Metabolomic differences between critically ill women and men

Sowmya Chary1, Karin Amrein2, Jessica A. Lasky-Su3, Harald Dobnig4 & Kenneth B. Christopher3,5*

Metabolism differs in women and men at homeostasis. Critically ill patients have profound dysregulation of homeostasis and metabolism. It is not clear if the metabolic response to critical illness differs in women compared to men. Such sex-specific differences in illness response would have consequences for personalized medicine. Our aim was to determine the sex-specific metabolomic response to early critical illness. We performed a post-hoc metabolomics study of the VITdAL-ICU trial where subjects received high dose vitamin D3 or placebo. Using mixed-effects modeling, we studied sex-specific changes in metabolites over time adjusted for age, Simplified Acute Physiology Score II, admission diagnosis, day 0 25-hydroxyvitamin D level, and 25-hydroxyvitamin D response to intervention. In women, multiple members of the sphingomyelin and lysophospholipid metabolite classes had significantly positive Bonferroni corrected associations over time compared to men. Further, multiple representatives of the acylcarnitine, androgenic steroid, bile acid, nucleotide and amino acid metabolite classes had significantly negative Bonferroni corrected associations over time compared to men. Gaussian graphical model analyses revealed sex-specific functional modules. Our findings show that robust and coordinated sex-specific metabolite differences exist early in critical illness.

Though inclusiveness of women subjects in clinical research was mandated by the National Institutes of Health (NIH) in 1993, most clinical research studies do not account for sex-specific differences1–3. The research that does exist shows that robust differences exist between women and men with respect to disease incidence, disease severity, metabolism and pharmacodynamics of interventions4,5. Although firm evidence exists for improved outcomes for female animals in experimental models of severe illness, such differences are not consistently observed in studies on critically ill patients6–9. Mechanistic understanding of sex-specific differences in the response to illness is essential if we are to progress to personalized medicine10.

Existing data show that metabolism differences are present in healthy women relative to men. At homeostasis, women incorporate free fatty acids into triglycerides whereas men oxidize circulating free fatty acids11. Circulating acylcarnitines which are reflective of energy metabolism, are generally lower in women12. Women also have less free fatty acid-induced insulin resistance13. Healthy women have increases in circulating lipid sphingomyelins which act in cell signaling and may reflect glucose metabolism14–17. Sex-specific differences in lipid and cholesterol metabolism are well established and likely due to sex chromosome and sex-specific hormone action18. The overall sex-specific metabolism differences at homeostasis are probably due to variation in metabolism related gene expression which contributes to sexual dimorphism12,19.

Metabolomics provides a window into the large number of circulating substrates and products of patient’s cellular metabolism20. A few large metabolomics studies on healthy individuals are notable for robust metabolite differences related to sex12,19,21–23. Data from healthy subjects has little relevance to critically ill patients where metabolic homeostasis is profoundly disturbed24. Heterogenous critical illness is not defined by a precise phenotypic framework and studies have provided limited mechanistic insights into pathophysiology25. Metabolomic studies performed early in critical illness can reflect illness severity and predict outcomes. But such work does not address sex-specific differences in the response to critical illness26–28. Therefore, to see whether sex-specific
metabolite modules exist that associate with sex. Further, with the metabolite change data we determined if regulated metabolism differences apply to the critically ill, we studied differences between women and men with regards to changes in metabolism during critical illness.

To test the hypothesis that significant sex-specific plasma metabolomic profile differences exist in the response to critical illness, we performed a metabolomics analysis of 1215 plasma samples from 428 subjects collected during the VITdAL-ICU trial. The VITdAL-ICU trial randomized 492 critically ill adults (166 of whom were women) with 25-hydroxyvitamin D \([25(OH)D]\) levels ≤ 20 ng/ml to high dose oral vitamin D3 or placebo. The VITdAL-ICU trial did not find significant differences in length of hospital stay or mortality outcomes. We assessed the effect of sex on changes in individual metabolites and plasma metabolite families over three time points early in the course of critical illness. Further, with the metabolite change data we determined if regulated metabolite modules exist that associate with sex.

Results
In the 428 subject analytic cohort, 35% of subjects were women. Baseline characteristics were balanced between subjects stratified by sex for C-reactive protein, Simplified Acute Physiology Score (SAPS) II, day 0 25(OH)D levels, intervention status and ICU type. Differences existed by sex with respect to age (see Table 1 and Supplementary Table S1). The overall 28-day mortality of the 428 subject analytic cohort was 22.2%. The 28-day mortality in women was 22.5% and in men was 22.0%.

### Single time point data.
In day 0 plasma samples (N = 428), significant differences exist in 12 individual metabolites (all multiple test-corrected threshold of \(P\)-value < 8.65 × 10\(^{-5}\), \(-\log_{10}(P) > 4.06\)) and in metabolomic profiles (CV-ANOVA \(P\)-value < 0.001) in female subjects relative to males (see Supplementary Table S2).

| Characteristic                   | Female | Male  | Total | \(P\)-value |
|----------------------------------|--------|-------|-------|-------------|
| No                               | 151    | 277   | 428   |             |
| Age years Mean (SD)              | 68.2 (13.3) | 62.0 (15.3) | 64.2 (14.9) | <0.001*     |
| Day 0 25(OH)D ng/ml Mean (SD)    | 13.2 (5.7) | 14.4 (10.1) | 13.9 (8.8) | 0.17*       |
| SAPS II Mean (SD)                | 34.6 (14.7) | 32.7 (15.8) | 33.4 (15.4) | 0.24*       |
| Day 0 C-reactive protein µg/mL Mean (SD) | 119.9 (96.4) | 127.6 (86.0) | 124.9 (89.8) | 0.40*       |
| Day 0 Procalcitonin ng/ml Median [IQR] | 0.45 [0.14, 1.98] | 0.77 [0.20, 3.02] | 0.66 [0.17, 2.79] | <0.001*     |
| Vitamin D\(_3\), Intervention No. (%) | 78 (51.7) | 134 (48.4) | 212 (49.5) | 0.52        |
| Change in 25(OH)D ng/ml Mean (SD) | 11.3 (18.0) | 10.0 (15.5) | 10.4 (16.4) | 0.43*       |
| ICU                              |        |       |       | 0.22        |

|                     | Female | Male  | Total | \(P\)-value |
|---------------------|--------|-------|-------|-------------|
| Anesthesia ICU No. (%) | 24 (15.9) | 59 (21.3) | 83 (19.4) |             |
| Cardiac Surgery ICU No. (%) | 42 (27.8) | 84 (30.3) | 126 (29.4) |             |
| Surgical ICU No. (%)   | 7 (4.6) | 16 (5.8) | 23 (5.4)  |             |
| Medicine ICU No. (%)   | 31 (20.5) | 59 (21.3) | 90 (21.0)  |             |
| Neurological ICU No. (%) | 47 (31.1) | 59 (21.3) | 106 (24.8) |             |

Table 1. Cohort characteristics. Data presented as No. (%) unless otherwise indicated. \(P\)-values determined by chi-square unless designated by (\(\dagger\)) then \(P\)-value determined by ANOVA or by (\(\ddagger\)) determined by Kruskal-Wallis test.

In linear regression models of metabolite data from single time points (day 0, 3 or 7), we find significant differences exist in 51 individual metabolites at 1 or more time point (all multiple test-corrected threshold of \(P\)-value < 8.65 × 10\(^{-5}\), \(-\log_{10}(P) > 4.06\)) in female subjects relative to males (see Supplementary Table S3).

In the repeated measures data, mixed-effects modeling of 1215 total day 0, 3 and 7 plasma samples from the analytic cohort (N = 432) shows 50 metabolites had significantly positive associations in women relative to men highlighted by increases in individual sphingomyelin species and decreased androgenic steroids in women relative to men (see Supplementary Table S3).

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Multiple time point data. In the repeated measures data, mixed-effects modeling of 1215 total day 0, 3 and 7 plasma samples from the analytic cohort (N = 432) shows 50 metabolites had significantly positive associations in women relative to men highlighted by increases in individual sphingomyelin species and decreased androgenic steroids in women relative to men. Decreases in androgenic steroids as well as bile acid and amino acid metabolism are found in women when compared to men. Decreases in androgenic steroids with a negative permutation \(Q^2\) intercept indicating model validity (see Supplementary Table S2)30,31. Day 0 differences are present with increased individual sphingomyelin species and decreased androgenic steroids in women relative to men (see Supplementary Table S3).

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acylcarnitine, androgenic steroid, bile acid, nucleotide and amino acid metabolite sub-pathways and individual sphingomyelin species that significantly increase or decrease in women relative to men over days 0, 3 and 7 (see Fig. 3).

Next, we explored the sex-specific associations of individual metabolites and 28-day mortality. We compared mixed-effects modeling of a total of 441 day 0, 3 and 7 plasma samples from 151 women in the analytic cohort to mixed-effects modeling of a total of 814 day 0, 3 and 7 plasma samples from 277 men in the analytic cohort. The data show that an increase in short chain acylcarnitines C4–C8 and branched-chain amino acids significantly associate with three fold higher 28-day mortality in women but not men (see Supplementary Table S6, Supplementary Fig. S1).

Metabolic networks and mediation. We investigated sex-specific metabolic networks by measuring pairwise correlations in metabolites which have similar effects via Gaussian graphical models (GGMs). The GGMs analysis revealed seven sex-specific functional modules at day 3 and seven at day 7 (see Supplementary Tables S7 & S8). Similar to the mixed-effects analyses, metabolism of branched chain amino acids, bile acids, androgenic steroids and lysophospholipids are prominently featured in the sex-specific GGM modules. Metabolites within in each functional module were either increased or decreased in women in unison and had biological or functional similarity. Of note, the sex-specific modules do include some individual metabolites that were not significantly associated with sex in our mixed-effects analysis (see Supplementary Tables S7 & S8: Modules B, E, H, I, K, M).

Finally, we focused on the potential mediation of the relationship between individual metabolite abundance and sex by inflammation status. Mediation analyses in day 3 data revealed no influence of Procalcitonin or of
C-reactive protein on the associations between sex and each of the individual 578 metabolites (all $P$-values were > 0.01 using 2000 bootstrap samples).

**Discussion**

Although previous work suggests that some sex-specific differences exist in the healthy at homeostasis, our data argue that distinct nuanced alterations in metabolism are present in women and men during critical illness. In our single time point data and our mixed-effects methods analysis, we consistently find robust increases and decreases in groups of metabolites along similar sub-pathways that have conserved function (see Fig. 3, Supplementary Tables S4 & S5). Further, we illustrate how groups of metabolites with similar sex-specific effects form GGM modules which highlight the same sub-pathways as our single time point data and our mixed-effects methods analysis. These modules serve to focus potential biological interpretation of our sex-specific metabolomics observations. All three analyses highlight the importance of sex-specific metabolism in critical illness related to branched chain amino acids, bile acids, androgenic steroids and lysophospholipids.

Critically ill patients preferentially catabolize fatty acids and amino acids for mitochondrial energy production. Sex-specific differences exist in the utilization of carbohydrates and lipids as energy. With increased energy needs during cell stress, women preferentially oxidize lipids over carbohydrates while men utilize carbohydrates. Firm evidence exists that female mitochondria have higher oxidative capacity, produce less reactive oxygen species, and preferentially utilize lipids for bioenergetics. Elevated circulating even-chain C4–C22 acylcarnitines are shown to be due to incomplete mitochondrial β-oxidation of fatty acids. Studies find that increases in circulating acylcarnitines are common in severe critical illness and are associated with adverse outcomes. To see whether these findings apply to critically ill women, we analyzed a total of 36 acylcarnitine species. We demonstrate that 15 circulating acylcarnitine species are significantly lower in women early in critical illness.
Our data are consistent with a more efficient fatty acid β-oxidation in critically ill women reflective of a sex-specific difference in mitochondrial response to critical illness. The circulating amino acid pool is supplied by dietary amino acids, endogenous amino acid synthesis and cellular protein turnover. Increases in circulating amino acids during critical illness are due to protein catabolism. Skeletal muscle protein is rapidly metabolized in response to severity of illness to provide substrate for liver gluconeogenesis, immune function support and immunoglobulin synthesis. Further, amino acid catabolism is a source for circulating C3, C4 and C5 acylcarnitines. Our findings of decreases in C3, C4 and C5 acylcarnitines as well as in multiple amino acid metabolite sub-pathways suggest sex-specific protein catabolism and energy substrate utilization during critical illness. Of particular interest, are the GGM modules B and H which highlight the importance of decrease in branched chain amino acid metabolites in women.

In women, we observe a combination of decreases in branch chain amino acid metabolites and in dicarboxylate fatty acids generated from fatty acid omega oxidation as well as in short-chain acylcarnitines derived from branch chain amino acids. Such decreases strongly suggest improved mitochondrial function and more complete fatty acid β-oxidation in women relative to men.

Our novel observations suggest that critically ill women have greater abundance of individual sphingomyelin species, plasmalogens and lysophospholipids compared to men. Studies show that during cell stress, specific sphingomyelin species regulate the initiation of apoptosis and autophagy. Plasmalogens are known to be endogenous antioxidants that protect the endothelium from oxidative stress injury by controlling toxic oxidation products. Lysophospholipids are signaling molecules that have chemocattractant effects and act to modulate the innate immune response. Our data are consistent with the hypothesis that the response to cell stress differs in women and men.
The liver is the essential organ for glucose, protein, amino acid, lipid and cholesterol metabolism. Sex-specific differences in liver metabolism at homeostasis are postulated to be an evolutionary consequence of the metabolic flexibility required for reproduction. Our data argue that bile acid metabolites are decreased in critically ill women. Though sex-specific differences in bile acid synthesis are reported, such differences in bile acid homeostasis are not well characterized. It is shown that cytochrome P450 enzymes are important for bile synthesis and regulated in a sex-specific manner. Bile acids activate the nuclear receptors farnesoid X receptor, pregnane X receptor and vitamin D receptor as well as the G-protein-coupled receptor TGR5. Such bile acid receptor activation results in gene expression which alters metabolism of bile acids, glucose, lipids, energy and inflammation. As elevation in blood bile acids are common in critical illness and the synthesis and pool composition of bile acids are sex-specific, such differences have widespread downstream metabolism pathway effects.

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Our novel study approach has several strengths. The use of a large number of plasma samples at multiple time points early in critical illness allows for a dynamic overview into sex-specific metabolomics (see Fig. 3). Mixed models are extremely useful for metabolomic data measured at multiple time points as they remove confounding variables with a fixed-effect (age, SAPS II, etc.) and also those with a random-effect (plasma sampling day)58,59. Importantly, by adjusting for the absolute change in 25(OH)D level at day 3, we mitigate the effect of the trial intervention on the observed sex-specific metabolomic changes which allow for study of the entire trial cohort increasing sample size and study power60,61. Further, our use of clinical trial data allows for modelling and normalization of metabolite abundance via adjustment for subject characteristics62. To account for multiple comparisons we utilized a conservative Bonferroni corrected $P\text{-value} < 8.65 \times 10^{-5}$63. Finally, some of the metabolism differences we observe are known to be sex-specific thus increasing the biological plausibility and relevance of our work.

### Table 3. Metabolites significantly decreased in Women relative to Men over time. Significant results presented following mixed-effects modeling of each of the 578 individual metabolites measured at day 0, 3 and 7. All estimates adjusted for age, SAPS II, admission diagnosis, 25(OH)D at day 0, absolute change in 25(OH)D level at day 3 and plasma day (as the random-intercept). A multiple test-corrected threshold of $P\text{-value} < 8.65 \times 10^{-5}$ was used to identify all significant associations. Negative $\beta$ coefficient values indicate lower abundance in females relative to males.

| Metabolite                        | $P\text{-value}$ | Bonferroni corrected $P\text{-value}$ | $-\log_{10}P$ | $\beta$ coefficient | Super pathway          | Sub pathway                        |
|-----------------------------------|------------------|---------------------------------------|----------------|---------------------|------------------------|-----------------------------------|
| N-acetylvaline                    | 2.55 E−05        | 1.47 E−02                              | 4.59           | −0.10               | Amino acid              | Leucine, isoleucine and valine metabolism |
| N-acetylleucine                   | 7.65 E−06        | 4.42 E−03                              | 5.12           | −0.12               | Amino acid              | Leucine, isoleucine and valine metabolism |
| Alpha-hydroxyisocaproate          | 9.27 E−07        | 5.36 E−04                              | 6.03           | −0.13               | Amino acid              | Leucine, isoleucine and valine metabolism |
| 3-hydroxyisobutyrate              | 1.37 E−06        | 7.90 E−04                              | 5.86           | −0.13               | amino acid              | Leucine, isoleucine and valine metabolism |
| 3-hydroxy-2-ethylpropionate       | 2.06 E−09        | 1.19 E−06                              | 8.69           | −0.15               | Amino acid              | Leucine, isoleucine and valine metabolism |
| 2-hydroxy-3-methylvalerate        | 5.00 E−08        | 2.89 E−05                              | 7.30           | −0.16               | Amino acid              | Leucine, isoleucine and valine metabolism |
| 2,3-dihydroxy-2-methylbutyrate    | 4.77 E−07        | 2.76 E−04                              | 6.32           | −0.17               | Amino acid              | Leucine, isoleucine and valine metabolism |
| Beta-hydroxyisovalerate           | 4.04 E−12        | 2.33 E−09                              | 11.39          | −0.19               | Amino acid              | Leucine, isoleucine and valine metabolism |
| Glycochenodeoxycholate            | 1.59 E−06        | 9.18 E−04                              | 5.80           | −0.20               | Lipid                  | Primary bile acid metabolism       |
| Taurocholate                      | 4.19 E−05        | 2.42 E−02                              | 4.38           | −0.21               | Lipid                  | Primary bile acid metabolism       |
| Glycochenodeoxycholate glucuronide| 2.94 E−12        | 1.70 E−09                              | 11.53          | −0.28               | Lipid                  | Primary bile acid metabolism       |
| Taurochenodeoxycholate            | 2.20 E−08        | 1.27 E−05                              | 7.66           | −0.28               | Lipid                  | Primary bile acid metabolism       |
| Glycochenodeoxycholate sulfate    | 4.26 E−11        | 2.46 E−08                              | 10.37          | −0.32               | Lipid                  | Primary bile acid metabolism       |
| Glycodeoxycholate sulfate         | 4.01 E−06        | 2.32 E−03                              | 5.40           | −0.23               | Lipid                  | Secondary bile acid metabolism     |
| Glycolithocholate sulfate         | 5.82 E−09        | 3.36 E−06                              | 8.24           | −0.27               | Lipid                  | Secondary bile acid metabolism     |
| Glycocholenate sulfate            | 6.22 E−14        | 3.59 E−11                              | 13.21          | −0.28               | Lipid                  | Secondary bile acid metabolism     |
| Taurolithocholate 3-sulfate       | 3.11 E−12        | 1.80 E−09                              | 11.51          | −0.34               | Lipid                  | Secondary bile acid metabolism     |
| Taurocholenate sulfate            | 1.03 E−16        | 5.94 E−14                              | 15.99          | −0.38               | Lipid                  | Secondary bile acid metabolism     |
We do acknowledge potential limitations to our approach. Our VITdAL-ICU trial subject population is heterogeneous with sex-specific imbalance in some admission diagnosis categories. Despite multivariable adjustment, our approach is subject to bias and confounding. Though our samples are derived from a randomized controlled trial, our study design is observational thus causal inference may be limited. Our subjects were all white with serum levels of 25(OH)D < 20 ng/ml, thus may not be representative of all critically ill. Our use of CRP as an indicator of inflammation is limited in the nine reproductive-aged women under study as CRP is associated with Progesterone and Estradiol levels. The single-center setting may limit generalizability of our findings. It is important to recognize that although the function and biological relevance of a metabolite may be characterized, the clinical significance may not be known. Finally, our study is a hypothesis generating exploratory analysis requiring subsequent confirmation and careful interpretation.

The importance of our study is that it offers a nuanced window into the differential metabolic response to critical illness between women and men. Beyond the known sex-dependent metabolism differences at homeostasis, we find that women respond to critical illness stressors in a dramatically different fashion than men. Our findings on sex-specific differences in metabolism pathways is an essential first step toward understanding how to provide patient-centered personalized medicine in the critically ill.

Methods
The details of the VITdAL-ICU trial as well as metabolomic processing and analysis are provided in Supplementary Methods. Briefly, the VITdAL-ICU trial randomized 475 critically ill adult patients to vitamin D3 or placebo once at a dose of 540,000 IU followed by 90,000 IU monthly. At VITdAL-ICU trial enrollment, written informed consent was obtained and included permission for plasma specimens to be saved for future research studies. The metabolomics study is considered post-hoc as it was designed following initiation completion of the VITdAL-ICU trial. The post-hoc study research protocol was approved by the Partners Human Research Committee Institutional Review Board at the Brigham and Women's Hospital. All research was performed in accordance with the Declaration of Helsinki.
To generate metabolomic data, a total of 1215 plasma samples from 428 VTidAL-ICU trial subjects at day 0, 413 subjects at day 3 and 374 subjects at day 7 were analyzed using four ultra high-performance liquid chromatography/tandem accurate mass spectrometry methods by Metabolon, Inc.\textsuperscript{65}. Metabolomic profiling identified 769 metabolites (Supplementary Data 2). We reduced baseline noise by removing metabolites with the lowest interquartile range of variability, leaving 578 metabolites\textsuperscript{66}. Metabolomic data underwent cube root transformation and Pareto scaling to normalize the distribution\textsuperscript{67}.

For univariate analysis of day 0 data, Student’s t-test was performed to determine if significant sex-specific differences exist using MetaboAnalyst\textsuperscript{48}. A Bonferroni multiple testing correction threshold of $P$-value $< 8.65 \times 10^{-5}$ was used to identify all significant differences\textsuperscript{63}. Day 0 data were also analyzed using orthogonal partial least square-discriminant analysis (OPLS-DA), a supervised method to assess the significance of classification discrimination (SIMCA 15.0 Umetrics, Umeå, Sweden). Permutation testing was performed for OPLS-DA model validation\textsuperscript{30,31}. Sevenfold cross-validation analysis of variance (CV-ANOVA) was utilized to determine OPLS-DA model significance\textsuperscript{31}.

For single time point data, correlations between individual metabolites and sex at day 0, 3 or 7 were separately determined utilizing linear regression models correcting for age, SAPS II, admission diagnosis, 25(OH)D at day 0 and absolute change in 25(OH)D level at day 3. A multiple test-corrected threshold of $P$-value $< 8.65 \times 10^{-5}$ was used to identify all significant associations in the single time point data\textsuperscript{68}. All linear regression models were analyzed using STATA 14.1MP\textsuperscript{69}. Rain plots were produced based on hierarchical clustering in R-3.6.2 adapted from source code published by Henglin et al.\textsuperscript{70}.

For repeated measures data, correlations between individual metabolites and sex over time (day 0, 3 and 7) were determined utilizing linear mixed-effects models correcting for age, SAPS II, admission diagnosis, 25(OH)D at day 0, absolute change in 25(OH)D level at day 3 and plasma day (as the random-intercept). This analysis was performed in the analytic-cohort ($N = 428$) with multiple test-corrected threshold of $P$-value $< 8.65 \times 10^{-5}$ was used to identify all significant associations. We repeated the analysis in only those subjects who received placebo ($N = 216$) with Benjamini–Hochberg adjustment of $P$-values\textsuperscript{31}. All mixed-effects models were analyzed using STATA 14.1MP\textsuperscript{69}. For data visualization purposes, a bipartite graph\textsuperscript{34} utilizing the Circos application (http://circos.ca/) in Perl was generated of metabolites which were significantly changed (increased or decreased) in females relative to males.

Mixed effects logistic regression was used separately in 151 women and in 277 men to estimate the odds of 28-day mortality of individual metabolites adjusted for age, SAPS II, admission diagnosis, 25(OH)D at day 0, absolute change in 25(OH)D level at day 3 and plasma day (as the random-intercept). A multiple test-corrected threshold of $P$-value $< 8.65 \times 10^{-5}$ was used to identify all significant associations in the repeated measures data\textsuperscript{68}. All mixed-effects models were analyzed using STATA 14.1MP\textsuperscript{69}. We used rain plots\textsuperscript{32} to separately visualize the mortality-dependent effect size and significance of individual metabolites in women and men.

As inflammation is important in response to critical illness, we evaluated a potential mediating effect of Procalcitonin or C-reactive protein on the association between sex and individual metabolite abundance adjusted for age, SAPS II, admission diagnosis, 25(OH)D at day 0, absolute change in 25(OH)D level at day 3 and plasma day (as the random-intercept). Analyses were performed on each of the 578 metabolites at day 3 using the R package mediation\textsuperscript{70} to obtain bootstrap $P$-values ($N = 2000$ samples)\textsuperscript{71,72}. Significant mediation was present if the $P$-value was $< 0.01$ and 10% or more of the association was mediated through Procalcitonin or C-reactive protein levels\textsuperscript{71,72}.

To identify sex-specific modules from metabolomics data, we estimated Gaussian graphical models (GGMs) for day 3 and 7. Modules serve to reconstruct pathway reactions from metabolomics data. GGMs are determined utilizing partial pairwise Pearson correlation coefficients following the removal of the effects of all other metabolites and covariates\textsuperscript{73}. We inferred a sex-specific network for relative metabolite abundance. We included age, SAPS II, admission diagnosis, 25(OH)D at day 0, absolute change in 25(OH)D level at day 3 and plasma day as covariates into the model\textsuperscript{74}. Edges between metabolites were allotted if both their Pearson correlations and partial correlations remained statistically significant at $P$-value $< 0.05$ following Bonferroni correction for 578 metabolites\textsuperscript{74}. GGMs were produced using the GeneNet R package, version 1.2.13 in R-3.6.2 adapted from published source code\textsuperscript{74}.

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Author contributions
S.C.: Conceptualization, Methodology, Formal analysis, Data Curation, Writing—Original Draft, Writing—Review & Editing, Visualization. K.A.: Conceptualization, Methodology, Investigation, Resources, Data Curation, Writing—Original Draft, Writing—Review & Editing, Funding acquisition. J.L.-S.: Conceptualization, Methodology, Software, Formal analysis, Investigation, Resources, Data Curation, Writing—Original Draft, Writing—Review & Editing, Visualization, Supervision, Project administration. H.D.: Conceptualization, Methodology, Investigation, Resources, Data Curation, Writing—Review & Editing, Visualization, Supervision, Project administration, Funding acquisition. K.C.: Conceptualization, Methodology, Software, Formal analysis, Investigation, Resources, Data Curation, Writing—Original Draft, Writing—Review & Editing, Visualization, Supervision, Project administration, Funding acquisition.

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Competing interests
Dr. Chary was a Masters student at Harvard Medical School during the implementation of the study. During her Masters, Dr. Chary was employed at Takeda Pharmaceutical Company and is currently employed by Biogen, Inc. Dr. Chary reports receiving salary and stock options from Takeda and from Biogen. Neither Takeda nor Biogen had any involvement in the VITdAL-ICU trial, the identification of metabolites, the post hoc study design, analysis of metabolomics, interpretation of the data, access to the data or writing of the manuscript. Dr. Lasky-Su served as a consultant and received personal fees from Metabolon, Inc., outside the submitted work. Dr. Amrein reports receiving lecture fees from Fresenius Kabi. Dr. Dobnig reports receiving lecture fees from Fresenius Kabi. Dr. Christopher reports no financial or other relationships that might lead to a conflict of interest.

Additional information
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