	0.001	1.61 × 10 ⁻⁵	0.664	-0.005	0.001	1.62 × 10 ⁻⁵	
Railway Lden	cg25201280 ^a	15	35838552	<i>ATPBD4</i>	TSS200	-0.001	0.0002	1.74 × 10 ⁻⁷	0.075	-0.001	0.0002	1.99 × 10 ⁻⁷	
	cg24653263 ^a	5	38258335	<i>EGFLAM</i>	TSS200	-0.003	0.001	8.97 × 10 ⁻⁷	0.193	-0.003	0.001	1.01 × 10 ⁻⁶	
	cg16825060 ^a	8	144242342	<i>LY6H</i>	TSS1500	-0.003	0.001	2.34 × 10 ⁻⁶	0.256	-0.003	0.001	2.56 × 10 ⁻⁶	
	cg23468045	5	12669584	NA	NA	0.001	0.0002	2.37 × 10 ⁻⁶	0.256	0.001	0.0002	2.19 × 10 ⁻⁶	
	cg19270309 ^a	17	77712853	<i>ENPP7</i>	3'UTR	0.002	0.0005	3.25 × 10 ⁻⁶	0.266	0.002	0.0005	3.56 × 10 ⁻⁶	
	cg07461273	7	99697172	<i>MCM7</i>	Body	-0.004	0.001	3.78 × 10 ⁻⁶	0.266	-0.004	0.001	3.70 × 10 ⁻⁶	
	cg01301319	7	27153580	<i>HOXA3</i>	5'UTR	-0.003	0.001	4.60 × 10 ⁻⁶	0.266	-0.003	0.001	4.32 × 10 ⁻⁶	
	cg23113715 ^a	22	25800663	NA	NA	-0.004	0.001	5.79 × 10 ⁻⁶	0.266	-0.004	0.001	5.73 × 10 ⁻⁶	
	cg13402217	1	151584375	<i>SNX27</i>	TSS1500	-0.003	0.001	6.03 × 10 ⁻⁶	0.266	-0.003	0.001	5.50 × 10 ⁻⁶	
	cg24047259 ^a	14	65347275	NA	NA	-0.001	0.0003	7.35 × 10 ⁻⁶	0.266	-0.001	0.0003	4.90 × 10 ⁻⁶	
Road traffic Lden	cg09129334	13	111837676	<i>ARHGEF7</i>	Body	-0.007	0.001	1.73 × 10 ⁻⁶	0.384	-0.007	0.001	2.12 × 10 ⁻⁶	
	cg17383236 ^a	7	100167504	NA	NA	-0.002	0.0005	2.76 × 10 ⁻⁶	0.384	-0.002	0.0005	2.74 × 10 ⁻⁶	
	cg01066220	6	31696240	<i>DDAH2</i>	Body	0.001	0.0001	3.48 × 10 ⁻⁶	0.384	0.001	0.0001	4.29 × 10 ⁻⁶	
	cg23910243 ^a	16	31484618	<i>TGFB111</i>	Body	-0.002	0.0005	4.32 × 10 ⁻⁶	0.384	-0.002	0.0005	7.13 × 10 ⁻⁶	
	cg06646021 ^a	1	229406520	<i>RAB4A</i>	TSS1500	-0.005	0.001	4.47 × 10 ⁻⁶	0.384	-0.005	0.001	4.78 × 10 ⁻⁶	
	cg03066594	20	10415919	<i>C20orf94</i>	TSS200	0.0005	0.0001	6.00 × 10 ⁻⁶	0.386	0.0004	0.0001	7.82 × 10 ⁻⁶	
	cg03966094 ^a	22	21058792	<i>TMEM191A</i>	Body	-0.003	0.001	7.06 × 10 ⁻⁶	0.386	-0.003	0.001	6.73 × 10 ⁻⁶	
	cg13948857	5	131763756	<i>C5orf56</i>	Body	-0.003	0.001	7.17 × 10 ⁻⁶	0.386	-0.003	0.001	7.22 × 10 ⁻⁶	
	cg08351004 ^a	2	172965650	<i>DLX2</i>	Body	-0.002	0.0005	9.53 × 10 ⁻⁶	0.456	-0.002	0.0005	1.23 × 10 ⁻⁵	
	cg13777730	1	234793300	NA	NA	-0.003	0.001	1.07 × 10 ⁻⁵	0.458	-0.003	0.001	1.10 × 10 ⁻⁵	
NO ₂	cg04337651 ^a	2	239344738	<i>ASB1</i>	Body	0.004	0.001	2.06 × 10 ⁻⁶	0.657	0.003	0.001	2.67 × 10 ⁻⁶	
	cg18776472	10	50732819	<i>ERCC6</i>	Body	-0.001	0.0002	8.20 × 10 ⁻⁶	0.657	-0.001	0.0002	1.00 × 10 ⁻⁵	
	cg18601596	6	39283313	<i>KCNK16</i>	Body	0.006	0.001	8.21 × 10 ⁻⁶	0.657	0.006	0.001	8.62 × 10 ⁻⁶	
	cg12392998	17	79550668	<i>NPLOC4</i>	Body	-0.002	0.0004	8.72 × 10 ⁻⁶	0.657	-0.002	0.0004	1.04 × 10 ⁻⁵	
	cg16550606	13	50160670	<i>RCBTB1</i>	TSS1500	0.004	0.001	1.33 × 10 ⁻⁵	0.657	0.004	0.001	1.44 × 10 ⁻⁵	
	cg25266109	19	12404608	<i>ZNF44</i>	Body	-0.0004	0.0001	1.38 × 10 ⁻⁵	0.657	-0.0004	0.0001	1.20 × 10 ⁻⁵	
	cg01746514 ^a	14	24520922	<i>LRRC16B</i>	TSS1500	-0.001	0.0002	1.43 × 10 ⁻⁵	0.657	-0.001	0.0002	1.58 × 10 ⁻⁵	
	cg15811902	15	75918385	<i>SNUPN</i>	5'UTR	-0.002	0.0005	1.61 × 10 ⁻⁵	0.657	-0.002	0.0005	1.48 × 10 ⁻⁵	
	cg26898336 ^a	17	15244519	<i>TEKT3</i>	5'UTR	0.002	0.0005	1.65 × 10 ⁻⁵	0.657	0.002	0.0005	1.89 × 10 ⁻⁵	
	cg21099332	5	39270715	NA	NA	0.004	0.001	1.66 × 10 ⁻⁵	0.657	0.004	0.001	1.88 × 10 ⁻⁵	
PM _{2.5}	cg26704043	6	5282702	<i>FARS2</i>	5'UTR	0.014	0.003	4.18 × 10 ⁻⁷	0.180	0.014	0.003	6.22 × 10 ⁻⁷	
	cg05157625	14	93153553	<i>RIN3</i>	Body	0.021	0.004	1.08 × 10 ⁻⁶	0.231	0.021	0.004	1.13 × 10 ⁻⁶	
	cg20099458 ^a	7	5272275	<i>WIPI2</i>	3'UTR	0.014	0.003	1.61 × 10 ⁻⁶	0.231	0.014	0.003	1.58 × 10 ⁻⁶	
	cg06587257	12	50452135	<i>ACCN2</i>	5'UTR	0.022	0.005	2.71 × 10 ⁻⁶	0.292	0.023	0.005	2.04 × 10 ⁻⁶	
	cg14531665	9	91058614	<i>SPIN1</i>	Body	0.012	0.003	5.91 × 10 ⁻⁶	0.398	0.011	0.003	7.07 × 10 ⁻⁶	
	cg06526020	6	34308880	<i>NUDT3</i>	Body	0.029	0.006	6.43 × 10 ⁻⁶	0.398	0.028	0.006	8.80 × 10 ⁻⁶	
	cg21058520	6	100914733	NA	NA	0.004	0.001	6.76 × 10 ⁻⁶	0.398	0.004	0.001	8.31 × 10 ⁻⁶	
	cg16259904	10	134146220	<i>LRRC27</i>	5'UTR	0.027	0.006	8.90 × 10 ⁻⁶	0.398	0.027	0.006	1.01 × 10 ⁻⁵	
	cg12770741	17	883776	<i>NXN</i>	TSS1500	0.018	0.004	9.15 × 10 ⁻⁶	0.398	0.018	0.004	1.01 × 10 ⁻⁵	
	cg26750893 ^a	2	38043481	NA	NA	0.016	0.004	1.05 × 10 ⁻⁵	0.398	0.016	0.004	1.28 × 10 ⁻⁵	

Note: Beta coefficients represent increase or decrease in DNA methylation per 10 dB increase in aircraft, railway or road traffic Lden or 10 µg/m³ increase in NO₂ or PM_{2.5}. All estimates were from multiexposure epigenome-wide linear mixed-effects models, with random intercept at the level of participant. Multiexposure models included all five exposures (aircraft, railway, road traffic Lden and respective truncation indicators, NO₂ and PM_{2.5}) at the same time. In a preliminary step, DNA methylation β-values were regressed on the Illumina control probe-derived first 30 principal components to correct for correlation structures and technical bias, and residuals of these regressions covering 430,477 CpGs were used as the technical bias-corrected methylation level at the CpG sites. Extreme values of the residuals (lying beyond three times the interquartile range below the first quartile and above the third quartile at each CpG site) were replaced with their corresponding detection threshold value ("modified winsorization"). The "winsorized" data were then used as the dependent variables in the EWAS. Model 1: adjusted for age; sex; educational level; area; neighborhood socioeconomic status; greenness index; smoking status; smoking pack-years; exposure to passive smoke; consumption of fruits, vegetables and alcohol; nested study; asthma status; noise truncation indicators; survey; and leukocyte composition (main model). Model 2: Model 1+body mass index and physical activity. CHR, chromosome; CpG, Cytosine-phosphate-Guanine; Lden, day-evening-night noise level; NA, not annotated; NO₂, particulate matter with aerodynamic diameter < 2.5 µm; SAPALDIA, Swiss Cohort Study on Air Pollution and Lung and Heart Diseases in Adults; SE, standard error.

^aLocation overlaps with significant differentially methylated region.

cg18776472 (*ERCC6*), cg12392998 (*NPLOC2*), and cg26898336 (*TEKT3*); and a stronger and genome-wide significant effect on PM_{2.5}-associated cg21058520 (Table S6). Results from the multiexposure main model of CpGs associated with aircraft, railway, and road traffic Lden, NO₂—at nominal *p*-value < 1.00 × 10⁻³—are presented in Tables S2–S6.

Post hoc analysis of overlap among the top 100 exposure-specific CpGs identified from the multiexposure EWAS showed minimal signal overlap between exposures (Figure 2A). We identified two overlapping signals between road traffic Lden and NO₂ [cg12439232 and cg15590912 (*CCSAP*)]. Yet, road traffic Lden was associated with decreased methylation, whereas NO₂ was

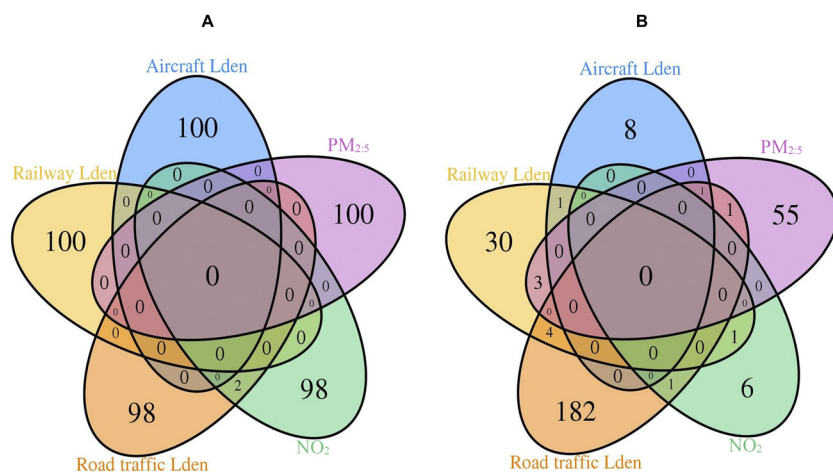


Figure 2. Overlap of top 100 CpG signals (A) and genes annotated to significant differentially methylated regions (B) in relation to aircraft, railway, and road traffic Lden, NO₂, and PM_{2.5} identified from multiexposure EWAS in the SAPALDIA study. Note: EWAS, epigenome-wide association study; Lden, day-evening-night noise level; NO₂, nitrogen dioxide; PM_{2.5}, particulate matter < 2.5 microns in diameter; SAPALDIA, Swiss Cohort Study on Air Pollution and Lung and Heart Diseases in Adults. CpGs were identified by multiexposure EWAS using multivariable linear mixed-effects models with random intercepts at the level of participants, and adjusted for age, sex, educational level, area, neighborhood socioeconomic status, greenness index, smoking status, smoking pack-years, exposure to passive smoke, consumption of fruits, vegetables and alcohol, nested study, asthma status, noise truncation indicators, survey and leukocyte composition. In a preliminary step, DNA methylation β -values were regressed on the Illumina control probe-derived first 30 principal components to correct for correlation structures and technical bias, and residuals of these regressions covering 430,477 CpGs were used as the technical bias-corrected methylation level at the CpG sites. Extreme values of the residuals (lying beyond three times the interquartile range below the first quartile and above the third quartile at each CpG site) were replaced with their corresponding detection threshold value (“modified winsorization”). The “winsorized” data were then used as the dependent variables in the present EWAS. CpGs (annotated gene) intersecting at the level of road traffic Lden and NO₂ were cg12439232 and cg15590912 (CCSAP). Genes (DMR) intersecting at the level of road traffic Lden and PM_{2.5} was *PRRT1* (chr6:32,115,964–32,117,401); and at the level of aircraft and road traffic Lden and PM_{2.5} was *HOXA2* (chr7:27,141,774–27,143,806). *VTRNA2-1* (chr5:135,415,129–135,416,613) intersected between aircraft and railway Lden, *ZFP57* (chr6:29,648,161–29,649,084), between railway Lden and NO₂, and *ZSCAN31* (chr6:28,303,923–28,304,451), between road traffic Lden and NO₂. *TRIM39*, *TRIM39-RPP21*, and *HCG18* (chr6:30,296,689–30,297,941) intersected between railway Lden and PM_{2.5}, whereas *SLC27A3* (chr1:153,746,588–153,747,856), *B3GALT4* (chr6:33,244,976–33,246,185), *EN2*, and *AC008060.8* (chr7:155,249,398–155,251,925) intersected between railway and road traffic Lden.

associated with increased methylation at both sites (Tables S4 and S5). Complete EWAS results from the single- and multiexposure main models of 430,477 CpGs in relation to aircraft, railway, and road traffic Lden; NO₂; and PM_{2.5} have been deposited in the Zenodo public online depository (<https://zenodo.org/record/3832253#.XsLLJEKxU2x>).

We identified independent DMRs (FDR = 0.05) across all exposures in the main model (Table 3). There were 14 (10), 48 (39), 183 (189), 8 (8), and 71 (60) DMRs (genes), associated with aircraft, railway, and road traffic Lden, NO₂, and PM_{2.5}, respectively. Among the top 10 CpGs identified in the multiexposure model, two aircraft [cg10975000 and cg25462190 (*N4BP3*)], six railway [cg25201280 (*ATPBD4*), cg24653263 (*EGFLAM*), cg16825060 (*LY6H*), cg19270309 (*ENPP7*), cg23113715, and cg24047259], and five road traffic Lden-associated CpGs [cg17383236, cg23910243 (*TGFBIII*), cg06646021 (*RAB4A*), cg03966094 (*TMEM191A*), and cg08351004 (*DLX2*)] were within the exposure-specific DMRs. Three NO₂ [cg04337651 (*ASB1*), cg01746514 (*LRRCL16B*) and cg26898336 (*TEKT3*)] and two PM_{2.5}-associated CpGs [cg20099458 (*WIP12*) and cg26750893] were also within the exposure-specific DMRs. Top DMRs independently associated with exposures, were annotated to *VTRNA2-1* (aircraft and railway Lden; Chr5:135,415,129–135,416,613), *OXT* (road traffic Lden; Chr20:3,051,954–3,053,196), *ZSCAN31* (NO₂; Chr6:28,303,923–28,304,451), and *TRIM39*, *HCG18*, and *TRIM39-RPP21* (PM_{2.5}; Chr6:30,296,689–30,297,941). Most of the CpGs within the DMRs associated with source-specific Lden showed decreased methylation (aircraft = 64%, railway = 69%, and road traffic = 93%), whereas those associated with air pollution showed increased methylation (NO₂ = 63% and PM_{2.5} = 93%). Tables S7 and S8 show the results from the multiexposure main model of DMRs associated with aircraft, railway, and road traffic Lden; NO₂; and PM_{2.5} at FDR < 0.05.

Post hoc analysis of overlap among the gene-annotated significant DMRs also showed minimal overlap between exposures (Figure 2B). *SLC27A3*, *B3GALT4*, *EN2*, and *AC008060.8* overlapped between railway and road traffic Lden; *TRIM39*, *TRIM39-RPP21*, and *HCG18* overlapped between railway Lden and PM_{2.5}; and *HOXA2* overlapped between aircraft and road traffic Lden and PM_{2.5}. Other overlapping genes include *VTRNA2-1* (aircraft and railway Lden), *ZFP57* (railway Lden and NO₂), and *ZSCAN31* (road traffic Lden and NO₂), and *PRRT1* (road traffic Lden and PM_{2.5}). Overlapping DMRs showed opposing effect direction between exposures, except for consistently increased methylation at *TRIM39*, *TRIM39-RPP21*, *HCG18* (railway Lden and PM_{2.5}), and *HOXA2* (aircraft Lden and PM_{2.5}), and decreased methylation at *SLC27A3*, *B3GALT4*, *EN2*, and *AC008060.8* (railway and road traffic Lden) (Tables S7 and S8).

Pathway Enrichment

For the candidate single phenotype and combined “allostatic load” pathways, multiple testing-corrected enrichment analyses showed varying enrichment across exposures (Table 4). Considering the results for single phenotypes, PM_{2.5}-related methylation was the most enriched, whereas NO₂-related methylation was the least enriched. Single pathways (*p*-values) enriched for PM_{2.5} included CRP (0.0004), glycemia (0.0947), WC (0.0004), BMI (0.0004), and eGFR (0.0004). Aircraft and road traffic Lden-related methylation were enriched for CRP (0.0038; 0.0395), BMI (0.0007; 0.0008), and eGFR (0.0015; 0.0031). Railway Lden-related methylation was enriched for eGFR (0.0562) and CAR (0.0229), whereas NO₂-related methylation was enriched only for eGFR (0.0058). Considering the global “allostatic load” pathway, we found consistent enrichment across all exposures, including

Table 3. Summary of EWAS-derived differentially methylated regions and enrichment in relation to transportation noise and air pollution exposure in the SAPALDIA study.

Exposure	DMRs [Genes (n)]	Average effect on DMRs	Top DMR (Gene)	CpGs in top DMR (n)	FDR <i>p</i> -value (top DMR)	Top enriched canonical pathway (TECP)	Genes in TECP	Top enriched disease networks
Aircraft Lden	14 (10)	↓ methylation (64%)	Chr5:135415129–135416613 (<i>VTRNA2-1</i>)	19	8.04×10^{-14}	NA	NA	Cell cycle, embryonic development, organismal development, cardiovascular system development and function, gene expression, organ development
Railway Lden	48 (39)	↓ methylation (69%)	Chr5:135415129–135416613 (<i>VTRNA2-1</i>)	19	3.61×10^{-5}	Type II diabetes mellitus signaling; diphthamide biosynthesis	<i>PRKAA1</i> , <i>SLC27A3</i> , <i>TNFRSF11B</i> , <i>DPH6</i>	cardiovascular system development and function, gene expression, organ development
Road traffic Lden	183 (189)	↓ methylation (93%)	Chr20:3051954–3053196 (<i>OXT</i>)	13	3.93×10^{-6}	Wnt/ β -catenin signaling; cholecystinin/gastrin-mediated signaling	<i>CDH1</i> , <i>C5NK1E</i> , <i>SOX2</i> , <i>SOX8</i> , <i>WNT16</i> , <i>MEF2D</i> , <i>PXN</i> , <i>SHC1</i> , <i>TNF</i>	Cell-mediated immune response, cell-to-cell signaling and interaction, cellular movement
NO ₂	8 (8)	↑ methylation (63%)	Chr6:28303923–28304451 (<i>ZSCAN31</i>)	11	2.57×10^{-6}	NA	NA	Cell cycle, cell-to-cell signaling and interaction, post-translational modification
PM _{2.5}	71 (60)	↑ methylation (93%)	Chr6:30296689–30297941 (<i>TRIM39</i> , <i>HCG18</i> , <i>TRIM39-RPP21</i>)	14	4.28×10^{-8}	β -alanine and 4-aminobutyrate degradation I; systemic lupus erythematosus signaling	<i>ABAT1</i> , <i>PRPF31</i> , <i>PRPF8</i> , <i>PTPN6</i>	Cell cycle, nervous system development and function, organ injury, and abnormalities

Note: Each DMR analysis had the corresponding multiexposure EWAS-derived parameters as input. Multiexposure EWAS derived from linear mixed-effects models, with random intercept at the level of participant, and adjusted for age; sex; educational level; area; neighborhood socioeconomic status; greenness index; smoking status; smoking pack-years; exposure to passive smoke; consumption of fruits, vegetables, and alcohol; nested study; asthma status; survey; noise truncation indicators; and leukocyte composition. In a preliminary step, DNA methylation β -values were regressed on the Illumina control probe-derived first 30 principal components to correct for correlation structures and technical bias, and residuals of these regressions covering 430,477 CpGs were used as the technical bias-corrected methylation level at the CpG sites. Extreme values of the residuals (lying beyond three times the interquartile range below the first quartile and above the third quartile at each CpG site) were replaced with their corresponding detection threshold value (“modified winsorization”). The “winsorized” data were then used as the dependent variables in the EWAS. Significant (FDR < 0.05) and annotated DMRs were used for canonical pathway and network enrichment in the Ingenuity Pathway Analysis software (Ingenuity Systems). ↓, decrease in methylation; ↑, increase in methylation; CpG, Cytosine-phosphate-Guanine; DMRs, limiting statistical power to detect enriched canonical pathways; EWAS, epigenome-wide association study; FDR, false discovery rate; Lden, day-evening-night noise level; NA, not applicable due to few significant and annotated; NO₂, nitrogen dioxide; PM_{2.5}, particulate matter with aerodynamic diameter < 2.5 μ m; SAPALDIA, Swiss Cohort Study on Air Pollution and Lung and Heart Diseases in Adults.

aircraft (0.0004), railway (0.0871), and road traffic Lden (0.0004); PM_{2.5} (0.0004); and NO₂ (0.0510).

Agnostic functional enrichment showed significantly enriched canonical pathways for railway, road traffic Lden and PM_{2.5}, with predominance of inflammatory and immune regulation-related pathways across exposures (Table S9). Some of the enriched canonical pathways included type 2 diabetes signaling, tight junction signaling, mTOR signaling, and lipopolysaccharide/interleukin-1 (IL-1)-mediated inhibition of retinoid X receptor (RXR) function (railway Lden); Wnt/ β -catenin signaling, cholecystinin/gastrin-mediated signaling, G-protein coupled receptor signaling, and Th1 and Th2 activation pathways (road traffic Lden); β -alanine, 4-aminobutyrate degradation; and systematic lupus erythematosus signaling, and IL-4 signaling (PM_{2.5}). Network analyses showed the enrichment for disease mechanisms related to cellular signaling/interaction and embryonic/organ development (all exposures), cell-mediated immune/inflammatory responses (road traffic Lden and PM_{2.5}) and diseases related to connective tissue development and function (railway and road traffic Lden). Pathways related to cardiovascular function, gene expression, and respiratory system disease (railway Lden); carbohydrate and small molecule transport/biochemistry, cancer, and auditory function (road traffic Lden); and hematological and nervous system function (PM_{2.5}) were enriched (Table S10).

Replication of previously reported NO₂ and PM_{2.5}-associated CpGs. Using the single-exposure model, corresponding to the previous EWAS studies, we replicated the increased methylation in cg08500171 (*BAT2*) associated with NO₂ ($p = 0.042$). This replication remained robust in the multiexposure model ($p = 0.019$). We observed a borderline significant decreased methylation in cg17629796 associated with PM_{2.5} ($p = 0.076$) in the single-exposure model, which became statistically significant in the multiexposure model ($p = 0.033$). Nine (41%) NO₂-associated CpGs and 5 (50%) PM_{2.5}-associated CpGs had the same direction of effect in the SAPALDIA study (Table 5).

Discussion

In this first multiexposure EWAS covering long-term aircraft, railway, and road traffic noise, as well as NO₂ and PM_{2.5} exposures, we found genome-wide significant signals—at the level of genomic regions—associated with source-specific transportation noise and air pollution exposures. We demonstrated some mutual confounding of the associations of exposures with DNA methylation, where single exposure models slightly overestimated the observed methylation effect. Overall, methylation signals minimally overlapped but showed common enrichment of inflammation and immune response-related pathways across exposures. In this study, we also validated air pollution-related CpG signals identified in external EWAS.

The railway noise-related decrease in methylation at cg25201280 (transcription start site 200 of *ATPBD4* gene on chromosome 15) and PM_{2.5}-related increase in methylation at cg26704043 (5' untranslated region of *FARS2* gene on chromosome 6) were the strongest single CpG signals identified in the study. *ATPBD4* (or *DPH6*) is a protein-coding gene that regulates adenosine triphosphate (ATP) binding and diphthamide synthase activity, within the protein metabolism pathway (Chertow 1981; Young et al. 2004). Polymorphisms in this gene were associated—in various genome-wide association studies (GWAS)—with adiposity (Kichaev et al. 2019; Pulit et al. 2019), cognitive function (Lee et al. 2018; Li et al. 2015), and Crohn's disease progression (O'Donnell et al. 2019). *FARS2* is a protein-coding gene that regulates the translation of mitochondrial proteins and gene expression (Bullard et al. 1999). Variants of this gene were associated—in GWAS—with extreme obesity (Wheeler et al. 2013) and acute myeloid leukemia (Lv et al. 2017), whereas in EWAS,

Table 4. Pathway enrichment tests (*p*-values) for transportation noise and air pollution exposures based on curated CpGs reported for selected cross-systemic outcomes.

Exposure	CRP	Metabolic syndrome	Lipids	FG/HbA1c	Insulin	WC	BMI	Blood pressure	eGFR	CAR	Allostatic load
N (CpGs)	256	10	14	9	158	168	893	20	296	7	1,626
Aircraft Lden	0.0038	0.5915	0.6326	0.4545	0.6630	0.2925	0.0007	0.1020	0.0015	0.9096	0.0004
Railway Lden	0.3163	0.4943	0.9533	0.5778	0.6284	0.3001	0.9023	0.6773	0.0562	0.0229	0.0871
Road traffic Lden	0.0395	0.2434	0.9969	0.3879	0.3461	0.5361	0.0008	0.2858	0.0031	0.8013	0.0004
NO ₂	0.2117	0.1237	0.7831	0.5733	0.3697	0.6873	0.8818	0.5355	0.0058	0.9728	0.0510
PM _{2.5}	0.0004	0.4517	0.1263	0.0947	0.3451	0.0004	0.0004	0.3759	0.0004	0.3814	0.0004

Note: Lipids includes triglycerides, high-, low- and very low-density lipoprotein cholesterol. Insulin includes measures of insulin secretion and resistance. WC also includes central obesity and adiposity. BMI also includes general obesity. Blood pressure includes systolic and diastolic blood pressure. eGFR also includes impaired renal function. CAR includes acceleration and deceleration capacity. Allostatic load combines all the phenotypes. Pathway enrichment *p*-values derived from Weighted Kolmogorov-Smirnov method using the absolute values of test statistics from multiexposure epigenome-wide association studies (EWAS), and comparing the EWAS-derived CpGs mapped to each pathway to the empirical null distribution derived by 10,000 permutation samples. The overall procedure included permutation-based multiple testing correction. EWAS was done using linear mixed-effects models, with random intercept at the level of participant, and adjusted for age; sex; educational level; area; neighborhood socioeconomic status; greenness index; smoking status; smoking pack-years; exposure to passive smoke; consumption of fruits, vegetables, and alcohol; nested study; asthma status; noise truncation indicators; survey; and leukocyte composition. In a preliminary step, DNA methylation β -values were regressed on the Illumina control probe-derived first 30 principal components to correct for correlation structures and technical bias, and residuals of these regressions covering 430,477 CpGs were used as the technical bias-corrected methylation level at the CpG sites. Extreme values of the residuals (lying beyond three times the interquartile range below the first quartile and above the third quartile at each CpG site) were replaced with their corresponding detection threshold value ("modified winsorization"). The "winsorized" data were then used as the dependent variables in the EWAS. BMI, body mass index; CAR, cardiac autonomic response; CpG, Cytosine-phosphate-Guanine; CRP, C-reactive proteins; eGFR, estimated glomerular filtration rate; FG, fasting glucose; HbA1c, glycated hemoglobin; Lden, day-evening-night noise level; NO₂, nitrogen dioxide; PM_{2.5}, particulate matter <2.5 microns in diameter; WC, waist circumference.

cg26704043 was associated with systemic lupus erythematosus (Imgenberg-Kreuz et al. 2018) and Grave's disease (Chen et al. 2019). These findings highlight the role of DNA methylation changes in these genes as potential mechanisms by which railway noise and PM_{2.5} contribute to the resulting inflammatory and neurological phenotypes.

Our observation of numerous independent DMRs (for all five exposures), despite paucity of single CpG signals, highlights the relevance of concurrent investigation of genomic regions in EWAS. Analyses of genomic regions contextualize CpGs to identify DMRs with possible functional relevance in regulation of gene transcription (Rakyan et al. 2011). It is interesting to note that transportation noise showed mostly decreased methylation, whereas air pollution showed mostly increased methylation at the respective DMRs. This difference in direction of association was also observed at the levels overlapping CpGs and annotated DMRs. The relevance of these findings remains to be clarified because DNA hypermethylation is often associated with transcriptional gene repression, whereas hypomethylation is associated with a chromatin configuration that allows transcription (Sawalha 2008). However, concordant hypermethylation in *TRIM39* (railway noise and PM_{2.5}) and *HOXA2* (aircraft noise and PM_{2.5}) and hypomethylation in *SLC27A3* and *EN2* (railway and road traffic noise) protein-coding genes indicate some synergistic pathways between exposures. *TRIM39* activates the apoptotic signaling pathway and plays a role in cellular signaling and response to stimuli (Zhang et al. 2012), and its hypermethylation was associated with depression (Crawford et al. 2018). *HOXA2* encodes a transcriptional regulator that controls cellular differentiation during development (Akin and Nazarali 2005). Hypermethylation at this gene was associated transcriptional suppression in various cancers (Li et al. 2013). *SLC27A3* is involved in lipid metabolism and brain development (Stahl 2004), and hypomethylation in this gene was associated with recurrent endometrial carcinoma (Hsu et al. 2013). *EN2* functions in the development of the central nervous system and is commonly associated with autism (Lupu et al. 2018; Márquez-Valadez et al. 2018). Aberrant methylation in *EN2* was recently linked to clear cell renal carcinoma (Lai et al. 2017). Although these highlight certain distinct and shared pattern of associations, they demonstrate the complex network in the mechanisms linking these exposures to disease. The integration of gene expression or transcriptomic data in future studies will improve our understanding of the mechanisms going from divergence in directions of associations to convergence on pathway levels across these exposures.

The results demonstrating enrichment for DNA methylation associated with phenotypes of "allostatic load" for both air pollution and noise exposure are novel but in line with previously hypothesized mechanisms. Chronic low-level exposure to air pollution and noise are established risk factors for cardio-metabolic disease and were previously linked to the single phenotypes making up the "allostatic load" pathway in this study (Münzel and Daiber 2018; Thomson 2019). Recent experimental evidence specifically linked PM and ozone pollution as well as transportation noise to alterations in the HPA axis (Jafari et al. 2017). The findings of "allostatic load" enrichment agree with the functional enrichment of pathways related to oxidative stress and immune responses across all exposures. Furthermore, the enriched disease networks identified for PM_{2.5} overlaps the networks that were recently reported for PM₁₀ in another study (Lee et al. 2019). Taken together, the evidence especially supports the recent links of transportation noise to markers of inflammation and oxidative stress (Bagheri Hosseinabadi et al. 2019; Münzel et al. 2017a, 2017b; Münzel and Daiber 2018). However, our findings on the inflammatory pathway might have been influenced by the higher prevalence of asthma in this sample (22%) in comparison with that of the entire SAPALDIA study (12%), but our models contained asthma status in addition to smoking and pack-years and should minimize this potential bias. In line with recent evidence on the early-life methylome effects of air pollution (Cai et al. 2017; Gruzjeva et al. 2017) and potentially noise exposures, we observed enrichment of pathways related to embryonic and organ development for all exposures. Altered DNA methylation due to these exposures might therefore explain the reported associations of PM and poor birth outcomes (Liu et al. 2019; Smith et al. 2020), as well as the early-life theory of age-related disease (Walhovd et al. 2016). This finding is even more relevant if these methylation changes are heritable and, when confirmed, improves our understanding of the relevance of this pathway in relation to these exposures.

The relevance of confounding by leukocyte composition on the association between exposures and peripheral blood DNA methylation has been of interest (Heiss and Brenner 2017; van Rooij et al. 2019). In post hoc analyses, we demonstrated the general weakening of top CpGs (with some signals not attaining nominal significance) when leukocyte composition was not considered. However, main effect sizes remained robust (Table S7). Source-specific transportation noise and air pollution exposures were also associated with various leukocyte types (Table S8). These associations indicate that changes in leukocyte composition probably drive

Table 5. Replication of previously reported EWAS signals for long-term exposure to NO₂ and PM_{2.5} in the SAPALDIA study, single- and multiexposure models.

Air pollutant; Cohort (Reference)	SAPALDIA (single-exposure model)										SAPALDIA (multiexposure model)									
	CpG ID	CHR	Location	Gene	Beta	SE	p-Value	Beta	SE	p-Value	Beta	SE	p-Value	Beta	SE	p-Value				
NO ₂ ; LifeLines, de FC Lichtenfels et al. 2018	cg04908668	6	32823941	PSMB9	-0.012	0.002	7.94 × 10 ⁻⁹	0.0003	0.0003	0.333	0.0004	0.0004	0.322	0.0004	0.0004	0.322				
	cg149338677	7	127231698	ATF5	0.023	0.004	1.05 × 10 ⁻⁸	0.0004	0.0004	0.619	0.0004	0.0004	0.663	0.0004	0.0004	0.663				
	cg00344801	22	46685728	TTC38	-0.028	0.005	2.38 × 10 ⁻⁸	0.001	0.001	0.209	0.001	0.001	0.282	0.001	0.001	0.282				
	cg18379295	14	52326155	GNG2	0.020	0.004	3.50 × 10 ⁻⁸	-0.001	-0.001	0.204	-0.001	-0.001	0.367	-0.001	-0.001	0.367				
	cg25769469	5	71643841	PTCD2	0.035	0.006	3.69 × 10 ⁻⁸	0.001	0.001	0.547	0.002	0.002	0.237	0.002	0.002	0.237				
	cg02234653	2	224625080	AP1S3	-0.017	0.003	4.70 × 10 ⁻⁸	0.0005	0.0005	0.001	0.001	0.001	0.752	0.001	0.001	0.752				
	cg08500171 ^a	6	31590674	BAT2	0.024	0.004	9.81 × 10 ⁻⁸	0.002	0.002	0.042	0.003	0.003	0.019	0.003	0.003	0.019				
	cg08120023	1	116947203	C1orf203	-0.003	0.0005	4.27 × 10 ⁻⁹	0.001	0.001	0.586	0.002	0.002	0.179	0.002	0.002	0.179				
	cg22856765	8	42693384	THAP1	-0.008	0.001	9.61 × 10 ⁻⁹	-0.0001	-0.0001	0.960	-0.0001	-0.0001	0.930	-0.0001	-0.0001	0.930				
	cg18164357	11	77534497	C11orf67	-0.009	0.001	1.02 × 10 ⁻⁸	0.0005	0.0005	0.042	0.003	0.003	0.016	0.003	0.003	0.016				
	cg13918628	9	35610380	CD72	-0.012	0.002	1.02 × 10 ⁻⁸	-0.0003	-0.0003	0.184	-0.0003	-0.0003	0.074	-0.0003	-0.0003	0.074				
	cg03870188	13	113717830	MCF2L	-0.004	0.001	1.02 × 10 ⁻⁸	0.0001	0.0001	0.670	0.001	0.001	0.650	0.001	0.001	0.650				
	cg20939320	3	132563279	NCRNA00119	-0.006	0.001	3.49 × 10 ⁻⁸	0.001	0.001	0.091	0.001	0.001	0.185	0.001	0.001	0.185				
cg13420207	7	81666278	CACNA2D1	-0.010	0.002	5.56 × 10 ⁻⁸	-0.0004	-0.0004	0.947	-0.0004	-0.0004	0.973	-0.0004	-0.0004	0.973					
cg04914283	1	23181832	EPHB2	-0.005	0.001	8.85 × 10 ⁻⁸	0.001	0.001	0.298	0.001	0.001	0.118	0.001	0.001	0.118					
cg16205861	12	54146572	RG9MTD2	0.010	0.002	6.03 × 10 ⁻⁸	-0.0002	-0.0002	0.941	-0.0002	-0.0002	0.731	-0.0002	-0.0002	0.731					
cg12790758	15	37369914	MEIS2	-0.004	0.001	6.57 × 10 ⁻⁸	-0.001	-0.001	0.788	-0.001	-0.001	0.882	-0.001	-0.001	0.882					
cg18201392	1	185023741	RNF2	-0.005	0.001	7.06 × 10 ⁻⁸	0.0006	0.0006	0.287	0.0005	0.0005	0.898	0.0005	0.0005	0.898					
cg05171937	12	27396765	STK38L	0.010	0.002	8.02 × 10 ⁻⁸	0.002	0.002	0.118	0.001	0.001	0.133	0.002	0.002	0.133					
cg06226567	20	22559676	C20orf56	0.003	0.001	3.50 × 10 ⁻⁸	0.0002	0.0002	0.863	0.001	0.001	0.514	0.001	0.001	0.514					
cg26583725	13	110397643	NA	-0.001	2.3 ⁻⁴	4.90 × 10 ⁻⁸	0.0001	0.0001	0.183	0.0001	0.0001	0.228	0.0001	0.0001	0.228					
cg23890774	19	36618841	NA	0.078	0.014	1.98 × 10 ⁻⁸	0.0001	0.0001	0.704	0.0003	0.0003	0.263	0.0003	0.0003	0.263					
cg12575202	10	133331128	NA	-0.467	0.080	5.40 × 10 ⁻⁹	-0.001	-0.001	0.786	-0.001	-0.001	0.529	-0.001	-0.001	0.529					
cg08630381	13	100612277	NA	0.461	0.073	2.58 × 10 ⁻¹⁰	-0.001	-0.001	0.415	-0.001	-0.001	0.394	-0.001	-0.001	0.394					
cg17629796 ^a	13	30707265	NA	-0.563	0.094	2.11 × 10 ⁻⁸	-0.002	-0.002	0.076	-0.002	-0.002	0.033	-0.002	-0.002	0.033					
cg07084345	15	61972967	NA	-0.512	0.075	7.26 × 10 ⁻¹²	0.002	0.002	0.816	0.008	0.008	0.769	0.002	0.002	0.769					
cg04319606	2	26785290	C2orf70	0.261	0.068	1.31 × 10 ⁻⁷	0.0002	0.0002	0.893	0.002	0.002	0.989	-0.0002	-0.0002	0.989					
cg09568355	2	45228633	NA	0.261	0.049	1.41 × 10 ⁻⁷	0.003	0.003	0.286	0.002	0.002	0.179	0.004	0.004	0.179					
cg03513315	2	30988383	PESI	0.307	0.058	1.35 × 10 ⁻⁷	-0.0003	-0.0003	0.631	0.001	0.001	0.837	-0.0001	-0.0001	0.837					
cg25489413	7	44794343	ZMIZ2	-0.365	0.068	6.48 × 10 ⁻⁸	0.001	0.001	0.712	0.002	0.002	0.933	-0.0002	-0.0002	0.933					
cg00005622	8	145180403	NA	-0.398	0.064	5.16 × 10 ⁻¹⁰	-0.004	-0.004	0.109	-0.004	-0.004	0.136	-0.004	-0.004	0.136					

Note: Beta coefficients represent increase or decrease in DNA methylation in relation to NO₂ or PM_{2.5} exposure. In a preliminary step, DNA methylation β-values were regressed on the Illumina control probe-derived first 30 principal components to correct for correlation structures and technical bias, and residuals of these regressions covering 430,477 CpGs were used as the technical bias-corrected methylation level at the CpG sites. Extreme values of the residuals (lying beyond three times the interquartile range below the first quartile and above the third quartile at each CpG site) were replaced with their corresponding detection threshold value ("modified winsorization"). The "winsorized" data were then used as the dependent variables in the present EWAS. The multiexposure model contained all five exposures at same time. All SAPALDIA estimates were derived from linear mixed-effects EWAS models, with random intercept at the level of participant, adjusted for age; sex; educational level; area; neighborhood socioeconomic status; greenness index; smoking status; smoking pack-years; exposure to passive smoke; consumption of fruits, vegetables, and alcohol; nested study; asthma status; noise truncation indicators; survey; and leukocyte composition. Multiexposure model included all five exposures (Aircraft, railway, road traffic, Lden and respective truncation indicators, NO₂ and PM_{2.5}) at the same time. CHR, chromosome; COPD, chronic obstructive pulmonary disease; CpG, Cytosine-phosphate-Guanine; EPIC, European Prospective Investigation into Cancer and Nutrition; EWAS, epigenome-wide association study; N.A., not annotated; NL, Netherlands; NO₂, nitrogen dioxide; PM_{2.5}, particulate matter with aerodynamic diameter <2.5 μm; SAPALDIA, Swiss Cohort Study on Air Pollution and Lung and Heart Diseases in Adults; SE, standard error.

^aValidated in the SAPALDIA study.

^bSAPALDIA estimates derived from never-smoker sample comparable to the EPIC-NL never-smoker estimates.

methylation at certain sites and should be considered potential confounders in this and similar studies. Our findings in this EWAS (adjusted for leukocyte composition) are more likely to reflect actual DNA methylation changes rather than the underlying leukocyte composition changes in relation to transportation noise and air pollution exposure. This finding would otherwise be difficult to interpret, given the observed associations of exposures with leukocyte types.

The strengths of our study include being the first EWAS to assess mutually independent effects of source-specific transportation noise and air pollution exposures. Our study considered in parallel the independent effects of these exposures on DNA methylation at single CpG sites and in genomic regions. The availability of individually assigned, source-specific noise and air pollution data, derived from validated models, allowed the exploration of mutual confounding of these exposures on DNA methylation. We were able to control for potential confounders given the detailed characterization of the participants in the SAPALDIA study. The multiexposure approach also allowed the investigation of the independent pathway enrichment using both candidate and agnostic approaches to improve mechanistic understanding of transportation noise and air pollution exposures. We used a novel approach to investigate indirectly the effect of these exposures on physiological stress systems. We could also explore the influence of leukocyte composition, among other sensitivity analyses. The availability of genome-wide methylation data allowed the validation of previously reported air pollution signals, contributing to their external validity in the SAPALDIA study.

Our study is limited in its cross-sectional design, which precludes causal inferences and differentiating short-term vs. long-term exposure effects. Our noise and air pollution estimates may have been biased by errors in input data, which could be exposure-specific and potentially account for variations in observed effects. Such bias, however, would most likely be nonsystematic and non-differential. The air pollution and noise levels in the SAPALDIA study are relatively low and may have reduced statistical power of identifying signals and limited the replication of previous findings in settings of higher exposure. Unlike road traffic noise, which was ubiquitous, railway and aircraft noise in the SAPALDIA study were less common, with only ~45% exposure to these sources. Nevertheless, we made some significant observations. We could not identify any EWAS on stress hormones that directly captured alterations in the HPA axis; thus, our definition of “allostatic load” may have been biased. However, we expect any bias to be minimal, given our inclusion of CAR and other downstream biomarkers of physiological stress. As demonstrated in a recent review (Johnson et al. 2017), there is yet no consensus on which markers best capture “allostatic load.” We find it interesting that “allostatic load” score was always associated with negative health outcomes regardless of constituent biomarkers or phenotypes (Castagné et al. 2018; Johnson et al. 2017; Ribeiro et al. 2019).

Conclusions

DNA methylation was independently associated with transportation-related noise and air pollution exposures in the SAPALDIA study, with enrichment for pathways related to inflammation and immune response. Differential methylation due to these exposures may therefore explain the link between these exposures and several age-related outcomes. More EWAS with combined exposures and gene expression or transcriptome data are needed, especially from more polluted areas (including low- and middle-income countries), to corroborate present findings and to better capture the full extent and relevance of DNA methylation changes associated with these exposures. In particular, it remains to be seen if DNA methylation related to the identified

pathways mediates the association between transportation noise and air pollution exposures and incident age-related phenotypes.

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