### Soil Microbial Functional Succession Over One

 $\begin{array}{c} 001 \\ 002 \\ 003 \\ 004 \\ 005 \\ 006 \\ 007 \end{array}$ 

 $008 \\ 009 \\ 010$ 

 $016 \\ 017 \\ 018$ 

 $020 \\ 021$ 

 $023 \\ 024 \\ 025 \\ 026$ 

 $042 \\ 043$ 

### Year of Human Decomposition

Allison R. Mason<sup>1</sup>, Lois S. Taylor<sup>2</sup>, Naomi Gilbert<sup>1</sup>, Steven W. Wilhelm<sup>1</sup>, Jennifer M. DeBruyn<sup>1,2\*</sup>

<sup>1</sup>Department of Microbiology, University of Tennessee-Knoxville, 1311 Cumberland Avenue, Knoxville, 37996.

<sup>2</sup>Department of Biosystems Engineering and Soil Science, University of Tennessee-Knoxville, 2506 E.J. Chapman Drive, Knoxville, 37996.

\*Corresponding author(s). E-mail(s): jdebruyn@utk.edu;

#### Abstract

Background The succession of microbial communities during vertebrate decomposition has been observed in various settings, documenting changes in taxa as decomposition progresses. These studies have predominantly employed phylogenetic markers (*i.e.*, rRNA genes), describing community composition and structure, but ultimately are not informative of which members are active or metabolic pathways they might be expressing. This has left a foundational knowledge gap regarding the functional roles of microorganisms in vertebrate decomposition, which ultimately impact ecosystem functioning. Here we present the first known study investigating gene expression in soil impacted by human decomposition in a terrestrial ecosystem. Total RNA was extracted and metatranscriptomes obtained from soil samples collected over the course of one year from below three decomposing human bodies.

**Results** Microbial gene expression profiles shifted in response to decomposition: decomposition impacted soils were most different from controls (*i.e.* nearby soils unimpacted by decomposition) at day 86, and profiles remained altered even

after one year. Shifts in gene expression were partially explained by environmental and soil physiochemical variables, including internal body accumulated degree hours (p=0.001), as well as soil temperature (p=0.045), pH (p=0.042) and electrical conductivity (p=0.037). Differential expression analysis revealed that microbes in decomposition soils displayed increased expression of stress response genes (mean fold change 3.48), particularly heat shock proteins (p<0.001), whose expression increased between days 0 and 58 and remained elevated through day 376. Further, we identified genes whose expression was altered at certain timepoints. This included increased expression of genes encoding hydroxylamine oxidoreductase (HAO) (85x), nitric oxide reductase (83x), and nitrous oxide reductase (19x) at day 86 when dissolved oxygen was ~85%, suggesting that microbial communities may be converting hydroxylamine to nitric oxide and reducing nitrous oxide to nitrogen gas during reduced oxygen conditions.

Conclusions Our results show that human decomposition alters soil microbial gene expression profiles providing evidence of altered microbial metabolisms (e.g., taurine metabolism, nitrogen cycling, and lipid metabolism) and reveal the potential of vertebrate decomposition to have both ephemeral and lasting effects on ecosystem processing in response to mortality events.

**Keywords:** Human Decomposition, Microbial Succession, Metatranscriptomics, Soil Microbial Ecology

#### Introduction

 $\begin{array}{c} 064 \\ 065 \end{array}$ 

 $073 \\ 074 \\ 075$ 

 $077 \\ 078$ 

 $\begin{array}{c} 079 \\ 080 \end{array}$ 

 $\begin{array}{c} 082 \\ 083 \end{array}$ 

 $084\\085$ 

 $087 \\ 088$ 

Soil microbial communities are important drivers of ecosystem processes in terrestrial environments. Many soil microbes are decomposers that are involved in degradation of complex organic matter and drive nutrient cycling in terrestrial ecosystems. Environmental disturbances can impact the presence and/or activity of soil microorganisms that are invovled in these cycles, ultimately affecting nutrient availability and the release of greenhouse gas emissions, such as  $CO_2$  and  $N_2O$  [1, 2]. Vertebrate death and subsequent carcass deposition in terrestrial ecosystems is one disturbance resulting in the deposition of large quantities of organic C and N [3–10], along with other elements (P, K, S, etc) [11], which collectively contribute to microbially-mediated

biogeochemical cycling. In additon to this, changes in pH, temperature, and fluctuations in soil oxygen provide abiotic filtering further impacting microbial metabolic strategies [7–9, 11, 12].

 $093 \\ 094$ 

 $\begin{array}{c} 095 \\ 096 \end{array}$ 

 $097 \\ 098$ 

 $099 \\ 100$ 

 $101 \\ 102$ 

103

 $104 \\ 105$ 

 $\begin{array}{c} 106 \\ 107 \end{array}$ 

108

 $109 \\ 110$ 

 $\begin{array}{c} 111 \\ 112 \end{array}$ 

113

 $114\\115$ 

 $\begin{array}{c} 116 \\ 117 \end{array}$ 

118

119 120

121 122

123

124 125 126

127 128

129

 $130 \\ 131$ 

132

133 134

While C and N transformations have been documented during decomposition, the functional response of microbes and their roles in nutrient cycles remain unclear. The composition and structure of decomposition-impacted soil microbial communities have been investigated using amplicon sequencing of marker genes (i.e., 16S rRNA, 18S rRNA, ITS), revealing successional dynamics [13]. This has allowed us to investigate changes in microbial biodiversity and composition in response to vertebrate decomposition, revealing patterns such as increases in the anaerobic taxa Firmicutes and Bacteroidetes. However, few studies have investigated soil biogeochemistry and microbial communities within the same study, which can further help to describe microbial ecology in human and animal decomposition systems. Taylor et al. (2024) [14] suggested that fungal community shifts were linked to changes in soil dissolved oxygen, highlighting interactions between soil microbes and the surrounding environment. While insightful for making potential connections between taxa and physiochemistry, these analyses cannot inform which taxa are active members of the community responsible for chemical transformations, which functional pathways/genes are expressed, and how these pathways are altered in response to decomposition.

Methods such as RNA sequencing (*i.e.*, metatranscriptomics) and metabolomics can be used to investigate microbial community functional succession in response to decomposition by measuring gene expression and metabolites, respectively. This can inform how ecological functions, including C and N cycling, are impacted by decomposition events in terrestrial ecosystems. To date, only two studies have applied metatranscriptomic approaches to assess mRNA in vertebrate decomposition samples [15, 16]:

Burcham et al. (2019) [15] examined gene expression of internal organ microbial communities during mouse decomposition, while Ashe et al. (2021) [16] examined gene expression of oral microbial communities during human decomposition. Both studies suggest that the host microbial community functionality is altered during decomposition, including differential expression of amino acid and carbohydrate metabolism in the heart [15] and shifts in gene transcripts across different taxa [16]. We expect that soil microbial community gene expression profiles are also altered; however, this has never been examined to our knowledge. The decomposition-impacted soil metabolome was assessed by DeBruyn et al. (2021) [17], showing changes in soil metabolites over time including increased prevalence of amino acids, however it is unclear which microbes are responsible for these shifts. Additionally, DeBruyn et al. (2021) [17] showed the soil metabolome was still altered compared to starting conditions at the end of the 21-week study, suggesting long-term impacts of decomposition on soil microbial functioning.

142

 $146 \\ 147$ 

 $148 \\ 149$ 

 $151 \\ 152$ 

 $156 \\ 157$ 

168

 $170 \\ 171$ 

 $175 \\ 176$ 

 $177 \\ 178$ 

The purpose of this study was to investigate soil microbial gene expression during a one-year period of human decomposition and address the following questions: (1) which genes are differentially expressed in soils impacted by human decomposition? (2) how does gene expression change over time in decomposition-impacted soils? (3) do microbial gene expression profiles return to pre-decomposition conditions after one year? The human body is comprised of nutrient-rich organic molecules, many of which are broken down during decomposition. We hypothesized that gene expression would change over time as resources are used and transformed and soil chemical and physical conditions change due to the influx of decomposition products [8, 9, 17]. For example, we expected to observe changes in the expression of genes encoding enzymes involved in nitrogen cycling, as changes in nitrogen pools have been previously described in decomposition soils [8]. Of the main macromolecules in the body (carbohydrates, proteins, lipids, and nucleic acids), we were particularly interested in

lipid metabolism, as we expect lipids from the body to enter the soil during decomposition and previous studies showed an increase of lipolytic organisms in decomposition soils [12, 18]. Finally, multiple studies have shown than soil chemistry [5, 8] and microbial community composition [19, 20] (via 16S rRNA gene amplicon sequencing) are still impacted after one year, therefore we did not expect soil expression profiles to return to pre-decomposition conditions.

 $185 \\ 186$ 

 $187 \\ 188$ 

189

 $190 \\ 191$ 

192 193

 $194 \\ 195$ 

196 197

 $198 \\ 199$ 

200

 $\begin{array}{c} 201 \\ 202 \end{array}$ 

 $\begin{array}{c} 203 \\ 204 \end{array}$ 

205

 $\begin{array}{c} 206 \\ 207 \end{array}$ 

 $\frac{208}{209}$ 

 $210 \\ 211 \\ 212$ 

 $\begin{array}{c} 213 \\ 214 \end{array}$ 

 $215 \\ 216 \\ 217$ 

218

 $\frac{219}{220}$ 

 $\frac{221}{222}$ 

223

 $\begin{array}{c} 224 \\ 225 \end{array}$ 

 $\begin{array}{c} 226 \\ 227 \end{array}$ 

228 229 230

To answer these questions, metatranscriptomes of soil samples collected at six key timepoints over one year of human decomposition were used to determine the active populations and expression of genes and pathways relevant to the enhanced biogeochemical cycling observed in decomposition hotspots. We compared gene expression between decomposition timepoints and control soils that were unexposed to decomposition products to identify functions or functional pathways of interest. This assessment of functional profiles within decomposition-impacted soils provided insight into the microbial response to vertebrate decomposition in terrestrial settings and biogeochemical cycling within these hotspots.

#### Results

#### Soil Physiochemistry

Soil chemistry was altered in response to human decomposition, with multiple parameters still impacted after one year [14]. Generally, soil pH decreased and remained low in decomposition soils of all but one individual. Soil electrical conductivity (EC) increased in response to decomposition, remaining elevated through approximately day 58 before gradually decreasing throughout the remainder of the study (Supplementary Material 1). Respiration (evolved  ${\rm CO_2}$ ) increased by an order of magnitude beginning at day 12, which corresponded to a reduction in soil dissolved oxygen (DO)

to 29% - 48.9%. Ammonium concentrations increased 78-fold, reaching maximum concentrations between days 12 and 58. This was followed by decreased ammonium and increased nitrate concentrations at day 86, with nitrate concentrations reaching a maximum at day 168 (Supplementary Material 1).

#### Sequencing

231

232

 $\begin{array}{c} 233 \\ 234 \end{array}$ 

 $\begin{array}{c} 235 \\ 236 \end{array}$ 

237 238

239 240 241

242

 $243 \\ 244$ 

 $\begin{array}{c} 245 \\ 246 \end{array}$ 

247

 $248 \\ 249$ 

 $\begin{array}{c} 250 \\ 251 \end{array}$ 

252

 $\begin{array}{c} 253 \\ 254 \end{array}$ 

 $\begin{array}{c} 255 \\ 256 \end{array}$ 

257

 $\begin{array}{c} 258 \\ 259 \end{array}$ 

 $\frac{260}{261}$ 

262

 $\begin{array}{c} 263 \\ 264 \end{array}$ 

 $\begin{array}{c} 265 \\ 266 \end{array}$ 

267 268 269

 $\begin{array}{c} 270 \\ 271 \end{array}$ 

272

 $\begin{array}{c} 273 \\ 274 \end{array}$ 

 $275 \\ 276$ 

Illumina sequencing of the 24 libraries yielded a total of 5,073,476,730 reads, or 2,536,738,365 paired reads, with a mean of 105,697,432 paired reads per sample. Removal of adapters and low-quality reads removed 4.7% of all reads, leaving 4,834,123,062 total reads. Filtering of ribosomal RNA further removed 7.3% of reads, leaving 4,479,804,360 reads for assembly. After co-assembly, a total of 6,257,674 proteins were identified by Prodigal. From this, 1,048,573 proteins were annotated by eggNOG-mapper (16.7%). Most of the annotated proteins were taxonomically annotated as bacteria (91.3%), followed by eukaryotes (7.6%), and archaea (0.81%). Of the 7.6% of eukaryotic proteins, 64.4% (4.9% of all proteins) were annotated as fungi. For this study, genes of interest included all bacterial, archaeal, and fungal proteins, therefore all non-fungal eukaryotic proteins (32,004) were removed prior to downstream analysis. The reference file of genes was then used to determine gene transcript counts in all samples using CLC genomic workbench. The percent of reads mapped to genes of interest ranged from 21% to 38%, with an average of 31% reads mapped. Gene counts were then combined in a single file and used for downstream analyses in R.

#### Microbial gene expression in response to human decomposition

Gene expression profiles in decomposition-impacted soils shifted away from controls and day zero samples as decomposition progressed (Fig 1A). Expression was most different from controls on study days 58, 86, 168 (Supplementary Material 2), before

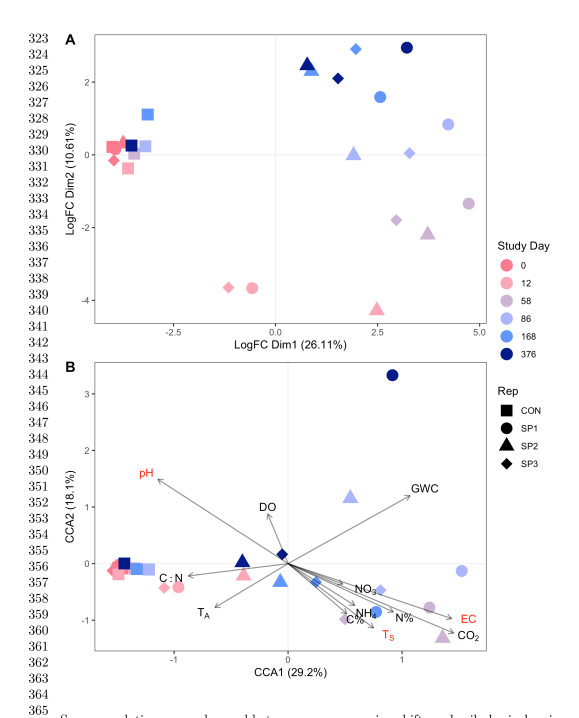
shifting back toward control conditions on study day 376. After one year of decomposition, soil gene expression profiles had not returned to pre-decomposition conditions, as evidenced by their clustering away from controls and day zero samples in the MDS plot (Fig 1A).

 $277 \\ 278$ 

 $\begin{array}{c} 279 \\ 280 \end{array}$ 

283 284

Figure 1: Microbial gene expression profiles are altered during human decomposition. Multidimensional scaling (MDS) shows gene expression within soils changed as decomposition progressed (A). Additionally, canonical correspondence analysis (CCA) shows that environmental variables explained 47.3% of the variation in gene expression profiles (B). Variables in bold red type significantly (p < 0.05) explained some of the variation in gene expression profiles as assessed by Permutational Analysis of Variance (PERMANOVA). In both panels soils from controls (CON) and the three donors (SP1, SP2, SP3) are denoted by symbol shape, while color represents study day. In B, soil physiochemical variable loadings are reporesented by arrows: Gravimetric water content (GWC), electrical conductivity (EC), pH (pH), dissolved oxygen (DO), respiration (evolved CO<sub>2</sub> µmol gdw<sup>-1</sup>), ammonium (NH<sub>4</sub>), and nitrate (NO<sub>3</sub>) concentrations (mg gdw<sup>-1</sup>), percent carbon (%C), percent nitrogen (%N), carbon:nitrogen ratio (C:N), ambient temperature ( $T_A$ ), and soil temperature ( $T_B$ ).



Some correlations were observed between gene expression shifts and soil physiochemical data at decomposition timepoints. Canonical correspondence analysis (CCA) was

 $\begin{array}{c} 366 \\ 367 \end{array}$ 

used to constrain gene expression data with soil physiochemical data (Fig 1B). CCA1 and CCA2 explained 29.2% and 18.1% of the variance in gene expression, respectively. Transcript profiles at day 12 were associated with an increase in soil carbon to nitrogen ratio (C:N). Gene expression profiles at days 58 to 86 were positively correlated with increased soil temperature, EC, and evolved  $\rm CO_2$ , while study day 168 was associated with elevated levels of soil  $\rm NO_3$ . Further, Permutational Analysis of Variance (PERMANOVA) revealed that internal accumulated degree hours (ADH), soil temperature, pH, and EC significantly explained some of the variation in gene expression profiles (p < 0.05). No other soil chemical variables were significant at  $\alpha = 0.05$  (Supplementary Material 3).

 $\frac{369}{370}$ 

 $\frac{371}{372}$ 

373

 $\frac{374}{375}$ 

 $\frac{376}{377}$ 

378

 $\frac{379}{380}$ 

 $\frac{381}{382}$ 

383

 $384 \\ 385 \\ 386$ 

 $\frac{387}{388}$ 

389

 $\frac{390}{391}$ 

 $\frac{392}{393}$ 

394

 $\frac{395}{396}$ 

397 398

399

 $400 \\ 401$ 

402 403

404

 $405 \\ 406$ 

407 408

409

410 411

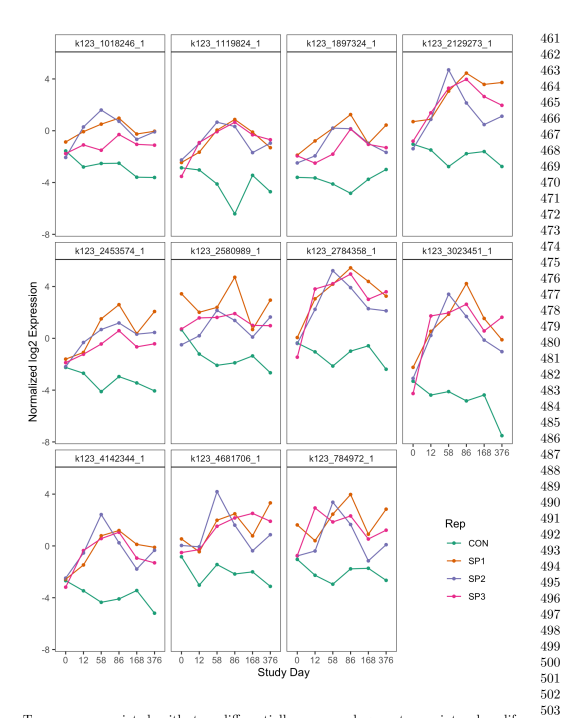
412 413

414

Overall, decomposition changed soil gene expression profiles over the one-year study relative to control soils. Differential expression analysis between decomposition and control soils identified 7,047 down-regulated and 38,425 up-regulated genes. Gene transcripts that were associated with control soils belonged to a wide variety of clusters of orthologous genes (COG) functional categories. Specifically, the top 20 genes whose expression was higher in control soils belonged to ten unique COG categories, including signal transduction mechanisms, transcription, and those of unknown function. In contrast, the top 20 genes whose expression was higher in decomposition soils only fell into four COG categories (Supplementary Material 4 A): 1) post-translational modification, protein turnover, and chaperones; 2) energy production and conversion; 3) cell motility; and 4) carbohydrate transport and metabolism. The most common COG category represented in decomposition soils (80% of the top 20 genes) was posttranslational modification, protein turnover, and chaperones. Within this category, several heat shock stress response genes were identified, including SSA2, HSP82, and clpB (Supplementary Material 5). Further investigation into these genes shows their expression increased in response to decomposition, typically reaching maximum transcript levels around study days 58 and 86 (Fig 2). This corresponded to elevated soil

415 temperatures below decomposing bodies between study days 12-80, with soil temperatures increasing to approximately 43°C [14], and maximum soil EC and minimum dissolved oxygen measurements between days 12 and 58 (Supplementary Material 1).

Figure 2: Normalized log2 expression of heat shock proteins identified by differential expression analysis comparing decomposition and control soils. Each panel represents a single heat shock transcript, labled with query ID. Symbol color denotes if the sample is a control (CON, green), or one of three individuals: SP1 (orange), SP2 (purple), or SP3 (pink).



Taxonomy associated with top differentially expressed gene transcripts also differed between control and decomposition soils. The top 40 significantly differentially

507 expressed gene transcripts in decomposition soils were associated with Fungi, Acti-508 nobacteria, and Xanthomonadales, while gene transcripts in controls were associated 510 with Acidobacteria, Cyanobacteria, Proteobacteria ( $\alpha$ ,  $\delta$ ,  $\gamma$ ), and Planctomycetes (Sup-512 plementary Material 4 B). The greatest number of differentially expressed genes 513 relative to control samples was observed at day 86, where we saw 145,460 and 124,883 up- and down-regulated genes, respectively.

# Fate of decomposition products as evidenced in gene expression profiles over time

521 522

 $526 \\ 527$ 

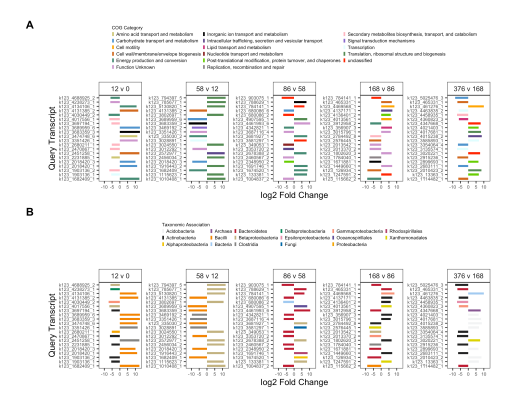
Differential expression analysis between respective sequential study days further revealed which genes were altered between decomposition timepoints. The top ten significantly up- and down-regulated genes, determined by the lowest p-values from differential expression analysis (< 0.05), are reported in Supplementary Material 6 and Fig 3.

530 531

Figure 3: Top twenty up- and down-regulated genes in decomposition soils comparing sequential study days (0, 12, 58, 86, 168, 376) colored by COG functional category (A) and taxonomic annotation (B). Positive values denote increased expression compared to the preceding timepoint, while negative values denote a decrease.

 $542 \\ 543$ 

 $544 \\ 545 \\ 546$ 



 $\begin{array}{c} 585 \\ 586 \end{array}$ 

 $\begin{array}{c} 591 \\ 592 \end{array}$ 

Expression of genes annotated with the COG categories cell wall/membrane/envelope biogenesis, inorganic ion transport and metabolism, and carbohydrate transport and metabolism increased from day 0 to 12. In contrast, expression of secondary metabolite biosynthesis, transport, and catabolism genes decreased during this period (Fig 3A). Transcripts from *Bacilli* and *Clostridia* increased, while transcripts from *Actinobacteria* decreased between study days zero and 12 (Fig 3).

Between days 12 and 58, 90% of the topmost upregulated genes were associated with the translation, ribosomal structure and biogenesis COG and all were taxonomically associated with *Betaproteobacteria* (Fig 3A,B). Many of these genes were annotated as ribosomal protein large (RPL), involved in ribosomal binding. Genes across multiple COG categories with taxonomic associations to *Bacilli* and *Clostridia* decreased

between study days 12 and 58, six of which were transcripts that previously increased between days zero and 12 (Fig 3B, Supplementary Material 6).

Multiple transcripts associated with the energy production and conversion COG, as well as transcripts annotated with the COGs inorganic transport and metabolism, and translation, ribosomal structure and biogenesis, increased between days 58 and 86 (Fig 3A). Two of the upregulated energy and production and conservation transcripts were associated with cytochrome c oxidase subunits in Betaproteobacteria, while another was annotated as hao, encoding the enzyme hydroxylamine dehydrogenase which is involved in conversion of hydroxylamine to nitrite during nitrification (Supplementary Material 6). Further investigation into hydroxylamine dehydrogenase showed a significant increase in hao transcripts at day 86 followed by subsequent decreases at days 168 and 376 (F = 4.183; p = 0.02). This increase corresponded to decreased soil ammonium levels and subsequent accumulation of nitrate (Supplementary Material 1). Half of the topmost downregulated genes between days 58 and 86 were not assigned to a COG (*i.e.*, unclassified) or were of unknown function.

Differential expression comparing study days 86 with 168 and 168 with 376 identified genes across a variety of functional categories, with many unclassified in the COG database or with unknown function (Fig 3A). Expression of carbohydrate transport and metabolism genes associated with *Bacilli* decreased between day 168 and 376. *Acidobacteria* transcripts increased in decomposition-impacted soils between study day 168 and 376, but were not associated with any single COG category (Fig 3B).

#### Carbon compound metabolism

599

600

 $601 \\ 602 \\ 603$ 

604

 $605 \\ 606$ 

 $607 \\ 608$ 

609

610 611

 $612 \\ 613$ 

614

 $615 \\ 616$ 

617 618

619

 $620 \\ 621$ 

 $622 \\ 623$ 

 $624 \\ 625$ 

 $626 \\ 627$ 

 $628 \\ 629$ 

630

 $631 \\ 632$ 

 $633 \\ 634$ 

635 636 637

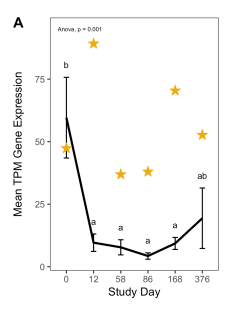
638

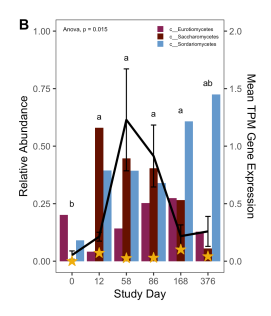
639 640

We expected to observe increased expression of lipid metabolizing genes during active and advanced decomposition as microbes degraded lipids deposited in the soil [18].

Therefore, we investigated changes in triacylglycerol lipase (enzyme commission number: 3.1.1.3) gene transcription in our soils. Generally, lipase transcripts decreased as decomposition progressed (HLM F = 6.564, p < 0.001), however we also observed a significant interaction between study day and taxonomic annotation (F = 8.786; p < 0.001). Specifically, lipase gene transcripts annotated as bacteria decreased with decomposition time (F = 10.392; p = 0.001), while fungal lipase transcripts increased, reaching a maximum at study day 58 (F = 4.509; p = 0.015) (Fig 4).

Figure 4: Mean transcript abundance, in transcripts per million (TPM), of all bacterial (A) and fungal (B) triacylglycerol lipase (EC 3.1.1.3) genes over time. Black lines (A, B) report mean and standard deviation of TPM from three individuals (black line), while gold stars denote mean TPM in control soils. P-values are the result of ANOVAs where average TPM and study day are the dependent and independent variables, respectively, while letters are the result of post-hoc Tukey tests between decomposition timepoints. In B, bars show the relative abundance of the fungal classes *Saccharomycetes*, *Sordariomycetes*, and *Eurotiomycetes*, reported in Taylor et al. (2024).





 $645 \\ 646$ 

 $647 \\ 648$ 

 $650 \\ 651$ 

 $652 \\ 653$ 

 $655 \\ 656 \\ 657$ 

 $679 \\ 680$ 

#### Nitrogen- and sulfur compound transformations

691 692 693

694

695 696

 $697 \\ 698$ 

699

 $700 \\ 701$ 

 $702 \\ 703$ 

704

 $705 \\ 706$ 

 $707 \\ 708$ 

709

710 711

 $712 \\ 713$ 

714

 $715 \\ 716$ 

 $717 \\ 718$ 

719

 $720\\721$ 

 $722 \\ 723$ 

724

 $725 \\ 726$ 

727 728 729

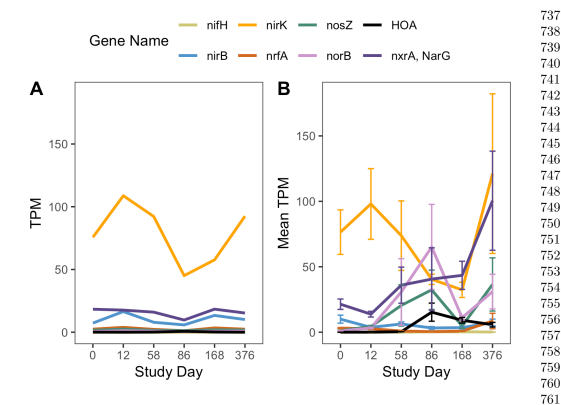
730

731

732

Expression of nitrogen cycling genes was impacted in response to human decomposition. Due to the detection of hao in our differential expression analysis, and our hypotheses predicting changes to nitrogen transformation processes, the expression of genes encoding common enzymes involved in nitrogen cycling (nifH, nirB, nirK, norB, nosZ, nrfA, nxrA, and amoA) were assessed using their enzyme commission numbers (Fig 5A,B). nifH, encoding a subunit of nitrogenase which is involved in nitrogen fixation, displayed little to no changes in gene expression between control and decomposition soils. Transcripts for two genes encoding enzymes contributing to the last two steps of denitrification, norB (encodes nitric oxide reductase) and nosZ (encodes nitrous oxide reductase), increased between study days 12 and 86, and decreased at study day 168 before increasing again at day 376. In contrast, expression of genes encoding nitrate reductase, narG, and NO-forming nitrite reductase, nirK, remained low until day 376 when transcripts for both genes increased. As noted above, expression of hao, encoding hydroxylamine dehydrogenase, increased at study day 86 before decreasing at remaining timepoints (Fig 3A, Fig 5B). Expression of amoA, encoding a subunit of ammonia monooxygenase, and nxrA, encoding a subunit of nitrite oxidoreductase, which are involved in nitrification, changed in response to decomposition. amoA transcripts initially decreased at day 12, remaining reduced until study day 376. Similarly, abundance of genes that encode for enzymes involved in dissimilatory nitrate reduction, nirB, and nrfA, was low for the first 168 days, with nrfA expression increasing at day 376 (Fig 5B).

Figure 5: Mean gene expression, in transcripts per million (TPM), of commonly used marker genes for enzymes involved in nitrogen cycling over time in controls (A) and decomposition (B) soils. Data in B represent mean and standard deviation of TPM from three individuals.



Expression of genes involved in metabolism of nitrogen and sulfur-containing compounds were also impacted by human decomposition. Specifically, four of the top ten genes whose expression decreased at day 12 were related to taurine metabolism, with their annotations associated with tauD, encoding taurine dioxygenase. (Supplementary Material 6). Further investigation into tauD showed that mean expression of these genes decreased steadily over one year, beginning at day 12 (Fig 6B); however, tauD expression in response to human decomposition was variable across taxonomic associations. Most tauD transcripts were associated with Gammaproteobacteria, Actinobacteria, Betaproteobacteria, Alphaproteobacteria, and fungi. While a majority of the tauD gene queries displayed reduced expression over time, expression of fungal-associated and a few Betaproteobacteria-associated tauD genes increased at day 58 (Supplementary Material 7). Sources of taurine in the human body include taurine

764

 $768 \\ 769$ 

 $771 \\ 772$ 

 $773 \\ 774$ 

 $776 \\ 777$ 

 $781 \\ 782$ 

absorbed from the diet and taurine produced from anaerobic microbial deconjugation of bile salts via bile salt hydrolase (BSH) enzymes [21]. Therefore, we also looked at expression of genes encoding BSH enzymes in decomposition soils. Expression of these genes was elevated at days 12, 58, and 86 before converging toward pre-decomposition levels at days 168 and 376 (Fig 6A). Hierarchical liner mixed effects (HLM) models showed that both tauD (HLM F = 7.356, p = 0.002) and BSH (F = 13.768, p < 0.001) gene expression was significantly different over time (Fig 6A,B).

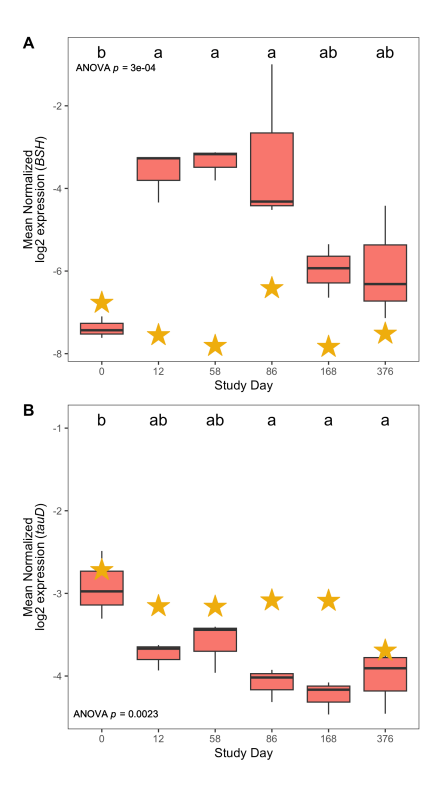
 $785 \\ 786$ 

 $790 \\ 791$ 

 $792 \\ 793$ 

 $794 \\ 795$ 

Figure 6: Mean bile salt hydrolase, BSH, (A) and tauD, taurine dioxygenase, (B) log2 normalized expression in controls (gold stars) and decomposition (boxplots) soils. Boxplots display the 25th and 75th quartiles and median log2 normalized values between all three individuals at each timepoint. ANOVA p-value is the result of a hierarchical linear mixed effects model accounting for repeated measures of each donor block, while letters denote the results of post-hoc Tukey test.



#### Discussion

 $878 \\ 879$ 

 $\begin{array}{c} 881 \\ 882 \end{array}$ 

884

 $\begin{array}{c} 891 \\ 892 \end{array}$ 

 $\begin{array}{c} 903 \\ 904 \end{array}$ 

906

 $910 \\ 911$ 

914

 $\begin{array}{c} 915 \\ 916 \end{array}$ 

The goal of this study was to assess soil microbial gene expression in response to human decomposition. Metatranscriptomics were applied to soil samples collected over one year from below three decomposing human bodies. From this, we found that microbial gene expression shifted over time, with samples reproducible between individuals. Additionally, we showed that gene expression profiles had not recovered to pre-decomposition conditions after one year. Comparison of control and decomposition expression profiles revealed that heat-shock proteins were elevated in response to decomposition. We also described expression patterns between decomposition time-points, noting changes in functional gene categories at certain timepoints, in particular with respect to lipid, nitrogen and sulfur metabolism.

# Decomposition impacted soil community gene expression, even after a year

Gene expression profiles remained altered after one year of decomposition. It is unclear if soil microbial communities, in terms of gene expression profiles, have reached a new steady state as a result of decomposition, or if they would eventually return to pre-decomposition conditions. The soil pH, EC, NH<sub>4</sub><sup>+</sup>, NO<sub>3</sub><sup>-</sup>, and total nitrogen (TN) exhibited differences (although not statistical) in these soils following a year of decomposition, however bacterial and fungal community structures, as assessed by rRNA amplicon libraries, were still altered [14]. This indicates that decomposition can continue to structure microbial communities and impact their function for extended periods of time. While nutrient pools and communities both demonstrate less rapid change at later time points in the study, there is not evidence suggesting an arrival at a steady-state post-disturbance microbial community. In some studies, human decomposition can result in elevated carbon and nutrients (organic nitrogen, ammonium, nitrate, and phosphate) for longer than a year [3], suggesting decomposition events

have long lasting effects on the local ecosystem. Together, this has implications for terrestrial ecosystem processing (e.g., nutrient cycling, emission of greenhouse gasses, etc.), as we show that decomposition alters functional metabolism pathways within soil microbial communities. Further work with extended sample collections beyond one year are needed to address how long microbial communities and their functions are impacted.

 $921 \\ 922$ 

 $923 \\ 924$ 

925

 $926 \\ 927$ 

 $928 \\ 929$ 

 $930 \\ 931$ 

 $932 \\ 933$ 

934 935

936

 $937 \\ 938$ 

 $939 \\ 940$ 

941

 $942 \\ 943$ 

 $944 \\ 945$ 

946

947 948

949 950

951

952 953

954 955

956

957 958

 $959 \\ 960$ 

Bacteria, fungi, and archaea were all represented in expressed genes throughout decomposition, suggesting that members of all three domains have the potential to contribute to decomposition processes and nutrient cycling. While a majority of annotated transcripts were identified as bacteria, fungal transcripts were the second most abundant group. Fungal transcripts made up almost half (seven of the top 15) of the significantly differentially expressed genes associated with decomposition-impacted soils. Additionally, with respect to expression shifts between decomposition timepoints, fungal transcripts were among the topmost upregulated genes at study day 86. The presence of fungal transcripts is not surprising as fungi are key decomposers, involved in the degradation of organic matter in terrestrial ecosystems [22]. It was interesting to see an increase in certain fungal transcripts, such as lipase, at study days 58 and 86 when soil oxygen began to recover. We would expect lipids to enter the soil as tissues are broken down during decomposition, so we were surprised to see bacterial lipase genes decrease during decomposition. This suggests that microbial activity in decomposition soils may be constrained by the changing chemical environment, potentially altered oxygen levels in the case of bacterial lipase gene expresssion. Prior work with these soils showed that soil oxgyen concentration was a key driver of changes in both bacterial and fungal community composition [14].

#### Increased stress responses during decomposition

Soil microbial communities expressed stress response genes in response to human decomposition. Differential expression analysis identified increased expression of multiple heat shock proteins associated with the taxa Xanthomonadales, Actinobacteria, and fungi. Upon further investigation, expression of these genes increased through day 58 and remained high for the remainder of the year. Soil temperature was elevated relative to controls between study days 8 and 80, with maximum temperatures >40°C, while soil electrical conductivity increased up to 663 μS/cm (16X higher than background) through day 58 before slowly decreasing through the remainder of the study. Soil electrical conductivity (correlates with ionic strength [23] and can indicate soil salinity) has previously been shown to increase in decomposition soils [8–10, 14]. As a result, we would expect these microbes to be experiencing both heat and osmotic stress during this period. Prior work has observed increased heat shock gene expression during salt stress in paddy soils [24] and the presence of both heat and osmotic stress genes in desert soils along a salt gradient [25], suggesting saline conditions can alter the expression of heat and/or osmotic stress genes. In our study we observed that stress response within soil microbial communities is stimulated during human decomposition, however, at this time, it is unclear if expression of these genes is in response to heat stress alone, or in combination with osmotic stress.

#### Increased expression of fungal lipase genes during decomposition

Human fat tissue contains lipids that are broken down during decomposition. There-fore, we assessed expression of triacylglycerol lipase genes in decomposition soils. Our 1008 results show that expression of triacylglycerol lipase genes was altered in response to decomposition, and these shifts differed between bacterial and fungal transcripts, specifically bacterial triacylglycerol lipase transcripts decreased in response to decomposition, while fungal triacylglycerol lipase transcripts increased. Further, expression of these genes corresponded to changes in relative abundance of the fungal classes Saccharomycetes, Sordariomycetes, and Eurotiomycetes [14]. These fungi have been previously associated with decomposition soils [26, 27] and are known to contain triacylglycerol lipase genes in their genomes [28, 29], suggesting that they play a role in lipid degradation in decomposition soils.

 $1013 \\ 1014$ 

 $\begin{array}{c} 1015 \\ 1016 \end{array}$ 

1017

1018 1019

 $1020 \\ 1021$ 

 $1022 \\ 1023$ 

 $\begin{array}{c} 1024 \\ 1025 \end{array}$ 

 $1026 \\ 1027$ 

1028

1029 1030

 $1031 \\ 1032$ 

1033

 $1034 \\ 1035$ 

 $1036 \\ 1037$ 

1038

 $1039 \\ 1040$ 

 $1041 \\ 1042$ 

1043

 $1044 \\ 1045$ 

 $1046 \\ 1047$ 

1048

 $1049 \\ 1050$ 

 $1051 \\ 1052 \\ 1053$ 

 $1054 \\ 1055$ 

 $\begin{array}{c} 1056 \\ 1057 \end{array}$ 

1058

Our observation of an overall decrease in triacylglycerol lipase transcripts contrasts with previous work by Howard et al. (2010) [18], who observed increased gene copy number of Group 1 lipase genes via qPCR during swine decomposition. Fatty acid composition differs in human compared to pig tissue [30], potentially altering the lipid profile available for microbes, leading to differences in decomposition products within the soil [17]. These products can then directly or indirectly alter community composition and/or activity of functional proteins via substrate availability or the chemical environment. Further, decomposition of humans and pigs resulted in increased pH in soils below pigs, and decreased pH below humans [17]. Altered pH and soil chemistry could result in a different functional potential and/or gene expression in decomposition-impacted soils. Many triacylglycerol lipases have a pH optimum that is neutral to basic [31–33], so cells may be decreasing expression under acidic conditions in human decomposition soils. Availability of lipid species and changes to pH may select for taxa that favor these substrates/pH conditions; for example, Mason et al. (2022) [12] suggested the abundance of the fungal taxa Saccharomycetes was related to antemortem BMI due to relative proportions of fat and muscle tissue.

#### Evidence for phased denitrification and nitrification

The human body is a concentrated source of nitrogen that is released into the surrounding soil during decomposition, therefore we also evaluated expression of genes involved 1059 in nitrogen cycling. Expression of common marker genes for nitrogen cycling was 1060 1061 altered in decomposition soil and suggested nitrogen transformations during human 1062 decomposition are driven by soil oxygen concentrations with hydroxylamine as an 1064 important intermediate. We observed low or reduced expression of nitrification genes 1065 1066 1066 1066 1066 1067 1068 1068 1069 maximum on day 12, and low nitrate conditions indicating that nitrification was inhib-1070 1066 1067 1068 1069 ited. This period of reduced soil oxygen constraining nitrification was also described 1072 in a decomposition experiment with beaver carcasses Keenan et al. (2018) [8].

1074

1104

1075 We observed increased expression of hao, which encodes the enzyme hydroxylamine dehydrogenase (HAO) at day 86 while oxygen was reduced (~85%). This corresponded 1077 1078 to simultaneous increases in expression of genes encoding nitic oxide reductase (norB) 1079 1080 and nitrous oxide reductase (nosZ). Traditionally HAO has been thought to process hydroxylamine to nitrite during nitrification, while NorB and NosZ are enzymes 1083 involved in the last two steps of denitrification converting nitric oxide (NO) to dini-1085 trogen gas  $(N_2)$ . However, recent work has suggested hydroxylamine can be converted to nitric oxide (NO), as well as can interact with multiple phases of the nitrogen cycle 1088 [34]. Even though amoA expression was shown to decrease during reduced oxygen 1090 conditions, amoA transcripts were still present and likely able to convert ammonium to hydroxylamine as soil oxygen was not completely depleted during decomposition. 1092 1093 Additionally, a previous study reported that the growth of the ammonia oxidizing 1094 1095 bacteria Nitrosomonas europaea under anoxic conditions lead to accumulation of hydroxylamine in a chemostat bioreactor [35], suggesting anaerobic ammonium oxi-1098 dation (anammox) may also be occurring in decomposition soils. However, we did 1100 not observe increases in nirK expression, which might suggest conversion of nitrite to NO for use in the anammox pathway. NO produced via HAO activity may be used

1103 for anammox in these soils; however, the role of hydroxylamine as an intermediate

in an ammox is still debated [34]. Therefore, our current hypothesis is that hydroxylamine accumulates under anaerobic conditions during decomposition, which can then be converted to NO by HAO. This NO would then be present for an aerobic denitrifying bacteria to convert to nitrous oxide ( $N_2O$ ) by NorB and finally to  $N_2$  by NosZ. Keenan et al. (2018) [8] also noted a brief increase in  $N_2O$  emissions, which suggests denitrification was occurring during this phase of reduced soil oxygen concentrations.  $1105 \\ 1106$ 

 $\begin{array}{c} 1107 \\ 1108 \end{array}$ 

1109

 $\begin{array}{c} 1110 \\ 1111 \end{array}$ 

1112 1113

 $\begin{array}{c} 1114 \\ 1115 \end{array}$ 

 $\begin{array}{c} 1116 \\ 1117 \end{array}$ 

1118 1119

1120

 $1121 \\ 1122$ 

1123 1124

1125

 $\frac{1126}{1127}$ 

 $\begin{array}{c} 1128 \\ 1129 \end{array}$ 

1130

1131 1132

1133 1134 1135

 $\begin{array}{c} 1136 \\ 1137 \end{array}$ 

 $1138 \\ 1139$ 

1140

 $1141 \\ 1142$ 

 $\begin{array}{c} 1143 \\ 1144 \end{array}$ 

1145

 $1146 \\ 1147$ 

 $\begin{array}{c} 1148 \\ 1149 \end{array}$ 

1150

As soils fully reoxygenated by day 168, we observed increased expression of genes encoding enzymes involved in aerobic nitrification, amoA and nxrR. Nitrification is an oxygen-dependent process which would be converting the accumulated ammonium to nitrate; the increase in nitrate concentrations may then serve as a substrate for denitrification. We observed increased expression of marker genes encoding all four enzymes in the complete dissimilatory denitrification pathway (narG, nirK, norB, and nosZ) at day 376. Increased expression of nitrification and denitrification marker genes is consistent with accumulation of nitrite, nitrate, and N<sub>2</sub>O after oxygen is reintroduced to soils described in Keenan et al. (2018) [3, 8]. Together, gene expression patterns in our study provide further insight into nitrogen transformations in during vertebrate decomposition, suggesting an important role of hydroxylamine.

#### Increased expression of bile salt hydrolases

Sulfur is present in various organic molecules, including taurine, a sulfur- and nitrogencontaining acid involved in bile acid formation [21]. Taurine is present in the human
body, where it can be absorbed from the diet or synthesized in the liver [36]. However, taurine is also produced as a byproduct of the deconjugation of bile salts via bile
salt hydrolases (BSH) present in the anaerobic gut taxa *Lactobacillus* and *Clostridium*[21]. In our study, we observed increased expression of genes encoding BSH enzymes
between days 12 and 86. Given that increased expression of BSH genes corresponded
to the beginning of active decomposition, when decomposition products were observed

1151 to enter the soil, and the period of reduced dissolved oxygen in our study, it is likely 1152 that taurine accumulation is the result of BSH enzyme activity by anaerobic microor-1154 ganisms. While we did not measure taurine concentrations in this study, our results 1156 correspond to previous decomposition studies that report accumulation of taurine 1157 in various organs and body regions [37–39] and soils [17, 40] during decomposition via metabolomics, and increased relative abundance of Clostridium and Lactobacillus 1161 within the body [41–43] and in decomposition soils [19] via DNA sequencing methods, 1162 including in these soils [14].

1164

1165 One pathway of taurine metabolism is through desulfurization via the  $\alpha$ -ketoglutarate-1167 dependent enzyme taurine dioxygenase (TauD). Specifically, this enzyme, encoded by the gene tauD, converts 2-oxoglutarate and taurine to produce aminoacetalde-1169 1170 hyde, succinate, sulfite, and  $CO_2$  [44]. Succinate and sulfite from this reaction can 1171 1172 then be used for the citric acid cycle and sulfur metabolism, respectively. Given increased BSH expression in our study and reported taurine accumulation in others, 1175 we would expect taurine to be present for microbial metabolism by TauD. However, 11761177 we observed a general decrease in tauD expression between days 12 through 376. This 1178 trend was driven by reduced expression of tauD transcripts associated with Proteobac-1179 1180 teria, Gammaproteobacteria, and Actinobacteria whose relative abundance have been 1182 shown to remain consistent or increase during human decomposition [19], suggesting 1183 that tauD expression is downregulated under decomposition conditions. However, we 1184 1185 noted that expression of tauD genes associated with fungi and a few Betaproteobac-1186 1187 teria displayed increased expression at day 58, corresponding to increased expression of bile salt hydrolases (BSH) between days 12 and 86. The reduction in tauD expres-1190 sion may be due to increased sulfur availability. We did not measure sulfur species 1191 1192 in this experiment; however, others have observed increased sulfur concentrations in 1193 decomposition-impacted soils [3, 7, 11]. Thus, sulfur scavenging pathways such as tau-1195 rine desulfurization by TauD [45], whose genes are expressed under sulfur-limiting 1196

conditions, likely display reduced expression under sulfur replete conditions. Additionally, taurine may be processed through other pathways. For example, taurine can be deaminated by taurine dehydrogenase to produce sulfite and acetyl-CoA for carbon metabolism [44, 46]. Overall, our results suggest that human decomposition has potential impacts on soil sulfur biogeochemistry through deposition of inorganic (sulfate) and organic (sulfur-containing amino acids) sulfur compounds.

1197 1198

 $1199 \\ 1200$ 

1201

 $\begin{array}{c} 1202 \\ 1203 \end{array}$ 

 $1204 \\ 1205$ 

 $1206 \\ 1207 \\ 1208$ 

 $1209 \\ 1210$ 

 $1211\\1212$ 

 $\begin{array}{c} 1213 \\ 1214 \end{array}$ 

1215

 $1216 \\ 1217$ 

 $1218 \\ 1219$ 

1220

 $\begin{array}{c} 1221 \\ 1222 \end{array}$ 

 $1223 \\ 1224$ 

1225

 $1226 \\ 1227$ 

 $1228 \\ 1229 \\ 1230$ 

 $1231 \\ 1232 \\ 1233$ 

1234 1235

 $1236 \\ 1237$ 

1238

1239 1240

 $1241 \\ 1242$ 

#### Conclusion

This study represents the first investigation of soil microbial gene expression during human decomposition. Metatranscriptomic analysis of soils from three human individuals over one year shows that decomposition impacted microbial community gene expression profiles, exhibiting functional shifts over time. This included altered expression of genes involved in lipid, N and S metabolism as microbes processed the nutrient-rich tissues of the human body. Additionally, we noted that functionality within decomposition-impacted soils was still affected after one year and had not returned to starting or background conditions. Together, these results show that vertebrate decomposition has lasting impacts on local soil ecosystems, including soil microbial communities. These results have important implications for understanding biogeochemical changes due to vertebrate mortality events in terrestrial ecosystems.

#### Materials and Methods

#### Study design

In February 2018, three deceased male human subjects (hereafter, "donors") were placed supine on the soil surface at the University of Tennessee Anthropology Research Facility (ARF) and allowed to decompose. Located in Knoxville, TN (35° 56′ 28″ N, 83° 56′ 25″ W) the ARF is a roughly 2-acre outdoor facility dedicated to studying

1243 human decomposition [47]. The soils at the ARF are comprised of the Loyston-Talbott1244
1245 Rock outcrop (LtD) and Coghill-Corryton (CcD) complexes. LtD soils are a silty clay
1246 loam and channery clay overlaying lithic bedrock, while CcD soils are comprised of
1247 clay from weathered quartz limestone [14, 47]. A site that had not been previously
1249 exposed to decomposition was used for this study.
1251

1252The decomposition field experiment is fully described in Taylor et al. (2024) [14]. 1253 1254 Briefly, experiments were conducted in a block design, where each block consisted of 1255one decomposition site and one control site [14]. In total three blocks, i.e., three donors 1256 1257paired with three respective control sites, were included in the study. Each control site 1258 1259 was chosen in a manner to ensure their location was uphill and roughly 2 m away from decomposition sites [14]. Donor internal temperatures were recorded by probes located 1261 1262 in the abdomen, while ambient air temperatures were monitored via sensors located 1263 1264 roughly 50 cm above the soil surface. Soil temperature and salinity were measured with sensors placed directly underneath each individual (Decagon Devices, GS3) [14]. 1266 1267 Donor ages ranged from 65 to 86 and were within 1 kg of each other with regard to 1268 1269 weight (90.7 to 91.6 kg); donor BMI varied between 27.7 to 29.6 [14].

#### 1272 Sampling and physiochemistry

 $1270 \\ 1271$ 

1273

1274 Decomposition of all subjects was observed for one year. During the one-year study 1275 period, soils were sampled at 20 timepoints chosen to correspond with morphological 1276 1277stages of decomposition as described by [48]. Once advanced decay was reached, soils 1278 1279 were collected at intervals of 350 accumulated degree days (ADD), calculated using ambient air temperatures, up to one year. All soil cores were taken using a 1.9 cm 1281 (3/4 inch) diameter soil auger to a depth of 16 cm. Soils were divided into two depth 1283 1284 fractions: 0-1 cm (interface) and 1-16 cm (core) for the analyses reported in Taylor et 1285 al. (2024) [14]; the entire 0 to 16 cm core was used for this current study. Decomposi-1286 1287 tion soils were taken from directly beneath the cadavers, taking care to not re-sample 1288

the same location more than once. At the time of sampling, soil dissolved oxygen was measured in triplicate using an Orion Star<sup>TM</sup> A329 pH/ISE/Conductivity/Dissolved Oxygen portable multiparameter meter (ThermoFisher) [14].

1289 1290

 $1291 \\ 1292$ 

1293 1294

 $\begin{array}{c} 1295 \\ 1296 \end{array}$ 

1297 1298

1299

 $1300\\1301$ 

 $1302 \\ 1303$ 

1304

 $1305 \\ 1306$ 

 $1307 \\ 1308$ 

1309

 $1310\\1311$ 

 $1312 \\ 1313$ 

1314

1315 1316

 $\begin{array}{c} 1317 \\ 1318 \end{array}$ 

 $1319 \\ 1320$ 

 $1321 \\ 1322$ 

1323

 $1324 \\ 1325$ 

 $1326 \\ 1327$ 

1328

1329 1330

A subset of 6 study timepoints were chosen for metatranscriptomics analysis. Study days 0, 12, 58, 86, 168, and 376 were chosen as they represented disctinct morphological and soil biogeochemical stages during decomposition. Study day 0 was chosen as a baseline sample prior to cadaver placement. Study day 12 was the start of active decomposition and corresponded to maximum soil ammonium concentrations and minimum soil oxygen (approximately 39%). Study day 58 was chosen as this sample represented the pH minimum, and respiration and soil temperature were at a maximum [14]. Additionally, ammonium concentrations began to decrease around day 58. Study day 86 was when soil oxygen started to recover and nitrate levels began to increase. Study day 168 was chosen as nitrate was at its maximum and soil dissolved oxygen had returned to 99%. Finally, day 376 was chosen to represent the end of the study, 1 year since cadaver palcement. Each study day was represented by four soil samples for RNA extraxtion: one pooled control sample which was a mix of the three control locations, plus one sample from each of the three donors, yielding a total of 24 samples for this study.

Soil samples were transported back to the University of Tennessee (Knoxville, TN) and processed within 24 hours of collection. Soils were homogenized by hand to remove insect larvae, roots, rocks, and other debris (> 2 mm). A subset of soils were used to measure pH, electrical conductivity (EC), and evolved  $CO_2$  as described in Taylor (2024). Soil nitrogen species ( $NH_4^+$ ,  $NO_3^-$ ) and total carbon (TC) and nitrogen (TN) were measured in all soil samples as described in [14]. Reported values for soil physiochemistry represent the full 16 cm core; estimated by summing interface and core

1335 values reported by Taylor et. al, (2024) [14] in 1:16 and 15:16 ratios, respectively. Con-1337 trol reported here are means of the three experimental controls that were unimpacted 1338 by decomposition. 1339

1340

1341 Roughly 10 g of soil was reserved for nucleic acid extraction, placed in a 4 oz. Whirl-Pak<sup>TM</sup> bag (Nasco), and flash frozen in liquid nitrogen. All samples were stored at  $1344\,\text{--}80^\circ\mathrm{C}$  until further analysis. Bacterial and fungal community composition was assessed 1346 via amplicon sequencing of the 16S rRNA gene and ITS2 region as described in Taylor et al. (2024). 1348

1349 1350

1351 1352

#### RNA Extraction and Sequencing

RNA was extracted from 2 g of soil using Qiagen's RNeasy® PowerSoil® Total RNA 1353 1354 kit. Manufacturer's instructions were followed with a few modifications. Soils became 1355 1356 saline during decomposition; therefore, we followed the manufacturer's suggestion and  $\overline{1358}$  incubated all extracts at -20°C following addition of solution SR4 (step 9) to decrease 1359 salt precipitation. All RNA samples were resuspended in 40  $\upmu{\rm l}$  of Solution SR7. RNA 1360 1361 concentrations were assessed fluorometrically using the Qubit® RNA HS assay (cat-1362 alog no. Q32852) with 1 µl of RNA. DNA contamination was removed by DNase 1363 1364treating RNA extracts twice using Qiagen's DNase Max® kit in  $50~\mu l$  reactions. RNA 1366 concentrations were remeasured after DNase treatment. PCR with V4 16S rRNA gene 1368 primers [49, 50] was conducted using RNA extracts as the template to confirm removal 1369 of all DNA prior to sequencing. RNA aliquots were shipped to HudsonAlpha Dis-1370 1371 covery (Huntsville, AL) for library preparation and RNA sequencing. Dual-indexed libraries were prepared using the Illumina® Stranded Total RNA prep with riboso-1373 1374mal RNA depletion via ligation with Ribo-Zero Plus. Libraries were then pooled and 13751376 sequenced on Illumina's NovaSeq 6000 v4 platform, resulting in demultiplexed fastq 1377 files for each sample. 1378

#### **Bioinformatics**

Read quality control (QC) was conducted in KBase [51] using Trimmomatic [52]. Paired fastq files were imported to KBase through Globus. Poor quality reads were removed, and adapters trimmed via Trimmomatic (v0.36) using default settings and the TruSeq3-PE-2 adapter file. After QC check with FastQC, trimmed libraries were exported as fastq files from KBase through Globus. Remaining ribosomal RNA was filtered using bbmap (maxindel = 20, minid = 0.93) from the Joint Genome Institute's (JGI) bbtools suite [53]. After this step, all non-ribosomal reads from all 24 samples were merged into one file. This file was then used to co-assemble reads into contigs using the de novo assembler MEGAHIT (v1.2.9) [54] (-12 -k-min 23, -k-max 123, -k-step 10).

1381 1382 1383

1384

 $1385 \\ 1386$ 

 $1387 \\ 1388$ 

1389

 $1390 \\ 1391$ 

 $1392 \\ 1393$ 

1394

 $1395 \\ 1396$ 

 $1397 \\ 1398$ 

 $1399 \\ 1400$ 

 $1401\\1402$ 

1403

 $1404\\1405$ 

 $1406\\1407$ 

1408

 $1409 \\ 1410$ 

 $1411 \\ 1412$ 

1413

1414 1415

 $\begin{array}{c} 1416 \\ 1417 \end{array}$ 

 $1418 \\ 1419 \\ 1420$ 

1421 1422

 $1423 \\ 1424$ 

 $1425 \\ 1426$ 

Gene identification and annotation from co-assembled contigs was performed using Prodigal [55] and eggNOG mapper [56], respectively. Briefly, the fastq containing all contigs was summitted to Prodigal (v2.6.3) for protein coding gene predication for a meta-sample (-p meta -f gff). Next, predicated genes were functionally and taxonomically annotated using eggNOG mapper (v2.1.6) using basic settings to perform a diamond blastp search [57]. Only genes that were both functionally and taxonomically annotated by one of the databases used by eggNOG mapper and identified as bacterial, archaeal, or fungal were chosen as genes of interest. Transcript counts for all genes of interest were obtained by mapping reads from each respective sample to genes of interest obtained from co-assembly using QIAGEN CLC Genomics Workbench 20.0 (https://digitalinsights.qiagen.com/).

#### Differential Expression

Transcript counts from all samples were combined in a single workable data file and imported into R for differential expression analysis using the R packages edgeR [58]

1427 and limma [59] following a modified pipeline by Phipson et al. (2020) [60]. The tran- $_{1429}$  script count table was imported into R and converted to a DGElist object. Genes 1430 without sufficient counts for statistical analysis were removed to increase power using 1431 1432 the edgeR function filterByExpr(), using study day as the comparison group. 1433 1434 Raw counts were then log2 normalized and gene expression profiles compared via 1435 1436 multidimensional scaling (MDS) and hierarchical clustering. Mutlidimendional scal-1437 1438 ing (MDS) using plotMDS() from the limma package was used to assess differences between samples. MDS values were extracted from the MDS object, and the first two 1440 1441 dimensions plotted using ggplot2 [61]. We also assessed the relationship bewteen gene 1443 expression profiles and changes in the soil environment using canonical correspondence analysis (CCA). Environmental variables of interest included decomposition 1445 1446 time in accumulated degree hours (ADH) based on ambient temperatures, ADH based 1447 1448 on internal gut temperatures, ADH based on soil temperatures, gravimetric moisture, pH, electrical conductivity (EC), dissolved oxygen (DO), CO<sub>2</sub> (mol gdw<sup>-1</sup>), NH<sub>4</sub> (mg gdw<sup>-1</sup>), NO<sub>3</sub> (mg gdw<sup>-1</sup>), N %, C %, and CN ratio. First, permutational multivari-1452 1453 ate analysis of variance (PERMANOVA) with adonis() (vegan v2.6.7) [62] was used 1454 to identify significant soil parameters. Then the vegan functions cca() and scores() 1455 1456 were applied to run the CCA and extract scores, respectively. Scores for the first two 1458 dimenstions were plotting using ggplot2, with loadings extracted from the CCA biplot. 1459 1460 For differential expression analysis, raw filtered reads were normalized using edgeR's 1461 1462 trimmed mean of M values (TMM) normalization using the function calcNormFac-1463 tors(). TMM normalized reads were then log2 transformed using limma's voom() and 1464 1465 differential expression assessed. Empirical Bayes shrinkage was used correct to p-1466 1467 values for false discovery rates. The topmost up and down regulated genes for each 1469 comparison, determined by  $\log 2$  fold change and adjusted p-values, were then reported. 1470Expression of certain genes were assessed after performing transcripts per million 1471

(TPM) normalization and statistical analyses with a combination of analysis of variance (ANOVA) and post-hoc Tukey tests. ANOVA across all timepoints were applied to hierarchical linear mixed effects models to account for repeated sampling within each donor block.

 $1473 \\ 1474$ 

 $1475 \\ 1476$ 

1477

 $1478 \\ 1479 \\ 1480$ 

 $1481 \\ 1482 \\ 1483$ 

 $1484 \\ 1485$ 

1486

 $1487 \\ 1488 \\ 1489$ 

1490 1491 1492

 $1493 \\ 1494$ 

1495 1496

 $1497 \\ 1498$ 

 $1499 \\ 1500$ 

 $1501 \\ 1502$ 

 $1503 \\ 1504$ 

 $\begin{array}{c} 1505 \\ 1506 \end{array}$ 

 $1507 \\ 1508$ 

 $1509 \\ 1510$ 

 $\begin{array}{c} 1511 \\ 1512 \end{array}$ 

 $1513 \\ 1514$ 

#### Availability of data and materials

RNA sequence files from the Novoseq intrument can be found at XXXX. The datasets supporting the conclusions of this article are available at the GitHub repository Mason\_MetaT\_XXX\_2024."

#### References

- [1] Benninger, L. A., Carter, D. O. & Forbes, S. L. The biochemical alteration of soil beneath a decomposing carcass. *Forensic Science International* **180**, 70–5 (2008).
- [2] Towne, E. G. Prairie vegetation and soil nutrient responses to ungulate carcasses. Oecologia 122, 232–239 (2000). URL https://doi.org/10.1007/PL00008851.
- [3] DeBruyn, J. M., Keenan, S. W. & Taylor, L. S. From carrion to soil: microbial recycling of animal carcasses. *Trends in Microbiology* (2024). URL https://doi. org/10.1016/j.tim.2024.09.003. Publisher: Elsevier.
- [4] Parmenter, R. R. & MacMahon, J. A. Carrion decomposition and nutrient cycling in a semiarid shrub–steppe ecosystem. *Ecological Monographs* **79**, 637–661 (2009).
- [5] Macdonald, B. C. T. et al. Carrion decomposition causes large and lasting effects on soil amino acid and peptide flux. Soil Biology and Biochemistry 69, 132–140 (2014).

- 1519 [6] Bump, J. K. et al. Ungulate carcasses perforate ecological filters and create
  1520
  1521 biogeochemical hotspots in forest herbaceous layers allowing trees a competitive
  1522
  1523 advantage. Ecosystems 12, 996–1007 (2009).
- 1524 1525 [7] Aitkenhead-Peterson, J. A., Owings, C. G., Alexander, M. B., Larison, N. & 1526 1527 Bytheway, J. A. Mapping the lateral extent of human cadaver decomposition 1528 with soil chemistry. Forensic Science International 216, 127–34 (2012).
- 1530
  1531 [8] Keenan, S. W., Schaeffer, S. M., Jin, V. L. & DeBruyn, J. M. Mortality hotspots:
  1532
  1533 nitrogen cycling in forest soils during vertebrate decomposition. Soil Biology and
  1534 Biochemistry 121, 165–176 (2018).

1535

1541

1547

1553

1557

- 1536
   1537 [9] Fancher, J. P. et al. An evaluation of soil chemistry in human cadaver decomposition islands: Potential for estimating postmortem interval (PMI). Forensic
   1539
   1540 Science International 279, 130–139 (2017).
- 1542 [10] Quaggiotto, M.-M., Evans, M. J., Higgins, A., Strong, C. & Barton, P. S.
  1544 Dynamic soil nutrient and moisture changes under decomposing vertebrate
  1545 carcasses. Biogeochemistry 146, 71–82 (2019).
- 1554 [12] Mason, A. R. et al. Body mass index (BMI) impacts soil chemical and microbial 1555 response to human decomposition. mSphere e0032522 (2022).
- 1558 [13] Mason, A. R., Taylor, L. S. & DeBruyn, J. M. Microbial ecology of vertebrate
   1559 decomposition in terrestrial ecosystems. FEMS Microbiology Ecology 99, fiad006
   1561 (2023). URL https://doi.org/10.1093/femsec/fiad006.

[14] Taylor, L. S. et al. Transient hypoxia drives soil microbial community dynamics and biogeochemistry during human decomposition. FEMS Microbiology Ecology 100, fiae119 (2024). URL https://doi.org/10.1093/femsec/fiae119.  $1565 \\ 1566$ 

 $\begin{array}{c} 1567 \\ 1568 \end{array}$ 

 $1569 \\ 1570$ 

 $1571 \\ 1572$ 

 $1573 \\ 1574$ 

 $\begin{array}{c} 1575 \\ 1576 \end{array}$ 

 $\begin{array}{c} 1577 \\ 1578 \end{array}$ 

1579

 $1580 \\ 1581$ 

 $1582 \\ 1583 \\ 1584$ 

1585

 $\begin{array}{c} 1586 \\ 1587 \end{array}$ 

 $1588 \\ 1589$ 

 $1590 \\ 1591$ 

1592 1593

1594 1595

 $\begin{array}{c} 1596 \\ 1597 \end{array}$ 

 $1598 \\ 1599$ 

 $\begin{array}{c} 1600 \\ 1601 \end{array}$ 

 $1602 \\ 1603$ 

1604

1605 1606 1607

 $\begin{array}{c} 1608 \\ 1609 \end{array}$ 

- [15] Burcham, Z. M. et al. Total RNA analysis of bacterial community structural and functional shifts throughout vertebrate decomposition. Journal of Forensic Sciences 64, 1707–1719 (2019).
- [16] Ashe, E. C., Comeau, A. M., Zejdlik, K. & O'Connell, S. P. Characterization of bacterial community dynamics of the human mouth throughout decomposition via metagenomic, metatranscriptomic, and culturing techniques. Frontiers in Microbiology 12, 689493 (2021).
- [17] DeBruyn, J. M. et al. Comparative decomposition of humans and pigs: soil biogeochemistry, microbial activity and metabolomic profiles. Frontiers in Microbiology 11, 608856 (2021).
- [18] Howard, G. T., Duos, B. & Watson-Horzelski, E. J. Characterization of the soil microbial community associated with the decomposition of a swine carcass. *International Biodeterioration & Biodegradation* 64, 300–304 (2010).
- [19] Cobaugh, K. L., Schaeffer, S. M. & DeBruyn, J. M. Functional and structural succession of soil microbial communities below decomposing human cadavers. *Plos One* 10, e0130201 (2015).
- [20] Singh, B. et al. Temporal and spatial impact of human cadaver decomposition on soil bacterial and arthropod community structure and function. Frontiers in Microbiology 8, 2616 (2018).
- [21] Urdaneta, V. & Casadesús, J. Interactions between Bacteria and Bile Salts in the Gastrointestinal and Hepatobiliary Tracts. Frontiers in Medicine 4 (2017).

- 1611 [22] van der Wal, A., Geydan, T. D., Kuyper, T. W. & de Boer, W. A thready affair:
- linking fungal diversity and community dynamics to terrestrial decomposition
- processes. FEMS Microbiology Reviews 37, 477–494 (2013).

1617 [23] Essington, M. E. Soil and water chemistry: an integrative approach (CRC press, 2015).

- [24] Peng, J., Wegner, C.-E. & Liesack, W. Short-term exposure of paddy soil micro-bial communities to salt stress triggers different transcriptional responses of key
- taxonomic groups. Frontiers in Microbiology 8 (2017).

1627 [25] Pandit, A. S. et al. A snapshot of microbial communities from the Kutch: one of the largest salt deserts in the World. Extremophiles 19, 973–987 (2015). 

1631 [26] Metcalf, J. L. et al. Microbial community assembly and metabolic function during mammalian corpse decomposition. Science 351, 158–62 (2016). 

[27] Fu, X. et al. Fungal succession during mammalian cadaver decomposition and potential forensic implications. Scientific Reports 9, 12907 (2019). 

[28] Dujon, B. et al. Genome evolution in yeasts. Nature **430**, 35–44 (2004). 

1642 [29] Haridas, S. et al. The genome and transcriptome of the pine saprophyte Ophios-toma piceae, and a comparison with the bark beetle-associated pine pathogen Grosmannia clavigera. BMC Genomics 14, 373 (2013).

[30] Notter, S. J., Stuart, B. H., Rowe, R. & Langlois, N. The initial changes of fat deposits during the decomposition of human and pig remains. Journal of Forensic Sciences **54**, 195–201 (2009).

[31] Kok, R. G. et al. Characterization of the extracellular lipase, LipA, of Acineto-bacter calcoaceticus BD413 and sequence analysis of the cloned structural gene. 

- Molecular Microbiology 15, 803–818 (1995).
- [32] Hasan, F., Shah, A. A. & Hameed, A. Influence of culture conditions on lipase production by Bacillus sp. FH5. *Annals of Microbiology* **56**, 247–252 (2006).

 $1657 \\ 1658 \\ 1659$ 

 $1660 \\ 1661$ 

 $\begin{array}{c} 1662 \\ 1663 \end{array}$ 

 $1664 \\ 1665$ 

1666

1667 1668 1669

 $\begin{array}{c} 1670 \\ 1671 \end{array}$ 

 $1672 \\ 1673$ 

 $\begin{array}{c} 1674 \\ 1675 \end{array}$ 

1676

 $1677 \\ 1678$ 

1679 1680 1681

1682

 $1683 \\ 1684$ 

 $\begin{array}{c} 1685 \\ 1686 \end{array}$ 

1687 1688

 $1689 \\ 1690$ 

 $1691 \\ 1692$ 

1693 1694

 $\begin{array}{c} 1695 \\ 1696 \end{array}$ 

1697 1698

 $1699 \\ 1700 \\ 1701 \\ 1702$ 

- [33] Zouaoui, B. & Bouziane, A. Production, optimization and characterization of the lipase from Pseudomonas aeruginosa. *Romanian biotechnological letters* 17, 7187–7193 (2012).
- [34] Soler-Jofra, A., Pérez, J. & van Loosdrecht, M. C. M. Hydroxylamine and the nitrogen cycle: A review. Water Research 190, 116723 (2021).
- [35] Yu, R., Perez-Garcia, O., Lu, H. & Chandran, K. Nitrosomonas europaea adaptation to anoxic-oxic cycling: Insights from transcription analysis, proteomics and metabolic network modeling. Science of the Total Environment 615, 1566–1573 (2018).
- [36] Seidel, U., Huebbe, P. & Rimbach, G. Taurine: A regulator of cellular redox homeostasis and skeletal muscle function. *Molecular Nutrition & Food Research* 63, 1800569 (2019).
- [37] Mora-Ortiz, M., Trichard, M., Oregioni, A. & Claus, S. P. Thanatometabolomics: introducing NMR-based metabolomics to identify metabolic biomarkers of the time of death. *Metabolomics* **15**, 37 (2019).
- [38] Locci, E. et al. A 1H NMR metabolomic approach for the estimation of the time since death using aqueous humour: an animal model. Metabolomics 15, 76 (2019).
- [39] Zelentsova, E. A. et al. Post-mortem changes in the metabolomic compositions of rabbit blood, aqueous and vitreous humors. Metabolomics 12, 172 (2016).

- 1703 [40] Hoeland Katharina, M. Investigating the potential of postmortem metabolomics
- in mammalian decomposition studies in outdoor settings. Ph.D. thesis, University
- of Tennessee-Knoxville, https://trace.tennessee.edu/utk\_graddiss/7000 (2021).

- 1709 [41] Javan, G. T. et al. Human thanatomicrobiome succession and time since death.
- Scientific Reports 6, 29598 (2016).

- 1713 [42] Javan, G. T., Finley, S. J., Smith, T., Miller, J. & Wilkinson, J. E. Cadaver
- thanatomicrobiome signatures: the ubiquitous nature of Clostridium species in
- human decomposition. Frontiers in Microbiology 8, 2096 (2017).

- 1719 [43] DeBruyn, J. M. & Hauther, K. A. Postmortem succession of gut microbial
- communities in deceased human subjects. Peerj 5, e3437 (2017).

- 1723 [44] Cook, A. M. & Denger, K. Metabolism of taurine in microorganisms. Taurine 6
- 3-13 (2006).

- [45] Kertesz, M. A. Riding the sulfur cycle – metabolism of sulfonates and sul-
- fate esters in Gram-negative bacteria. FEMS Microbiology Reviews 24, 135-175
- (2000).

- 1733 [46] Brüggemann, C., Denger, K., Cook, A. M. & Ruff, J. Enzymes and genes of
- taurine and isethionate dissimilation in Paracoccus denitrificans. Microbiology
- (Reading, England) 150, 805–816 (2004).

- 1739 [47] Keenan, S. W. et al. Spatial impacts of a multi-individual grave on microbial
- and microfaunal communities and soil biogeochemistry. PLoS One 13, e0208845
- (2018).

- [48] Payne, J. A. A summer carrion study of the baby pig Sus Scrofa Linnaeus.
- Ecology 46, 592–602 (1965).

[49] Apprill, A., McNally, S., Parsons, R. & Weber, L. Minor revision to V4 region SSU rRNA 806R gene primer greatly increases detection of SAR11 bacterioplankton. Aquatic Microbial Ecology 75, 129–137 (2015).  $1749 \\ 1750$ 

 $1751 \\ 1752$ 

1753 1754

 $1755 \\ 1756$ 

1757 1758

1759

1760 1761 1762

1763

1764 1765 1766

1767 1768

 $1769 \\ 1770$ 

1771 1772 1773

1774

 $1775 \\ 1776$ 

 $1777 \\ 1778$ 

1779 1780

1781 1782 1783

1784

 $1785 \\ 1786$ 

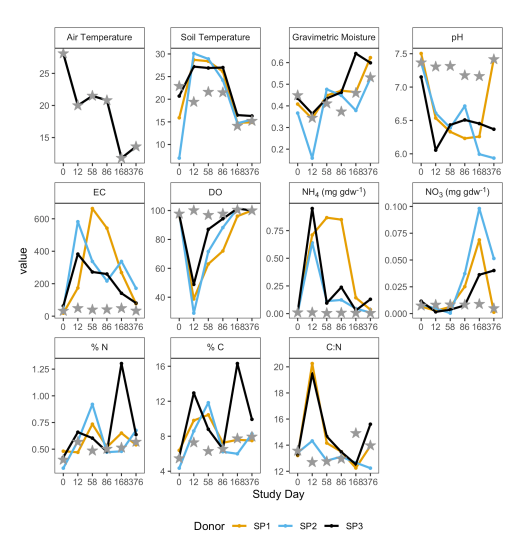
1787 1788

 $1789 \\ 1790$ 

 $1791 \\ 1792$ 

- [50] Parada, A. E., Needham, D. M. & Fuhrman, J. A. Every base matters: assessing small subunit rRNA primers for marine microbiomes with mock communities, time series and global field samples. *Environmental Microbiology* 18, 1403–14 (2016).
- [51] Arkin, A. P. et al. KBase: The United States Department of Energy Systems Biology Knowledgebase. Nature Biotechnology 36, 566–569 (2018).
- [52] Bolger, A. M., Lohse, M. & Usadel, B. Trimmomatic: a flexible trimmer for Illumina sequence data. *Bioinformatics* 30, 2114–2120 (2014).
- [53] Bushnell, B. BBMap. URL sourceforge.net/projects/bbmap/.
- [54] Li, D., Liu, C.-M., Luo, R., Sadakane, K. & Lam, T.-W. MEGAHIT: an ultra-fast single-node solution for large and complex metagenomics assembly via succinct de Bruijn graph. *Bioinformatics* 31, 1674–1676 (2015).
- [55] Hyatt, D. et al. Prodigal: prokaryotic gene recognition and translation initiation site identification. BMC Bioinformatics 11, 119 (2010).
- [56] Cantalapiedra, C. P., Hernández-Plaza, A., Letunic, I., Bork, P. & Huerta-Cepas, J. eggNOG-mapper v2: functional annotation, orthology assignments, and domain prediction at the metagenomic scale. *Molecular Biology and Evolution* 38, 5825–5829 (2021).
- [57] Buchfink, B., Reuter, K. & Drost, H.-G. Sensitive protein alignments at tree-of-life scale using DIAMOND. *Nature Methods* 18, 366–368 (2021).

1795 [58] Robinson, M. D., McCarthy, D. J. & Smyth, G. K. edgeR: a Bioconduc-tor package for differential expression analysis of digital gene expression data. Bioinformatics 26, 139–140 (2010). 1801 [59] Smyth, G. K. in limma: Linear Models for Microarray Data (eds Gentleman, R., Carey, V. J., Huber, W., Irizarry, R. A. & Dudoit, S.) Bioinformatics and Computational Biology Solutions Using R and Bioconductor 397–420 (Springer New York, New York, NY, 2005). [60] Phipson, B. et al. Differential expression analysis (2020). URL https://combine-australia.github.io/RNAseq-R/06-rnaseq-day1.html#References. [61] Wickham, H. qqplot2: Elegant Graphics for Data Analysis (Springer-Verlag New York, 2016). URL https://ggplot2.tidyverse.org. [62] Oksanen, J. et al. vegan: Community Ecology Package (2024). URL https:// vegandevs.github.io/vegan/.  ${f Acknowledge ments}$ 1824 We would like to thank the Forensic Anthropology Center at the University of Tennessee-Knoxville for their help in setting up field experiments. We would like to  $\frac{1827}{1828}$  thank Mary Davis for her help in managing the field site and helping to obtain donors 1829 for this work. This research was funded by a National Institute of Justice Award (DOJ-NIJ-2017-R2-CX-0008) to LST and JMD. Supplementary Information 

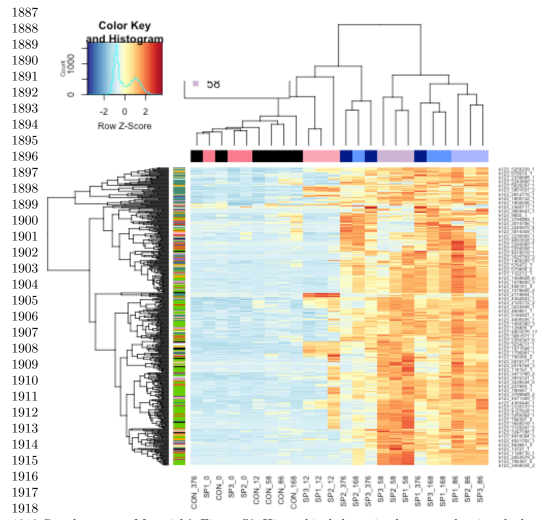


 $1871 \\ 1872$ 

 $1873 \\ 1874$ 

 $\begin{array}{c} 1875 \\ 1876 \end{array}$ 

Supplementary Material 1: Figure S1. Soil physiochemical parameters in decomposition soils during the one-year study. Data is shown for each individual donor: SP1 (gold), SP2 (blue), and SP2 (black). Values for the full 16 cm core samples were estimated by summing values interface (0-1 cm) and core (0-16 cm) reported by Taylor et al, (2024) in 1:16 and 15:16 ratios, respectively. Controls reported here are means of three experimental controls that were unimpacted by decomposition and are represented by stars.



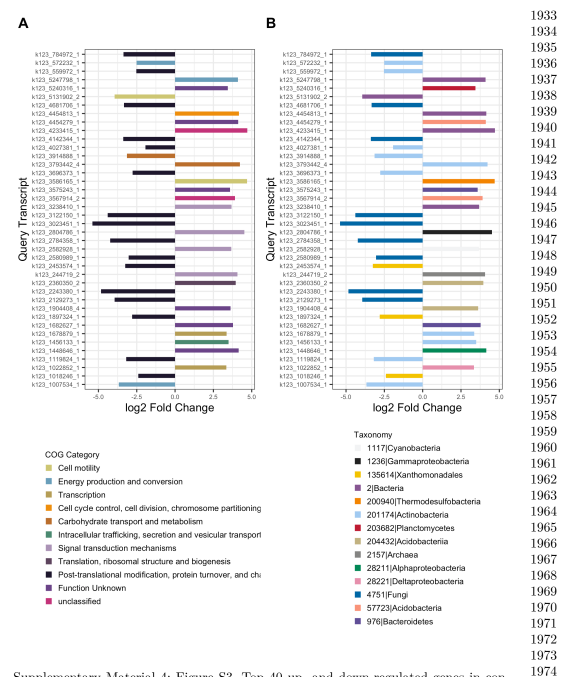
1919 Supplementary Material 2: Figure S2. Hierarchical clustering heatmap showing the log 1920 counts per million (CPM) of the top 500 most variable genes across samples. Variable 1921 genes were determined by selecting genes with the highest variance in gene expression. 1922 Samples are clustered along the x-axis using Euclidean distances between samples and 1923 colored by study day.

1925

1931 1932

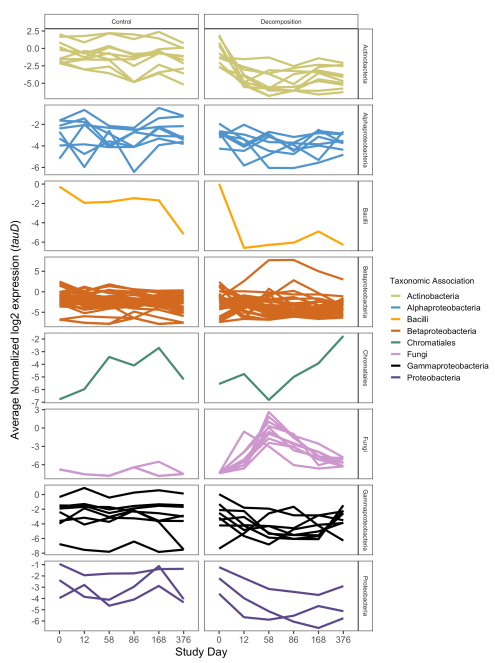
 $\begin{array}{c} 1926 \\ 1927 \\ 1928 \\ \end{array} \\ \begin{array}{c} \text{Table S1. Permutational analysis of variance (PERMANOVA) results identifying} \\ 1928 \\ \text{significant environmental parameters which explain some of the variation in soil gene} \\ 1929 \\ \text{expression profiles. Environmental parameter data is from Taylor et al. (2024).} \\ \text{Variables with p} < 0.05 \text{ are indicated in bold.} \end{array}$ 

Supplementary Material 3



Supplementary Material 4: Figure S3. Top 40 up- and down-regulated genes in controls relative to decomposition soils across all study days, colored by COG functional category (A) and taxonomic annotation (B). Positive values denote higher expression in controls, while negative values are higher in decomposition soils.

1980 1981 1982 1983 1984 1985 1986 Table S2. Top 20 most up- and down-regulated gene queries, determined by change and adjusted p-values, in control relative to decomposition soils. log2 fold change values represent genes whose expression was higher in conwide while negative log2 fold change values were higher in decomposition soils. To annotation, COG categories, gene description, gene names, and EC were via eggNOG-mapper. Supplementary Material 5 1996 1997 1998 1999 2000 2001 2002 2003 2004 2006 2007 2008	Positive atrol soils, Caxonomic
Table S3. Top 10 most up- and down-regulated genes, determined by lo change and adjusted p-values, for each sequential timepoint comparison. log2 fold change values represent genes whose expression was higher in to decomposition timepoint soils, while negative log2 fold change values are earlier decomposition timepoint soils. Taxonomic annotation, COG categor names, and EC were assigned via eggNOG-mapper. The comparison of distinguishes each timepoint comparison.  Supplementary Material 6  Supplementary Material 6  2017  2018  2020  2021  2022  2023  2024	Positive he later higher in ries, gene



 $2030 \\ 2031 \\ 2032 \\ 2033 \\ 2034$ 

 $\begin{array}{c} 2035 \\ 2036 \end{array}$ 

 $\begin{array}{c} 2040 \\ 2041 \end{array}$ 

 $2048 \\ 2049 \\ 2050$ 

 $2051 \\ 2052 \\ 2053 \\ 2054 \\ 2055 \\ 2056 \\ 2057$ 

Supplementary Material 7: Figure S4. Mean normalized  $\log 2$  expression of tauD genes by taxonomic association (color) in control and decomposition soils at each study day. Each line represents one tauD gene query, while color denotes taxonomic association as determined by eggNOG-mapper.