Urinary metabolic phenotyping for Alzheimer’s disease

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Finding early disease markers using non-invasive and widely available methods is essential to develop a successful therapy for Alzheimer’s Disease. Few studies to date have examined urine, the most readily available biofluid. Here we report the largest study to date using comprehensive metabolic phenotyping platforms (NMR spectroscopy and UHPLC-MS) to probe the urinary metabolome in-depth in people with Alzheimer’s Disease and Mild Cognitive Impairment. Feature reduction was performed using metabolomic Quantitative Trait Loci, resulting in the list of metabolites associated with the genetic variants. This approach helps accuracy in identification of disease states and provides a route to a plausible mechanistic link to pathological processes. Using these mQTLs we built a Random Forests model, which not only correctly discriminates between people with Alzheimer’s Disease and those who were not. Further annotation of top-ranking metabolic features nominated by the trained model revealed the involvement of cholesterol-derived metabolites and small-molecules that were linked to Alzheimer’s pathology in previous studies.

Unmet medical need and the repeated failure of clinical trials in Alzheimer’s disease (AD) have together resulted in a surge of research seeking to understand disease mechanisms and generate novel therapeutic approaches. In order for such therapies to succeed it is widely accepted that trials will need to be performed early in the disease process1,2. Currently, the optimal biomarkers used to detect AD processes early in the course of the disease are Positron Emission Tomography (PET) imaging and cerebrospinal fluid (CSF) markers3. However, PET imaging is not universally available and obtaining CSF is a relatively invasive procedure. Progress has been made in the attempt to supplement these relatively specific biomarkers with other biomarkers that might be more applicable to larger populations using, for example, blood biomarkers. Putative markers in blood have been identified using proteomics, transcriptomics, metabolic and lipidomic phenotyping platforms. Efforts are now underway...
to replicate these and to generate single markers or biomarker panels that might be used as part of a process identifying people with early disease4,26.

There are relatively few studies examining the potential of urine as a biomarker fluid in AD4–6. probably because being separated from the brain not only by the blood–brain barrier but also by glomerular filtration, urine seems inherently unlikely to possess a signature of neurodegeneration. Most of the studies reported to date are small both in terms of the numbers of molecular targets and numbers of individuals examined. However, urine is a complex fluid with metabolites that reflect a response to injury11 and oxidative stress12 amongst other biological events at the systems level and might, therefore, be useful as a target fluid in neurodegeneration and other brain diseases13. An added advantage is that urine carries information on the metabolites arising from the gut microbiome14, an area of research that is gaining greater focus in neurodegeneration and AD15,16. In mouse models a range of methods to characterize the urine metabolome have been employed, with some success in identifying markers that differ between transgenic animals and controls17–20, but these models do not encompass the full systems effect and metabolic progression of AD in humans. Therefore deep exploration of the urinary metabolome for biomarkers relevant to AD may yield valuable mechanistic information. We report here the application of in-depth urinary metabolic phenotyping in a large multi-centre study consisting of four groups of participants. One group consisted of individuals clinically diagnosed with AD. A second group consisted of individuals with no apparent clinical symptoms of cognitive decline or dementia (control group—CTL). The final two groups consisted of individuals diagnosed with Mild Cognitive Impairment (MCI) who either remained cognitively stable throughout the follow-up term of the study (sMCI) or converted to a clinical AD diagnosis at a later study visit (cMCI). All samples that underwent metabolic phenotyping were collected at a single time point (baseline assessment).

Metabolic phenotyping was completed using a novel dual-platform approach consisting of both proton nuclear magnetic resonance spectroscopy (1H-NMR) and ultra-high-performance liquid chromatography coupled with mass spectrometry (UHPLC-MS) to ensure comprehensive metabolite coverage. Critically, in order to address the challenge of rich metabolic datasets, where analytical variables far exceed the numbers of samples available to study, we applied a novel approach consisting of metabolomic dataset normalisation followed by a feature reduction process that only selected those metabolites which associated with a metabolic regulatory genetic element—a technique known as Quantitative Trait Loci for metabolites (mQTL). In this way, we identified a panel of metabolites that might accurately classify the participant groups. Of particular interest was the ability of the classification model to differentiate between baseline MCI participants, who later either remained cognitively stable (sMCI) or converted to clinical AD (cMCI) which heralds promise for non-invasive scoring for early biomarkers of AD.

The classification model was used to prioritise metabolites for annotation according to their importance to predict AD. The analysis of the annotated metabolites revealed multiple direct and a few indirect links to the etiopathological processes in AD.

**Results**

We report here the largest study to date of metabolic phenotyping analysis of urine as a potential marker of AD. Metabolic phenotyping was performed on urine obtained from the AddNeuroMed/Dementia Case Registry (ANM/DCR) cohorts15-21 using two analytical platforms: UHPLC-MS (n = 561 samples) and 1H-NMR (n = 575 samples). The cohort of participants used in the analysis includes participants with normal cognition (control group—CTL), stable mild cognitive impairment (sMCI), mild cognitive impairment converting to dementia (cMCI) and participants with Alzheimer’s disease (AD) (Table 1). The number of available samples and metabolic features is summarised in Table 2. Using an innovative approach to the challenge of very high dimensionality data analysis, we first prioritise a set of metabolites using mQTL mapping, ensuring that features with a degree of genetic regulation were selected. From this reduced feature selection, we were then able to select metabolites that are associated with AD and were able to predict conversion to dementia from MCI.

**Dimensionality reduction of metabolic features.** Given that the number of metabolic features found in the cohort samples was orders of magnitude higher than the number of samples (specifically 55,675 features), as the first step we developed a novel feature reduction method. We utilised the availability of genetic data, in particular, 12 million SNPs obtained in the previous study24 and looked for an association between these SNPs and the metabolic features, performing metabolic quantitative trait locus analysis. We hypothesized that an association between a metabolite and a disease state is more likely to be relevant to an etiopathological mechanism if this metabolite was also associated with a genetic variant previously linked to a relevant disease phenotype.

The mQTL analysis resulted in a total of 1542 individual metabolic features relating to either a chemical shift in the case of 1H NMR (233 metabolic features) or a chromatographic retention time and mass to charge ratio (m/z) feature in the UHPLC-MS data (1309 features). The resultant metabolic features were associated with 6932 SNPs at a q-value < 0.01 (Table 3, Fig. 1). Of these, 6047 unique SNPs were linked to features from the UHPLC-MS data, 876 SNPs to features from the 1H NMR data, with 838 SNPs common to both. Given the 12 million SNPs tested, the probability of observing 838 or more SNPs in the intersection between UHPLC-MS and 1H NMR results by chance is vanishingly small (p-value < 2.23E–308). Previously, a total of 276 metabolomic QTLs have been reported25–30, and 83% of these SNPs were reproduced in the current study (Supplementary Materials Table S2), thereby, validating our pipeline.

**Ranking of metabolic features for annotation.** Metabolite annotation remains the bottleneck and limitation of metabolic phenotyping studies12,26. After reducing the number of metabolic features to 1542, we aimed to prioritise them according to their relative importance of correctly predicting the AD state of the sam-
We first built and tested Random Forests classification models on different combinations of features and diagnostic classes (for details see “Methods”). Briefly, we hypothesised that genomics data in combination with metabolic features and sample covariates could improve the prediction quality and tried combinations of following features: (1) 1542 metabolic features after the mQTL filtering, (2) 6932 SNPs associated with metabolic features and (3) sample covariates—age, sex and study site. Also, to account for the imbalance between the diagnostic classes, with the prevalence of AD and CTL classes over sMCI and cMCI classes, we applied re-sampling. First, we analysed all data in relation to the four original diagnostic categories (AD/CTL/cMCI/sMCI). Then, hypothesising that cMCI would be most similar to AD and sMCI most similar to control, we performed binary over-sampling by creating AD + cMCI and CTL + sMCI groups. Finally, we applied the under-sampling by using AD and CTL groups only. As the model trained on 1542 metabolic features together with sample covariates and only two diagnostic classes (AD and CTL) gave the lowest prediction error, it was selected as the final model (Fig. 2).

To further validate the performance of the final model, MCI samples were used. The model proved effective in separating MCI samples from individuals who subsequently converted to dementia (cMCI) from those who remained stable (sMCI), finding that 82.96% of cMCI samples were predicted as AD and 77.78% of sMCI samples as CTL (Fig. 3). This provided additional evidence that our feature reduction method resulted in a meaningful set of metabolites enabling the detection of early AD patients, along with the previous validation of 83% of associated SNPs in AD-related literature. The AUROC value of the model was 0.99 (Table 4 and Fig. 3), showing that the final model was quite robust.

Next, we used the developed Random Forest model for prioritisation of the metabolic features by computing the permutation importance score for each metabolic feature (Supplementary Materials Table S3). This method gave us 235 metabolic features with a score of at least $10^{-4}$. Out of these, 32 features were successfully annotated

### Table 1. Overview of study participants. CTL normal cognition (control) participants, sMCI stable mild cognitive impairment, cMCI mild cognitive impairment converting to dementia, AD participants with Alzheimer’s disease, MMSE Mini-Mental State Examination.

|                           | CTL | sMCI | cMCI | AD  | Total |
|---------------------------|-----|------|------|-----|-------|
| Number of participants    | 214 | 200  | 55   | 197 | 666   |
| Age                       |     |      |      |     |       |
| Mean ± sd                 | 76.1 ± 5.1 | 76.7 ± 5.5 | 77.9 ± 7.9 | 76.6 ± 5.8 |       |
| Sex                       |     |      |      |     |       |
| Male                      | 103 | 99   | 20   | 101 | 323   |
| Female                    | 111 | 101  | 35   | 96  | 343   |
| MMSE score                |     |      |      |     |       |
| Mean ± sd                 | 28.9 ± 1.1 | 27.3 ± 1.7 | 26.3 ± 1.8 | 20.0 ± 4.3 |       |
| Metabolic UHPLC-MS data obtained | 172 | 167 | 45  | 177 | 561 |
| Metabolic NMR data obtained | 174 | 173 | 46  | 182 | 575 |
| Metabolic data obtained (UHPLC-MS and NMR) | 132 | 140 | 36  | 162 | 470 |
| Genetic and metabolic UHPLC-MS data available | 119 | 80  | 24  | 122 | 345 |
| Genetic and metabolic NMR data available | 120 | 83  | 23  | 126 | 352 |
| APOE genotype available   | 154 | 155 | 44  | 174 | 527 |

### Table 2. Summary of samples and metabolic features available for the analysis. Details of used metabolomic platforms and assays are available in “Methods” section. Number of metabolic QTL samples—genetic and metabolic data availability.

| Platform      | Assay  | Abbreviation | # metabolic features | # samples | # metabolic QTL samples |
|---------------|--------|--------------|----------------------|-----------|------------------------|
| UHPLC-MS      | HILIC ESI+ | UHPOS | 6851                 | 561       | 345                    |
|               | RPC ESI− | URNEG | 16,961               | 561       | 345                    |
|               | RPC ESI+ | URPOS | 13,217               | 561       | 345                    |
| NMR           | 1H NMR | NMR | 18,646               | 575       | 352                    |
Figure 1. Manhattan plots presenting significant QTL associations with metabolic features across the metabolite phenotyping datasets. The x-axis shows each SNP that was analysed, sorted by chromosome and position. The y-axis shows the $-\log_{10}$ of the p-value for association with metabolic features concentration. Four sections correspond to four different metabolomic assays presented in our study: UHPOS, URNEG, URPOS and NMR.
by spectral interpretation. Since multiple analytical signals can correspond to the same metabolite, the annotation process resulted in a total list of 23 metabolites (Table 5).

### Analysis of annotated metabolites

We found three broad groups of annotated metabolites differentially present in the disease state: (1) conjugated metabolites of cholesterol derived compounds, (2) small molecule metabolites and (3) metabolites of exogenous sources.

Following the annotation, we quantified the differences in the levels of assigned metabolites across study groups using ANOVA (Table 5). A heatmap demonstrating inter-annotation correlations is shown in Fig. 4.

Having assigned chemical identity, we then investigated whether the associated SNPs were linked to any previously reported GWAS traits to gain potential insight into the relationship between metabolomic and genomic data.
variations. We mapped any polymorphisms associated with our annotated metabolites to the GWAS Catalog’s SNPs by genomic regions and summarised them in Table 6.

We speculate that all endogenous annotated metabolites are linked to AD processes through alteration of DNA methylation, gut microbiota malfunction and possibly altered metabolism of polyamines, cholesterol and sugar (Table 6, Fig. 5). The key observations from these analyses are reported in the sections below.

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**Figure 3.** Receiver Operating Characteristic (ROC) curves for RF model discriminating AD vs CTL and then applied to cMCI and sMCI study groups. The model was trained with 1542 prioritised metabolic features and three covariates (age, sex, study site) identified from the AD vs CTL comparison only. The area under the ROC curve (AUROC) value for the AD vs CTL is 0.99. The AUROC value for cMCI vs sMCI classes is 0.88.

**Figure 4.** The heatmap showing concentrations of the annotated metabolic features. Note (*) indicates metabolite conjugation with N-acetylglucosamine, note (**) indicates metabolite conjugation with N-acetylglucosaminide. Columns sharing the same metabolite names are isomers of each other.
Cholesterol metabolism. There are three unique metabolites derived from cholesterol metabolism, all in the conjugated form: tauro(chole)deoxycholic acid, pregnenolone and pregnanediol. These metabolites have higher levels in the AD and cMCI study groups (Table 5).

We found the two hormones, pregnenolone and pregnanediol, conjugated with sulfate and N-acetylglucosamine, both biotransformations were observed in human urine previously49. Annotated tauro(chole)deoxycholic bile acid (an isomer of taurochenodeoxycholic or taurodeoxycholic acid) is conjugated with an N-acetylglucosamine, known biotransformation prior to bile acid renal excretion50. There is an increased level of taurochenodeoxycholic or taurodeoxycholic acid conjugates in AD patients (see “Discussion” section).

Sugar metabolism and gut microbiota. We identified sucrose and galactose sugars with sucrose levels significantly higher in AD study group.

The following annotated metabolites are produced in or derived from gut microbiota processes: N,N,N-trimethyl-L-alanyl-L-proline betaine, 3-aminoisobutyrate, trimethylamine, lysine, butyrylcarnitine, and mentioned above, taurodeoxycholic acid. These metabolites have increased levels in AD patients (Table 5).

DNA methylation and polyamine metabolism. Evidence of DNA methylation in AD patients’ urine is found in the observation of two methylcytidine metabolites: 5-methylcytidine, and 2-O-methylcytidine. Another metabolite that shared the pattern of alterations with two methylated cytidine metabolites was annotated as “unknown

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**Table 5.** Annotated metabolites. Note (*) signifies isomers that cannot be differentiated using mass spectrometry fragmentation data. Column headers: Assay—metabolomic assay; #—a number of metabolic features in the dataset; PIC—Permutation Importance Score from Random Forest algorithm showing the importance of metabolite for classification purpose; F (3, 471)—ANOVA results (MANOVA in case of multiple metabolic features) presented as F-statistic; p-value—ANOVA (MANOVA in case of multiple metabolic features) results presented as adjusted p-value; CTL vs AD, CTL vs cMCI and CTL vs sMCI—post hoc tests results presented as adjusted p-value. In the last four columns, Scientific Notation is used due to the presence of very small numbers.

| Metabolite family          | Metabolite annotation                                      | Assay   | # | PIC | F (3, 471) | p-value | CTL vs AD | CTL vs cMCI | CTL vs sMCI |
|----------------------------|------------------------------------------------------------|---------|---|-----|------------|---------|-----------|------------|------------|
| Exogenous metabolites      | N-Desmethyl O-desacetyl diltiazem glucuronide              | UHPOS   | 1 | 0.04943 | 112.8      | <2E-16  | 1.54E-11  | 1.54E-11   | 5.94E-02   |
|                            | N-Desmethyl O-desacetyl hydroxy diltiazem glucuronide      | UHPOS   | 1 | 0.02553 | 80.3       | <2E-16  | 1.54E-11  | 1.54E-11   | 3.32E-02   |
|                            | N-Desmethyl hydroxy diltiazem glucuronide                  | URPOS   | 1 | 0.00741 | 40.13      | <2E-16  | 1.54E-11  | 1.54E-11   | 1.27E-01   |
|                            | Paracetamol                                                | UHPOS   | 1 | 0.03214 | 59.34      | <2E-16  | 1.54E-11  | 9.98E-01   | 7.30E-09   |
|                            | Paracetamol sulphate                                       | URPOS   | 1 | 0.003   | 26.66      | 6.19E-16| 1.54E-11  | 9.26E-01   | 9.66E-01   |
|                            | 3-Methoxy-paracetamol sulphate                             | URPOS   | 1 | 0.00105 | 11.97      | 1.45E-07| 4.00E-07  | 1.26E-01   | 7.85E-01   |
|                            | Quinine                                                    | UHPOS   | 1 | 0.00028 | 13.26      | 2.56E-08| 3.66E-07  | 2.66E-03   | 1.02E-06   |
| Cholesterol derived metabolites | Hydroxylated pregnenolone sulphate N-Acetylglucosamine isomer 2* | URNEG   | 1 | 0.0051 | 34.75      | <2E-16  | 1.53E-11  | 5.61E-02   | 9.86E-01   |
|                            | Hydroxylated pregnenolone sulphate N-Acetylglucosamine isomer 1* | URNEG   | 1 | 0.01073 | 46.28      | <2E-16  | 1.53E-11  | 8.84E-03   | 8.26E-02   |
|                            | Pregnenolone sulphate N-acetylglucosamine                  | URNEG   | 5 | 0.00463 | 26.42      | 8.39E-16| 1.54E-11  | 4.87E-03   | 8.22E-02   |
|                            | Pregnanediol sulphate N-acetylglucosamine                  | URNEG   | 1 | 0.00025 | 13.06      | 3.34E-08| 3.02E-07  | 9.49E-01   | 9.30E-01   |
|                            | Taurochenodeoxycholic or taurodeoxycholic acid NAcetylglucosamin* | URNEG   | 1 | 0.00052 | 20.31      | 2.18E-12| 3.14E-10  | 9.99E-01   | 9.03E-01   |
| Nucleosides, amines, carnitines, glycines | 3-Aminoisobutyrate                                         | NMR     | 2 | 0.00015 | 2.18       | 8.99E-02| 9.44E-01  | 6.25E-01   | 3.75E-01   |
|                            | N,N,N-Trimethyl-L-alanyl-L-proline betaine                  | URPOS   | 2 | 0.00299 | 34.09      | <2E-16  | 1.53E-11  | 8.65E-02   | 5.57E-02   |
|                            | Butyryl or isobutyryl carnitine*                           | UHPOS   | 1 | 0.00054 | 14.54      | 4.57E-09| 1.16E-08  | 7.25E-03   | 5.33E-01   |
|                            | Trimethylamine                                              | NMR     | 1 | 0.00014 | 1.68       | 1.71E-01| 2.81E-01  | 9.48E-01   | 1.67E-01   |
|                            | L-Lysine                                                   | NMR     | 1 | 0.00011 | 1.33       | 2.56E-01| 9.90E-01  | 9.07E-01   | 4.65E-01   |
|                            | 5-Methylcytidine                                           | UHPOS   | 2 | 0.00075 | 18.56      | 2.16E-11| 2.67E-11  | 1.79E-01   | 9.89E-02   |
|                            | 2-O-Methylcytidine                                         | UHPOS   | 1 | 0.00033 | 20.54      | 1.62E-12| 1.93E-09  | 7.12E-01   | 9.79E-01   |
|                            | Unknown nucleoside with adenosyl moiety                    | UHPOS   | 1 | 0.00026 | 9.24       | 5.99E-06| 2.90E-06  | 1.71E-01   | 3.18E-01   |
|                            | N-Acetylisoputreanine-gamma-lactam                         | URPOS   | 2 | 0.00046 | 13.42      | 2.06E-08| 4.55E-09  | 1.74E-02   | 5.36E-03   |
| Sugars                     | Sucrose                                                    | NMR     | 1 | 0.00027 | 0.17       | 9.20E-01| 6.78E-04  | 5.99E-01   | 1.56E-01   |
|                            | Galactose                                                  | NMR     | 2 | 0.0001  | 0.17       | 9.20E-01| 9.05E-01  | 9.77E-01   | 9.74E-01   |
| Metabolic pathway | Metabolite annotation | Chr | Genes/genomic region | Phenotypic traits | Relationship to AD |
|-------------------|-----------------------|-----|----------------------|------------------|-------------------|
| Metabolic pathway | N-Desmethyl O-desacetyl diltiazem glucuronide | 1 | Regulatory feature: ENSR0000006069 | Anxiety and major depressive disorder, Obesity-related traits | SORCS2 belongs to the Vps10 receptor family that has previously been linked to neurodegeneration and AD<sup>**34</sup>, and is known to play functional roles in protein transport. In addition, the receptor family includes the SORL1 gene that encodes protein SorLA—a key protein in amyloid-beta precursor protein (APP) processing<sup>**35</sup>. |
| Metabolic pathway | N-Desmethyl O-desacetyl hydroxy diltiazem glucuronide | 15 | MESP2 | Coronary artery aneurysm in Kawasaki disease |  |
| Paracetamol | Paracetamol sulphate | | | | |
| | 3-Methoxy-paracetamol sulphate | 4 AND 17 | SORCS2 AND CNTROB | Biopolar disorder, Interleukin-10 levels | The mQTL association links aldehyde oxidase 1 (AOX1) gene and quinine. AOX1 gene has a previously reported GWAS trait “Late-onset Alzheimer’s disease”<sup>37</sup>. |
| | Quinine | 2 | AOX1 | Late-onset Alzheimer's disease | Beta-chimaerin (CHN2) gene plays a role in neural development by regulating Rac1 activity<sup>38</sup> and is known to be downregulated with age. Through Rac1 activation, gene CHN2 is linked with Alzheimer's disease<sup>39</sup>. |
| Cholesterol metabolism (CM) | Hydroxylated pregnenolone sulphate N-acetylglucosamine isomer 2<sup>*</sup> | 7 | CHN2 | Age at onset, Alzheimer's disease, Obesity-related traits, Psychosis |  |
| | Hydroxylated pregnenolone sulphate N-acetylglucosamine isomer 1<sup>*</sup> | | | |  |
| | Pregnenolone sulphate N-acetylglucosamine | | | |  |
| | Pregnenediol sulphate N-acetylglucosamine | | | |  |
| | Tauro(cheno) deoxycholic acid N-acetylglucosaminide <sup>*</sup> | 5 | UGT3A1 | Blood metabolite levels, Primary biliary cholangitis (PBC) | Neither the gene UGT3A1 nor the PBC has a known relationship to AD, although we note that a progressive cognitive impairment different to delirium is a feature of PBC, independently of liver pathology<sup>**40,**41</sup>. In animal models of biliary cirrhosis that has led to memory impairment, hippocampal pregnenolone sulphate infusion resulted in a memory-enhancing effect<sup>**42</sup>. |
| CM, gut microbiota | 3-Aminoisobutyrate | 5 | AGXT2 | Metabolite levels, Asymmetrical dimethylarginine levels, Symmetrical dimethylarginine levels |  |
| | N,N,N-Trimethyl-l-alanyl-l-proline betaine | 11 AND 21 | Regulatory feature: ENSR00000961656 AND intergenic variant | General cognitive ability, Obesity-related traits, Metabolite levels |  |
| Gut microbiota | Butyryl or isobutyryl carnitine <sup>*</sup> | 15 | intergenic variant | |  |
| | Trimethylamine | 10 | PYROXD2 | Estimated glomerular filtration rate, Creatinine levels |  |
| | l-Lysine | 19 | SLC7A9 | |  |

Continued
nucleoside with adenosyl moiety. We were not able to identify its structure definitively in this analysis but the mass spectrometric data provided the molecular formula C10H11N5O3, and fragmentation data indicated the presence of adenosyl moiety in the molecule (some details regarding structure elucidation effort are presented in the "Supplementary Document Metabolite_Annotation_details.docx").

Another annotated metabolite is N-acetylisoputreanine-gamma-lactam, a catabolic product of spermidine. This metabolite levels alter not only in AD but also in cMCI and sMCI groups (Table 5).

Exogenous metabolites. Amongst exogenous metabolites, we found two prescribed medications, paracetamol and the calcium channel blocker diltiazem, in addition to the metabolite quinine. All three exogenous metabolites have associations with genetic variants (Table 6) and their levels significantly alter in study groups, as shown in Table 5.

Discussion

Previous studies have reported data suggesting that a panel of metabolites circulating in blood was able to predict incipient AD with very high degrees of accuracy\(^\text{51-52}\), raising considerable hopes for finding pre-clinical AD biomarkers in blood. Subsequent studies using a similar, if not identical approach, in multiple, larger cohorts failed to replicate these findings\(^\text{53-55}\). Other studies have reported metabolic and lipidomic differences in blood from people with disease compared to age-matched controls with various degrees of power, success and outcome\(^\text{56-59}\). However, none of these studies have been unequivocally replicated. One of the limitations of studies with high dimensionality datasets and relatively small numbers of samples is susceptibility to over-fitting. Additionally, because of the inherent problems of heterogeneity in neurodegenerative disease and the diversity of analytical platforms used in metabolic profiling (\(^1\)H-NMR, GC-MS, UHPLC-MS, CE-MS etc.), it is perhaps not surprising that there has been relatively little replication of metabolic phenotyping studies that seek biomarkers of disease. Similar problems plagued early genetic studies seeking susceptibility factors, but these have been largely overcome by the introduction of studies based on tens of thousands of individuals. In genetics it was possible to combine data from different cohorts using imputation techniques. This approach is more challenging for metabolic phenotyping approaches, which often utilize heterogeneous technologies and independent assays, the results from which are more difficult to build a comprehensive picture of the metabolic landscape. With increased throughput and lower cost of metabolic assays, such larger studies will become possible in future.

In the absence of studies with large sample size, one approach to combat the limitation of high dimensionality in molecular studies is to reduce the dimensionality of the data. To achieve this, here we used the mQTL approach. In doing so, we provide for the possibility of a degree of validation in other, much larger dataset derived from genetic associations studies, also enabling the inference of a degree of causality when an association is discovered. While this work requires replication, this finding holds promise for biomarkers in urine—arguably the most readily available biomarker fluid. Using this mQTL targeting approach, we show a highly significant association of a relatively small set of 32 metabolic features with AD. A model generated from these features not only accurately predicts the disease state, but more importantly, the same model when applied to samples from participants with the clinical diagnosis of mild cognitive impairment (MCI), distinguishes those subjects that...
subsequently progress to dementia (cMCI) from those that remain stable (sMCI). Note that, MCI is commonly referred to as a prodromal condition, but in fact, is a highly heterogeneous state defined by impaired cognitive function relative to age-adjusted norms. An easy to perform test to identify people in the prodromal phase of AD, distinguishing the subjects with MCI that are likely to convert to AD from those remaining stable, would be an important advance for the field. Our results suggest that the analysis of urinary metabolites might provide such a test.

Given that the sample size in our study is small, in the absence of a larger replication cohort, it is important to provide as much additional evidence as possible showing that the method we developed is reliable. One such type of evidence comes from the analysis of the annotated metabolites. One set of metabolites that we found is related to cholesterol—a critical biological precursor, required for the biosynthesis of downstream metabolites such as bile acids, hormones and steroids, which are commonly found in their conjugated form in the urine. Hormones and related steroidal structures derived from cholesterol are known to have a role in brain function.

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Figure 5. Annotated metabolites and their linkage to AD. Red colour indicates metabolites annotated in the study. Up arrow next to the metabolite’s name indicates increased levels in AD patients samples. Down arrow shows decreased levels in AD patients samples.
Observed in this study pregnenolone sulfate is a known neuro-steroid species\textsuperscript{60,61} reported to influence cognition in rodent models\textsuperscript{62} and patient studies, perhaps through its role in modulating gamma-aminobutyric acid subunit A (GABA\textsubscript{A}) and N-methyl d-aspartate (NMDA) receptors\textsuperscript{63,64}. The other hormone we annotated—pregnenediol, a metabolite of pregnenolone, was previously reported to be lower in the urine of older men\textsuperscript{65}, however, this has not been linked to AD. Concentrations of non-conjugated bile acids were observed to be altered in AD in human blood and brain samples and in transgenic models of the disease\textsuperscript{66,67}. A recent study on mouse models discovered that bile acids strongly inhibit the cysteine dioxygenase type-1-mediated (CDO1-mediated) cysteine oxidation was prevented following injections of 17-β estradiol, suggesting a potential interlinked role for disruptions in sugar and sex hormone metabolism, alternations of both are reported in our results. However, it is necessary to highlight that we have very limited known covariates of this retrospective data cohort that was collected ten years ago. Original study participants exclusion criteria included other neurological or psychiatric diseases, significant unstable systemic illness or organ failure and alcohol or substance misuse\textsuperscript{68}. That knowledge gives us some assurance that study participants did not have other diagnosed medical conditions like diabetes mellitus type 2 and renal disease.

Evidence of gut microbiota malfunction is a noticeable trait in our findings. We annotated trimethylamine, N,N,N-trimethyl-l-alanyl-l-proline betaine, 3-aminoisobutyrate, l-lysine and butyrylcarnitine metabolites with different levels of concentrations amongst study groups. We detected several metabolites closely linked to betaine, though not betaine directly. Betaine is a source for trimethylamine production and an alternate methyl source for converting homocysteine to methionine\textsuperscript{69}, increasing DNA methylation and altering gene expression\textsuperscript{70}. We detected trimethylamine, a metabolite released in gut microbiota from trimethylamine-containing dietary phospholipid components such as choline, lecithin, l-carnitine and mentioned above betaine. The oxidation of trimethylamine generates trimethylamine N-oxide that was reported to be elevated in Alzheimer’s patients\textsuperscript{71}. In addition, observed here N,N,N-trimethyl-l-alanyl-l-proline betaine is a recently discovered plasma biomarker of kidney function. Plausibly this metabolite is a product of betaine and myoinos light chain degradation\textsuperscript{72}, though this hypothesis has yet to be confirmed. Another detected product of gut microbiota processes is butyrylcarnitine—a product of l-carnitine processing in the human body. Previous research conducted in the mouse brain has shown that in old age, the AD genetic load significantly increase levels of butyrylcarnitine\textsuperscript{73}. Changes in butyrylcarnitine concentrations together with the observation of elevated levels of lysine in AD patients suggest alterations of carnitine in AD, since lysine is one of the sources for carnitine production in humans. These complex metabolic processes in gut microbiota require further investigation.

Exogenous metabolites found in this study and their linkage with genomic variants (paracetamol and quinine) or with possible dementia protective effect (diltiazem metabolomic profile) are significant findings for further investigation. Given that the prescription of quinine preparations is recommended only for malaria prophylaxis, this is an unlikely explanation for the quinine finding in this European population of older people. However, its prevalence in dietary sources may explain its presence in the population. This finding, along with the reporting of altered levels of diltiazem and paracetamol in AD is not obviously explicable, although reverse causality, where the disease state is resulting in a change in either prescription or compliance with medication, is an obvious possible explanation of the finding.

While the strengths of the study lie in the mQTL approach and the acquisition of a large amount of metabolic phenotyping data, there are undoubtedly limitations to consider. First, although large compared to previously reported urine studies, the analysis remains vulnerable to over-fitting and bias as the number of features is three times that of the number of samples, even following the mQTL based feature reduction. Clearly, larger and independent datasets are necessary to replicate the findings we report here. This could be achieved using a targeted quantitative mass spectrometry assay, specifically designed to quantitate the metabolic pathways of the key metabolites identified here. This would both help validate the findings and provide greater insight into pathway mechanisms and the networks involved. Secondly, the cohort used here, lacks specific diagnostic biomarkers (such as PET or CSF measures) indicative of pathological load and the diagnostic categorisation rests on experienced clinician assessment together with systematic assessment by a research worker, albeit using a widely tested and proven methodology. Ideally, data would also be collected on diet and life-style factors such as exercise and other environmental exposures together with detailed assessments of co-morbidity and prior medical history and medication use. Such exploration of factors beyond diagnostic category that might influence the metabolic profile will be important topics for future studies. Lastly, by focusing on the metabolites with an associated genetic variant, we may exclude metabolites that have a significant association with AD but are
largely influenced by the environment, lifestyle and not as strongly by genetics. The further research of exogenous metabolites, such as medicines and dietary compounds, as well as effects of environmental factors on metabolic processes in AD is necessary.

Methods

We confirm that: all experiments were performed in accordance with relevant guidelines and regulations; all methods were carried out in accordance with relevant guidelines and regulations; all experimental protocols were approved by the European Union AddNeuroMed program.

Study participants.

The data and biospecimens were collected (AddNeuroMed/Dementia Case Registry (ANM/DCR) cohort) in European, multi-site study, public-private partnership21-23. The cohort includes participants with established AD, MCI and normal cognition who were evaluated using well-established systematic interviews for diagnosis. Formally, the diagnoses are not pathologically confirmed or supported by specific biomarkers of pathology, but it has been shown that this diagnostic method is highly predictive using MRI scans24 and subsequent post-mortem classification25. In addition, all disease categories, and conversion to dementia from MCI, were diagnosed by an experienced clinician according to criteria as described below. Briefly, the inclusion and exclusion criteria for study groups were as follows. Inclusion criteria for AD study group: (1) ADRDA/NINCDS and DSM-IV criteria for probable Alzheimer’s disease; (2) mini Mental State Examination score ranged from 12 to 28; (3) age 65 years or above. Inclusion criteria for MCI and CTL study groups: (1) mini Mental State Examination score range between 24 and 30; (2) geriatric Depression Scale score less than or equal to 5; (3) age 65 years or above; (4) medication stable; (5) good general health. The distinction between MCI and controls was based on two criteria: (1) subject scores 0 on Clinical Dementia Rating Scale = control; (2) Subject scores 0.5 on Clinical Dementia Rating scale = MCI. For the MCI subjects it was preferable that the subject and informant reported occurrence of memory problems. All AD subjects had a CDR score of 0.5 or above. The distinction between cMCI (converted MCI) and sMCI (stable MCI): based on follow-up interviews and tests. Exclusion criteria (all study groups): (1) significant neurological or psychiatric illness other than AD; (2) significant unstable systematic illness or organ failure; (3) alcohol or substance misuse.

All samples were collected under human participant research protocols, including informed consent containing the clause necessary for allowing bio-specimens and data sharing abroad. Blood and urine were collected at baseline and stored at −80°C. Genetic data from blood samples were obtained using the Illumina 610-Quad chip in two different batches as previously described24, while the third batch was obtained using HumanOmniExpress 24 v1.1. For this study morning (non-first void) urine samples were collected at baseline assessment and were subjected to metabolic profiling analysis via UHPLC-MS and by 1H NMR spectroscopy (561 and 575 samples respectively) (Table 1).

1H NMR metabolic phenotyping.

Samples were analyzed by 1H-NMR spectroscopy in-line with previously published standard protocols for the study of human urine samples76. In brief, samples were prepared using a Gilson 215 liquid handling robot and transferred to 4 in. length × 5 mm outer diameter NMR tubes in batches of 80 patient samples and four QC pooled samples. Racks of 96 prepared NMR tubes were transferred to a refrigerated SampleJet sample handler robot (Bruker Co) working at 5°C. One dimensional 1H-NMR general profile and two-dimensional J-resolved (Jres) experiments were acquired on a Bruker Avance III HD 600 spectrometer following the set up previously described76. Experiments were acquired and processed in automation using TopSpin 3.2 and ICON NMR. Phasing, baseline correction and calibration to TSP were also carried out in automation after each acquisition. Spectra quality was assessed using an in-house developed bioinformatics tool nPYC27 following the quality criteria previously described76.

UHPLC-MS metabolic phenotyping.

UHPLC-MS analysis of urine samples was performed as previously described84 utilizing a combination of reversed-phase chromatography (RPC) and hydrophilic interaction chromatography (HILIC). UHPLC was performed using Waters Acquity UPLC systems (Waters Corp., Milford, MA, USA), coupled to Waters Xevo G2-S QTOF mass spectrometers (Waters Corp., Wilsolm, UK) via Z-spray electrospray ionization (ESI) sources. RPC separations were paired with both positive and negative ion mode detection (generating URPOS and URNEG datasets respectively), while the HILIC separation was paired with positive ion mode detection only (UHPOS). All UHPLC-MS datasets underwent feature extraction using Progenesis QI 2.1 software (Nonlinear Dynamics, Newcastle, UK) as previously described, and feature filtering was performed using previously described quality control materials79. Preprocessing including batch and run order correction was performed using an in-house developed bioinformatics tool nPYC27. Features were removed from the data sets where their analytical variation, assessed by repeated measurement of a pooled quality control sample (study reference), exceeded 30% (relative standard deviation) or where the analytical variation exceeded the total observed variation among all study samples. Features were also inspected for correlation between their observed intensity and sample dilution within a pooled QC dilution series and features with Pearson correlation coefficients less than 0.7 were removed.

Metabolic QTL mapping.

The goal of QTL mapping is to identify associations between genetic markers and phenotypic variation79, in the case of metabolic QTL, the genome-wide contribution of individual alleles to a metabolic feature concentration27. The R package MatrixEQTL80 was used for QTL mapping by modelling the effect of genotype as additive linear including covariates to account for: age, sex, data collection centre and cohort. The null-hypothesis, "there is no QTL effect for the metabolic feature concentration", was tested against the alternative, "there is QTL effect between SNP and metabolic feature concentration", for each pair of meta-
bolic feature and SNP. We corrected results for multiple comparisons by calculating q-value using the Benja-
mimi–Hochberg procedure. The number of metabolic features and SNPs found to be associated using stringent q-value threshold 0.01 is presented in Table 3.

Data preparation For mQTL analysis. UHPLC-MS metabolic datasets UHPOS, URPOS and URNEG were normalised with EigenMS method that removes bias of unknown complexity from this type of metabolomics experiment at the same time preserving known bias, diagnoses in the case of this study. Subsequently, both EigenMS processed UHPLC-MS data, and raw one-dimensional 1H NMR metabolic data were normalised using the quantile normalisation method to make metabolomics data suitable for the QTL mapping.

Imputation and quality control of genomics data. After quality control using PLINK, genomics data from separate batches were assembled and remapped to “hg19-build37” reference genome. The imputation was performed using IMPUTE2 software. Since SNPs with low minor allele frequency are non-informative and have a potential of creating spurious findings, the imputation results were converted into matrix form and filtered using minor allele frequency threshold 10%. The final genomics matrix includes 12,105,785 SNPs. The results of population stratification indicated biases by cohort and data collection centres (Supplementary materials Figures S1 and S2).

Model selection. We used the Random Forests algorithm to prioritise metabolic features that can potentially be used as Alzheimer’s disease biomarkers. The main focus of this study is on metabolic features. However, there are also 6923 SNPs that metabolic features are associated with and covariates available for the samples: age, gender and data collection centre. To find the best prediction model, we considered the following sets of features:

A. Metabolic features only;
B. Metabolic features and SNPs;
C. Metabolic features, SNPs and covariates;
D. Metabolic features and covariates.

As discussed earlier, we tested three different ways of classifying diagnostic groups: four original diagnostic categories (AD/CTL/cMCI/sMCI), binary over-sampling AD + cMCI/CTL + sMCI, and binary under-sampling AD/CTL by removing cMCI and sMCI data. We explored different classifications approaches and feature sets by using the R implementation of RF to find the best combination. As the comparison criteria, we used the Out-Of-Bag (OOB) errors which is a standard approach for Random Forests. OOB error is a method of measuring the prediction error utilising bootstrap aggregation to subsample data used for training. It helps to avoid the need for an independent validation dataset. Results in the form of OOB errors of the RF models built for the different combinations of feature set and diagnostic groups are shown in Fig. 2. Each model was repeated ten times, using an increasing number of trees per run. The binary classification AD/CTL gives the best OOB errors for all four feature sets: the feature set A—0.0325, B—0.0404, C—0.0363 and D—0.0292, correspondingly. The feature set D, metabolic features and covariates, and binary classification AD/CTL, Alzheimer’s disease and healthy control samples only, after the tuning of RF parameters, gave the OOB predicted error 0.0241. The value of the area under the receiver operating characteristic curve (AUROC) for the model is 0.99 (Table 4 and Fig. 3). This model is our final RF model used in further analysis.

Tuning of random forests parameters. There are two parameters to tune for the RF algorithm: the number of trees used in the forest—ntree, and the number of variables used in each tree—mtry. We applied the alternating iterative procedure to find the best possible parameters values for ntree and mtry. The results plotted on Supplementary materials Figure S3 shows that with the ntree value equal to 680 the out-of-bag error rate stabilises and reaches its minimum 0.0254 with standard deviation 0.0056. The best mtry value we found is 90 (Supplementary materials Figure S4). It gives minimal OOB error rate 0.0241 with standard deviation equals to 0.0046.

Ranking of metabolic features. We ranked metabolic features using the permutation importance score obtained from the final RF model. This score is based on the idea that if the feature is not essential, then rearranging the values of that variable does not degrade classification accuracy. The list of ranked metabolic features with calculated permutation importance score and added metabolite annotations are presented in the Supplementary materials Table S3.

Metabolite annotation. For metabolite identification features of interest derived from the UHPLC-MS datasets first underwent correlation analysis using an R script developed in-house. This enabled the observation of co-eluting adducts and in-source fragments that are characteristic of metabolites. Further structural data were obtained via the use of high-resolution accurate mass to charge ratio (m/z) values and collision-induced dissociation (CID) fragmentation patterns. CID experiments were completed at a range of collision voltages at 5 V step intervals (5–45 V). The front quadrupole of the QTOF MS system was engaged to select for the feature of interest prior to CID. Chromatographic retention time matching to in-house standards was also completed where the standard was available for purchase. Online databases such as METLIN, and the Human Metabolome Database (HMDB) were also used to assist with metabolite identification. Where available, analytical standards were purchased and spiked into representative samples to increase confidence in the annotation.
Tentative annotation of features of interest derived from $^1$H NMR analysis was completed by searching on literature and using in-house databases to match possible patterns of interest with the relevant spectra of standard compounds. The multiplicity of the signals of interest was confirmed using the corresponding Jres spectrum and a case of 2D spectra of representative samples including $^1$H,$^1$H-COSY, $^1$H,$^1$H-TOCSY, $^1$H, $^3$C-HSQC.

**Data availability**
Genomics data in European Nucleotide Archive PRJNA266531.

**Code availability**
Github repository.

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Author contributions
The study was conceived and designed by S.L., E.H. and J.K.N. carried out the metabolic profiling design, N.K. designed the analytical pipeline; experimental NMR profiling assays were carried out by B.J., UPLC-MS analysis was conducted by M.G.R., E.C., L.W. and M.R.L., Statistical analysis of the metabolomic data was carried out by T.K., J.P., A.N.H. and A.H.; Data integration and analysis was undertaken by N.K. M.G. repeated the analysis with updated inputs. Interpretation of the metabolomics data was carried out by E.H., J.K.N., L.W., S.S. and C.L.Q. A literature search and linkage of annotated metabolites to AD pathology was done by N.K. in conjunction with S.L. The manuscript was written by N.K. and S.L. in conjunction with A.B., M.G., L.W. All authors reviewed the manuscript.

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Competing interests
The authors declare no competing interests.
Additional information

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