Metabolomics Study of Urine in Autism Spectrum Disorders Using a Multiplatform Analytical Methodology

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ABSTRACT: Autism spectrum disorder (ASD) is a neurodevelopmental disorder with no clinical biomarker. The aims of this study were to characterize a metabolic signature of ASD and to evaluate multiplatform analytical methodologies in order to develop predictive tools for diagnosis and disease follow-up. Urine samples were analyzed using $^1$H and $^1$H–$^{13}$C NMR-based approaches and LC–HRMS-based approaches (ESI+ and ESI− on HILIC and C18 chromatography columns). Data tables obtained from the six analytical modalities on a training set of 46 urine samples (22 autistic children and 24 controls) were processed by multivariate analysis (orthogonal partial least-squares discriminant analysis, OPLS-DA). The predictions from each of these OPLS-DA models were then evaluated using a prediction set of 16 samples (8 autistic children and 8 controls) and receiver operating characteristic curves. Thereafter, a data fusion block-scaling OPLS-DA model was generated from the 6 best models obtained for each modality. This fused OPLS-DA model showed an enhanced performance ($R^2_Y$(cum) = 0.88, $Q^2$(cum) = 0.75) compared to each analytical modality model, as well as a better predictive capacity (AUC = 0.91, p-value = 0.006). Metabolites that are most significantly different between autistic and control children (p < 0.05) are indoxyl sulfate, N-acetyl-l-arginine, methyl guanidine, and phenylacetylglutamine. This multimodality approach has the potential to contribute to find robust biomarkers and characterize a metabolic phenotype of the ASD population.

KEYWORDS: metabolomics, autism spectrum disorder, ASD, NMR, LC–HRMS, data fusion

INTRODUCTION

Autism spectrum disorder (ASD) refers to a group of complex neurodevelopmental disorders present since early childhood and persisting lifelong.1 The prevalence of ASD was recently estimated in France to 36.5/10 000 children with a sex ratio of 4.1 boys for 1 girl.2

Autism is typically diagnosed before 3 years of age,3 and ASD children are characterized by deficits in social communication and social interaction, as well as restricted and repetitive behaviors and interests, as listed in the American Psychiatric Association’s Diagnostic and Statistical Manual of Mental Disorders, 5th ed. (DSM-V).4 Diagnosis is made clinically by using different scale tests that evaluate the behavior of the patient,5 and to date, there is no reliable biochemical marker of this disorder. Although etiologies of autism remain unknown, studies have found implications of genetic, environmental, and metabolic factors.6 ASD has several suspected causes, including dysfunctions of the neurologic, immunologic, and/or gastrointestinal systems, with some markers showing ubiquitous distribution. Due to the social and communication impairments of patients with ASD, identifying gastrointestinal problems remains difficult. However, some studies have linked an imbalance of the gut microbiota with ASD.7

Some metabolic disorders have been found to be more frequent in the autistic population compared to the general population. These abnormalities include phenylketonuria, creatine deficiency syndromes, adenylosuccinate lyase deficiency, S-nucleotidase, and metabolism of purine pyrimidine disorders. The metabolic abnormalities that contribute to the etiology of autism is still unknown, but these findings suggest that ASD phenotypes may be associated with metabolic pathways imbalance. In order to explore this hypothesis, metabolomics studies which have already shown their potential in biomarker research in central nervous system disorders have been performed.8

Metabolomics is the study of the metabolome, which represents the whole content of low-molecular-weight com-

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Analytical platforms most commonly used to identify and quantify metabolites are mass spectrometry coupled to separation techniques such as gas chromatography (GC–MS) or liquid chromatography (LC–MS) and nuclear magnetic resonance spectroscopy (NMR).

To date, few metabolomics studies have been described for biomarkers exploration in ASD. These studies have been primarily based on urine screenings, since urine can be obtained in large quantities by noninvasive sampling. Moreover, repeated sampling is easy to achieve, a major consideration in the case of ASD children. These studies have been performed using a single analytical platform, based on either NMR or MS technologies. Although they have made the proof of concept that exploration of the metabolome allows us to classify ASD children compared to control, each analytical platform cannot cover the whole diversity of metabolites in body fluids (molecular diversity and expression levels). Using a single platform also results in partial information and difficulties in confirming identified metabolites as reliable biomarkers. In this study, we take full advantage of NMR and MS complementarity to explore the urine metabolome, using the combination of 1H NMR, 13C heteronuclear single quantum coherence (HSQC) NMR, and liquid chromatography coupled to high-resolution mass spectrometry (LC–HRMS). One of the challenges of this approach is to associate data from different analytical platforms in order to generate a statistical model that better represents urinary metabolic differences between children with or without ASD. In order to achieve this, we first built independent supervised multivariate models for each analytical modality in order to select the most discriminant metabolites, which were then concatenated in a new single matrix for a block-scaling model analysis. This multiprofiling approach, combined with data fusion, gave us the opportunity to better classify children with or without ASD compared to a unique analytical platform approach.

## EXPERIMENTAL SECTION

### Patients and Controls

Patients who met ASD diagnostic criteria according to the World Health Organization’s International Statistical Classification of Diseases and the American Psychiatric Association’s Diagnostic and Statistical Manual of Mental Disorders, 4th edition, were included in the study after medical consultations at the Regional Center for Autism in Tours, France, between 2011 and 2012. All parents of participants and participants provided informed consent. Urine samples were collected from 30 children with ASD and 32 control children living in France.

The 62 patients urine samples (30 ASD and 32 controls) were split into two sets: a training set of 46 samples (22 ASD and 24 controls), and an independent validation set of 16 samples (8 ASD and 8 controls). Information including sex, diagnosis, gastrointestinal disturbances, and age at sampling were collected for each participant. Urinary samples were collected in sterile polypropylene tubes untreated with preservatives. After centrifugation at high speed, each urine sample was aliquoted in a 1.5 mL sterile Eppendorf tube and stored at −80 °C immediately after collection until analysis.

### NMR Study

**Sample Preparation.** Urine samples were thawed at room temperature and centrifuged at 3000g for 10 min as described previously. The supernatant (500 µL) was then added with 100 µL of phosphate buffer (pH = 7.4 ± 0.5) and 100 µL of D2O solution for 1D analysis or 100 µL of D2O with internal reference [3-trimethylsilylpropionic acid (TSP), 0.05 wt% in D2O] for 2D analysis. Samples were then transferred into conventional 5 mm NMR tubes for NMR analysis.

**NMR Spectroscopy Experiments.** 1H NMR spectra were obtained as previously described using a Bruker DRX-500 spectrometer (Bruker SADIS, Wissembourg, France) operating at 500 MHz, using a “cpmg” pulse program.

All sensitivity-enhanced 1H–13C HSQC spectra were adapted from a previously described method on a Bruker DPX Avance spectrometer operating at 300 MHz, using an “hsqcqpppr” pulse program in the Bruker library.

### Data Preprocessing for NMR Analysis

Spectra were processed using TopSpin version 2.1 software (Bruker Daltonik, Karlsruhe, Germany), then aligned using the work software. The aligned spectra were then displayed in R, and zones with no peaks were removed from the file. Finally, intensities of points corresponding to the same peak or nearby peaks were added. These buckets corresponded to either single metabolites or a range of overlapped metabolites. The signal intensity in each bucket was normalized by the total sum of peak intensities and gathered in the 1H NMR matrix for further statistical analysis.

2D spectra were processed using MestReNova version 7.1.0 software (Mestrelab Research, S.L., Santiago De Compostela, Spain) as previously described. Each urine spectrum was normalized with an external reference TSP that served as a chemical shift reference set at 0 ppm and as a quantitative reference signal. The final 2D matrix contained 677 different 1H–13C cross-peaks between 10 and 150 ppm.

Peaks with low variability (relative standard deviation [RSD] lower than 15%) were excluded from all data tables since they would not be good predictive biomarkers.

### LC–HRMS Study

**Sample Preparation.** Samples were prepared from 20 µL of urine, diluted at 1/10 in water or acetonitrile (ACN) depending on the type of analytical column used (for HILIC or C-18, respectively). After vortexing for 10 min and centrifugation for 10 min at 10000g, 150 µL of the supernatant was transferred into a 96-well plate. Quality controls samples (QCs) were obtained from a pooled mixture of equal volumes of all urine samples. QCs followed the same pre-analytical and analytical steps described above. Fifteen QCs were injected to equilibrate the chromatographic system before each analytic batch. The running order of samples was randomized, and QCs were analyzed every 10 samples.

**Liquid Chromatography–High-Resolution Mass Spectrometry Analysis.** **MS Analysis.** LC–HRMS analysis was performed on a UPLC Ultimate 3000 system (Dionex), coupled to a Q-Exactive mass spectrometer (Thermo Fisher Scientific, Bremen, Germany) and operated in positive and negative electrospray ionization (ESI+) and ESI− modes (one run for each mode). The system was controlled by Xcalibur 2.2 (Thermo Fisher Scientific). Four untargeted LC–HRMS methods were conducted for better metabolome coverage: C18 and HILIC chromatography coupled to ESI in both positive and negative ion polarities. Each sample analysis resulted in four separate data acquisitions. Chromatography was carried out with a Phenomenex Kinetex 1.7 µm XB-C18 (150 mm × 2.10 mm) column with a Waters Cortecs 1.6 µm HILIC (150 mm × 2.10 mm) column kept at 40 °C. For C18
chromatography, a multistep gradient (followed by a 2 min equilibration time) had a mobile phase A of 0.1% formic acid in water, and a mobile phase B of ACN acidified with 0.1% formic acid; the gradient operated at a flow rate of 0.2 mL/min over a run time of 30 min for both negative and positive modes.

The multisteps gradient was programmed as follows: 0–3 min, 0% B; 3–8 min, 0–15% B; 8–15 min, 15–50% B; 15–20 min, 50–100% B; 20–25 min, 100% B; 25–28 min, 100–0% B. For HILIC chromatography, a multistep gradient (followed by a 2 min equilibration time) had a mobile phase A of ammonium formate in water (10 mM), and a mobile phase B with ACN containing ammonium formate (10 mM); the flow rate was 0.4 mL/min over a run of 22 min for both positive and negative modes. The multisteps gradient was programmed as follows: 0–5 min, 0% A; 5–12 min, 0–20% A; 12–18.5 min, 20–60% A; 18.5–19.5 min, 60% A; 19.5–20 min, 60–0% A. The autosampler temperature (Ultimate WPS-3000 UHPLC system, Dionex, Germany) was set at 4 °C, and the injection volume for each sample was 5 μL for C18 and 10 μL for HILIC.

Heated ESI source parameters were, for both modes, a spray voltage of 3 kV, capillary temperature of 380 or 325 °C, sheath gas flow of 40 arbitrary units (AU) or 35 AU, auxiliary gas flow of 20 or 10 AU, spare gas flow of 2 or 1 AU, and tube lens voltage of 50 or 60 V for C18 or HILIC, respectively. During the full-scan acquisition, which ranged from 66.7 to 1000 m/z, the instrument operated at 70 000 resolution (m/z = 200), with an automatic gain control (AGC) target of $1 \times 10^6$ charges and a maximum injection time (IT) of 250 ms.

**MS² Analysis of VIPs.** First, a pool sample was injected for molecular ion mass determination at a resolution of 140 000. The most discriminant metabolites (variable importance parameter, or VIPs) obtained from mass spectrometry were then further investigated for their identification. Targeted tandem mass spectrometry (MS²) experiments were conducted with an inclusion list of ions mass-selected from HILIC and C18 analysis using an isolation window for the quadrupole of 0.5 m/z and a resolution of 35 000 (m/z = 200) for the fragmentation spectrum, with an AGC target of $2 \times 10^4$ charges, a maximum IT of 100 ms, and a normalized energy collision (Supporting Information, Figure S-3).

**Data Preprocessing for LC–HRMS Analysis.** XCMS software was implanted in the Galaxy platform (http://workflow4metabolomics.org/the-galaxy-environment/) was used to process raw data for peak alignment and framing. This step produced a table of detected features, characterized by sample retention time, m/z ratio, and intensity (i.e., peak area). We normalized each peak area to the total peak area of each chromatogram. For chromatograms obtained with HILIC column in negative ionization mode, intensities of signals were corrected with Loess after we observed an analytical deviation of signals before normalization to total peak area.

The CAMERA package was used to group isotopes and adducts in order to annotate and to identify features. The stability of signals intensities across batches was evaluated. Extracted ion chromatograms (EICs) were checked in order to review consistency of integration across samples, to analyze peak shapes, and to exclude background noise. Variability of features passing this EIC quality review process was then evaluated. QCs variability, given by RSD of each feature, was assessed. Features with RSD in QCs higher than in samples were excluded. We only kept features with RSD in QCs below 30% for further multivariate analysis. Features greater than 30% variance in QCs were not considered, except if significant variance was observed between groups, meaning that biological variability may exceed analytical variability. Similarly to NMR experiments, we excluded peaks with RSD in samples lower than 15%.

**Data Processing**

**Multivariate Data Analysis.**

**Quality Control Analysis for LC–HRMS.** Clustering of QCs was assessed by principal component analysis (PCA) according to total peak area data in order to compare analytical variability with biological variability.

**Samples Batch Data Analysis.** The 62 patients samples (30 ASD and 32 controls) were split into two sets: a training set of 46 samples (22 ASD and 24 controls) for the identification of the most discriminants metabolites between ASD and control urine samples, and an independent validation set of 16 samples...
Table 1. ASD and Control Groups’ Characteristics

<table>
<thead>
<tr>
<th></th>
<th>training set</th>
<th></th>
<th>independent validation set</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>ASD (n = 22)</td>
<td>control (n = 24)</td>
<td>ASD (n = 8)</td>
<td>control (n = 8)</td>
</tr>
<tr>
<td>age, years</td>
<td>8.64 ± 3.62</td>
<td>8.08 ± 3.67</td>
<td>9.24 ± 3.79</td>
<td>9.37 ± 4.07</td>
</tr>
<tr>
<td>no. of males (%)</td>
<td>19 (86.36%)</td>
<td>21 (87.50%)</td>
<td>7 (87.50%)</td>
<td>6 (75%)</td>
</tr>
<tr>
<td>no. of females (%)</td>
<td>3 (13.64%)</td>
<td>3 (12.50%)</td>
<td>1 (12.50%)</td>
<td>2 (25%)</td>
</tr>
<tr>
<td>diagnostic</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>autism disorder</td>
<td>9</td>
<td>0</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Asperger’s syndrome</td>
<td>2</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>PDD-NOS</td>
<td>11</td>
<td>0</td>
<td>6</td>
<td>0</td>
</tr>
<tr>
<td>neurotypical</td>
<td>0</td>
<td>24</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td>gastrointestinal</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>disturbance</td>
<td>yes</td>
<td>NA</td>
<td>5</td>
<td>NA</td>
</tr>
<tr>
<td>no</td>
<td>12</td>
<td>NA</td>
<td>3</td>
<td>NA</td>
</tr>
<tr>
<td>NA</td>
<td>1</td>
<td>24</td>
<td>0</td>
<td>8</td>
</tr>
</tbody>
</table>

“Data are expressed as mean.

(8 ASD and 8 controls) to evaluate the performance of the classification models. This was accomplished by randomizing samples. The training and independent validation sets were matched by age and sex.

The general workflow for the training set is shown in Figure 1.

Each analytical method generated a data table with detected features presented in columns and urine samples presented in rows. The preprocessed data sets were used as input for Simca P+ version 13.0 (Umetrics, Umeå, Sweden), and data analysis was preceded by log transformation and unit variance scaling. The training sets of these six data sets were tested individually in order to find the best orthogonal partial least-squares discriminant analysis (OPLS-DA) model. Model development was performed in order to (i) select a minimum set of discriminant metabolites (VIP > 1.0) that are the most implicated in the difference between ASD and control urine samples and (ii) test performance of the optimal model with receiver operating characteristic (ROC) curve analysis and the validation set data. The SIMCA prediction score (Ypred) on the independent validation set was used to build the ROC curves. The main benefit of OPLS-DA compared to PLS-DA is its ability to separate the systemic variation in variables X into two parts: variation related to class membership to variation unrelated to class membership (orthogonal). This partitioning of the X-data will facilitate model interpretation and prediction of new samples.30 OPLS-DA models performed on training set were then evaluated by cross-validation by withholding 1/7 of the samples in seven successive simulations, so that each sample was omitted once in order to prevent against overfitting. The set of multiple models resulting from cross-validation was used to calculate jackknife uncertainty measures.27 We set a maximum number of iterations at 200 in order to ensure convergence of the OPLS algorithm.28 A Pearson correlation test was performed between discriminant variables for each analytical method to remove the redundancy of information due to VIPs that correspond to the same metabolites. The performance of each of the six OPLS-DA models was assessed using a validation set data and ROC curves analysis.

A data fusion block-scaling model was then generated by combining data tables coming from the previous six OPLS-DA models. Block scaling, in the context of data fusion provides a way to balance influence of blocks of variables in relation to their size.30 Block scaling allows each group of variables to be considered independently as an entity with a specific variance.30 This final data fusion block-scaling OPLS-DA model was also evaluated using a ROC curve analysis.

ROC curves were performed using GraphPad Prism version 6.00 for Windows (GraphPad Software, La Jolla, CA, USA, http://www.graphpad.com/).

**Univariate Data Analysis**

Univariate analysis focused on the Variable Importance in Projection (VIP > 1) obtained from the data fusion block-scaling model using Wilcoxon test. A statistical correction for multiple tests was applied in order to adjust the p-value for significance by accounting for the number of metabolites evaluated (Bonferroni adjustment). So differences were deemed significant when p < 0.05/n (where n is the number of VIPs). Statistical analyses were performed with JMP statistical software version 7.0.2 (SAS Institute, Cary, NC, USA).

**Variable Importance Parameter Annotation**

Most discriminant VIPs in multivariate analyses were investigated to be annotated or identified. The VIP assignments for molecular formula elucidation were made with the help of seven golden rules in mass spectrometry.31 From the molecular formula, free access databases queries using ChemSpider (http://www.chemspider.com/), Human Metabolome Database (http://www.hmdb.ca/), and MassBank (http://www.massbank.jp/) were used to annotate compounds. Finally, each VIP was analyzed at high resolution and by MS2. From HRMS (ion mass and isotopic abundances), one or several parent structures are generated. To determine the structure of these compounds, Mass Frontier software’s fragment comparator (Thermo Scientific) gives the opportunity to compare fragments derived from different compounds. First, Mass Frontier software predicted comprehensive pathways based on a set of general ionization, fragmentation, and rearrangement rules. Fragmentation spectra were then compared to ab initio fragmentation spectra of potential identified metabolites using Mass Frontier.32 Experimental MS2 mass spectra with fragment structure annotations are given in the Supporting Information (Figure S-3, which illustrates this compound annotation workflow).

For VIPs obtained from 2D NMR (1H–13C HSQC), chemical shifts were submitted to the MetaBoliner database (http://wishart.biology.ualberta.ca/metabominer/). In MetaBoliner, a compound is considered to be present if the requirements of minimal signatures are met. A minimal
Table 2. Summary of Statistical Values of OPLS-DA Models and ROC Curve Analysis of Training Set and Independent Validation Set Obtained with the Different Methodologies

<table>
<thead>
<tr>
<th>Methodology</th>
<th>no. of variables</th>
<th>R²Y(cum)</th>
<th>Q²(cum)</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
<th>p-value</th>
<th>AUC</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>¹H NMR</td>
<td>8</td>
<td>0.52</td>
<td>0.37</td>
<td>81.8</td>
<td>91.7</td>
<td>6 x 10⁻⁴</td>
<td>0.83</td>
<td>0.02</td>
<td>62.5</td>
</tr>
<tr>
<td>¹H−¹³C NMR</td>
<td>7</td>
<td>0.59</td>
<td>0.51</td>
<td>86.4</td>
<td>75.0</td>
<td>2 x 10⁻⁷</td>
<td>0.84</td>
<td>0.03</td>
<td>75.0</td>
</tr>
<tr>
<td>HILIC ESI+</td>
<td>9</td>
<td>0.51</td>
<td>0.47</td>
<td>86.4</td>
<td>91.7</td>
<td>1 x 10⁻⁴</td>
<td>0.70</td>
<td>0.17</td>
<td>100</td>
</tr>
<tr>
<td>HILIC ESI−</td>
<td>4</td>
<td>0.40</td>
<td>0.34</td>
<td>77.3</td>
<td>79.2</td>
<td>1 x 10⁻³</td>
<td>0.64</td>
<td>0.34</td>
<td>62.5</td>
</tr>
<tr>
<td>C18 ESI+</td>
<td>7</td>
<td>0.64</td>
<td>0.53</td>
<td>90.9</td>
<td>91.7</td>
<td>3 x 10⁻⁸</td>
<td>0.78</td>
<td>0.06</td>
<td>62.5</td>
</tr>
<tr>
<td>C18 ESI−</td>
<td>11</td>
<td>0.48</td>
<td>0.39</td>
<td>86.4</td>
<td>75.0</td>
<td>2 x 10⁻³</td>
<td>0.83</td>
<td>0.03</td>
<td>87.5</td>
</tr>
<tr>
<td>block model</td>
<td>46</td>
<td>0.88</td>
<td>0.75</td>
<td>100</td>
<td>100</td>
<td>9 x 10⁻¹²</td>
<td>0.91</td>
<td>0.006</td>
<td>100</td>
</tr>
</tbody>
</table>

“Y (cum)” is the cumulative predicted Y values, “Y (cum)⁰” is the cumulative observed Y values, “Sensitivity (%)” is the % of healthy children correctly identified as healthy, “Specificity (%)” is the % of diseased children correctly identified as diseased. AUC = Area under the ROC curve, 95% CI = 95% Confidence Interval. 

The different cumulative modeled variations in Y matrices [R²Y(cum)] on spectral datasets and predictability of the model (Q²) are given [observations (N) = 46]. Sensitivity = the number of diseased subjects that are correctly identified as diseased. Specificity = the number of healthy subjects that are correctly identified as healthy in the training set and validation set. p-value of ROC curve analysis.

Results and Discussion

The 62 patient urine samples (30 ASD and 32 controls) were split into two sets: a training set of 46 samples (22 ASD [mean 8.64 years] and 24 controls [mean 8.08 years]) and an independent validation set of 16 samples (8 ASD [mean 9.24 years] and 8 controls [mean 9.37 years]). Information collected on each participant, including age, sex, age at diagnosis, medication, and age at sampling, is summarized in Table 1. Differences of age and sex between groups of training sets are not significant (p-value = 0.52 for age and p-value = 1 for sex) likewise for validation set (p-value = 0.73 for age and p-value = 1 for sex).

NMR Experiments

¹H NMR. ¹H NMR spectroscopy, a rapid, robust, and reliable analytical tool with high reproducibility, has shown its potential to explore the urine metabolome in ASD patients. 34,35 ¹H NMR spectra (0–9.5 ppm) were divided in 147 buckets. After removing buckets with low variability in patients (RSD < 15%), the data table of the training set (138 buckets) was analyzed by OPLS-DA. Performances of OPLS-DA internal cross-validated model obtained with 8 discriminant variables, were R²Y(cum) = 0.52, Q² = 0.37 (Table 2). The sensitivity (percentage of ASD children correctly identified) obtained from the OPLS-DA model using the training set was 81.8%, and specificity (percentage of healthy children correctly identified) was 91.7%. OPLS-DA model p-value was significant (p-value = 6 x 10⁻⁴). This OPLS-DA model was then externally evaluated using the independent validation set and ROC curve analysis. Area under curve (AUC) was 0.83, and p-value was 0.03 (Supporting Information, Figure S-2). The sensitivity and specificity obtained from the ROC curve analysis using the independent validation set were lower, with respective values of 62.5% and 87.5% (Table 2). This result shows that the internal validation during OPLS-DA model construction overvalued prediction’s capacity due to overfitting of data. Assessing classification performances of each model by an independent set appears to be necessary in order to minimize this overvaluation.

¹H−¹³C NMR. ¹H NMR signals can be disturbed by spectral overlap, resulting in a potential lack in spectral resolution and making difficult the identification step. To improve the resolution of urine components, a two-dimensional NMR (¹H−¹³C HSQC NMR) acquisition may be valuable. ¹H−¹³C HSQC-based NMR avoids spectral overlap by dispatching the overall information in two dimensions. Compared to the data table of 138 buckets of ¹H NMR, the ¹H−¹³C HSQC NMR list containing 677 different ¹H−¹³C cross-peaks between 10 and 150 ppm was established as previously described due to less overlap of the signals.

The model obtained by ¹H−¹³C HSQC NMR gave a better information on prediction with a Q² = 0.51 (Table 2) obtained from 7 cross peaks in the optimized OPLS-DA model compared to ¹H NMR OPLS-DA model (Q² = 0.37). The OPLS-DA model sensitivity for the training set was found higher compared to the ¹H NMR (86.36% vs 81.82%). However, specificity was found lower compared to the ¹H NMR (75% vs 91.7%). The p-value of the ¹H−¹³C HSQC NMR OPLS-DA model was significant (p-value = 2 x 10⁻⁵). Regarding the validation set, AUC ROC curve gives results similar to those obtained with ¹H NMR (0.84 vs 0.83), and the p-value of the ¹H−¹³C HSQC NMR ROC curve was 0.03.

The sensitivity of ROC curve analysis for independent validation was found better compared to the ¹H NMR (75% vs 62.5%) with an equal specificity between them (87.5%). There again, these results show that overfitting of data in OPLS-DA models could be attenuated by the use of a validation set, and a better spectral separation results in a better sensitivity in discriminant analysis.

LC–HRMS Analysis

Since only a few untargeted metabolomics studies by LC–MS have been realized with urines of ASD patients, and since they have been done using a single chromatographic column type (C18), 36 we decided, in order to better cover the urine metabolome, to analyze each sample using LC–HRMS coupled to a HILIC or C18 column in both positive (ESI+) and negative (ESI−) ionization modes. Four independent models were built depending on column separation and ionization mode.
Quality Control

Fifteen QCs were injected to equilibrate the chromatographic system before each analytical batch. PCA scatter plots of QCs and samples were analyzed in order to compare analytical and biological variabilities for each batch. As shown in Figure S-1 in the Supporting Information, the clustering of QCs compared to samples clearly shows that biological variability exceeds the analytical one. This QCs step validates all batches series.

HILIC Analysis

HILIC provides a good separation for polar metabolites that are of major abundance in urine. This is the first study performed with HILIC chromatography in order to explore ASD urine metabolome. After data preprocessing (see above), data tables contained 1067 and 845 features in ESI+ and ESI−. Multivariate analysis gave the similar ability to separate the two populations of urine samples with slightly better statistical values ($R^2Y$(cum) = 0.51 and $Q^2 = 0.47$) for ESI+ compared to ESI− ($R^2Y$(cum) = 0.4 and $Q^2 = 0.34$).

The OPLS-DA models’ sensitivities (built with 9 and 4 variables from ESI+ and ESI−, respectively) were 86.4% and 77.3% for ESI+ and ESI−, respectively. Likewise, the OPLS-DA models’ specificities were 91.7% and 79.2% for of ESI+ and ESI−, respectively. AUCs curves for the validation set (ESI+ and ESI−, respectively) were 0.70 and 0.64 (Supporting Information, Figure S-2). Model curves were not found to be significant ($p > 0.05$), unlike $p$-values of OPLS-DA models that were significant (Table 2). Reasons for these discrepancies could be an overfitting of the OPLS-DA models and/or too small a number of samples for the validation set.

C18 Analysis

Reverse-phase separations are ideal for relatively nonpolar metabolites. A total of 1555 and 1226 features in ESI+ and ESI−, respectively, were selected as repeatable. From C18 ESI− data, an OPLS-DA model built with 11 discriminant variables gave statistical parameters $R^2Y$(cum) = 0.48 and $Q^2 = 0.39$ (Table 2). The sensitivity and specificity of this OPLS-DA model were 86.36% and 75%, respectively, with a significant $p$-value = $2 \times 10^{-5}$. The performance of this model was also evaluated using the ROC curve, and we found sensitivity and specificity similar to those of the OPLS-DA model (Table 2). AUC was found to be 0.83, with a significant $p$-value = 0.03 (Supporting Information, Figure S-2). The significant $p$-value of ROC curve confirms here that the OPLS-DA model is able to classify ASD patients and controls with minimum overfitting.

The OPLS-DA model obtained with 9 discriminant variables from C18-ESI+ data gave better information on the class descriptor, with $R^2Y$(cum) = 0.64. Moreover, compared to other LC−HRMS modalities, ESI+ model gave the highest predictive ability, with $Q^2 = 0.53$ and $p$-value = $3 \times 10^{-6}$. OPLS-DA model sensitivity and specificity were 90.9% and 91.7%, respectively. Despite these models results, the ROC curve’s AUC value was not found to be significant ($p$-value = 0.06).

Multivariate Statistical Analysis of the Combined $^1$H NMR, $^1$H−$^{13}$C HSQC NMR, ESI+/− C18 Column, and ESI+/− HILIC Column

The fusion and extraction of information from multiple data tables has become a decisive issue. Several strategies can be used to associate data from different analytical platforms. The simplest approach is to concatenate the different data sets in a low-level fusion approach where the data matrix results in a fused data table that is used for multivariate analysis. This approach is greatly affected by disparate signal intensities in the range and size of the data matrix obtained from the different analytical platforms. In order to address this issue, the selection of the most relevant/predictive variables from each data tables may solve the problem of dimensionality. Such strategies are called intermediate or midlevel data fusion. The model responses are combined in order to produce a final “fused” response that provides a meaningful synthesis. Following a midlevel data fusion strategy, the most discriminant variables of each analytical platform ($^1$H NMR, $^1$H−$^{13}$C HSQC, C18 ESI+/−, and HILIC ESI+/−) were selected and combined in order to build a data fusion block-scaling model. Since a subset of the most valuable variables can be selected from each data sources, the prediction performance can be increased when compared with individual analysis. While some signals are expected to be common to different blocks, the remainder would be specific and should give complementary information. Since a data fusion model could be dominated by the largest matrices for numerical reasons, the fairness between blocks was ensured by unit variance scaling normalization.

We first built six independent OPLS-DA models in order to select the most discriminant variable (VIP > 1.0) for each analytical platforms: 8, 7, 7, 11, 9, and 4 variables from $^1$H NMR, $^1$H−$^{13}$C HSQC, C18 ESI+−, C18 ESI−−, HILIC ESI+, and HILIC ESI−−, respectively. An equal weight was then assigned to each block corresponding to each analytical platform in the data table.

The data fusion block-scaling model was obtained from 46 features giving performances parameters $R^2Y$(cum) = 0.88, $Q^2 = 0.96$.
Dihydroxy-1H-indole glucuronide I and desaminotyrosine, which are related to tyrosine metabolism, were also found in the most discriminant metabolites. These results, associated with perturbations in the levels of valine, tyrosine, and N-acetylaspartic acid, are in agreement with perturbations of amino acids levels published by Tu et al. and may be interpreted as abnormal amino acids metabolism affecting neurotransmitters levels such as dopamine, noradrenaline, and epinephrine as observed in ASD children.

Level of dihydroxyuracil, an intermediate breakdown product of uracil, was found altered in ASD children’s metabolotypes. Purines and pyrimidines disorders, such as dihydropteridine reductase deficiency or dihydropyrimidine dehydrogenase or dihydroxyphenylalanine deficiencies and adenosine deaminase deficiencies, have been linked to autistic features.

Deficiencies of dihydropteridine dehydrogenase, the enzyme which catalyzes the conversion of uracil to dihydroxyuracil, have been linked to ASD. Indeed, symptoms of this pyrimidine disorder include psychomotor retardation, epileptic encephalopathy, and autistic features.

Recent studies have shown that an antipurinergic therapy could reverse the behavioral and metabolic disturbances in the maternal immune activation mouse model and in the Fragile X (Fmr1 knockout) mouse model. From these two mechanically distinct examples of ASD mouse models, the purinergic pathway is a neurochemical hypothesis that triggers the evolutionary conserved cell danger to stress that may be associated with ASD.

Indoxyl and indoxyl sulfate, which are produced by tryptophan metabolism in gut bacteria, were found here as metabolite candidates (different from analytical modality: NMR 1D and 2D and HILIC ESI+) and confirmed our previous results. Since indoxyl sulfate has been found as a VIP in more than one analytical platform, it may be assumed that this metabolite expression is modified between our two groups of children.

Desaminotyrosine, which can also be the result of deamination of tyrosine by intestinal microflora, was found increased in the ASD group. Moreover, a recent study by Noto et al. using GC/MS as an analytical platform points out a gut microbiota dysfunction, including tyrosine metabolism perturbation. Our study also points out perturbations of phenylacetylglutamine (PAG) and p-cresol sulfate concentrations, which are also produced by the microbiota respectively from tyrosine and phenylalanine. These results which confirmed those previously reported underline the importance of mammalian-microbial cometabolites in ASD, supporting emerging evidence for a gut-brain connection in autism, wherein gastrointestinal microbiota may contribute to the ASD symptoms. It has to be noticed that half of our autistic cohort is clinically diagnosed as suffering of gastrointestinal disturbances that include diarrhea, constipation, and colitis.

Whether it is the cause or the consequence of autism’s physiopathology, our results confirm that gut microbiome seems to be associated with this disorder.

It has been recently shown that there are gender differences in emotions and sociability in children with autism spectrum disorders. This suggests that, in addition to phenotypic differences, there are metabotype differences linked to gender.

Table 3. Putatively Annotated Metabolites of Data Fusion OPLS-DA Model (VIP > 1)

<table>
<thead>
<tr>
<th>Analytical Platform</th>
<th>Potential Assignment</th>
<th>P-value</th>
<th>Differentiation for ASD Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>C18 ESI</td>
<td>Dihydroxy-1H-indole glucuronide I</td>
<td>0.005</td>
<td>↑†</td>
</tr>
<tr>
<td>C18 ESI</td>
<td>Dihydroxyuracil</td>
<td>0.03</td>
<td>↓</td>
</tr>
<tr>
<td>C18 ESI</td>
<td>N-acetyl-l-arginine</td>
<td>0.009</td>
<td>↑</td>
</tr>
<tr>
<td>C18 ESI</td>
<td>Unknown</td>
<td>0.009</td>
<td>↓</td>
</tr>
<tr>
<td>C18 ESI</td>
<td>Unknown</td>
<td>0.006</td>
<td>↑</td>
</tr>
<tr>
<td>C18 ESI</td>
<td>N-acetylaspartic acid</td>
<td>0.03</td>
<td>⊥</td>
</tr>
<tr>
<td>C18 ESI</td>
<td>Desaminotyrosine</td>
<td>0.006</td>
<td>⊥</td>
</tr>
<tr>
<td>C18 ESI</td>
<td>Guanidinosuccinic acid</td>
<td>0.007</td>
<td>⊥</td>
</tr>
<tr>
<td>HILIC ESI</td>
<td>Indoxyl</td>
<td>0.01</td>
<td>↑</td>
</tr>
<tr>
<td>HILIC ESI</td>
<td>Unknown</td>
<td>0.003</td>
<td>⊥</td>
</tr>
<tr>
<td>HILIC ESI</td>
<td>Valine, norvaline, 5-aminopentanoic acid</td>
<td>0.007</td>
<td>⊥</td>
</tr>
<tr>
<td>HILIC ESI</td>
<td>Unknown</td>
<td>0.0004</td>
<td>↑</td>
</tr>
<tr>
<td>HILIC ESI</td>
<td>α-N-phenylacetyl-l-glutamine</td>
<td>0.004</td>
<td>⊥</td>
</tr>
<tr>
<td>HILIC ESI</td>
<td>p-Cresol sulfate</td>
<td>0.02</td>
<td>⊥</td>
</tr>
<tr>
<td>NMR 1D</td>
<td>Indoxyl sulfate</td>
<td>0.01</td>
<td>↑</td>
</tr>
<tr>
<td>NMR 1D</td>
<td>Methylnitroguanidine</td>
<td>0.003</td>
<td>⊥</td>
</tr>
<tr>
<td>NMR 2D</td>
<td>Valine</td>
<td>0.06</td>
<td>↑</td>
</tr>
<tr>
<td>NMR 2D</td>
<td>Indoxyl sulfate</td>
<td>0.003</td>
<td>⊥</td>
</tr>
<tr>
<td>NMR 2D</td>
<td>Glucuronic acid</td>
<td>0.009</td>
<td>⊥</td>
</tr>
</tbody>
</table>

“↓p-value of univariate analysis (Wilcoxon test). ↑↑ denotes higher level in ASD urines, ↓↓ denotes lower level in ASD urines. "Significant p-value after Bonferroni correction (p < 0.003).”

First, we found higher levels of N-acetylaspartic acid in the ASD group compared to controls. Arginine has been described to be higher in plasma samples of ASD children. Excessive arginine is thought to induce oxidative stress via NO production. We also found perturbations of guanidinosuccinic acid and methylguanidine, which are produced from oxidation of argininosuccinic acid and creatine, respectively, by free radicals. The exact relationship between arginine pathway and oxidative stress in neuropsychiatric disorders remains unclear; however, a common susceptibility gene for ASD and schizophrenia, NOS1, has been suggested to be involved in the arginine-NO pathway. This metabolic profile is consistent with impaired oxidative stress in children with autism. The exact relationship between the guanidino compounds pathway and autism needs further investigation.
enrolled. Since only a few females have been included in this study, this tends to decrease the biological significance of our conclusions. However, significant metabolic differences found in this work between ASD and control children may underline a typical general metabolic signature of ASD.

**CONCLUSION**

The discovery of biomarkers for an early diagnosis of ASD, as well as a follow-up of its evolution for improved patient care, is still a challenge to achieve. Because of its noninvasive accessibility and since it has already shown potential for containing discriminative metabolites, we decided to explore the metabolome of children’s urine as deeply as possible, using six complementary analytical platforms: 1H NMR, 1H-13C NMR, and ESI+ and ESI− LC–MS with C18 and HILIC chromatographic support.

First, our results highlighted and confirmed that several metabolic pathways associated with amino acids, including tyrosine, asparagine, phenylalanine, tryptophan, and arginine, seem to be involved in ASD. In addition, as other studies have previously shown, a gut dysbiosis is likely to be associated with ASD. It is important to notice that all these metabolism perturbations could be observed in a single study, thanks to the use of a multiplatform strategy that enabled a deep exploration of the metabolome. Second, our results showed that OPLS-DA model construction based on data fusion and block scaling by criteria and classification. Bio. J. Psychiatry 1971, 118, 381–384.


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A. 1H NMR, GC-EI-TOFMS, and Data Set Correlation for Fruit


