Potential of Urinary Metabolites for Diagnosing Multiple Sclerosis

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Supporting Information

Abstract: A definitive diagnostic test for multiple sclerosis (MS) does not exist; instead physicians use a combination of medical history, magnetic resonance imaging, and cerebrospinal fluid analysis (CSF). Significant effort has been employed to identify biomarkers from CSF to facilitate MS diagnosis; however, none of the proposed biomarkers have been successful to date. Urine is a proven source of metabolite biomarkers and has the potential to be a rapid, noninvasive, inexpensive, and efficient diagnostic tool for various human diseases. Nevertheless, urinary metabolites have not been extensively explored as a source of biomarkers for MS. We demonstrate that urinary metabolites have significant promise for monitoring disease progression, and response to treatment in MS patients. NMR analysis of urine permitted the identification of metabolites that differentiate experimental autoimmune encephalomyelitis (EAE)-mice (prototypic disease model for MS) from healthy and MS drug-treated EAE mice.

Multiple sclerosis (MS) is a demyelinating disease of the central nervous system (CNS) characterized by selective loss of myelin sheath encapsulating the neuronal axons.1 Treatments of MS are more effective during the early course of the disease when symptoms are mild.2 Thus, the early diagnosis of MS is critical in order to quickly initiate treatments that slow the progression of the disease and improve the quality of a patient’s life.3 Unfortunately, MS is a very challenging disease to properly diagnose,4 where a misdiagnosis is a common occurrence that inevitably leads to a delay in the correct treatment.5 Significant effort has been employed to identify biomarkers from cerebrospinal fluid (CSF) to facilitate MS diagnosis, but this endeavor has proven to be challenging and has not been successful (see supplementary discussion).6 The analysis of urine samples for MS biomarkers has only been minimally investigated but holds significant promise.7 The pathology of MS is associated with widespread demyelination, glial scarring, and an inflammatory response that will inevitably result in a systematic change in the excreted metabolome. Thus, urinary metabolites may be a valuable approach for diagnosing MS and evaluating the in vivo efficacy of MS drug candidates.8 MS is believed to be an immune-mediated (autoimmune) disease requiring the mediation of T cells and/or B cells. Thus, experimental autoimmune encephalomyelitis (EAE) can be induced by immunizing mice with myelin antigens or their peptide fragments emulsified in complete Freund’s adjuvant (CFA).9 Because of similarities with respect to genetic susceptibility, environmental triggers, disease pathology, and clinical signs and disease course, the EAE model has also been used extensively in MS drug discovery research and as a disease model for MS.10 In fact, the majority of drugs being tested now in the phase II and phase III clinical trials were first examined in EAE.11 This includes fingolimod (Gilenya, Novartis), a first-in-class orally administered drug approved for MS therapy.12,13 Fingolimod is more efficacious than other MS treatments.14,15 Importantly, the mechanism of action for fingolimod has been shown to be similar in both humans and the EAE mouse model.16 Specifically, fingolimod suppresses the disease-inducing abilities of myelin-reactive T cells by multiple mechanisms.17,18 Thus, the analysis of changes in urinary metabolites in mice resulting from fingolimod treatment is likely to translate to similar human studies.

Our studies involve the use of 6-to 8-week-old female C57BL/6 mice, which develop a chronic progressive form of paralysis when immunized with the myelin oligodendrocyte glycoprotein (MOG) 35–55 (EAE mice).19 To monitor a response to drug treatment, a group of EAE mice were treated with fingolimod. Establishment of EAE and the efficacy of fingolimod treatment were confirmed by clinical scoring for paralysis and histological evaluation of brains and spinal cords (Figure 1a and Supplementary Table 1). The experimental design consisted of seven treatment groups (n = 13) that include healthy, saline, CFA, EAE, saline plus fingolimod (saline-treated), CFA plus fingolimod (CFA-treated), and EAE.
plus fingolimod (EAE-treated). Fingolimod (1 mg/kg body weight) completely prevented clinical EAE, except for one mouse with EAE score of 2 (mild EAE). Expectedly, MOG-specific T cells in EAE mice treated with fingolimod expanded comparably with those of untreated mice, but they did not induce the disease (Supplementary Figure 1a).

Individual urine samples were collected daily from all the mice belonging to the seven treatment groups, where sample collection began on day 7 and continued until day 30 postimmunization. Ninety one-dimensional (1D) $^1$H NMR spectra were acquired for the urine samples collected on day 17, when the EAE severity reached a peak (Figure 1a).

Representative NMR spectra are shown in Figure 1b, where a visual comparison shows a clear difference between EAE and healthy mice and, importantly, a similarity in the pattern between healthy and EAE-treated mice. A set of NMR peaks (labeled A) were significantly decreased in the spectra for EAE mice relative to healthy mice. Conversely, two sets of NMR peaks (labeled B) were increased in the EAE mice relative to healthy mice. These spectral differences are potential biomarkers for MS. The NMR spectra for the other control groups were essentially identical to the NMR spectra for the healthy mice and EAE-treated mice (Supplementary Figure 2) suggesting that evidence of inflammatory changes in the CNS can be captured by analyzing urinary metabolites.

Principal component analysis (PCA) of the 1D $^1$H NMR spectra further defines a difference in the urine profiles between healthy and EAE mice (Figure 1c–f). The two-dimensional (2D) PCA scores plot generated from 86 of the 1D $^1$H NMR spectra (4 spectra were rejected during the analysis) exhibited two distinct clusters. The rejected spectra fell significantly outside the 95% confidence limit for the EAE mice cluster and the PCA model (not shown) and were randomly distributed throughout the scores plot. The ellipses that correspond to the 95% confidence limits from a normal distribution for each cluster clearly define two statistically distinct classes (Figure 1).

**Figure 1.** NMR metabolomics of EAE mice. (a) Clinical scores of EAE and EAE-treated mice. Groups of mice were immunized with MOG 35–55 in CFA, and the animals were treated with or without fingolimod daily (1 mg/kg body weight) from day 7 postimmunization through day 30. The animals were monitored for EAE signs and the disease-severity was scored. (b) Examples of 1D $^1$H NMR spectra of urine samples collected from EAE, healthy, and EAE-treated mice. Major spectral differences are highlighted. (c) 2D PCA and (d) 2D OPLS-DA scores plot generated from the 1D $^1$H NMR spectra acquired for the urine samples from the seven treatment groups, namely, healthy (dark blue +), saline (black ▲), CFA (brown (*)), EAE (orange ■), saline-treated (light blue □), CFA-treated (green ○), and EAE-treated mice (purple ●). The OPLS-DA used one predictive component and one orthogonal component to yield a $R^2$X of 0.690, $R^2$Y of 0.938, and $Q^2$ of 0.841. The CV-ANOVA validation of the OPLS-DA class distinctions yielded a $p$ value of $1.6 \times 10^{-31}$. The ellipses correspond to the 95% confidence limits from a normal distribution for each cluster. (e,f) Metabolomics tree diagrams determined from the PCA and OPLS-DA scores plots, respectively. The coloring scheme for each group in the tree diagram correlates with the data point colors in the scores plot. The $p$ value for each node is indicated on the tree diagram.
Correspondingly, this result indicates that the urinary metabolites for EAE mice are distinct from the healthy mice. Importantly, the EAE-treated mice cluster together with the healthy mice. This shift in the urinary metabolite profile provides a proof-of-concept for two key events: (i) fingolimod was effective in suppressing the disease severity in EAE mice, and (ii) the therapeutic efficacy of fingolimod was expected to change the metabolite pattern of EAE group toward the healthy group. This is consistent with the clinical scoring and histological evaluation observed for the EAE-treated mice (Figure 1a and Supplementary Table 1).

Orthogonal partial least-squares discriminant analysis (OPLS-DA) of the 1D 1H NMR metabolomics data was performed to further substantiate the observed difference in urinary metabolites between EAE, healthy, and EAE-treated mice (Figure 1d). Strikingly, the OPLS-DA 2D scores plot also contains two statistically distinct clusters. Importantly, a CV-ANOVA test validated the OPLS-DA model with a resulting $p$ value of $1.6 \times 10^{-31.20}$ This provides statistical verification that the urinary metabolites from EAE mice are distinct from both healthy and EAE-treated mice and that there is a corresponding similarity between healthy and EAE-treated mice. Further validation of the statistical significance of the clustering pattern in the PCA and OPLS-DA 2D scores plot was determined by generating metabolomics tree diagrams (Figures 1e,f) using our PCAtoTree program, where we recently replaced bootstrap values with a Mahalanobis metric ($p$ values).$^{21,22}$ Again, the high $p$ values (most $>0.1$) indicate that only the nodes that separate the EAE mice from the healthy and the EAE-treated mice ($p$ value $<5.0 \times 10^{-17}$) are statistically significant. PCA and OPLS-DA scores plot generated from the NMR analysis of urine samples collected on days 23 and 30 are presented in the Supporting Information (Supplementary Figures 3 and 4) and yielded similar results. It is also important to note that the healthy and EAE-treated mice also clustered together and were indistinguishable from the four other negative controls, further establishing that cluster separation is a result of EAE.

The group distinction based on the observed changes in urinary metabolites is disease related as opposed to other environmental factors. The C57Bl/6 mice are inbred animals, were obtained simultaneously, were randomly distributed between each group, and were maintained under identical conditions besides the described group-specific treatments. The only other variable between the groups was the hydration and nutrient supplement (DietGel) provided to EAE and EAE-treated mice with a clinical score of 3 or higher. This is based on accepted protocols for the ethical treatment of animals. Importantly, both EAE and EAE-treated mice received the DietGel supplement, but separation was still observed in the PCA and OPLS-DA scores plot. Critically, the EAE mice treated with fingolimod still clustered with the healthy mice and other control mice, which did not receive the DietGel supplement (Figure 1c,d).

A diet control study comparing healthy mice with or without access to a DietGel supplement resulted in urinary metabolomes distinct from EAE mice (Supplementary Figure 5). Mice received the DietGel supplements for 5 to 7 days, similar to the EAE mice in the EAE induction and treatment study shown in Figures 1 and 2 (days 12 to 17). Clearly, the DietGel supplement is not the source of the observed changes in the

![Figure 2](https://example.com/fig2.png)

**Figure 2.** Potential metabolite biomarkers for EAE. (a) S-plot generated from the OPLS-DA model presented in Figure 1d. Each NMR bin with a covariance of greater than 0.10 or less than −0.10 were identified as major contributors to class separation. These bins are highlighted as red triangles and are labeled with the assigned metabolite. (b) The normalized average peak intensities or bin integrals identified in the S-plot from panel a as major contributors to class separation are plotted as a function of treatment class and metabolite assignment. The mean standard deviation in the normalized average peak intensities is indicated on the bar graphs.
EAE mice urine. A corresponding 2D PCA scores plot comparing urine samples collected over three days from healthy mice with and without the DietGel supplement resulted in a larger group variation for mice receiving the DietGel supplement (Supplementary Figure 5b). In fact, the distribution appears bimodal, potentially distinguishing between individual animals and their preference for the DietGel supplement over food-pellets. Correspondingly, the mice that only received the Teklad food pellets fall within the ellipse defining the 95% confidence interval for the mice receiving the DietGel supplement. Presumably, the pellet-only fed mice overlap with the mice that prefer the Teklad food pellets over the DietGel supplement. Variations were also observed between the healthy mice samples collected from day 5 to day 7 and from those of day 17 (Supplementary Figure 5a). Nevertheless, the 2D PCA scores plot (Supplementary Figure 5c) once again yields a clear separation between healthy and EAE mice. Furthermore, the scores plot indicates that the healthy mice with or without access to the DietGel supplement are more similar to each other than to the EAE mice. This clearly indicates that metabolite changes due to EAE are more pronounced than perturbations from dietary variations. This bodes well for being able to detect similar urinary biomarkers for MS despite an expected wider diversity in the diets of human patients.

The class distinction observed in the PCA and OPLS-DA scores plot suggest the presence of a set of metabolites that could be used as potential biomarkers to differentiate EAE mice from healthy mice. The S-plot generated from the OPLS-DA model identifies the major spectral features that contribute to the class separation observed in the scores plot, where the corresponding metabolite assignments are labeled in Figure 2a. To further validate the statistical relevance of these metabolites to differentiate between healthy and EAE mice, normalized average peak intensities were calculated for each bin and then compared between the seven treatment groups using a standard Student’s t test (Figure 2b). These normalized average peak intensities are proportional to metabolite concentrations. Statistically significant changes were observed in the relative metabolite concentrations between healthy and EAE mice. Specifically, hippurate (p value = 1.3 × 10^{-5}) and fructose (p value = 2.1 × 10^{-5}) were up-regulated in EAE mice compared to healthy mice. Conversely, citrate (p value = 1.3 × 10^{-6}), oxoglutaric acid (p value = 7.0 × 10^{-7}), taurine (p value = 4.9 × 10^{-7}), and urea (p value = 5.8 × 10^{-8}) were down-regulated in EAE mice compared to healthy mice. In addition, the effect of fingolimod treatment was also assessed by a similar comparison between the EAE mice with and without fingolimod treatment. Hippurate (p value = 2.7 × 10^{-5}) and fructose (p value = 2.2 × 10^{-5}) were up-regulated in EAE mice compared to EAE-treated mice. Conversely, citrate (p value = 1.2 × 10^{-5}), oxoglutaric acid (p value = 2.4 × 10^{-5}), taurine (p value = 2.5 × 10^{-5}), and urea (p value = 2.4 × 10^{-5}) were down-regulated in EAE mice compared to EAE-treated mice (Figure 2b).

Two-dimensional 1H−13C HSQC NMR spectra were also acquired to provide a further, in-depth analysis of EAE-induced metabolite changes in urine samples (Figure 3a). The improved resolution and correlated 1H and 13C chemical shifts increase the accuracy and the number of metabolite assignments, but the analysis of the low naturally abundant 13C-labeled metabolites (only 1.1%) is a significant challenge that required large urine volumes (∼500 µL) to obtain acceptable spectral signal-to-noise. Unfortunately, obtaining this much urine from completely paralyzed EAE mice is not practically feasible. Thus, it was necessary to pool an equal volume of urine samples from a group of 4 to 5 mice of identical clinical scores within a cage. Pooling samples does have an advantage since it will minimize within-group variations and maximize between-

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**Figure 3.** Detailed analysis of EAE-dependent urine metabolites. (a) An overlay of 2D 1H−13C HSQC NMR spectra acquired from healthy (black) and EAE (red) mice urine samples. Metabolite assignments are indicated on the spectrum. (b) Heat map generated from the normalized relative intensity of the corresponding metabolite peak obtained from the 2D 1H−13C HSQC NMR spectrum. The dendrogram represents a hierarchical clustering of the metabolites according to relative concentration changes between healthy, EAE, and EAE-treated mice. Metabolites significantly altered in the urine from EAE, but not in EAE-treated mice, which are in italics (p < 0.05 *, p < 0.01 ***, p < 0.001****).

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**Table 1.** Comparison of normalized peak intensities of significant metabolites between healthy and EAE mice. The Student’s t test (Figure 2b) was used to determine the statistical significance of the changes. Metabolites that are significantly different (p < 0.05) are indicated in bold.

<table>
<thead>
<tr>
<th>Metabolite</th>
<th>Healthy (Mean ± SD)</th>
<th>EAE (Mean ± SD)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hippurate</td>
<td>1.2 ± 0.3</td>
<td>10.6 ± 2.1</td>
<td>1.3E-5</td>
</tr>
<tr>
<td>Oxoglutaric acid</td>
<td>4.9 ± 0.7</td>
<td>0.7 ± 0.2</td>
<td>7.0E-7</td>
</tr>
<tr>
<td>Citric acid</td>
<td>1.5 ± 0.2</td>
<td>0.2 ± 0.1</td>
<td>4.9E-7</td>
</tr>
<tr>
<td>Taurine</td>
<td>2.8 ± 0.4</td>
<td>0.5 ± 0.2</td>
<td>5.8E-8</td>
</tr>
<tr>
<td>Urea</td>
<td>6.0 ± 1.2</td>
<td>0.6 ± 0.1</td>
<td>5.0E-9</td>
</tr>
<tr>
<td>Fructose</td>
<td>2.0 ± 0.3</td>
<td>0.3 ± 0.1</td>
<td>2.7E-5</td>
</tr>
<tr>
<td>Acetate</td>
<td>1.4 ± 0.2</td>
<td>0.2 ± 0.1</td>
<td>2.2E-5</td>
</tr>
<tr>
<td>Taurine sulfate</td>
<td>1.3 ± 0.2</td>
<td>0.3 ± 0.1</td>
<td>1.2E-5</td>
</tr>
</tbody>
</table>

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**Figure 2.** (a) An overlay of 2D 1H−13C HSQC NMR spectra acquired from healthy (black) and EAE (red) mice urine samples. Metabolite assignments are indicated on the spectrum. (b) Heat map generated from the normalized relative intensity of the corresponding metabolite peak obtained from the 2D 1H−13C HSQC NMR spectrum. The dendrogram represents a hierarchical clustering of the metabolites according to relative concentration changes between healthy, EAE, and EAE-treated mice. Metabolites significantly altered in the urine from EAE, but not in EAE-treated mice, which are in italics (p < 0.05 *, p < 0.01 ***, p < 0.001****).
group differences by effectively averaging NMR peak intensities (metabolite concentrations) within a group.

Urine samples were collected from triplicate sets of EAE, EAE-treated, and healthy mice and then used to acquire the corresponding 2D $^1$H−$^{13}$C HSQC NMR spectra. Normalized differences in average peak intensities relative to healthy mice are summarized in the heat-map depicted in Figure 3b. Over 32 metabolites were identified with different concentrations between EAE and healthy mice, which also contained the metabolites identified from the OPLS-DA S-plot analysis. Interestingly, some of the identified metabolites were previously implicated in the pathogenesis of MS and EAE (histamine and glutamate)$^{23,24}$ or identified as markers of metabolic diseases with associated neurological problems ($3$-hydroxyisobutyric acid, $3$-ureidopropionate, and guanidinoacetate).$^{25−27}$ Also, one of the identified metabolites, indoxyl sulfate, is likely to originate from the gut microbiota.$^{28}$ This is consistent with the previous observation that EAE induction and progression is influenced by gut microorganisms as opposed to attenuation of EAE in germ-free mice.$^{29}$ Other potential metabolites identified in our NMR analysis of urine samples have also been associated with EAE, MS, and other neurological diseases (see Supplementary Discussion).

MS is accompanied by various pathological features like inflammation, demyelination, and axonal damage. This inherent complexity of MS makes it challenging to identify a single biomarker to monitor the progression and treatment of the disease. Alternatively, our NMR analysis of urine samples demonstrates that a diverse set of metabolites can be used to differentiate between healthy and EAE mice. These metabolite changes can also be used to monitor the recovery of EAE mice upon treatment with fingolimod. Also, the 1D $^1$H NMR analysis of urine samples takes only about 10 min per sample and completely lacks any of the risks or side effects associated with the analysis of CSF. Thus, the NMR analysis of urine holds the promise of being an easy, fast, and safe diagnostic tool for MS. Additionally, the NMR analysis of urine may be a valuable approach for evaluating the in vivo efficacy of drug leads and for designing patient-specific treatments.

**METHODS**

**Peptide Synthesis.** MOG 35−55 (MEGVGYRSPFSRVVHLNYRNGK) was synthesized on 9-fluorenylmethoxycarbonyl chemistry (Neopeptide, Cambridge, MA) to a purity of more than 90% as verified by HPLC and mass spectroscopy. The peptide was dissolved in 1x phosphate buffered saline and stored at −20 °C until used. The MOG 35−55 peptide was used to induce EAE in mice.

**Mice.** Ninety 6 to 8 week-old female C57Bl/6 (H-2b) mice were obtained from the Jackson Laboratory (Bar Harbor, ME). C57Bl/6 (H-2b) mice are inbred animals and individual variations in the gut flora are not expected. Furthermore, the animals are maintained in the same environment and received the same food. All the mice were located in the same room within the University of Nebraska—Lincoln (UNL) Life Sciences Annex, managed and maintained by the Institutional Animal Care Program in accordance with the animal protocol guidelines of UNL. The mice were provided with the Teklad global 16% protein rodent diet (Harlan Laboratories, Indianapolis, IN). The mice were randomly classified into 21 individually ventilated cages. Fifteen of the cages had four animals, while the other six cages had five animals. The mice were acclimatized for three days before the start of the experiment.

During the experiment, all the mice were given *ad libitum* access to food and water. Correspondingly, the amount of food and water consumed by an individual mouse was not regulated or quantified. Also, the mice were not fasted during urine collection or at any other time during the experiment. In this manner, the metabolomics experiment captures the natural variation in nutrient consumption per animal and provides a more realistic metabolomic background. Additionally, if a cage containing either an EAE group or EAE group under fingolimod treatment with a mouse exhibiting a clinical score of 3 or above, the entire cage was supplemented with a nutrient fortified hydration gel (moisture, 73%; DietGel 76A, ClearH2O, Portland, ME). The DietGel combines hydration and standard nutrients (76A maintenance diet formulation)$^{30}$ in a palatable form for compromised rodents. Importantly, all the mice in the cage still have access to the Teklad global 16% protein rodent diet in addition to the nutrient fortified hydration gel. The EAE and EAE-treated mice received supplemental DietGel on day 12. The EAE mice received the supplemental DietGel for the remainder of the experiment; whereas, the EAE-treated mice stopped receiving the supplemental DietGel from day 15. All of the EAE mice treated with fingolimod had adequately recovered by day 15.

**EAE Induction and Treatment.** The experimental design consisted of a healthy group (n = 12) and six treatment groups (n = 13). These include saline, CFA, EAE, saline treated, CFA-treated, and EAE-treated. Each group of mice was divided into three cages, respectively, containing 4, 4, and 5 (or 4 for healthy) animals. The cages for each experimental group were selected randomly. To induce EAE, peptide emulsions were prepared by mixing MOG 35−55 in CFA containing *Mycobacterium tuberculosis* H37RA extract (Difco Laboratories, Detroit, MI) to a final concentration of 5 mg mL$^{-1}$. Each animal received 200 μg of peptide-emulsion subcutaneously in the inguinal and sternal regions. In addition, pertussis toxin (List Biological Laboratories, Campbell, CA) was administered (200 ng/mouse) intraperitoneally on day 0 and day 2 postimmunization.$^{31−33}$ Seven days postimmunization, fingolimod dissolved in 1x sterile normal saline (working dilution of 0.2 mg mL$^{-1}$), was administered intraperitoneally to the animals corresponding to treated groups as indicated above at 1 mg/kg body weight daily until day 30.

**Urine Collection and Clinical Scoring.** Urine samples were collected both prior to and after disease induction. Urine collections occurred three times daily (10−11 a.m.; 2−3 p.m.; and 10−11 p.m.) from each animal by expressing the bladder. The samples collected from each batch of animals were pooled on a daily basis and preserved at −80 °C until further analysis. In addition, the samples collected from individual animals on days 17, 23, and 30 postimmunization were also preserved as separate aliquots. The immunized mice were monitored for clinical signs of EAE and scored as described previously:$^{31,34}$ 0, no signs of disease; 1, limp tail or hind limb weakness; 2, limp tail and hind limb weakness; 3, partial paralysis of hind limbs; 4, complete paralysis of hind limbs; and 5, moribund or dead.

**NMR Sample Preparation.** The samples for the 1D $^1$H NMR experiments were prepared by adding 600 μL of a 50 mM phosphate buffer in 99.8% D$_2$O (Isotec, St. Louis, MO) at pH 7.2 (uncorrected) to 25 μL of urine collected from each mouse. The six treatment groups contained 13 mice per group and the healthy group contained 12 mice for a total of 90 NMR samples. The samples for the 2D $^1$H−$^{13}$C HSQC experiments were prepared by adding 100 μL of a 50 mM phosphate buffer in 99.8% D$_2$O at pH 7.2 (uncorrected) and 500 μL of urine pooled from a group of 4 to 5 healthy or EAE mice. Each pooled group of mice was assigned the same clinical assessment score, and the animals were placed in the same cage. The urine was collected at the same time on a daily basis. A triplicate set of NMR samples were prepared for each group for the 2D $^1$H−$^{13}$C HSQC experiments.

**NMR Data Collection and Analysis.** All NMR experiments were performed with Bruker AVANCE DRX 500 MHz spectrometer equipped with 5 mm triple-resonance cryogenic probe ($^1$H, $^13$C, and $^{15}$N) with a Z-axis gradient. A BACS-120 sample changer with Bruker ICON-NMR software was used to automate the NMR data collection. The 1D $^1$H NMR data was collected at 298 K with 32 K data points, a spectrum width of 5483 Hz, 128 scans, and 16 dummy scans using an excitation sculpting pulse sequence to remove the solvent peak.$^{35}$ The 2D $^1$H−$^{13}$C HSQC NMR spectrum was collected at 298 K with 512 scans, 32 dummy scans, and a 1.5 s relaxation delay. The spectrum was
collected with 2 K data points and a spectrum width of 4734 Hz in the direct dimension and 64 data points and a spectrum width of 18 864 Hz in the indirect dimension. ACD/1D NMR manager version 12.0 (Advanced Chemistry Development, Inc.) was used to process the 1D 1H NMR spectra. Intelligent binning was used to integrate each region with a bucket size of 0.025 ppm. The buckets were normalized by the intensity of the 3-(trimethylsilyl)propionic acid-2,2,3,3-d4 (TMSP) peak. Each NMR spectrum was mean centered and autoscaled by the standard deviation as described in Zhang et al.36

NMRPipe37 was used to process the 2D 1H−13C HSQC spectra. Peak-picking and peak-matching were accomplished using NMRViewJ Version 8.0.38 Peak-intensities were normalized for each 2D 1H−13C HSQC NMR spectrum by dividing each peak-intensity by the average peak-intensity for a given spectrum. Each NMR peak for each metabolite from the triplicate set of 2D1H spectrum was assigned with SIMCA P+ 12 (UMETRICS). The tree diagram and the dendrograms were generated using a new updated version of our RR015468-01).

ACKNOWLEDGMENTS

The authors declare no competing financial interest.

REFERENCES


