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Toledo, Jon B Arnold, Matthias Kastenmüller, Gabi <u>et al.</u>

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Metabolic network failures in Alzheimer's disease: A biochemical road map

Jon B. Toledo^{a,b,**}, Matthias Arnold^c, Gabi Kastenmüuller^{c,d}, Rui Chang^e, Rebecca A. Baillie^f, Xianlin Han^g, Madhav Thambisetty^h, Jessica D. Tenenbaum^j, Karsten Suhre^{c,k}, J. Will Thompson^l, Lisa St. John-Williams^l, Siamak MahmoudianDehkordi^m, Daniel M. Rotroff^m, John R. Jack^m, Alison Motsinger-Reif^m, Shannon L. Risacher^{n,o}, Colette Blach^{i,j}, Joseph E. Lucas^p, Tyler Massaro^p, Gregory Louie^{q,r}, Hongjie Zhu^{q,r}, Guido Dallmann^s, Kristaps Klavins^s, Therese Koal^s, Sungeun Kim^{n,o}, Kwangsik Nho^{n,o}, Li Shen^{n,o}, Ramon Casanova^h, Sudhir Varma^h, Cristina Legido-Quigley^t, M. Arthur Moseley^l, Kuixi Zhu^e, Marc Y. R. Henrion^e, Sven J. van der Lee^u, Amy C. Harms^v, Ayse Demirkan^u, Thomas Hankemeier^{u,v}, Cornelia M. van Duijn^{u,v}, John Q. Trojanowski^a, Leslie M. Shaw^a, Andrew J. Saykin^{n,o}, Michael W. Weiner^w, P. Murali Doraiswamy^{q,r}, Rima Kaddurah-Daouk^{q,r,x,*}, and for the Alzheimer's Disease Neuroimaging Initiative and the Alzheimer Disease Metabolomics Consortium¹

^aDepartment of Pathology & Laboratory Medicine, University of Pennsylvania, Philadelphia, PA, USA ^bDepartment of Neurology, Houston Methodist Hospital, Houston, TX, USA ^cInstitute of Bioinformatics and Systems Biology, Helmholtz Zentrum Müunchen, German Research Center for Environmental Health, Neuherberg, Germany ^dGerman Center for Diabetes Research (DZD), Neuherberg, Germany eDepartment of Genetics and Genomic Science, Icahn School of Medicine at Mount Sinai, New York, NY, USA ^fRosa & Co LLC, San Carlos, CA, USA ^gSanford Burnham Prebys Medical Discovery Institute, Orlando, FL, USA ^hClinical and Translational Neuroscience Unit, Laboratory of Behavioral Neuroscience, National Institute on Aging, National Institutes of Health, Baltimore, MD, USA ⁱDuke Molecular Physiology Institute, Duke University, Durham, NC, USA ^jDepartment of Biostatistics and Bioinformatics, Duke University, Durham, NC, USA ^kDepartment of Physiology and Biophysics, Weill Cornell Medical College, Qatar, Doha, Qatar ¹Duke Proteomics and Metabolomics Shared Resource, Center for Genomic and Computational Biology, Duke University, Durham, NC, USA "Bioinformatics Research Center, Department of Statistics, North Carolina State University, Raleigh, NC, USA ⁿDepartment of Radiology and Imaging Sciences, Indiana University School of Medicine, Indianapolis, IN, USA °The Indiana Alzheimer Disease Center, Indiana University School of Medicine, Indianapolis, IN, USA PInstitute for Genome Sciences and Policy, Duke University, Durham, NC, USA ^qDepartment of Psychiatry, Duke University, Durham, NC, USA 'Duke Institute for Brain Sciences, Duke University, Durham,

^{*}Corresponding author. Tel.: 919-684-2611; Fax: 919-681-7668, kaddu001@mc.duke.edu (R.K-D.). **Corresponding author. Tel: 713-363-8184. jbtoledoatucha@houstonmethodist.org (J.B.T.).

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NC, USA ^sBIOCRATES Life Sciences AG, Innsbruck, Austria ^tIPS, Faculty of Life Sciences and Medicine, King's College London, London ^uDepartment of Epidemiology, ErasmusMC, Rotterdam, The Netherlands ^vAnalytical Biosciences, Leiden Academic Centre for Drug Research, Leiden University, Leiden, Netherlands ^wDepartment of Radiology, Center for Imaging of Neurodegenerative Diseases, San Francisco VA Medical Center/University of California San Francisco, San Francisco, CA, USA ^xDepartment of Medicine, Duke University Medical Center, Durham, NC, USA

Abstract

Introduction—The Alzheimer's Disease Research Summits of 2012 and 2015 incorporated experts from academia, industry, and nonprofit organizations to develop new research directions to transform our understanding of Alzheimer's disease (AD) and propel the development of critically needed therapies. In response to their recommendations, big data at multiple levels are being generated and integrated to study network failures in disease. We used metabolomics as a global biochemical approach to identify peripheral metabolic changes in AD patients and correlate them to cerebrospinal fluid pathology markers, imaging features, and cognitive performance.

Methods—Fasting serum samples from the Alzheimer's Disease Neuroimaging Initiative (199 control, 356 mild cognitive impairment, and 175 AD participants) were analyzed using the AbsoluteIDQ-p180 kit. Performance was validated in blinded replicates, and values were medication adjusted.

Results—Multivariable-adjusted analyses showed that sphingomyelins and ether-containing phosphatidylcholines were altered in preclinical biomarker-defined AD stages, whereas acylcarnitines and several amines, including the branched-chain amino acid valine and α -aminoadipic acid, changed in symptomatic stages. Several of the analytes showed consistent associations in the Rotterdam, Erasmus Rucphen Family, and Indiana Memory and Aging Studies. Partial correlation networks constructed for A β_{1-42} , tau, imaging, and cognitive changes provided initial biochemical insights for disease-related processes. Coexpression networks interconnected key metabolic effectors of disease.

Discussion—Metabolomics identified key disease-related metabolic changes and diseaseprogression-related changes. Defining metabolic changes during AD disease trajectory and its relationship to clinical phenotypes provides a powerful roadmap for drug and biomarker discovery.

Keywords

Metabolomics; Metabonomics; Pharmacometabolomics; Pharmacometabonomics; Biomarkers; Serum; Metabolism; Systems biology; Biochemical networks; Precision medicine; Alzheimer's disease; Dementia; Branched-chain amino acids; Sphingomyelins; Phospholipids; Acylcarnitines

1. Introduction

Alzheimer's disease (AD) is the most common cause of dementia. An anticipated 136 million people will be affected by dementia by 2050, presenting major global health and

economic challenges. There are currently no treatments that modify AD; hence, AD remains the largest unmet medical need within neurological disorders [1,2].

Many biochemical processes are affected in AD, including amyloid precursor protein metabolism, phosphorylation of tau protein, oxidative stress, impaired energetics, mitochondrial dysfunction, inflammation, membrane lipid dysregulation, and neurotransmitter pathway disruption [3,4]. Impaired cerebral glucose uptake occurs decades before the onset of cognitive dysfunction in AD [5], and neurotoxicity associated with $A\beta$ is thought to participate in impaired neuronal energetics including mitochondrial dysfunction and release of reactive oxygen species. Growing evidence supports the concept that insulin resistance can contribute to AD pathogenesis, and therefore, AD could be regarded as a metabolic disease mediated in part by brain insulin and insulin-like growth factor resistance [3]. Mapping the trajectory of biochemical changes in AD is therefore becoming a priority as filling knowledge gaps about disease mechanisms and their link to metabolic processes can lead to developing much-needed biomarkers and therapies [3]. How does peripheral metabolism, diet, gut microbiome, and exposome impact the metabolic heath of the brain, and thus cognitive function? Which pathways are affected by genes that have been implicated in AD, such as presentiin 1 (PSENI) and PS2 or apolipoprotein E (APOE) genes? Biochemical information elucidating these questions is critical for developing drugs that target enzymes and transporters which regulate metabolism.

Metabolomics provides powerful tools for mapping global biochemical changes in disease and treatment [6-10]. In contrast to classical biochemical approaches that focus on single metabolites or reactions, metabolomics and lipidomics approaches simultaneously identify and quantify hundreds to thousands of metabolites [11–19]. Measurement of large numbers of metabolites enables network analysis approaches and provides means to identify critical metabolic drivers in disease pathophysiology [20]. Initial small-scale metabolomics studies in AD have highlighted metabolic alterations including ceramide-sphingomyelin pathways [10], glycero-phosphatidylcholines (aa = diacyl, ae = acyl-alkyl) [PC] [15,21], PE plasmalogens [22,23], amines [24], and mitochondrial defects [25] among others [13,14]. Metabolic networks have linked central perturbations in norepinephrine and purines with elevated cerebrospinal fluid (CSF) tau, and changes in tryptophan and methionine to decreased A β levels [18]. More recently, the ARIC Neurocognitive Study identified PC aa C36:1 as being linked to lower risk of dementia; however, no metabolite from the panel measured added significantly to prediction of dementia beyond routine clinical variables [26]. A recent plasma-pathology correlative study found that plasma ceramides C16:0, C18:1, C20:0, and C24:1 and monohexosylceramides C18:1 and C24:1 were elevated in those with autopsy-confirmed AD pathology [27], but these metabolites did not differentiate AD from dementia with Lewy bodies. An autopsy study of frontal cortex metabolites from AD patients versus controls showed six central metabolic pathways were altered along with glycerophospholipid metabolism and aspartate metabolism. A metabolomics study in an AD mouse model (APPswe/PS1deltaE9 double transgenic) found abnormalities in polyamine metabolism, essential amino acids, branched-chain amino acids (BCAAs), and serotonin, as well as phospholipid and acylcarnitine homeostasis with brain changes preceding those in the blood [28]. Although these studies highlight specific metabolic underpinnings of AD, not all metabolomics findings have been replicated. For example, a metabolomics study of two

separate cohorts—the Baltimore Longitudinal Study of Aging, and the Age, Gene/ Environment Susceptibility-Reykjavik Study—did not replicate an earlier finding [11,12,15].

Earlier metabolomics studies had major limitations, including not accounting for important confounds such as impact of medications use; small studies that lacked evaluation across data sets; limited ability to connect peripheral metabolic changes with central changes to define what might be related, and lack of attempts to connect metabolic changes within a pathway and network context. Network biology and "network medicine" approaches have become important tools to dissect molecular mechanisms triggering neurodegeneration [29]. This approach accounts for the fact that complex diseases arise from alterations in multiple genes, proteins, and metabolites, and a network may be described as an interaction map among the wide range of biological entities which contribute to disease. As many of the metabolites that are associated with AD are interconnected through metabolic pathways, cofactors, and common intermediates, changes to one metabolite can entail several others, as well as have downstream effects on other coregulated pathways. A systems biology approach integrating metabolites and their interrelations (for instance quantified by partial correlations) in metabolic networks can provide important mechanistic insights about how biochemical reactions are dysregulated during different stages of disease. In contrast to looking at single dysregulated metabolite at a time, the visualization of changes in the metabolic network captures the totality of influences on interconnected biochemical reactions in far more informative ways and allows one to follow these changes over disease stages.

In this large study, we profiled baseline serum samples from the Alzheimer's Disease Neuroimaging Initiative–1 (ADNI-1) cohort where vast data exist on each patient including cognitive decline and imaging changes over many years, information on CSF markers, genetics, and other omics data. We used CSF biomarkers to define early metabolic changes in cognitively normal participants who have CSF pathology and to evaluate metabolic signatures that might be related to $A\beta_{1-42}$ and tau pathology. Using partial correlation networks, we defined progressive metabolic changes that accompany changes in CSF $A\beta_{1-42}$, CSF tau, brain structure, and cognition [30], whereas coexpression networks were used to connect key metabolic changes implicated in disease. The relationship of metabolites with longitudinal cognitive and imaging changes helped us define metabolic signatures correlated with disease progression. Key associations were also present in multiple independent cohorts. We believe that the systems approach taken in our study to elucidate metabolic changes along different stages during the progression of AD will transform our understanding of disease mechanisms and lead to valuable peripheral biomarkers that can inform and accelerate clinical trials.

2. Methods

2.1. Study cohorts and samples

2.1.1. ADNI-1 baseline samples—ADNI shipped 831 samples with unique identifiers belonging to 807 subjects. These initial identifiers were different from the ADNI subject identifiers. There were duplicate aliquots from the same CSF draw for 24 subjects to help us

evaluate analytical performance. Only after the final raw data were submitted to ADNI, the information was obtained to link the samples identifier to the subject RID and identify the duplicates.

Data were obtained from the ADNI database in September 2015 (http://adni.loni.usc.edu). ADNI-1 was launched in 2004 by the National Institute on Aging (NIA), the National Institute of Biomedical Imaging and Bioengineering, the Food and Drug Administration, private pharmaceutical companies, and nonprofit organizations [31]. ADNI-1 patients underwent extensive clinical and cognitive testing, including the Alzheimer's Disease Assessment Scale–Cognition (ADAS-Cog13), which was used as a measure of general cognition in this analysis. AD dementia diagnosis was established based on the NINDS-ADRDA criteria for probable AD. Mild cognitive impairment (MCI) participants did not meet these AD criteria and had largely intact functional performance, meeting predetermined criteria for amnestic MCI [31]. Controls were cognitively normal (CN) (Supplementary Table 1). Additional details of participant selection criteria and protocol are available at http://www.adni-info.org and in the methods in the Supplementary Materials. The study was approved by institutional review boards of all participating institutions, and written informed consent was obtained from all participants and/or authorized representatives before study commencement.

2.1.2. Rotterdam and Erasmus Rucphen Family cohorts—Participants from the Erasmus Rucphen Family (ERF) study (N= 905) were metabolically profiled from fasting blood samples using the Biocrates AbsoluteIDQ-p150 kit platform (Supplementary Table 2) [32], which measures a subset of metabolites from the P180 and excludes many of the amines. A previously described quality control (QC) protocol was applied [32]. Valine was measured in fasting blood samples using the Brainshake platform [33] in 2752 participants from the Rotterdam large prospective cohort study [34]. Participants of the ERF study underwent a standardized cognitive test battery at the study center on the same day blood was drawn (Supplementary Table 2) [35]. Participants of the Rotterdam study underwent cognitive tests at the time of valine measurement, and all participants were followed up for AD clinical diagnosis [36], as previously described in detail [37].

The general cognitive ability or "g-factor" was calculated using previously described methods in dementia-free participants with available cognitive tests in the ERF study (N= 905) and Rotterdam Study (N= 2480) [38]. In short, the g-factor is a general cognitive function phenotype created by principal component analysis of multiple cognitive tests. A higher g-factor is associated with a higher general cognitive function, in contrast to the cognitive measure used for analysis of the ADNI-1 cohort, the ADAS-Cog13.

2.1.3. The Indiana Memory and Aging Study—The Indiana Memory and Aging Study (IMAS) is an ongoing longitudinal study investigating multimodal neuroimaging, cognition, fluid biomarkers, and genetics in early prodromal stages of AD with follow-up visits every 18 months [39–42]. IMAS participants included CN participants, euthymic older adults with subjective cognitive decline in the absence of significant psychometric deficits, and patients with amnestic MCI or probable AD (Supplementary Table 3). Because of limited sample size compared to other cohorts, analyses were limited to assessment of [¹¹C] Pitts-burgh

compound B (PiB) positron emission tomography (PET) amyloid status. Thirty-four participants had PET scans to measure brain Aâ load; 30 participants underwent [11C]PiB PET scans on a Siemens HR + PET scanner; and 4 participants underwent [¹⁸F]Florbetapir PET scans on a Siemens mCT. For the [¹¹C]PiB PET, participants underwent either a 90minute dynamic scan starting at time of tracer injection or a 50-minute dynamic scan after a 40-minute uptake period after injection of approximately 10 mCi of $[^{11}C]PiB$. The $[^{18}F]$ Florbetapir PET scans were collected as a 30-minute dynamic scan after a 40-minute uptake period after an injection of approximately 10 mCi of [¹⁸F]Florbetapir. [¹¹C]PiB and ^{[18}F]Florbetapir scans were motion-corrected and normalized to Montreal Neurologic Institute space using parameters from a same time point structural magnetic resonance imaging (MRI) scan. For the [¹¹C]PiB PET images, a 40- to 90-minute standardized uptake value ratio (SUVR) image was created by averaging the appropriate frames and intensity normalizing to mean cerebellar gray-matter uptake. For the [¹⁸F]Florbetapir PET, a 40- to 70-minute SUVR image was created by averaging the appropriate frames and intensity normalizing to mean whole cerebellar uptake. Finally, amyloid positivity was defined as a mean [¹¹C]PiB PET SUVR of 1.37 or a mean [¹⁸F]Florbetapir SUVR of 1.20 from a cortical gray-matter region of interest. These cutoffs were determined by simultaneous processing of the ADNI [¹¹C] PiB and [¹⁸F]Florbetapir PET images using the same pipeline and adjusting the locally derived cutoffs to best match either the previously reported ^{[11}C]PiB PET cutoff of mean cortical SUVR 1.5 [43] or the [¹⁸F]Florbetapir PET cutoff of SUVR 1.10 [44], respectively. A side-by-side comparison of the three cohorts, including sample sizes, baseline cognitive diagnoses, and studied outcomes in each cohort, is offered in Supplementary Table 4.

2.2. AbsoluteIDQ-p180 kit metabolite measurements

Metabolites were measured with a targeted metabolomics approach using the AbsoluteIDQp180 kit (BIOCRATES Life Science AG, Innsbruck, Austria), with an ultra-performance liquid chromatography (UPLC)/MS/MS system [Acquity UPLC (Waters), TQ-S triple quadrupole MS/MS (Waters)] which provides measurements of up to 186 endogenous metabolites quantitatively (amino acids and biogenic amines) and semiquantitatively (acylcarnitines, sphingomyelins, PCs, and lyso-glycero-phosphatidylcholines (a = acyl) [lysoPCs] across multiple classes) (see methods in Supplementary Materials). The AbsoluteIDQ-p180 kit has been fully validated according to European Medicine Agency Guidelines on bioanalytical method validation. In addition, plates include an automated technical validation to approve the validity of the run and provide verification of the actual performance of the applied quantitative procedure including instrumental analysis. The technical validation of each analyzed kit plate was performed using MetIDQ software based on results obtained and defined acceptance criteria for blank, zero samples, calibration standards and curves, low/medium/high-level QC samples, and measured signal intensity of internal standards over the plate. This is a highly useful platform that was used in hundreds of publications, including several studies in AD [11,12,15].

Deidentified samples were analyzed following the manufacturer's protocol, with metabolomics laboratories blinded to diagnosis and pathological data. Serum samples from all 807 ADNI-1 participants were analyzed, but after QC, a smaller number of participants

were included in the analysis (Supplementary Fig. 1). Three participants were excluded because of incomplete clinical data, 70 samples were excluded because of nonfasting status, and two samples were excluded during the multivariate outlier detection step (see the following), leaving 732 participants included in the final analyses. Each assay plate included two sets of replicates: (1) A set of duplicates obtained by pooling the first 72 samples in the study (QC pool duplicates) and (2) 20 blinded analytical duplicates (blinded duplicates).

2.3. P180 QC

Metabolites with >40% of measurements below the lower limit of detection (LOD) were excluded from the analysis. Metabolite values were scaled across the different plates using the QC pool duplicates. LOD values were imputed using each metabolite's LOD/2 value. Using the blinded duplicates, we selected metabolites with a coefficient of variation <20% and an intraclass correlation coefficient >0.65. Based on the QC process, 32 of the flow injection analysis metabolites and 14 of the UPLC metabolites were excluded from further analysis (Supplementary Table 5). We checked for the presence of multivariate outlier participants by evaluating the first and second principal components in each platform. Two multivariate outliers were beyond 7 standard deviations and were therefore excluded. For the participants with duplicated measurements, we used the average values of the two measured values in further analyses.

2.4. CSF $A\beta_{1-42}$ and tau biomarkers

Lumbar puncture was performed in the mornings after an overnight fast. $A\beta_{1-42}$, total tau (ttau), and tau phosphorylated at threonine 181 (p-tau₁₈₁) were measured using the multiplex xMAP Luminex platform (Luminex Corp, Austin, TX) with Innogenetics immunoassay kitbased reagents (INNO-BIA AlzBio3; Ghent, Belgium; for research use– only reagents) (methods in Supplementary Materials) [45]. CSF samples were available and measured for 48.8% of the CN, 52% of the MCI, and 54.9% of the AD participants. $A\beta_{1-42}$ -defined groups were classified as normal or pathological based on the previously published concentration (192 pg/mL) [46].

2.5. MRI measures

A 1.5-T MRI nonaccelerated sagittal volumetric 3D magnetization-prepared rapid gradientecho MRI images were acquired at each performance site for the ADNI-1 participants (http://www.adni-info.org; http://adni.loni.usc.edu). Only images that passed QC evaluations were included. Cortical gray-matter volumes were processed using the FreeSurfer version 4.4 image processing framework (http://surfer.nmr.mgh.harvard.edu/) [47]. FreeSurfer ventricular volume of MRI scans that passed the QC was adjusted for total intracranial volume and used for longitudinal analyses. The Spatial Pattern of Abnormality for Recognition of Early Alzheimer's Disease (SPARE-AD), an index that captures brain atrophy related to AD and has shown association with AD CSF biomarker and clinical measures, and was calculated for the baseline visit of ADNI-1 participants, was assessed in the present analysis [48,49].

2.6. Medication adjustment

In the ADNI and IMAS cohort, 41 major medication classes used to treat psychiatric (including different categories of benzodiazepines, antipsychotics, and antidepressants) and cardiovascular conditions (including different categories of antihypertensives, cholesterol treatment, and antidiabetics), as well as dietary supplements (Co-Q10, fish oil, nicotinic acid, and acetyl L-carnitine), were systematically coded and available for model-based evaluations of the influence of each drug type on metabolite levels (methods in Supplementary Materials). Intake of any medication within a category was coded as present or absent. Dose effect was not evaluated. The list of the studied medication categories and the percentage of subjects taking these medications in each of the diagnostic categories for the ADNI cohort is listed in Supplementary Table 6.

2.7. Statistical analysis

Metabolites with a skewness >2 [50] that showed a departure of the normality distribution (D'Agostino test *P*-value <.05) [50,51] were \log_{10} transformed to normalize their distribution. We implemented a two-stage regression approach, whereby metabolites were first adjusted for confounding medications and dietary supplements in a linear regression model. For each metabolite, medications were backward-selected via Bayesian information criteria to select an optimal combination of medications for preventing confounding while limiting model complexity. One AD medication class, that is, anticholinesterases, were excluded from this process due to highly significant correlations (Spearman's $\rho = 63.4\%$, $P = 1.28 \times 10^{-83}$) of these medications with diagnosis (for details, see methods in Supplementary Materials). The residuals for each metabolite were then carried forward to test associations with clinical outcomes.

The cross-sectional association with categorical outcomes (clinical diagnosis and CSF $A\beta_{1-42}$ group) was studied using a logistic regression model. For the cross-sectional quantitative outcomes (t-tau/A β_{1-42} ratio, SPARE-AD, and ADAS-Cog13), a linear regression model was applied. Age and gender were forced covariates in all the models associating with clinical variables, and education was also forced into the models for ADAS-Cog13 and clinical diagnosis, whereas APOE £4 was backward-selected based on Bayesian information criteria for each outcome (see methods in Supplementary Materials and Supplementary Table 7). Diagnosis was not included as a covariate in the models in the primary analyses that studied A β_{1-42} , t-tau/A β_{1-42} ratio, SPARE-AD, and ADAS-Cog13 associations. The P-values were Bonferroni corrected to adjust for multiple comparisons and a corrected .05 two-tailed *P*-value was considered significant. A Cox hazard model including age, gender, APOE ɛ4 presence, and education as covariates was used to evaluate the association of metabolite levels with progression from MCI to AD with a median follow-up of 3.0 years (interquartile range [IQR]: 2.0–6.1). A mixed-effects model that included age, gender, education, APOE e4 presence, time, and metabolite level as independent variables was used to study longitudinal associations between the metabolites and volumetric MRI changes (transformed to normalized distribution) during follow-up in the MCI participants (AD participants were excluded because of short follow-up) [52]. A mixed-effects model was also used to evaluate the association of metabolites with change in ADAS-Cog13 (transformed to normalized distribution) and included education as an additional covariate.

Both models accounted for baseline cognitive and MRI measures for each participant. Median follow-up times for the MRI and cognitive analyses were 3.0 years (IQR: 2.0–5.0). An interaction with time was included in all mixed-effects models for the studied metabolites.

In the Rotterdam study, a linear regression model was fitted for the cross-sectional analysis with *g*-factor as the outcome and value as the determinant, adjusting for age, gender, lipid-lowering medication, and education. *P*-values and effect estimates of the significant metabolites are reported. [¹¹C]PiB PET analysis for IMAS samples included age, gender, and *APOE* ε 4 presence, along with the A β_{1-42} status on PET, as independent predictors of target metabolite measures using a linear regression model. All analyses were performed using the R software package [53].

2.7.1. Co-expression network construction and module analysis—We

investigated the global baseline cross-sectional correlation structure of metabolites and their correlation with a subset of clinical and biomarker measures at baseline (A β_{1-42} , tau/A β_{1-42} ratio, and ADAS-Cog13). We built the p180 coexpression network based on baseline-normalized data adjusted for age, education, gender, and *APOE* ϵ 4 presence using the WGCNA R package [54].

2.7.2. Partial correlation analysis—Biochemically related metabolites and propagation patterns of effects on the clinical variables were investigated from a network perspective. A Gaussian graphical model (GGM) calculation was performed as described in Krumsiek et al. [55] using the GeneNet R package with default parameters. To illustrate effect propagation on clinical variables, we colored the resulting network similar to that in Mittelstrass et al. [56]. In brief, a GGM is an undirected graphical model based on partial correlation coefficients, that is, pairwise correlation coefficients conditioned against correlations with all other included variables. GGMs, contrary to correlation networks, thus can reveal the direct relations between metabolites. To account for correlations between metabolites and clinical or other potentially predictive variables, we used metabolite residuals that accounted for effects of medication and dietary supplements (as described previously) and additionally included age, gender, APOE e4 presence, and education as covariates in the GGM generation process. To obtain significant partial correlations, we used a significance threshold of 0.05 after Bonferroni correction for all possible edges in the model (0.05/10,296 $=4.86 \times 10^{-6}$). For each clinical variable, we colored the network representation of the GGM using the results of our regression analyses using $sign(\beta)^*(-log10(P))$ to visualize both strength of association and direction of effect.

3. Results

Metabolomic analyses were performed in the ADNI-1 cohort, and key findings were further tested in the Rotter-dam, EFR, and IMAS cohorts. Overall descriptions of sample size, composition, and studied outcomes across the different cohorts are shown in Supplementary Table 4. The results are presented for each cohort in the following.

3.1. ADNI-1 cohort

In ADNI-1, CN, MCI, and AD participants did not differ in mean age but, as expected, differed in *APOE* ϵ 4 frequency, baseline cognition, MRI atrophy index, and CSF levels of T-tau and A β_{1-42} . The heat map (Fig. 1) (and the later described coexpression networks in Supplementary Fig. 2) shows that the global (i.e., direct and indirect) correlation structure between metabolites is formed into biochemical classes, illustrating that the metabolites with significant findings can be seen as proxies for the group of their correlating metabolites (Supplementary Material File 2 presents the correlation values).

3.2. ADNI-1: Metabolites associated with cross-sectional clinical, MRI, and CSF biomarker measures

The metabolites that remained in the analyses after the QC steps showed different correlation strengths, indicating groups of metabolites that may be involved in similar processes (Fig. 1). After applying Bonferroni multiple comparison correction, 13 metabolites showed significant associations (Bonferroni-adjusted P-value <.05) with cognitive scores and CSF and MRI biomarker measures (Table 1). Six metabolites were associated with CSF AB1-42 positivity (ether-containing PC [PC ae] C36:2, PC ae C40:3, PC ae C42:4, PC ae C44:4, SM (OH) C14:1, SM C16:0), four were associated with t-tau/A β_{1-42} ratio (C18, PC ae C36:2, SM C16:0, SM C20:2), five were associated with ADAS-Cog13 scores (C14:1, C16:1, SM C20:2, a-aminoadipic acid [a-AAA], and valine), and 6 were associated with SPARE-AD scores (C12, C16:1, PC ae C42:4, PC ae C44:4, α-AAA, and valine). In all analyses, higher acylcarnitine, PC, and sphingomyelin (SM) values were associated with worse clinical and biomarker measures, whereas the opposite direction of associations was observed for valine and α-AAA values. The complete results for the 138 studied metabolites are listed in Supplementary Table 8, where many amines (including isoleucine, glutamate, tyrosine, tryptophan, glycine, proline, histidine, T4OH proline) and other metabolites within PC and SM classes showed significant non-comparison-corrected associations with clinical markers and outcomes but did not survive Bonferroni multiple comparison correction. Supplementary Tables 9 and 10 present the analyses adjusted by clinical diagnosis and stratified by each of the three clinical diagnostic categories, respectively. All significant correlations were in the same directions in the clinical diagnostic groups.

We next evaluated differences in levels of key metabolites associated with cognitive or biomarker measures from the analyses reported previously between the three diagnostic groups (CN, MCI, and AD) subclassified by CSF $A\beta_{1-42}$ positivity status. Metabolites showed three different patterns of associations with the CSF AD biomarkers. PC ae C44:4, PC ae C36:2, and C18 represented the most significant examples of each of these patterns, and the values in the six groups are shown in Fig. 2. Of note, CN participants (red boxes) with pathological CSF $A\beta_{1-42}$ values showed significant metabolic changes in a specific group of metabolites compared with CN with no pathological CSF $A\beta_{1-42}$ values (Fig. 2A). Some of the changes associated with CSF $A\beta_{1-42}$ values appeared in clinical stages of disease (MCI and AD; Fig. 2B). Other metabolic changes were only observed in comparing CN participants to clinically impaired subjects (Fig. 2C) but showed no associations with

pathological CSF A β_{1-42} status. Fig. 2D illustrates value correlation with cognition in the ADNI-1 study.

3.3. Metabolites associated with longitudinal outcomes in the ADNI-1 cohort

We evaluated whether levels of metabolites at baseline were associated with (1) ADAS-Cog13 changes up to 5 years; (2) ventricular volume changes up to 5 years; or (3) progression from MCI to AD (Table 2). Regression coefficients of six metabolites (PC ae C40:3, PC ae C42:4, PC ae C44:4, SM (OH) C14:1, SM C16:0, and SM C20:2) showed a positive association with all three longitudinal outcomes. In addition, lower valine and α -AAA values were associated with faster cognitive decline, similarly the coefficient for valine was negatively associated with ventricular volume changes. Fig. 3 shows some of these associations as examples, including Fig. 3A which shows the Cox hazards model of the association between baseline concentration of SM 20:2 (presented as tertiles) and longitudinal cognitive (ADAS-Cog13) and MRI (brain ventricular volume) change.

3.4. Evaluation of findings in the Rotterdam and ERF studies

In the Rotterdam and ERF studies, only a subset of metabolites were measured from the panel of P180 metabolites evaluated in the ADNI-1 study (P150 panel, Supplementary Table 11). Using a targeted approach, we tested whether the metabolites that showed a significant association in the ADNI-1 study were also correlated with cognition (general cognitive ability: g-factor) in the Rotterdam Study or ERF.

For the cross-sectional analysis, eight metabolites were available in the ERF study. Two of these metabolites (PC ae C40:3 and SM C20:2) were associated with cross-sectional general cognitive ability in the expected direction based on the discovery ADNI-1 cohort. Notice that higher general cognitive ability levels indicate better cognition as opposed to ADAS-Cog13. Valine was strongly associated with a higher general cognitive ability (P=.00035) in the Rotterdam study (Fig. 2E), which is in line with the association with ADAS-Cog13 in ADNI-1 (Fig. 2D). Longitudinally, 342 participants developed AD in the Rotterdam study after a median follow-up time of 9.7 years (IQR 5.6–10.5). A Cox proportional hazard model was fitted adjusting for age at baseline, gender, education, and lipid-lowering medication and indicated that a 1-SD increase in valine concentration was also associated with a decreased risk of AD (P=.044).

3.5. Evaluation of $A\beta_{1-42}$ signature in the IMAS cohort

Three of the six metabolites (PC ae 42:4, PC ae 44:4, and SM (OH) C14:1) that showed an association with CSF A β_{1-42} positivity in the ADNI-1 cohort were also associated with amyloid positivity on PET in the IMAS cohort (*n* = 34; Supplementary Table 12).

3.6. Partial correlation networks for A β_{1-42} , T-tau, SPARE-AD, ADAS-Cog13—metabolic trajectory for disease

We strived to define insights into the trajectory of biochemical changes that correlate with established models for disease [30], A β pathology, tau, imaging changes, and cognitive decline, building partial correlation networks for each clinical measure.

The partial correlation networks established direct connections between the measured metabolites, which have been shown to depict direct biochemical relationships between metabolites while omitting mediated correlations [55,57]. Fig. 4 integrates the strength of the partial correlations between metabolites and overlays on these networks the associations with the studied outcomes $A\beta_{1-42}$ t-tau, SPARE-AD, and ADAS-Cog13 (partial correlation networks for p-tau and t-tau/A β_{1-42} ratio are not shown). The networks showing the direct links between metabolites (nodes) identified through their strong partial correlations (edges) expand the heat map information (Fig. 1). Through coloring of the metabolite nodes by their association to CSF, imaging, and cognitive markers, respectively (where bright colors indicate strong associations and blue and red color indicate upregulation and downregulation of metabolites), these networks demonstrate how the effects of clinical variables propagate along the edges within the network suggesting that the results follow biochemically plausible pathways. The studied outcomes cover the different stages of AD, matching known biomarkers of disease [30].

The network for $A\beta_{1-42}$ (Fig. 4A) highlighted direct correlations with short- and mediumchain SMs and PC with ether bonds, suggesting a role for membrane structure and function, contact sites, and membrane signaling in amyloid pathology. The correlation pattern for t-tau (Fig. 4B) highlighted metabolites among long-chain acylcarnitines and SMs implicated in lipid metabolism. The SPARE-AD and ADAS-Cog13 (Fig. 4B) partial correlation networks were very similar, suggesting associations of brain atrophy and cognitive decline with metabolic changes in BCAAs and short-chain acylcarnitines implicated in mitochondrial energetics as well as additional changes in lipid metabolism.

3.7. Coexpression network—direct and indirect connections for key metabolites

The partial correlation networks mentioned previously evaluated only direct connections among metabolites. To capture both indirect and direct correlations, we built coexpression networks to evaluate the number of modules in our data set and evaluate additional connections between key metabolites identified as related to cognitive or biomarker measures in ADNI-1. The full coexpression network, which identified seven metabolic modules, can be found in Supplementary Fig. 2, and Supplementary Material File 3 presents each of the individual correlations.

We investigated the correlation structure of the three metabolites in the ERF and Rotterdam data sets that significantly associated with cognition, namely PC ae C40:3, SM C20:2, valine as shown in Fig. 5. The subnet-work shows these three metabolites to have high correlations (marked as red edges) to other functional metabolic modules via direct and indirect links. Valine was highly correlated with isoleucine and α -AAA, whereas SM C20:2 was highly correlated with a subset of the SMs including SM C16:0. Finally, PC ae C40:3 was highly correlated with PCs and SMs, but not amines and acylcarnitines. These SMs and PCs were significantly associated with cognitive scores, CSF biomarkers, and MRI measures (Table 1).

4. Discussion

The Alzheimer Disease Metabolomics Consortium (ADMC) funded by the NIA under the AMP-AD and MOVE-AD initiatives and in partnership with ADNI has as its goal to create a comprehensive metabolomics database for AD. These data will fill in biochemical knowledge about disease mechanisms that can be used as a roadmap for novel drug discovery and establishment of blood-based biomarkers. Eight complementary, targeted and nontargeted, metabolomics platforms are currently in the process of generating data on ADNI participants to define the metabolic trajectory of disease connecting central and peripheral metabolic failures in a pathway and network context. We seek to replicate earlier findings and test hypotheses but also expand on biochemical coverage to better understand disease pathogenesis by using complementary data unique to ADNI-1. The unique opportunity of having longitudinal cognitive and imaging data on each subject for close to a decade enables identification of peripheral biomarkers that are disease related.

This is the first report from ADMC on use of a targeted, highly validated metabolomics platform with the analysis guided by CSF markers and imaging data. Using 732 baseline serum samples from the ADNI-1 cohort, we systematically evaluated the relationships between metabolomics data and cross-sectional clinical, CSF, and MRI measures, as well as their association with longitudinal cognitive and brain volume changes. We demonstrate a novel approach to the analysis of neurodegenerative disease mechanisms. Multiple comparisons and covariate-adjusted analyses, that included relevant medications, identified sets of metabolites that became altered at specific disease stages (preclinical AD with biomarker-defined AD pathology vs. symptomatic stages) [30]. Using partial correlation networks, we integrate our findings of metabolic effects on AD pathogenesis, linking central and peripheral metabolism in a way that consistently reconciles biochemical trajectories of disease with this established temporal sequence of pathophysiological stages of AD. In the following, we therefore discuss our findings in their temporal order along AD pathogenesis.

4.1. Aβ pathology

Changes observed earliest in AD, namely in biomarker-defined preclinical stages [58] in CN participants, were higher levels of a specific set of PCs (PC ae C36.2, PC ae C40.3, PC ae C42.4, and PC ae C44.4) and SMs (SM (OH) C14.1, SM C16.0). These metabolites were associated with abnormal CSF $A\beta_{1-42}$ values in CN participants to a similar degree as observed in MCI, indicating an early role of ether-containing PC species and SM in the development of the disease. Interestingly, most of these metabolites were also associated with later cognitive decline and global brain atrophy changes in the MCI group (Table 1). The use of information on AD CSF biomarkers was extremely useful and guided our metabolomics data. Only after substratification of the diagnostic groups (CN, MCI, and AD) based on CSF biomarkers, we were able to detect the increased values of these metabolites in participants with CSF pathology stressing the importance of using biomarkers to detect early biochemical changes [59]. Our findings along with those of several others [11,15] all point to problems with phospholipid metabolism that happen early in the disease process. These early changes by themselves are not sufficient predictors of disease or progression, but they might provide extremely valuable mechanistic insights.

Partial correlation networks showed that the pathological CSF A β_{1-42} values were associated with two groups of lipids, composed primarily of ether-containing PCs and relatively short-chain SMs. Ether-containing PC (PC ae) metabolites are PC species with an ether linkage of an aliphatic chain to the first hydroxyl position of glycerol. These lipids are not completely characterized and may represent a mixture of lipid metabolites including plasmalogens, acyl-alkyl PC, or PC containing an odd-numbered fatty acyl chain. When measured in the serum, the ether-containing lipids are derived from liver metabolism and are possible indicators of peroxisomal function and lipid oxidation status [60,61]. Plasmalogens and SMs are enriched in membrane rafts where they facilitate signal transduction and serve as a source for lipid secondary messengers [62]. The association of PCs and SMs in our study and others [14,63] with early changes in AD and with pathological CSF A β_{1-42} levels (Figs. 4B and 5) may be indicative of early neurodegeneration and loss of membrane function. Ether-linked PC metabolites are found in high abundance in plasma membranes and are a source for signaling molecules [64], particularly platelet-activating factor and arachidonic acid. Similarly, they are found in high abundance in immune cells, are regulatory factors, and may be part of the link between inflammation and AD [65]. Both SMs and ether-linked PCs may be located in membrane rafts, supporting the hypothesis that lipid rafts are directly associated with regulation of amyloid precursor protein processing, the production of $A\beta_{1-42}$, and facilitate its aggregation [66]. All of these lipid biological functions related to $A\beta_{1-42}$ might provide great insight about mechanisms of $A\beta_{1-42}$ pathology, but at this early stage, we cannot assign cause or effect. Because this study was conducted using peripheral blood, we cannot directly associate the changes in neuronal lipid rafts with changes in signaling and accumulation of cerebral A β_{1-42} .

Previous smaller studies relying on clinical measures identified only associations between these metabolite categories and dementia diagnosis or clinical progression [12,15,67], indicating that these metabolic pathways may contribute significantly to AD pathophysiology. Differences in the specific metabolites identified in the published studies [12,15,67] may be due to differences in sample selection criteria, disease severity, or for the absence or presence of controlling for different confounders in the data analysis.

4.2. Tau pathology

In this study, pathological CSF $A\beta_{1-42}$ shows an association with ether-linked PC, and shorter chain SM, but not amines, lysoPC, or acylcarnitines. $A\beta_{1-42}$ changes happen early in disease, followed by accumulation of tau protein in the CSF [30]. In our analysis, tau-related metabolites were very different both from those that correlate with $A\beta_{1-42}$ as well as from metabolites associated with brain atrophy and cognitive changes. Tau-related metabolites thus appear to belong to an intermediate stage between $A\beta_{1-42}$ accumulation and changes in imaging and cognitive function (Fig. 4B), further supporting our hypothesis that different metabolic events occur at different disease stages.

Long-chain acylcarnitines, PC ae C36:2, and SM.C20:2 were higher only in cognitively impaired participants with AD-like CSF $A\beta_{1-42}$ values, indicating that changes in these metabolites are more specific to AD-related neurodegeneration reflecting possible changes in multiple cellular processes. Specifically, accumulation of acylcarnitine species containing

long fatty acyl chains indicates malfunction of fatty acid transport and/or β -oxidation in mitochondria, inefficient utilization of fatty acids as energy substrates [68], or alterations in tau metabolism [69]. In the present study, we revealed that the levels of several acylcarnitine species were increased either at the MCI stage or in clinical AD [70] (Table 1). Acylcarnitines have important functions in the brain [70] such as mitochondrial function, energetics, and neurotransmission that need to be further explored and connected to peripheral function.

4.3. Brain volume changes and cognitive decline

In our study, partial correlation networks show a pattern of inverse associations between brain volume changes (measured by SPARE-AD) and cognition (ADAS-Cog13) and long and short acylcarnitines, value, and α -AAA, indicating a shift in energy substrate utilization in later stages of AD (Fig. 4). By using a second type of networking analysis, a coexpression network, our data show the relationship between valine and short acylcarnitines (Fig. 5, dark green cluster), reinforcing the hypothesized changes in energetics. The association of the long-chain acylcarnitines, odd-numbered acylcarnitines, and amino acids in relation with ADAS-Cog scores supports a switch of utilization from fatty acids to amino acids and glucose. In the network analysis, the amines and short-chain acylcarnitines did not link to PCs and SMs, rather they clustered together in smaller groups. This may indicate that the short-chain acylcarnitines are associated in energy and amino acid metabolism rather than lipid metabolism in AD participants. This novel finding indicates a disease-associated transition in pathways for utilization of energy substrates. Whether such a switch is associated specifically with AD pathogenesis or is a function of aging [71] remains to be tested. Interestingly, ADAS-Cog13 does not show the associations with SMs or ether-linked PCs found with pathological CSF A β_{1-42} . Therefore, it is possible that metabolic pathways other than those associated with lipid rafts are dys-regulated later in the disease process.

It is not clear if insulin resistance is a cause, an effect, or some combination of cause and effect in AD [72]. Insulin resistance, obesity, and diabetes are risk factors for AD [73], with lipid metabolism disorder (and inflammation) being a common link between metabolic disease, vascular disease, and AD. Several ether-linked PC metabolites have been associated with the risk of diabetes [74], insulin resistance promotes aminoacidemia and the use of amino acids for energy [68], and BCAA and α -AAA have been identified as predicting diabetes risk [75,76]. BCAAs (valine, leucine, and isoleucine) play central roles in metabolism and have been implicated in insulin resistance, type-2 diabetes mellitus, and obesity. Our findings in ADNI related to low levels of valine and its correlation with cognitive changes were confirmed in the large Rotterdam study, pointing to an important role for this BCAA in cognitive changes in AD. Low levels of BCAAs have been implicated in hepatic insulin resistance in liver disease and may have a broader role in insulin resistance in the brain [77]. The seemingly paradoxical directionality difference in correlation of BCAAs with diabetes and cognition needs to be further evaluated in longitudinal studies taking into account weight changes, tissue type, and differences in human and animal model systems [28,78]. Our understanding of the biochemical crossroads between diabetes and AD could be greatly enhanced by metabolic profiling of both central and peripheral tissues in both diseases and over time.

4.4. Concluding remarks

In summary, by using metabolomics and network approaches, this study has revealed lipid metabolic changes related to early stages of disease, as well as later changes related to mitochondrial energetics and energy utilization. We hypothesize that the lipid changes measured in this study reflect alterations in membrane structure and function early in the disease process and suggest a change in lipid rafts, which in turn, cause alterations in $A\beta$ processing [66]. We hypothesize that over time, the changes in lipid membranes, particularly mitochondrial membranes, results in increased lipid oxidation, loss of membrane potential, and changes in membrane transport [79,80]. All of these lipid membrane changes might be reflected as disruption in BCAA as an energy source, production of acylcarnitines, and altered energy substrate utilization. The link between the altered energy metabolism, glycolysis, use of ketone bodies, and development of insulin resistance are all subject of ongoing longitudinal studies using complementary metabolomics platforms that enable the study of those biochemical pathways. The specific interactions between the peripheral metabolic network changes, central changes, and the timeline of AD pathophysiology reveal that peripheral metabolic changes can impact long-term brain health and function. This study thus provides an approach to define viable disease biomarkers based on understanding of whole-body AD pathophysiology at a systems level.

Previous studies have shown seemingly conflicting results regarding the diagnostic and prognostic classification of metabolomics data in AD [11,15]. Our findings show that using a network approach, many of these findings might be related such as early changes in phospholipid metabolism. Large meta-analysis across many metabolomics studies using our network approaches is ongoing.

By using the unique data available as part of ADNI, we were able to establish robust associations accounting for several confounders that lead to a better understanding of metabolic changes present in AD. Gender showed the strongest associations with many metabolites, but also age, BMI, and several medications and food supplements, such as statins and fish oils, had significant effects on several metabolites. The fact that medication use varied among the groups (e.g., fish oil intake was more frequent in CN and MCI subjects while use of selective serotonin reuptake inhibitors was lower in CN subjects) and that sometimes studied groups are unbalanced in terms of demographics makes it imperative to account for all possible confounders. Our results thus suggest that future studies should account for associations of metabolites with both gender and medications. This is an important outcome, as prior studies have failed to address effects of medications on the metabolome; a limitation that can cause misleading conclusions. Many of the drugs used by AD patients were shown to have profound effects on metabolism (for reviews on drug effects see Kaddurah-Daouk and Weinshilboum [8,9]). A detailed analysis of medications as confounds and meta-analysis of previously reported metabolomics studies addressing confounds will be published separately.

Limitations of our study are several. At this early stage, we cannot determine which of the metabolic changes we see are causative and which are a consequence of disease. Future mechanistic studies in model systems are needed to test hypothesis generated along with building of predictive metabolic networks. Gender metabolic differences are clear from this

data set and although adjusted for in this study ongoing analysis and modeling of gender differences will be key. Profiling blood samples across the trajectory of disease will provide valuable information and is currently ongoing. In addition, analysis of CN cohort with normal CSF A β_{1-42} values will be useful to evaluate within subject variability. Linking peripheral and central metabolic changes in blood, CSF, and brain tissue is critical for better defining peripheral influences like diet and environment on brain health and disease. The effect of confounds like medications impacts metabolomics findings in significant ways and must be addressed carefully. In our study, we had medication data for two of the cohort (ADNI and IMAS), but this information was not available for the Rotterdam and ERF studies. Therefore, there are differences in how the data were processed across cohorts. Finally, neuropsychological assessments evaluated in the ADNI and Rotterdam and ERF scores were not the same. ADNI analyses were based on a global cognitive scale, whereas in the Rotterdam and ERF studies, a composite measure, the g-factor. Therefore, these scales might differently represent the summary of the diverse cognitive functions.

Over the past decade, we have gained significant knowledge about effects of commonly used medications and have tried to address their confounds in studies such as ADNI. More sophisticated modeling approaches will be needed to address complex interactions with medications so this study presents early effort. Defining genetic factors that influence metabolic changes is key and is ongoing. Broad biochemical coverage is needed to better define mechanism and trajectory of disease. The ADMC is producing metabolomics data sets from eight targeted and nontargeted platforms on the large ADNI cohort. After these and other cohort data sets are available, we look forward to working with the international scientific community to build a comprehensive model for AD using integrated multiomic network approaches.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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RESEARCH IN CONTEXT

- Systematic review: We searched PubMed and reviewed all publications reporting on the use of metabolomics technologies for defining signatures in plasma, serum, or cerebrospinal fluid (CSF) in Alzheimer's disease (AD), mild cognitive impairment (MCI) when compared to healthy control subjects. Few small cohort studies have reported on findings with no replication and validation and results have been conflicting. Previous studies have not accounted for medication effects and did not include imaging or Aβ biomarkers for connecting peripheral and central changes.
- 2. Interpretation: Studying a large cohort of patients (AD, MCI, and cognitively normal) at different disease stages with guiding data on CSF and magnetic resonance imaging AD biomarkers, we identified peripheral metabolic changes in preclinical stages of disease and additional metabolic changes associated with imaging changes and cognitive decline. Results were validated in three additional cohorts (Rotterdam, Erasmus Rucphen Family, and Indiana Memory and Aging Study). Partial correlation networks and coexpression networks revealed progressive interconnected biochemical changes in disease.
- **3.** Future directions: (1) Expand biochemical coverage and the number of metabolic processes studied, (2) continue to link peripheral and central metabolic changes, and (3) test which changes are causative and which are secondary to disease processes.



Fig. 1.

Clustering of pairwise metabolite correlations and association results with clinical variables. (A) Heat map of Spearman correlations between the residuals of metabolite concentrations on the single metabolites. Metabolites are clustered using hierarchical clustering using the Euclidean distance metric. The clustering assigns metabolites to their biochemical class: amino acids, biogenic amines, short-chain and long-chain acylcarnitines, lyso-lipids, PC, and SM. Significant clusters of acylcarnitines are outlined in blue and amines outlined in brown. (B) Association results of the regression analyses. The distribution of association

results of metabolites with clinical variables mirrors the correlation structure of the metabolites. Abbreviations: α -AAA, α -aminoadipic acid; AD, Alzheimer's disease; C0, free carnitine; C*x:y*, acylcarnitines; C*x:y*-OH, hydroxylacylcarnitines; C*x:y*-DC, dicarboxylacylcarnitines; CN, cognitively normal; lysoPC, lyso-glycero-phosphatidylcholines (*a* = acyl); MCI, mild cognitive impairment; Path. A β_{1-42} , pathological A β_{1-42} ; PC, glycero-phosphatidylcholines (*a* = diacyl, *ae* = acyl–alkyl); SDMA, symmetric dimethylarginine; SM, sphingomyelin; SM*x:y*, sphingomyelins; SM (OH) *x:y*, N-hydroxylacyloylsphingosyl-phosphocholine; T4-OH-Pro, trans-4-hydroxyproline.



Fig. 2.

Relationship between serum metabolites, clinical diagnosis, and $A\beta_{1-42}$ status. Serum PC ae 44:4 (A), PC ae 44:4 (B), and C18 (C) concentrations stratified by clinical diagnosis and CSFA β_{1-42} -defined groups. The concentration of each metabolite is shown for each diagnosis red: CN, green: MCI, blue: AD and by *N*. Abeta: normal concentrations of $A\beta_{1-42}$ (>192 pg/mL), and Path. Abeta: pathological concentrations of $A\beta_{1-42}$ (<192 pg/mL), Y- axes are values for each metabolite. Scatter plot for ADAS-Cog13 and serum values (D). Black line and shading are the regression line and 95% confidence interval. (D and E)

Correlations between valine levels and cognitive decline in ADNI-1 and Rotterdam, respectively. Abbreviations: α -AAA, α -Aminoadipic acid; ADAS-Cog13, Alzheimer's Disease Assessment Scale–Cognition; ADNI-1, Alzheimer's Disease Neuroimaging Initiative–1; C0, free carnitine; C*x:y*, acylcarnitines; C*x:y*-OH, hydroxylacylcarnitines; C*x:y*-DC, dicarboxylacylcarnitines; lysoPC, lyso-glycero-phosphatidylcholines (*a* = acyl); PC, glycero-phosphatidylcholines (*aa* = diacyl, *ae* = acyl–alkyl); SDMA, symmetric dimethylarginine; SM*x:y*, sphingomyelins; SM (OH) *x:y*, N-hydroxylacyloylsphingosylphosphocholine; T4-OH-Pro, trans-4-hydroxyproline.

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SM C20:2 25th percentile 50th percentile 75th percentile



Fig. 3.

Longitudinal associations for SM C20:2. (A) Cox hazards model of the association of conversion from MCI to AD. Black line: 1st tertile, red line: 2nd tertile, green line: 3rd tertile. Analysis was conducted using quantitative values, and stratification by tertiles was used only for graphical representation. (B) Association between baseline concentrations of SM 20:2 and longitudinal cognitive (ADAS-Cog13) and imaging (MRI: brain ventricular volume) changes during follow-up. Lines represent trajectories on subjects on the 25th percentile (black line), 50th percentile (red line), 75th percentile (green line) of baseline SM 20:2. Y-axes are ADAS-Cog13 score (left) and ventricular volume (right). Trajectories for these values are calculated based on the studied mixed-effects models. Abbreviations: AD, Alzheimer's disease; ADAS-Cog13, Alzheimer's Disease Assessment Scale–Cognition; MCI, mild cognitive impairment; MRI, magnetic resonance imaging.



Fig. 4.

Network model showing metabolic pathways correlated with the temporal evolution of biomarkers and clinical variables in AD. (A) Partial correlation network. Gaussian graphical model of metabolite concentrations showing reconstructed metabolic pathways and highlighting of the different modules involved in the steps along the temporal evolution of biomarkers and clinical variables in AD. Nodes in the network represent the metabolites, and edges (lines) illustrate the strength and direction of their partial correlations. Only partial correlations significant after Bonferroni correction for all possible edges are included.

Labels show the major classes of metabolites included in our study. Gray circles outline the modules highlighted in panel B. (B) Schematic diagram of the model of temporal evolution of biomarkers in AD, modified from Jack and Holtzman [30], augmented with colored versions of the network from panel A. In these networks, nodes are highlighted according to the strength and direction of the metabolite's association with the respective clinical trait with blue as positive and red as negative (networks in temporal order from left to right: pathological A β_{1-42} , T-tau, SPARE-AD, and ADAS-Cog13). Significant associations are colored in dark blue/bright red, and weaker (but at least nominally significant at 0.05) associations are displayed in fainter colors. Modules of metabolites implicated in the respective trait are highlighted by circles colored by their first occurrence in the temporal order following the color scheme of the time sequence on the bottom. The partial correlation network for $A\beta_{1-42}$ (panel A) highlighted direct correlations with short- and medium-chain SM and PC with ether bonds suggesting a role for membrane structure and function, contact sites, and membrane signaling in amyloid pathology. There was a different pattern for tau (panel B) with highlighted metabolites with long-chain acylcarnitines and SM implicated in lipid metabolism showing association with T-tau level. The SPARE-AD and ADAS-Cog13 partial correlation networks were very similar suggesting associations of brain atrophy and cognitive decline with metabolic changes in BCAAs and short-chain acylcarnitines that have been implicated in mitochondrial energetics as well as additional changes in lipid metabolism. Abbreviations: AD, Alzheimer's disease; ADAS-Cog13, Alzheimer's Disease Assessment Scale-Cognition; BCAA, branched-chain amino acid; PC, glycerophosphatidylcholines (aa = diacyl, ae = acyl-alkyl); SM, sphingomyelin; SPARE-AD, Spatial Pattern of Abnormalities for Recognition of Early AD.

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Fig. 5.

Coexpression subnetwork with direct and indirect interconnections between select metabolites. A coexpression subnetwork focused on three metabolites also identified in the Rotterdam data set (PC ae C40:3, valine, and SM C20:2) was generated from the primary network (Supplementary Fig. 2). The subnetwork shows these three metabolites have high correlations (red edges lines) and lower correlations (green edges lines) to multiple modules via direct and indirect interconnections. Each module is denoted by a color representing a robust set of coregulated metabolites in interconnected biochemical pathways, for example, orange module contained a subset of amines, green module consists of long-chain acylcarnitines; teal, brown, and blue modules contained exclusively PC and lysoPC; red module contained SM and PC; gray module contained short-chain acylcarnitines and other amines. Each node represents a metabolite. The edge (line) opacity is proportional to the Pearson correlation, that is, lighter means weaker correlation value and darker means stronger correlation. The intermodule edges represent correlations and potentially indirect interactions among metabolites and biochemical pathways. The coexpression network captures all significant associations between metabolites and reveals a global correlation structure and interconnections among different modules that adds to our understanding of the disease network. Abbreviations: lysoPC, lyso-glycero-phosphatidylcholines (a = acyl);

PC, glycero-phosphatidylcholines (*aa* = diacyl, *ae* = acyl–alkyl); PC ae, ether-containing PC; SM, sphingomyelin.

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Metabolites	MCI	AD	$A\beta_{1-42}$	T-tau/Aβ _{1−42}	ADAS-Cog13	SPARE-AD
C12	(0.1)(0.0)	-1.62 (1.0)	1.22 (1.0)	0.26 (.33)	5.88 (.073)	0.87 (.041)
C14:1	10.79 (1.0)	-12.25 (1.0)	12.93 (1.0)	2.46 (.05)	52.21 (.037)	6.8 (.1)
C16:1	1.25 (1.0)	-2.098 (1.0)	1.62 (1.0)	0.38 (.091)	9.4 (.0037)	1.2 (.020)
C18	14.62 (1.0)	-19.27 (1.0)	21.62 (1.0)	4.64 (.0055)	64.31 (.5)	10.0095 (.2)
PC ae C36:2	0.085 (.33)	-0.082(1.0)	0.16 (.007)	0.018 (.013)	0.23(1.0)	0.027~(1.0)
PC ae C40:3	0.98 (1.0)	-3.27 (1.0)	5.76 (.017)	0.49 (.55)	2.72 (1.0)	0.26(1.0)
PC ae C42:4	1.62 (.063)	-1.51 (.88)	2.32 (.017)	0.19 (.75)	3.63(1.0)	0.79 (.049)
PC ae C44:4	3.029 (1.0)	-3.37 (1.0)	6.11 (.016)	0.6 (.089)	11.24 (.64)	2.059 (.037)
SM (OH) C14:1	0.06 (1.0)	-0.054(1.0)	0.24 (.044)	0.027 (.081)	0.2~(1.0)	0.016(1.0)
SM C16:0	0.0065(1.0)	-0.0074(1.0)	0.015 (.016)	0.0017 (.013)	0.024 (1.0)	0.0037 (.57)
SM C20:2	0.66~(1.0)	-1.082 (.22)	0.74 (1.0)	0.18 (.047)	4.57 (<.0001)	0.4(.48)
α-AAA	-0.46(1.0)	0.67~(1.0)	-0.68 (1.0)	-0.13 (.098)	-3.7 (.0025)	-0.61 (<.0001)
Valine	-0.0038(1.0)	0.0073 (.079)	-0.004(1.0)	-0.0006(1.0)	-0.028 (< 0.001)	-0.0039 (< 0001)

Abbreviations: MRI, magnetic resonance imaging; CSF, cerebrospinal fluid; MCI, mild cognitive impairment; AD, Alzheimer's disease; ADAS-Cog13, Alzheimer's Disease Assessment Scale-Cognition; SPARE-AD, Spatial Pattern of Abnormalities for Recognition of Early AD; α-AAA, α-aminoadipic acid.

NOTE. The cells include the logistic (MCI and AD) and linear (A β 1-42, T-tau/A β 1-42, ADAS-Cog13, SPARE-AD) regression coefficients and, in parenthesis, the Bonferroni corrected *P*-value. All model included age and gender as covariates. APOE e4 presence included in A\b1-42 model and education was included in the MCI, AD, and ADAS-Cog13 models. A complete list is found in Supplementary Table 2. Bold text indicates significant results.

Table 2

Association of metabolites with longitudinal cognitive and MRI changes in MCI

Analytes	ADAS-Cog13 change	Ventricle volume change	Progression MCI to AD dementia
C12	0.091 (.26)	0.11 (.73)	1.37 (.4)
C14:1	1.39 (.034)	7.085 (.006)	2.11 (.22)
C16:1	0.15 (.13)	0.67 (.092)	1.9 (.19)
C18	-0.16 (.87)	1.94 (.64)	2.41 (.18)
PC ae C36:2	0.0075 (.094)	0.031 (.096)	1.056 (.012)
PC ae C40:3	0.38 (.02)	1.5 (.020)	5.98 (.027)
PC ae C42:4	0.15 (.04)	0.72 (.013)	1.96 (.042)
PC ae C44:4	0.49 (.0076)	2.33 (.0012)	5.89 (.027)
SM (OH) C14:1	0.015 (.04)	0.075 (.01)	1.08 (.025)
SM C16:0	0.0009 (.025)	0.0037 (.023)	1.004 (.029)
SM C20:2	0.11 (.0078)	0.48 (.0035)	1.9 (.0023)
a-AAA	-0.093 (.022)	-0.29 (.087)	0.68 (.061)
Valine	-0.0006 (.035)	-0.0027 (.026)	1.0 (.27)

Abbreviations: MRI, magnetic resonance imaging; MCI, mild cognitive impairment; ADAS-Cog13, Alzheimer's Disease Assessment Scale-Cognition; AD, Alzheimer's disease; PC ae, ether-containing PC; a-AAA, a-aminoadipic acid.

NOTE. Table depicts the association between selected metabolites and longitudinal ADAS-Cog13 (column 2) and ventricular volume (column 3) in mixed-effects models that were age, gender, and *APOE* adjusted. In addition, the ADAS-Cog13 model was adjusted for education. Boxes contain the coefficients and, in parenthesis, the *P*-values. The last column (column 4) presents the associations of the metabolites with progression from MCI to AD in Cox hazards models that included age, gender, education, and *APOE* as covariates. Values represent hazard ratio and, in parenthesis, the *P*-values. Significant associations are bolded for an easier visualization. All *P*-values were not multiple comparison corrected.