Integrative multi-omics analysis of genomic, epigenomic, and metabolomics data leads to new insights for Attention-Deficit/Hyperactivity Disorder

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Abstract
The evolving field of multi-omics combines data and provides methods for simultaneous analysis across several omics levels. Here, we integrated genomics (transmitted and non-transmitted polygenic scores [PGSs]), epigenomics, and metabolomics data in a multi-omics framework to identify biomarkers for Attention-Deficit/Hyperactivity Disorder (ADHD) and investigated the connections among the three omics levels. We first trained single- and next multi-omics models to differentiate between cases and controls in 596 twins (cases = 14.8%) from the Netherlands Twin Register (NTR) demonstrating reasonable in-sample prediction through cross-validation. The multi-omics model selected 30 PGSs, 143 CpGs, and 90 metabolites. We confirmed previous associations of ADHD with glucocorticoid exposure and the transmembrane protein family TMEM, show that the DNA methylation of the MAD1L1 gene associated with ADHD has a relation with parental smoking behavior, and present novel findings including associations between indirect genetic effects and metabolites.

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and CpGs of the STAP2 gene. However, out-of-sample prediction in NTR participants (N = 258, cases = 14.3%) and in a clinical sample (N = 145, cases = 51%) did not perform well (range misclassification was [0.40, 0.57]). The results highlighted connections between omics levels, with the strongest connections between non-transmitted PGSs, CpGs, and amino acid levels and show that multi-omics designs considering interrelated omics levels can help unravel the complex biology underlying ADHD.

**KEYWORDS**
ADHD, DNA methylation, genetic nurture, metabolites, multi-omics, polygenic scores

1 | INTRODUCTION

Attention-Deficit/Hyperactivity Disorder (ADHD) in children is a highly heritable (75%–91%) complex neurodevelopmental disorder that is characterized by high levels of hyperactivity, inattention, and impulsiveness (Roehr, 2013). The prevalence of ADHD in children is 5%–8%, and a majority of these children continue to display symptoms in adulthood (Derks et al., 2008; Faraone et al., 2018; Levy et al., 1997; Luo et al., 2019). Persons with ADHD are more likely to be extroverted and entrepreneurs but also have an increased lifetime risk of harmful outcomes (Faraone et al., 2021; Martel et al., 2010). Multiple psychiatric conditions are comorbid with ADHD, including borderline personality disorder, anxiety, and conduct/oppositional defiant disorder, resulting in challenges with the diagnosis, referral, and treatment of ADHD (Caye et al., 2019; Demontis et al., 2019; Distel et al., 2011; Joseph et al., 2015; Reale et al., 2017).

Several single-omics studies involving genomics, epigenomics, and metabolomics have been performed to unravel the etiology of ADHD. Large (N = 55,374) Genome-Wide Association (GWA) studies initially identified 12 (Demontis et al., 2019) and recently 27 genomewide significantly associated loci for ADHD (Demontis et al., 2023). Demontis et al. (2019) compared these results with those for continuous measures of ADHD symptoms in the general population (Middeldorp et al., 2016), concluding that the same genetic variants that give rise to an ADHD diagnosis also affect inattention and impulsivity in the general population.

The biological basis of ADHD has also been explored in Epigenome-Wide Association (EWA) and metabolomics studies. Epigenomics constitutes a link between genetic and environmental exposures through changes in gene expression with processes such as DNA methylation (Barros & Offenbacher, 2009). Methylation level variation is associated with exposures such as stressful life events, nutritional factors, and toxins, and is partly due to genetic factors (Moore et al., 2013). Interestingly, it has been hypothesized that the decrease in hyperactivity with age might be due to epigenetic factors regulating the expression of genes involved in hyperactivity (Elia et al., 2012). EWA studies that measured DNA methylation in cord blood at birth, in saliva from older children, or in blood from adults start to provide evidence for DNA methylation sites associated with ADHD symptom trajectories and ADHD clinical diagnosis in children and adults (Goodman et al., 2020; Mooney et al., 2020; Neumann et al., 2020; Rovira, Sanchez-Mora et al., 2020; Walton, 2019). EWA studies based on blood samples from adolescents and adults identified multiple new candidate loci involved with immune and neuronal functioning associated with adult ADHD symptoms in population-based samples (van Dongen et al., 2019) and loci involved in cholesterol signaling in a comparison of participants with persistent ADHD and controls (Meijer et al., 2020).

Metabolomics studies utilize a profile of small molecules derived from cellular metabolism (X. Liu & Locasale, 2017). The first studies of metabolomics and ADHD identified multiple metabolites associated with ADHD in plasma (Wang et al., 2021). For instance, metabolites involved in oxidative stress pathway(s) were positively associated with ADHD in children. Meta-analyses showed that across all ages, proteins involved in dopamine metabolism and the fatty acid docosahexaenoic acid in different tissues were negatively associated with ADHD (Bonvicini et al., 2016, 2018). However, sample sizes of the metabolomics studies concerning ADHD are relatively small and it is likely that other metabolites and biological pathways influencing ADHD are still to be identified (Bonvicini et al., 2018).

The growing numbers of single-omics studies for ADHD indicate that large cohorts increasingly collect information across omics levels, including genomics, epigenomics, and metabolomics in relation to health and behavior. Associations and correlations have been observed between the genome and epigenome (Min et al., 2021; van Dongen et al., 2016), the genome and the metabolome (Hagenbeek et al., 2020; Hagenbeek et al., 2023), and the epigenome and the metabolome (Gomez-Alonso et al., 2021). Single-omics analyses alone fail to account for the interactions between these omics levels. To address this limitation, multi-omics approaches are the next step to understand how specific phenotypes like ADHD are linked to different omics levels and their interconnectedness (Hasin et al., 2017). Optimal analysis of multi-omics data in association and etiological studies requires dedicated statistical treatment of simultaneous omics influences (Duruffle et al., 2020). Such approaches complement single-omics techniques and may result in novel insights and uncover new biological pathways underlying traits and diseases (Rajasundaram & Selbig, 2016). For example, in cancer research, multi-omics findings
show promise in advancing personalized medicine by identifying disease markers (Chakraborty et al., 2018).

We applied an integrative multi-omics approach to elucidate biological mechanisms underlying ADHD and identify potential biomarkers. We integrated genetic data comprising genome-wide methylation data, and urinary metabolomic data with transmitted and non-transmitted polygenic scores (PGSs) for 15 traits, which are genetically correlated with ADHD. Here, the PGS based on alleles transmitted from parents to offspring represent the offspring’s own genetic liability, whereas the non-transmitted PGS represent the environmental influences that are associated with the parental with alleles that were not transmitted (e.g., genetic nurture; Kong et al., 2018). The study included 854 twins originating from the Netherlands Twin Register (NTR; Ligthart et al., 2019) and 145 non-twin children from a youth psychiatry clinic. We included non-transmitted PGSs to explore the environmental influences that children are exposed to that are created by the parents’ genetics. To our knowledge, this is one of the first studies to include non-transmitted PGSs in a multi-omics design.

2 | MATERIALS AND METHODS

2.1 | Study population and procedures

Participants were selected from the Young Netherlands Twin Register (YNTR; van Beijsterveldt et al., 2013) and from a youth psychiatry clinic (LUMC-Curium, the Netherlands), as part of the ACTION (Aggression in Children: Unraveling gene–environment interplay to inform Treatment and InterveniON strategies) Biomarker Study (Bartels et al., 2018; Hagenbeek et al., 2020). The ACTION project collected first-morning urine samples and buccal-cell swabs from 1494 twins (747 complete pairs) and 189 children from a clinical cohort with standardized protocols (see http://www.action-euproject.eu/content/data-protocols). The first-morning urine samples were stored in a freezer at the child’s home (T ≈ −18°C) and were transported to the lab in a mobile freezer unit at T = −18°C. In the lab, urine samples were stored at T = −80°C until further processing. The buccal swabs were collected on two consecutive days: twice in the morning (before breakfast) and twice in the evening (before dinner). In the twin cohort, buccal-cell swabs were also collected from the parents and siblings of the twins.

ADHD status in the twin cohort was based on sex- and age-specific T-scores from the mother-rated ADHD DSM-oriented scale of the ASEBA-CBCL as completed around the time of biological sample collection. T-scores were calculated in the entire NTR for children with CBCL data available at 9–10 years of age (N = 23,858). Twins with ADHD CBCL T-scores of 65 or higher were classified as cases (N = 125) and twins with T-scores below 65 as controls (N = 729). In the LUMC-Curium cohort, parents (90% mothers, 10% fathers) completed a CBCL as part of a standardized clinical assessment of a maximum of 6 months before or after biological sample collection. T-scores were calculated via CBCL software. Children with T-scores of 70 or higher were classified as cases (N = 74), children with T-scores below 65 as controls (N = 71), and children with T-scores between 65 and 69 were excluded (N = 40).

NTR and LUMC-Curium participants were excluded if no ADHD status at the time of biological sample collection was available (N = 209), if they were assigned a control status while their co-twin was assigned a case status for ADHD (N = 51), if the collected urine sample was not the first-morning urine (e.g., parent-reported time of urine collection was after 12:00 in the afternoon, N = 13), or if any of the omics data were not available because of poor quality (N = 367). From the twin cohort, 854 out of 1494 participants, and from the clinical cohort, 185 participants out of 189 had complete phenotype and omics data across all omics levels (genomics, epigenomics, and metabolomics).

2.2 | Genotyping and polygenic scores

2.2.1 | Genotyping and imputation

Genotyping in the NTR and LUMC-Curium cohorts was performed simultaneously on Affymetrix AXIOM or Illumina GSA arrays (Beck et al., 2019; Ehli et al., 2017), and genome-wide single nucleotide polymorphism (SNP) data were available for 3334 participants, including 1702 parents and siblings of twins (AXIOM = 909, GSA = 2425).

For each genotyping platform, methods were removed based on the following criteria: (1) DNA sex did not match self-reported sex; (2) Plink heterozygosity F statistic (<−0.10) or >0.10; and (3) genotyping call rate (>0.90). Criteria for excluding SNPs included: (1) minor allele frequency (MAF) ≤0.005; (2) Hardy–Weinberg Equilibrium (HWE) p-value (<1 × 10−5); (3) call rate (>0.95); (4) number of Mendelian errors (>2); and (5) palindromic AT/GC SNPs (0.4 ≤ MAF ≤ 0.5; Purcell et al., 2007). For each platform, data were aligned with the GoNL reference set (V4; Boomsma et al., 2014). SNPs with an allele frequency difference of larger than 0.10, or mismatching alleles with the reference panel were removed and samples were excluded if DNA identity by Descent (IBD) state did not match the expected familial relations (PLINKv1.9). Genotypes were then re-aligned, for each platform separately, to the 1000G Phase 3 version 5 reference panel by the PERL based “1000G Imputation preparation and checking” tool v4.3 (https://www.well.ox.ac.uk/~wrayner/tools/). The two platforms were subsequently phased (EAGLE v2.4.1) and imputed (MINIMAC3 v2.0.1) following the Michigan Imputation server protocols (Das et al., 2016; Delaneau & Zagury, 2012; Loh et al., 2016). After imputation, the resulting per-platform chromosomal files were merged into a single best-guess Plink file (PLINKv1.9). Data were available for 3334 participants: 3149 NTR participants (1447 twins, 1702 parents/siblings of twins), and 185 LUMC-Curium participants.

2.2.2 | Computation of principal components

From the best-guess 1000-genomes imputed data, a complete list of all platform-genotyped SNPs on both AXIOM and Illumina GSA were
extracted from the NTR and LUMC-Curium cohorts. This creates a single dataset without large SNP missingness where most genotypic information from both platforms is included. The same SNPs were extracted from the 1000G reference panel genotype data with additional filters for MAF (>0.05), and call rate (>0.98). In the 1000G population alone, the SNPs were then linkage disequilibrium (LD) pruned (PLINK1.9; –indep 50 5 2) and long-range LD regions were removed (Abdellaoi et al., 2013). Next, the NTR, LUMC-Curium, and 1000 genomes sets were merged for all SNPs that passed the above Quality Control (QC) criteria and were present in all sets. Subsequently, 20 principal components (PCs) were calculated for the 1000 genomes populations, and our sample was placed in the same dimensional PC space based on genetic similarity of the SNPs, to control the PGSs for stratification due to ancestry (SMARTPCav7; Price et al., 2006).

2.3 Transmitted and non-transmitted alleles and calculation of polygenic scores

The NTR cohort included 1447 twins with genotyped parents, for whom allele transmission could be established based on the imputed best guess data. Allele transmission could not be established in the LUMC-Curium cohort, as parental genotype data were not assessed. SNPs were excluded when: (1) MAF < 0.01; (2) HWE p-value < 1 x 10⁻⁵; (3) call rate < 0.98; (4) SNPs had duplicate positions; (5) SNPs had three or more alleles; and (6) non-ACGT SNPs on the autosomes. The non-transmitted allele status was defined by generating a single transmission-disequilibrium test (TDT) pseudo-control genotype for each child (given the two parents; Plink-tucc option) after defining all children as being cases (Clayton, 1999). To determine the maternal and paternal transmission of haplotypes, the transmitted and non-transmitted alleles datasets were phased (SHAPEITv2.r904). The resulting haplotypes were converted into mother and father non-transmitted homozygous haplotype genotypes for polygenic score analyses of non-transmitted alleles per parent.

In an earlier study, we applied integrative multi-omics models to childhood aggression (Hagenbeek et al., 2023). There, we included transmitted and non-transmitted PGSs for childhood aggression and 14 traits with significant (p < 0.02) large genetic correlations (≤−0.40 or ≥0.40) with aggression (Ip et al., 2021). As childhood aggression (Ip et al., 2021) had a genetic correlation with ADHD of 1.0028 (se = 0.0732, p = 9.39 x 10⁻⁴), we included PGSs for the same traits in the current study. PGSs were calculated for transmitted and non-transmitted alleles based on 15 discovery GWAS meta-analyses, which omitted NTR and LUMC-Curium from the discovery meta-analysis. The 15 traits comprised childhood aggression (Ip et al., 2021), ADHD (Demontis et al., 2019), Major Depressive Disorder (MDD; Wray et al., 2018), Autism Spectrum Disorder (ASD; Grove et al., 2019), loneliness (http://www.nealelab.is/uk-biobank/), insomnia (Jansen et al., 2019), self-reported health (http://www.nealelab.is/uk-biobank/), smoking initiation (ever/never smoked; M. Liu et al., 2019), age of smoking initiation (Watanabe et al., 2019), number of cigarettes per day (M. Liu et al., 2019), childhood IQ (Benyamin et al., 2014), educational attainment (EA; Lee et al., 2018), age at first birth (Barban et al., 2016), wellbeing spectrum (Baselmans et al., 2019), and intelligence (Savage et al., 2018; also see Hagenbeek et al., 2023). We calculated the PGSs in LDpred (Vilhjalmsson et al., 2015) which infers the posterior mean effect size of each marker by using a prior on effect sizes and LD information from an external reference panel. In 2500 second-degree unrelated individuals randomly selected from the NTR (the reference panel) the LD weighted betas for all traits were estimated (LDpred v0.9, fraction F of variants with nonzero effect F = 0.50). For all phenotypes we obtained a transmitted and two non-transmitted (one non-transmitted for the father and one for the mother) PGSs by scoring the LD corrected betas using PLINKv1.9. Thus, in total, 45 PGS were specified for each child. The effects of sex, age at biological sample collection, genotype platform, and the first 10 genetic PCs were regressed on the standardized (mean of 0 and standard deviation of 1) PGSs. The residuals of the above regressions were included in the analyses.

2.4 DNA methylation

Genome-wide DNA methylation for the NTR and LUMC-Curium cohorts were measured, according to the manufacturer’s specification, on the Infinium MethylationEPIC BeadChip Kit (Illumina, San Diego, CA, USA; Moran et al., 2016) by the Human Genotyping Facility (HugeF) of ErasmusMC (the Netherlands; http://www.glindna.org/), using the ZymoResearch EZ DNA Methylation kit (Zymo Research Corp, Irvine, CA, USA) for bisulfite treatment of 500 ng of genomic DNA obtained from buccal swabs. QC and normalization of the methylation data were carried out with pipelines developed by the Biobank-based Integrative Omics Study (BIOS) consortium as was previously described (Sinke et al., 2019). In short, samples were removed if they failed to pass all five quality criteria of MethylAid (van Iterson et al., 2014) if samples had incorrect relationships (omicsPrint; van Iterson et al., 2018), or sex mismatches (DNAArray; Min et al., 2018). Methylation probes were set to missing in a sample if they had an intensity value of zero, a bead count < 3, or a detection p-value >0.01. DNA methylation probes were excluded if they overlapped with a SNP or Insertion/Deletion (INDEL), mapped to multiple locations in the genome, or had a success rate <0.95 across all samples. After QC, 785,311 out of 865,859 sites were retained for analysis for 1424 samples, and cellular proportions were predicted in epithelial tissues using the cell-type deconvolution algorithm Hierarchical Epigenetic Dissection of Intra-Sample-Heterogeneity (HepiDISH) with the reduced partial correlation (RPC; Zheng et al., 2018). Missing methylation β-values were imputed (missMDA; Sinke et al., 2019), and two duplicate samples of NTR participants were excluded. One sample was excluded from the LUMC-Curium dataset because this participant and co-twin were also included in the NTR. The effects of sex, age, percentages of epithelial and natural killer cells, EPIC array row, and bisulfite sample plate were regressed from the methylation β-values of the top 10% most variable methylation sites (78,772 in total) that survived QC (see Hagenbeek
et al., 2023). The resulting residual methylation levels were included in the analyses.

2.5 | Metabolomics

Urinary metabolomics data were generated by the Metabolomics Facility of Leiden University (Leiden, the Netherlands) on three platforms: (1) a liquid chromatography-mass spectrometry (LC–MS) platform targeting amines; (2) an LC–MS platform targeting steroid hormones; and (3) a gas chromatography–mass spectrometry (GC–MS) platform targeting organic acids. Subjects were randomized across batches, while retaining twin pairs on the same plate. Each batch included a calibration line, QC samples (pooled aliquots from all urine samples; every 10 samples), sample replicates, and blanks. In-house developed algorithms were applied, using the pooled QC samples, to compensate for shifts in the sensitivity of the mass spectrometer across batches. Metabolites were reported as "target area/area of internal standard" after QC correction, and metabolites with a relative standard deviation of the QC samples larger than 15% were excluded (six amines, three steroids, and one organic acid). Metabolite measurements that fell below the limit of detection/quantification were imputed with half of the value (with internal standards) urine to extract the organic acids and remove urea. Online derivatization procedures were performed in two steps: (1) oxidation with methoxamine hydrochloride (MeOX, 15 mg/mL in pyridine); and (2) N-Methyl-N-(trimethylsilyl)-trifluoroacetamide (MSTFA) silylation. Chromatographic separation was performed on a 25 m (HP–5MS UI) film thickness 30 × 0.25 m ID column, with helium as a carrier gas (1.7 mL/min). The mass spectrometer (Agilent Technologies, Waldbronn, Germany), using a single quadrupole with electron impact ionization (70 eV) was operated in SCAN mode (mass range 50–500), using 1 μL of the sample. In total, 21 organic acids were successfully measured.

2.6 | Statistical analyses

To examine molecular variation associated with ADHD for PGSs, DNA methylation, and metabolomics, and to build a predictive model for ADHD, we applied an integrative multi-omics method called sparse multi-block supervised analysis (Rohart et al., 2017; Singh et al., 2019). All analyses were performed in R (version 4.1.1), mainly in the mixOmics package (version 6.16.3; Rohart et al., 2017). The analytical design consisted of three steps (Hagenbeek et al., 2023): (1) three single-omics analyses; (2) two sets of pairwise cross-omics analyses; and (3) a multi-omics analysis (Figure 1). The twin sample was randomly split at twin pair level to create two subgroups with 70% of the data for model training (training data), and 30% of the data for model testing (test data; Table 1). The performance of final single- and multi-omics models was explored in a follow-up clinical sample. All single- and multi-omics models employed five-fold cross-validation (CV) with 50 repeats to determine the optimal number of components for each model (the mixOmics perf function), for variable selection in each model (the mixOmics perf function), and for assessment of final model performance in the training data (the mixOmics perf function). For these multi-omics analyses there is a strong requirement of no missing data at any omics level. To keep the input data the same for the single- and multi-omics models, we included participants with complete omics data in all analyses.

2.6.1 | Step 1: single-omics analyses

Principal Component Analysis (PCA) was run separately for a total of 45 transmitted and maternal and paternal non-transmitted PGSs, 78,772 CpGs in the DNA methylation data, and all 90 metabolites in the training data to obtain a first insight into the dimensionality in each omics level (Figure S1). Next, Partial Least Square Discriminant Analyses (PLS-DA) and sparse PLS-DA (sPLS-DA) were applied to the training data to assess the ability of each of the three omics levels to correctly classify ADHD status. CV determined the optimal number of components to include in each PLS-DA model (Table S1; Figure S2), determined the optimal number of components, and reduced the number of variables in each omics layer that contribute to each component in the sPLS-DA models (Table S1; Figure S3), and assessed the
Analyses

Demographics of the training, test, and follow-up data.

Workflow of the analyses performed in this study. We employed an analytical design consisting of three phases: (1) single-omics analyses; (2) pairwise cross-omics analyses; and (3) multi-omics analyses. First, we built single-omics biomarker panels in the twin cohort, with 70% of the twin data for model training (training data), 30% of the twin data for model testing (test data), and the clinical cohort (follow-up data). Second, the overall pairwise cross-omics correlations among the loading scores of the PLS components for all omics traits and the selected omics variables in the single-omics models were examined in the training data. Third, using the same data split for model training, testing, and follow-up, we built multi-omics biomarker panels and described the multi-omics connections of the selected omics traits. The details about the analyses are provided in Section 2. PLS, Partial Least Squares.

![Table 1](https://onlinelibrary.wiley.com/doi/10.1002/ajmg.b.32955)

**Table 1** Demographics of the training, test, and follow-up data.

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<thead>
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<tr>
<td></td>
<td>• Principal Component Analysis (PCA)</td>
<td>• 70% twin cohort (N = 596)</td>
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<tr>
<td></td>
<td>• Partial Least Squares Discriminant Analysis (PLS-DA)</td>
<td>• Cases: 88 (14.8%)</td>
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<tr>
<td></td>
<td>• Sparse Partial Least Squares Discriminant Analysis (sPLS-DA)</td>
<td>• Controls: 508 (85.2%)</td>
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<td></td>
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<td></td>
<td>• Predict out-of-sample case-control status</td>
<td>• 30% twin cohort (N = 258)</td>
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<tr>
<td></td>
<td>• Receiver Operating Characteristic (ROC) analysis</td>
<td>• Cases: 37 (14.3%)</td>
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<td></td>
<td></td>
<td>• Controls: 221 (85.7%)</td>
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<td><strong>All omics traits:</strong></td>
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<td>• Selected omics traits:</td>
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<td>• Partial Least Squares (PLS) analysis</td>
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<td></td>
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<td>• 70% twin cohort (N = 596)</td>
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<td></td>
<td>• Multi-block sparse Partial Least Squares Discriminant Analysis (MB-sPLS-DA)</td>
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<tr>
<td>3. Multi-omics</td>
<td><strong>Model testing &amp; validation:</strong></td>
<td><strong>Test data:</strong></td>
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<td></td>
<td>• Predict out-of-sample case-control status</td>
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**Figure 1** Workflow of the analyses performed in this study. We employed an analytical design consisting of three phases: (1) single-omics analyses; (2) pairwise cross-omics analyses; and (3) multi-omics analyses. First, we built single-omics biomarker panels in the twin cohort, with 70% of the twin data for model training (training data), 30% of the twin data for model testing (test data), and the clinical cohort (follow-up data). Second, the overall pairwise cross-omics correlations among the loading scores of the PLS components for all omics traits and the selected omics variables in the single-omics models were examined in the training data. Third, using the same data split for model training, testing, and follow-up, we built multi-omics biomarker panels and described the multi-omics connections of the selected omics traits. The details about the analyses are provided in Section 2. PLS, Partial Least Squares.

**Abbreviations:** ADHD, Attention-Deficit/Hyperactivity Disorder; MZ, monozygotic.

*Measured with the mother-rated Attention Deficit/Hyperactivity Problems DSM-oriented scale of the Achenbach System of Empirically Based Assessment (ASEBA) Child Behavior Checklist (CBCL). The ASEBA CBCL Attention Deficit/Hyperactivity Problems DSM-oriented scale scores in the clinical cohort include 90% mother report and 10% father report.
performance of the final sPLS-DA model (Table S1; Figure S4). Three types of distance measurements were assessed: (1) centroid distance, (2) Mahalanobis distance, and (3) maximum distance. The distance that showed the best predictive performance in the training data was selected for the final models.

We evaluated the out-of-sample predictions in the test and follow-up data (mixOmics predict function) based on the best-performing model (Table S1). We applied three methods to evaluate how well the final models classified case–control status. First, we employed a balanced misclassification rate, the balanced error rate (BER; false-negative + false-positive rate), that corrects for imbalances in the number of cases and controls. Second, we calculated the sensitivity, specificity, and accuracy of the single-omics models by comparing the predicted cases and controls with the true cases and controls. Third, we assessed the area under the curve (AUC) for each component in both the test and follow-up data with receiver operating characteristic (ROC) analysis.

2.6.2 | Step 2: Pairwise cross-omics analyses

To investigate pairwise cross-omics connections (i.e., PGSs-DNA methylation, PGSs-metabolomics, and DNA methylation-metabolomics) we constructed PLS regression models in canonical mode in the training data. For each combination of omics data, we constructed PLS models including different numbers of components. Two components were included for the PGS-DNA methylation model, two for the PGS-metabolomics model, and five for the DNA methylation-metabolomics model. This resulted in a total of two sets of PLS models, each containing three models corresponding to the pairwise combinations of omics data. The first set of models included all 45 PGSs, 78,772 CpGs, or 90 metabolites, allowing us to examine the correlations of the loading scores among all omics traits. In the second set of models, we focused on the 17 PGSs, 486 CpGs, and 90 metabolites selected by the single-omics sPLS-DA models (Data S1), which were found to be the most relevant for classifying ADHD cases and controls. By performing hierarchical clustering using the Ward linkage algorithm on Euclidean distances of the PLS variates, we identified biologically meaningful clusters (‘dendextend’ R-package; Galili, 2015). We specifically extracted the two largest clusters for each of the omics levels considered in the PLS models.

2.6.3 | Step 3: Multi-omics analyses

The multi-omics analysis was conducted through multi-block sPLS-DA (MB-sPLS-DA) in the training data including an empirical design matrix, based on the correlations among the loading scores from the pairwise cross-omics PLS models including all omics traits. The design matrix comprised of correlations of 0.25 between the PGSs and DNA methylation, 0.29 between the PGSs and the metabolites, and 0.23 between DNA methylation and the metabolites (based on pairwise cross-omics analyses). CV determined the optimal number of components to include in the MB-PLS-DA models, the number of traits to include per component per omics layer, and the performance of the final MB-sPLS-DA model (Table S2; Figures S5–S7). The accuracy of out-of-sample case–control status prediction was again evaluated in the test and follow-up data, with final multi-omics models evaluated by their BER and AUCs (ROC curves were calculated per component for each omics layer), and the model sensitivity, specificity, and accuracy were calculated from the confusion matrices.

2.7 | Biological characterization

We describe the correlations among the loading scores of the PLS components of the PGSs, CpGs, and metabolites that were selected by the single-omics sPLS-DA and the multi-omics MB-sPLS-DA models to facilitate high correlation patterns suitable for biological interpretation. We performed enrichment analysis for the CpGs selected across our analyses for all phenotypes (618) in the EWAS atlas on December 10, 2021 (Xiong et al., 2022). We performed enrichment on (1) the CpGs that were selected by the single-omics sPLS-DA model, (2) the CpGs in the clusters identified by the pairwise cross-omics analyses, (3) the CpGs that were selected for the multi-omics MB-sPLS-DA model, and (4) the CpGs that were included in the high correlation patterns of the multi-omics MB-sPLS-DA. When fewer than 20 CpGs were selected, the trait associations with the CpGs were manually retrieved from the EWAS atlas. We focused on the enrichment analysis on the CpGs since there is no similar tool to the EWAS atlas for the metabolites.

3 | RESULTS

3.1 | Single-omics models for ADHD

Based on the sPLS-DA analyses for PGSs, DNA methylation, or metabolomics data we built single-omics prediction models for ADHD case–control status. The single-omics models included 17 PGSs, 486 CpGs, and all 90 metabolites (Table S1; Data S1). The 17 selected PGSs in the sPLS-DA model comprised the transmitted PGS for ADHD, smoking initiation, MDD, EA, number of cigarettes per day, and insomnia. The non-transmitted maternal PGSs were selected for insomnia, self-reported health, childhood aggression, age of smoking initiation, and childhood IQ. The non-transmitted paternal PGSs were selected for ADHD, intelligence, ASD, EA, number of cigarettes per day, and the age of smoking initiation (Data S1). Trait enrichment analyses for all 486 selected CpGs in the sPLS-DA model showed the strongest enrichment for glucocorticoid exposure (i.e., based on an EWAS of administration of corticosteroid medication and DNA methylation measured in buccal cells [Braun et al., 2019]; N Overlap Differently Methylated CpGs (DMC) = 13, OR = 7.66, \( p = 1.12 \times 10^{-14} \)), and ancestry (DMC = 15, OR = 2.61, \( p = 5.73 \times 10^{-6} \); Table S3). Model performance was sub-optimal in the test and follow-up data for all three omics levels (AUC range = [0.45, 0.60], Table S4).
3.2 | PGS-DNA methylation analysis

An average correlation of \( r = 0.23 \) (\( q = 8.92 \times 10^{-16} \)) was found between the PGSs and DNA methylation in the loadings scores of the pairwise PLS cross-omics model. For the pairwise model, we applied hierarchical clustering which identified two clusters for the PGS and DNA methylation data (Figure 2a; Data S2). Cluster 1 contains eight of the PGSs (47.1%) selected by the sPLS-DA model for ADHD, and comprised the transmitted PGSs for ADHD, insomnia, MDD, smoking initiation, and the number of cigarettes per day, the non-transmitted paternal PGSs for insomnia and childhood aggression, and the non-transmitted paternal PGS for ADHD. The nine PGSs (52.9%) included in cluster 2, comprised the transmitted PGS for EA, the non-transmitted maternal PGSs for age at smoking initiation, childhood IQ, and self-reported health, and the non-transmitted paternal PGSs for age at smoking initiation, ASD, the number of cigarettes per day, and intelligence. The DNA methylation cluster 1 contains 226 (46.5%) of the CpGs selected by the sPLS-DA models for ADHD and showed the strongest trait enrichments for glucocorticoid exposure (DMC = 13, \( OR = 12.94, \ p = 6.64 \times 10^{-31} \)), and trihalomethanes (THM) exposure (DMC = 2, \( OR = 33.89, \ p = 1.85 \times 10^{-11} \) Table S5). The 260 (53.5%) CpGs included in cluster 2 showed the strongest trait enrichment for ancestry (DMC = 13, \( OR = 4.32, \ p = 3.30 \times 10^{-7} \)), and childhood stress (DMC = 3, \( OR = 20.69, \ p = 7.20 \times 10^{-7} \)). The correlations of the CpGs in clusters 1 and 2 with the CpGs ranged from \(-0.15\) to \(0.13\) and from \(-0.20\) to \(0.21\), respectively, and the correlations of the CpGs in clusters 1 and 2 with the CpGs ranged from \(-0.20\) to \(0.21\) and from \(-0.17\) to \(0.16\), respectively (Figure 2a; Data S2 and S3).

3.3 | PGSs-metabolomics analysis

Hierarchical cluster of the pairwise PGSs-metabolomics model, with an average correlation between the loading scores of \( r = 0.26 \) (\( q = 1.02 \times 10^{-17} \)), identified two clusters of PGSs and two clusters of metabolites (Figure 2b; Data S2). The PGS cluster 1 contains nine of the PGSs (52.9%) selected by the sPLS-DA model for ADHD, comprising the transmitted PGS for ADHD, insomnia, and MDD, the non-transmitted maternal PGSs for insomnia, childhood aggression, and self-reported health, and the non-transmitted paternal PGSs for ADHD, ASD, and intelligence. The eight PGSs (47.1%) included in cluster 2 comprises the transmitted PGSs for the number of cigarettes per day, EA, smoking initiation, the non-transmitted maternal PGSs for smoking initiation, and the number of cigarettes per day, the non-transmitted paternal PGSs for age at smoking initiation, ASD, and self-reported health, and the non-transmitted paternal PGSs for EA, and age at smoking initiation. Metabolite cluster 1 contains 48 metabolites (53.3%), all of which are amines (80% of all amines). The remaining 12 amines, as well as all steroids and organic acids, have been included in metabolite cluster 2 (\( N = 42, 46.7\% \)). Overall, we observed correlations ranging from \(-0.17\) to \(0.25\) for the PGSs in cluster 1 with metabolites, and from \(-0.22\) to \(0.16\) for the cluster 2 PGSs with metabolites (Figure 2b; Data S2 and S4). For the metabolites in cluster 1 the correlation ranged between \(-0.22\) and \(0.16\) with PGSs, and for cluster 2 between \(-0.17\) and \(0.25\).

3.4 | DNA methylation-metabolomics analysis

The pairwise cross-omics DNA methylation-metabolomics model showed an average correlation of \( r = 0.23 \) (\( q = 3.95 \times 10^{-27} \)) and hierarchical clustering identified two clusters of CpGs and of metabolites (Figure 2c; Data S2). The DNA methylation cluster 1 contains 241 (49.6%) of the CpGs selected by the sPLS-DA models for ADHD, and the strongest trait enrichments for these CpGs were observed for glucocorticoid exposure (DMC = 13, \( OR = 12.08, \ p = 7.63 \times 10^{-30} \)), and THM exposure (DMC = 2, \( OR = 31.79, \ p = 3.22 \times 10^{-11} \) Table S5). The 245 (50.4%) CpGs included in cluster 2 show the strongest trait enrichment for ancestry (DMC = 13, \( OR = 4.60, \ p = 9.96 \times 10^{-10} \)), childhood stress (\( OR = 21.97, \ p = 5.12 \times 10^{-7} \)), and household socioeconomic status in childhood (DMC = 3, \( OR = 18.34, \ p = 7.20 \times 10^{-7} \)). Metabolite cluster 1 contains 43 (47.8%) of the metabolites, including 42 amines and 1 organic acid, while metabolite cluster 2 contains 47 metabolites (52.2%), including 18 amines, 19 organic acids, and all 10 steroids. The correlation of the CpGs included in cluster 1 with metabolites range from \(-0.23\) to \(0.29\) (Figure 2c; Data S2 and S5). The CpGs included in cluster 2 have correlations ranging from \(-0.27\) to \(0.28\) with metabolites. For the metabolites in cluster 1, the correlations with CpGs ranged from \(-0.26\) to \(0.25\), and for cluster 2 from \(-0.27\) to \(0.29\).

3.5 | Multi-omics model for ADHD

We built a multi-omics panel for ADHD based on a multi-block sPLS-DA (MB-sPLS-DA) model including PGSs, DNA methylation, and metabolomics data. The four-component model included in total 30 PGSs, 143 CpGs, and all 90 metabolites (Table S2; Data S6). The number of components and number of variables per omics layer were determined by the single-omics models. The included variables in the multi-omics model were selected based on CV. The multi-omics model comprised the transmitted PGS for ADHD, aggression, cigarettes per day, insomnia, loneliness, childhood IQ, self-reported health, age at first birth, intelligence, smoking initiation, EA, MDD, and wellbeing. The non-transmitted maternal PGSs were selected for aggression, cigarettes per day, insomnia, loneliness, childhood IQ, self-reported health, and age smoking initiation. The non-transmitted paternal PGSs were selected for ADHD, aggression, cigarettes per day, insomnia, loneliness, age at first birth, intelligence, smoking initiation, age smoking initiation, and ASD. Trait enrichment analyses for all selected CpGs in the MB-sPLS-DA model showed the strongest enrichment for childhood stress (DMC = 3, \( OR = 38.00, \ p = 2.25 \times 10^{-8} \)), and respiratory allergies (AR; DMC = 2, \( OR = 28.48, \ p = 1.58 \times 10^{-5} \); Table S6). The multi-omics prediction in the test data showed a better prediction (0.42 ≤ BER ≤ 0.51; 0.43 ≤ AUC ≤ 0.62) as compared to single-omics models, while prediction in the follow-up data was similar to those.
FIGURE 2  Clustered heatmaps of the correlations among the loading scores of the Partial Least Squares (PLS) components including only the omics traits as selected by the single-omics sparse Partial Least Squares Discriminant Analyses (sPLS-DA). The hierarchical clustering was generated using the Ward linkage algorithm on Euclidean distances of the PLS variates. Two clusters have been identified for each dendrogram (cluster 1 = pink, cluster 2 = blue). Positive correlations among the omics traits have been depicted in red and negative correlation in blue. For the polygenic scores (PGSs), the “_NTm” suffix denotes PGSs non-transmitted by mother, the “_NTf” suffix denotes the PGSs non-transmitted by father, and childhood aggression is abbreviated as “aggression,” Attention-Deficit Hyperactivity Disorder as “ADHD,” Major Depressive Disorder as “MDD,” Autism Spectrum Disorder as “Autism,” and Educational Attainment as “EA.” For the metabolites, the “amines.” prefix indicates these metabolites were measured on the liquid chromatography-mass spectrometry (LC–MS) amines platform, the “steroids.” prefix indicates these metabolites were measured on the LC–MS steroids platform, and the “OA.” prefix indicates these metabolites were measured on the gas chromatography–mass spectrometry (GC–MS) organic acids platform. (a) Correlation among the 17 PGSs and 486 CpGs included in the two-component PGS-DNA methylation PLS model, where the selected CpGs are represented in the rows and the selected PGSs in the columns. The cluster assignments and the full correlation matrix are included in Data S2 and S3. (b) Correlations among the 17 PGSs and 90 metabolites included in the two-component PGS-metabolomics PLS model, where the selected metabolomics traits are represented in the rows and the selected PGSs in the columns. The cluster assignments and the full correlation matrix are included in Data S2 and Data S4, respectively. (c) Correlation among the 486 CpGs and 90 metabolites included in the five-component DNA methylation-metabolomics PLS model, where the selected metabolomics traits are represented in the rows and the selected CpGs in the columns. The cluster assignments and the full correlation matrix are included in Data S2 and Data S5.
observed for the single-omics models (0.49 ≤ BER ≤ 0.55; 0.50 ≤ AUC ≤ 0.53; Table S7; Table S4).

The average correlations among the loading scores of the PLS components between each omics layer in the multi-omics model were $r = 0.17$ ($q = 1.41 \times 10^{-16}$) for PGS-DNA methylation, $r = 0.14$ ($q = 1.24 \times 10^{-11}$) for PGSs-metabolomics, and $r = 0.20$ ($q = 1.1 \times 10^{-21}$) for DNA methylation-metabolomics. We observed high absolute correlations ($r \geq 0.60$) between 4 selected PGSs, 19 CpGs, and 10 metabolites, which can be summarized in three sets of correlational patterns (Figure 3; Data S7; Data S8). Correlation

**FIGURE 3** High cross-omics correlations among the loading scores of the PLS components of the multi-omics traits identified in the four-component multi-block sparse Partial Least Squares Discriminant Analysis (MB-sPLS-DA) model. The outer ring depicts the polygenic scores (PGSs), CpGs, and metabolites in yellow, pink, and green, respectively. For the PGSs, Educational Attainment is abbreviated as “EA,” and the “_NTm” suffix denotes the PGSs non-transmitted by mother. The inner plot depicts the correlations among the omics traits. For the metabolites, the “amines.” prefix indicates these metabolites were measured on the liquid chromatography-mass spectrometry (LC–MS) amines platform, and the “OA.” prefix indicates these metabolites were measured on the gas chromatography–mass spectrometry (GC–MS) organic acids platform. Here, only high absolute correlations ($r \geq 0.60$) between traits of at least two omics levels are depicted, with blue lines reflecting negative correlations and red lines positive correlations. Correlations are averaged across all components in the MB-sPLS-DA model. The full correlation matrix is included in Data S7 and the correlational patterns are included in Data S8.
pattern 1 comprises correlations of CpGs located in the STAP2 gene with the transmitted ADHD PGS (r range: −0.62 to −0.61), the transmitted self-reported health PGS (r range = [0.63, 0.66]), and 11 amino acids (r range = [−0.71, −0.60]). Correlation pattern 2 comprises correlations between eight CpGs and the non-transmitted paternal cigarettes per day PGS (r range = [−0.70, −0.61]). Two of these CpGs (cg15914316 and cg25352924) are located in the MAD1L1 gene (chromosome 7). Correlation pattern 3 comprises correlations between five CpGs and the transmitted childhood IQ PGS (r range = [−0.60, 0.67]). Three of these five CpGs are located in genes coding for transmembrane proteins (C7orf49, TMEM100, TMEM135, and TMEM140). The other two CpGs (cg10864680 and cg06843189) are located in TNFSF13B (chromosome 13) and NR3C2 (chromosome 4), respectively.

4 | DISCUSSION

We present an integrative multi-omics analysis of childhood ADHD based on genomics, epigenomics, and metabolomics data to identify multi-omics biomarkers and to investigate the connections among the different omics levels. The single-omics models identified 17 PGSs, 486 CpGs, and 90 metabolites as relevant variables, while the multi-omics model selected 30 PGSs, 143 CpGs, and 90 metabolites. Several of these selected variables showed support for known associations with ADHD. However, the out-of-sample predictions from both the single- and multi-omics models were sub-optimal, with BER ranging from 0.40 to 0.57 and 0.42 to 0.55, respectively. We also evaluated the pairwise associations between variables and found that the averaged correlations range from 0.23 to 0.29. In the multi-omics model, the connections among the ADHD biomarkers highlighted several genomic regions that may be involved in the disorder. Additionally, our findings further supported previously identified markers such as the gene MAD1L1, childhood stress, and glucocorticoid exposure.

The single-omics models selected 17 PGS based on 12 phenotypes, while the multi-omics model selected 30 PGSs comprising all 15 phenotypes. Transmitted PGSs were selected for ADHD, insomnia, MDD, EA, smoking initiation, and cigarettes per day. These traits are genetically correlated with ADHD (Demontis et al., 2019, 2021). One study that looked at the intergenerational transmission of EA and ADHD found that transmitted PGSs influenced ADHD behavior at school (de Zeeuw et al., 2020). For traits such as aggression, self-reported health, age of smoking initiation, intelligence, childhood IQ, and autism only non-transmitted PGSs were selected. This may represent an important observation: the genotype of the parents can help predict the ADHD status of their offspring when we consider their non-transmitted scores across a broad range of traits. We note that the predictive accuracy of the model is currently low but may improve when PGSs improve with increasing GWAS sample size.

The CpGs in the single-omics model were enriched for multiple traits and disorders previously associated with ADHD, such as glucocorticoid exposure, Klinefelter syndrome, and childhood stress (Friedrichs et al., 2012; Grimm et al., 2020; Harpin, 2005; Lo-Castro et al., 2011; Xiong et al., 2022). In the single- and multi-omics models, we observed enrichment for traits involved in diet exposure, such as glucocorticoid and vitamin B12 levels. We also found associations for THM exposure, a class of chemical compounds that can be present when water is disinfected with chlorine (Hood, 2005). Still, we note that the number of overlapping DMCs with all known DMCs in the EWAS atlas was usually relatively small. The multi-omics model selected two novel CpGs located in the gene MAD1L1 previously linked to ADHD in blood and saliva tissue (Goodman et al., 2020; Mooney et al., 2020; Rovira, Sanchez-Mora, et al., 2020). Another finding for rs11982272 located in MAD1L1 has previously been reported as a shared risk locus between ADHD and disruptive behavior disorders (DBD) but was only moderately associated with ADHD in a large GWAS (Demontis et al., 2021). MAD1L1 was also identified in studies focusing on schizophrenia and bipolar disorder, suggesting that MAD1L1 is a risk gene for several psychiatric disorders (Ikeda et al., 2018; Ripke et al., 2014).

The CpGs in the multi-omics models were connected with the non-transmitted paternal cigarettes per day PGS in our study. The connection suggests that the association between ADHD and the methylation around MAD1L1 might be caused by the effects of paternal smoking or that paternal smoking is a confounder between ADHD and the methylation around MAD1L1 (e.g., parental smoking leads to both ADHD and DNA methylation, but there is no connection between the DNA methylation and ADHD). Still, to our knowledge this is the first observation of paternal smoking effects on ADHD and paternal smoking should merit investigation in future studies. An earlier multi-omics study identified several genes whose expression or DNA methylation level in the (fetal) brain mediates the effects of genetic variants on ADHD, including a gene for a transmembrane protein TMEM125 (Hammerschlag et al., 2020). We observed four CpGs located in three genes of this same gene family (TMEM100, TMEM135, and TMEM140) had a connection with the transmitted childhood IQ PGS. The findings of MAD1L1 and the TMEM family support the results of the previous studies and imply an effect of these genomics regions on ADHD.

Both the single- and multi-omics models selected all 90 metabolites, which is not unexpected since the data were generated on targeted platforms, covering metabolites previously associated with aggression (Hagenbeek et al., 2016), and aggression is the trait which is one of the strongest associated variables with ADHD in childhood (Bartels et al., 2018). Each metabolite by itself has a small effect and single and multi-omics models are not equipped to identify these small effects, but a combination of all metabolites discriminates better between ADHD cases and controls.

We observed three connection patterns in our multi-omics analysis where the omics variables showed correlation coefficients of \( r \geq 0.60 \). As a result, we can make several observations about these patterns. First, only amines (mostly essential amino acids) showed high negative correlations with CpGs (\( r \) range = [−0.71, −0.60]) and we observed no high correlations between the organic acids and the steroids, and the other omics levels. Recent studies have linked several of these essential amino acids, such as glycine, serine, leucine, and...
valine, to childhood ADHD (Anand et al., 2021; Yu et al., 2021). Essential amino acids, are amino acids that that come from dietary intake. This suggests a relationship between diet, DNA methylation, and metabolites involved in ADHD and that DNA methylation mediates the effect of diet on metabolite levels. (Dekkers et al., 2016). A previous study suggested IGF2 methylation mediates the effect of prenatal diet on ADHD symptoms in childhood (Rijlaarsdam et al., 2017). Our observation concerning the amino acids supports hypotheses that essential amino acids affect childhood ADHD, and that diet might help to improve the symptoms of ADHD (Anand et al., 2021; Hamza et al., 2019; Yu et al., 2021). We would like to point out that, even though we highlight these associations, we cannot make claims about cause and effect. Our study identified associations between ADHD and amino acids, which indicate that it is worthwhile to further examine the possibility of causal relations among diet, DNA methylation, and amino acids.

Another multi-omics connection pattern comprised the transmitted PGSs for ADHD and self-reported health with 6 CpGs and 10 amino acids. The six CpGs are located in the STAP2 gene (Signal Transducing Adaptor Member 2) on chromosome 19. No SNPs or CpGs in or around this gene were associated with ADHD in GWA or EWA studies, suggesting STAP2 is a novel genomic region that could be involved in ADHD (Demontis et al., 2019, 2021; Goodman et al., 2020; Mooney et al., 2020; Neumann et al., 2020; Rovira, Demontis, et al., 2020; Rovira, Sanchez-Mora, et al., 2020; Walton, 2019; Walton et al., 2017). The six CpGs and the STAP2 gene have been associated with other traits such as smoking behavior, aging, and type 2 diabetes in blood (Xiong et al., 2022; Data 58). Although we observed no connection between these CpGs and the smoking-related PGSs, it should be taken into account that smoking PGSs are an imperfect measure of overall smoking exposure.

A challenge in machine learning via CVs is the risk of leaky preprocessing: data processing that inadvertently causes information to leak from the training set to the test set (Whalen et al., 2021). We preselected the top 10% most variable methylation sites before the split in training and test set to limit the computational burden of our method. We selected these methylation sites solely based on the criterion of variability, regardless of any association of these probes with a phenotype. Since we randomly selected test and training set samples, we may assume the same distributions in both sub-samples.

A source of confounding in the ACTION cohort are the effects of puberty. We do not, however, expect puberty to affect the results in any major way, given the ages of the subjects and the fact that only three females had had their first period.

A major limitation of most multi-block multi-omics methods, including MB-sPLS-DA, is that these methods do not allow for any missing values in the training data on any of the included omics levels. For our study, this meant we had to exclude 367 participants (almost one third of the original sample) because of missing or poor-quality data in at least one of the omics levels. Allowing missing data is an important improvement that multi-omics methods need to address in the future.

In contrast to previous multi-omics studies for ADHD, which relied on sequential integration of summary statistics (Cabana-Dominguez et al., 2022; Cheng et al., 2020; Hammerschlag et al., 2020), our multi-block integration approach allowed simultaneous modeling of multiple omics levels and gain insights into the relationships among the different omics levels (Wörheide et al., 2021). Unsupervised multi-omics methods, such as PLS structural equation modeling (PLS-SEM; Csala et al., 2020) or Multi-Omics Factor Analysis (MOFA, Argelaguet et al., 2018), constitute powerful alternatives to explore the relationships among omics levels, but are not appropriate for classification of groups, as was one of the aims of this study. Another asset of our study is that by employing an integrative multi-omics design with harmonized sample collection and measurement protocols in the twin and the clinical groups, we reduced the detrimental heterogeneity due to, for example, differences in sample size for different omics layers, or study protocols as far as possible (Wörheide et al., 2021). Problems with design and technical heterogeneity are not uncommon in sequential integration methods. In the current study, such undesirable sources of technical heterogeneity were minimized by our choice of analytical design and by streamlining the collection protocol for biological samples across twins from the NTR and participants from LUMC-Curium. We collected the biological samples in both cohorts in parallel and generated the omics data for both cohorts together. Other than the difference in cohort type, population versus clinical, the main difference between these cohorts is the absence of genotyping information for the parents of the children included in LUMC-Curium and non-transmitted PGSs could not be assessed in the clinical cohort.

5 | CONCLUSION

Our study underscores the complexity of ADHD, as it remains a challenge to detect multi-omics biomarkers and indicates that clinically useful prediction models based on omics data are not viable yet. We showed how different omics levels involved in ADHD are connected to each other, leading to new hypotheses about cross-omics interactions. The connection patterns reproduced previously identified associations with ADHD, including the MAD1L1 gene and glucocorticoid exposure, while also highlighting new interesting genomic regions such as the STAP2 gene. Multi-omics methods are expected to become increasingly important in the translation from association studies to clinical applications and it is of value to continue to use them. The integrated approach we applied in this study requires raw data analysis and demands that all participants are measured across all omics levels. Solving the challenge of missing data in multi-omics approaches will lead to increased sample sizes and increased power in future projects. Multi-omics might be especially promising for complex traits such as ADHD, where clinicians have been struggling to find fitting treatments for patients and struggle to predict the persistency of ADHD (Caye et al., 2019). By improving methods and enhancing models, as well as by including more omics levels such as transcriptomics, proteomics, and exposome (Cheng et al., 2020; Liu et al., 2021), the identification of multi-omics biomarkers will eventually aid in diagnosing and improving personalized treatment plans for persons diagnosed with ADHD.
**AUTHOR CONTRIBUTIONS**


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**CONFLICT OF INTEREST STATEMENT**

Erik A. Ehli was employed by the Avera Institute for Human Genetics (Sioux Falls, SD, USA). The remaining authors declare no conflicts of interest.

**DATA AVAILABILITY STATEMENT**

The standardized protocol for large scale collection of urine and buccal-cell samples in the home setting was developed for the ACTION Biomarker Study in children available at http://www.action-euproject.eu/content/data-protocols. The data of the Netherlands Twin Register (NTR) ACTION Biomarker Study may be accessed, upon approval of the data access committee, through the NTR (https://tweelingenregister.vu.nl/information_for_researchers/working-with-ntr-data).

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**REFERENCES**


SUPPORTING INFORMATION

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