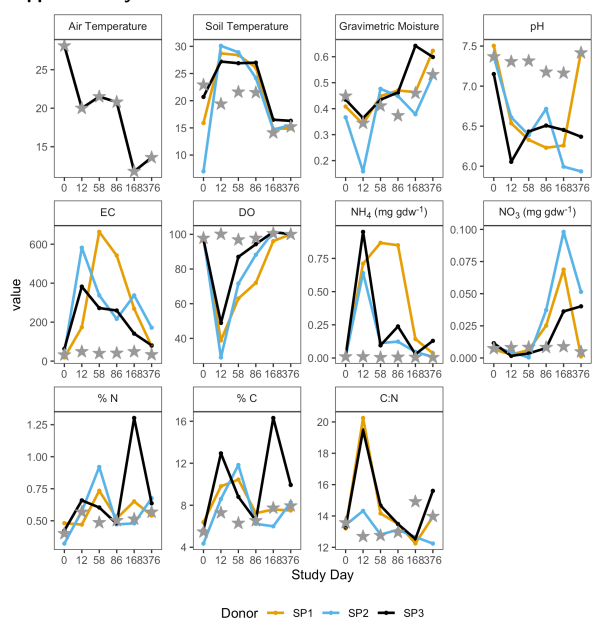
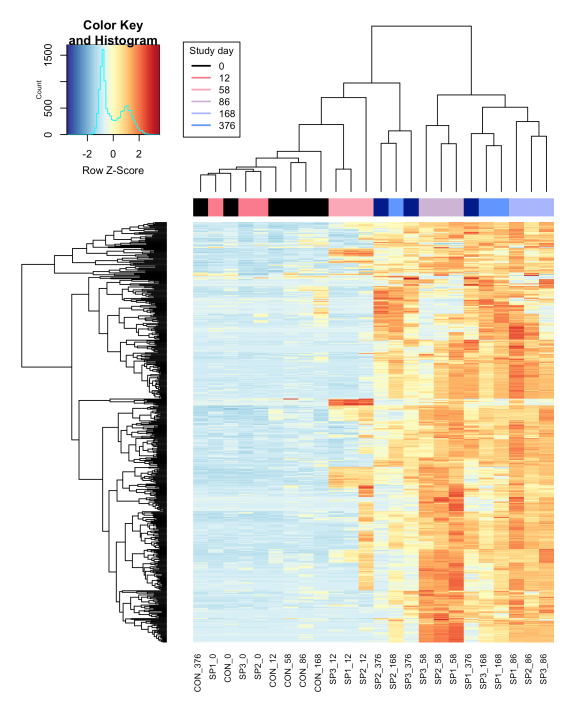
# **Supplementary Information**



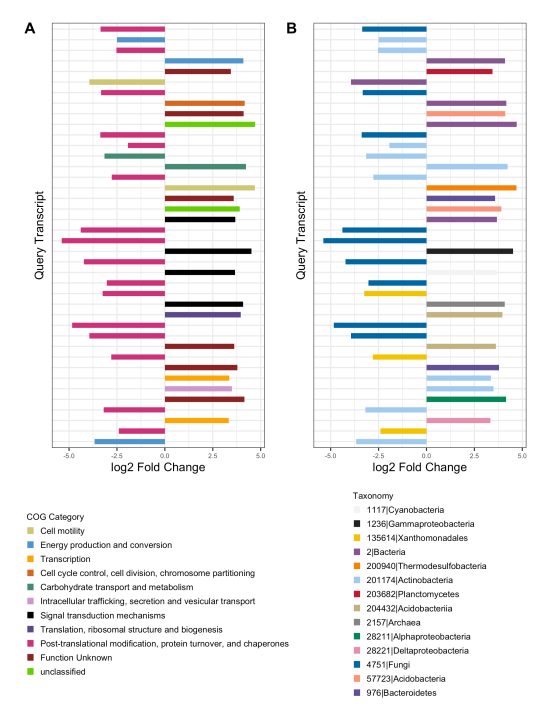
Supplementary Fig 1. Soil physiochemical parameters in decomposition soils during the one-year study. Data is shown for each individual donor: SP1 (gold), SP2 (blue), and SP2 (black). Values for the full 16 cm core samples were estimated by summing values interface (0-1 cm) and core (0-16 cm) reported 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.



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

Supplementary Table 1. Permutational analysis of variance (PERMANOVA) results identifying significant environmental parameters which explain some of the variation in soil gene expression profiles. Environmental parameter data is from Taylor et al. (2024). Variables with p < 0.05 are indicated in bold.

|                             | Df | SumsOfSqs | MeanSqs | F.Model | R2    | Pr(>F) |
|-----------------------------|----|-----------|---------|---------|-------|--------|
| $\overline{\mathrm{ADH_A}}$ | 1  | 0.287     | 0.287   | 3.061   | 0.082 | 0.047  |
| $\mathrm{ADH_{I}}$          | 1  | 0.536     | 0.536   | 5.712   | 0.152 | 0.001  |
| $\mathrm{ADH_S}$            | 1  | 0.154     | 0.154   | 1.640   | 0.044 | 0.158  |
| $\mathrm{T}_{\mathrm{A}}$   | 1  | 0.185     | 0.185   | 1.970   | 0.053 | 0.123  |
| $\mathrm{T_{S}}$            | 1  | 0.258     | 0.258   | 2.749   | 0.073 | 0.039  |
| GWC                         | 1  | 0.117     | 0.117   | 1.245   | 0.033 | 0.285  |
| $_{ m pH}$                  | 1  | 0.258     | 0.258   | 2.751   | 0.073 | 0.033  |
| EC                          | 1  | 0.294     | 0.294   | 3.139   | 0.084 | 0.031  |
| DO                          | 1  | 0.086     | 0.086   | 0.920   | 0.025 | 0.483  |
| $CO_2$                      | 1  | 0.131     | 0.131   | 1.399   | 0.037 | 0.240  |
| $\mathrm{NH_4}$             | 1  | 0.127     | 0.127   | 1.352   | 0.036 | 0.205  |
| $NO_3$                      | 1  | 0.081     | 0.081   | 0.861   | 0.023 | 0.479  |
| N %                         | 1  | 0.101     | 0.101   | 1.072   | 0.029 | 0.332  |
| C%                          | 1  | 0.092     | 0.092   | 0.980   | 0.026 | 0.403  |
| C:N                         | 1  | 0.059     | 0.059   | 0.630   | 0.017 | 0.727  |
| Residuals                   | 8  | 0.751     | 0.094   | NA      | 0.213 | NA     |
| Total                       | 23 | 3.517     | NA      | NA      | 1.000 | NA     |



Supplementary Fig 3. 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.

| Query         | Coef | fficient | p-Value | Taxonomic Annotation  | COG<br>Category | Description   | Gene<br>Name | EC       |
|---------------|------|----------|---------|-----------------------|-----------------|---|--------------|----------|
| k123_3586165_ | _1   | 4.699    | 0       | Thermodesulfobacteria | NU              | Type II secretion<br>system (T2SS),<br>protein E,<br>N-terminal<br>domain | -            | -        |
| k123_5240316_ | _1   | 3.444    | 0       | Planctomycetes        | S               | TIGRFAM type<br>VI secretion<br>protein,                                  | -            | -        |
|               |      |          |         | <b>.</b>              |                 | VC_A0111 family   |              |          |
| k123_4233415_ |      | 4.708    | 0       | Bacteria              | -               | -   | -            | -        |
| k123_5247798_ | _1   | 4.101    | 0       | Bacteria              | CO              | amine<br>dehydrogenase<br>activity  | -            | -        |
| k123_1022852_ | _1   | 3.340    | 0       | Deltaproteobacteria   | K               | ROS/MUCR<br>transcriptional<br>regulator protein                          | -            | -        |
| k123_4454813_ | _1   | 4.159    | 0       | Bacteria              | D               | protein conserved in cyanobacteria  | -            | -        |
| k123_3793442_ | _4   | 4.229    | 0       | Actinobacteria        | G               | pfkB family<br>carbohydrate<br>kinase                                     | kdgK         | 2.7.1.45 |
| k123_3567914_ | _2   | 3.910    | 0       | Acidobacteria         | -               | -   | -            | _        |
| k123_3575243_ | _1   | 3.586    | 0       | Bacteroidetes         | S               | Protein of<br>unknown function<br>(DUF1501)                               | -            | -        |

| Query                     | Coefficient | p-Value | Taxonomic Annotation | COG<br>Category | Description  | Gene<br>Name | EC      |
|---------------------------|-------------|---------|----------------------|-----------------|--|--------------|---------|
| k123_1456133              | _1 3.496    | 0       | Actinobacteria       | U               | TadE-like protein  | -            | -       |
| k123_1678879              | _1 3.364    | 0       | Actinobacteria       | K               | DNA binding  | -            | _       |
| k123_1448646              | _1 4.154    | 0       | Alphaproteobacteria  | S               | Type VI secretion<br>system effector,<br>Hcp                         | -            | -       |
| k123_2582928              | _1 3.665    | 0       | Cyanobacteria        | T               | Signal<br>Transduction<br>Histidine Kinase                           | -            | -       |
| k123_4454279 <sub>-</sub> | _1 4.115    | 0       | Acidobacteria        | S               | Type VI secretion<br>system effector,<br>Hcp                         | -            | -       |
| k123_2804786              | _1 4.518    | 0       | Gammaproteobacteria  | Т               | Domain present in phytochromes and cGMP-specific phosphodiesterases. | -            | -       |
| k123_1682627              | _1 3.786    | 0       | Bacteroidetes        | S               | Protein of<br>unknown function<br>(DUF1800)                          | -            | -       |
| k123_3238410              | _1 3.678    | 0       | Bacteria             | Т               | phosphoprotein<br>phosphatase<br>activity                            | -            | 6.5.1.3 |

| Query                     | Со | efficient | p-Value | Taxonomic Annotation | COG<br>Category | Description   | Gene<br>Name | EC       |
|---------------------------|----|-----------|---------|----------------------|-----------------|---|--------------|----------|
| k123_2360350              | _2 | 3.954     | 0       | Acidobacteriia       | J               | One of the early assembly proteins it binds 23S rRNA. One of the proteins that surrounds the polypeptide exit tunnel on the outside of the ribosome. Forms the main docking site for trigger factor binding to the ribosome | rplW         | -        |
| k123_244719_              | _2 | 4.077     | 0       | Archaea              | T               | Histidine kinase-,<br>DNA gyrase B-,<br>and HSP90-like<br>ATPase  | -            | 2.7.7.65 |
| k123_1904408              | _4 | 3.615     | 0       | Acidobacteriia       | $\mathbf{S}$    | ASPIC and UnbV  | -            | -        |
| k123_4681706 <sub>-</sub> | _1 | -3.333    | 0       | Fungi                | O               | Belongs to the heat shock protein 70 family   | SSA2         | -        |
| k123_2129273 <sub>.</sub> | _1 | -3.939    | 0       | Fungi                | O               | Belongs to the heat shock protein 70 family   | SSA2         | -        |
| k123_2784358              | _1 | -4.226    | 0       | Fungi                | O               | Heat shock<br>protein   | HSP82        | -        |

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| Query        | Coefficient | p-Value | Taxonomic Annotation  | COG<br>Category | Description  | Gene<br>Name | EC               |
|--------------|-------------|---------|-----------------------|-----------------|--|--------------|------------------|
| k123_2453574 |             | 0       | Xanthomonadales Fungi | O               | Part of a stress-induced multi-chaperone system, it is involved in the recovery of the cell from heat-induced damage, in cooperation with DnaK, DnaJ and GrpE MreB/Mbl protein | clpB         | -                |
| k123_784972_ | _1 -3.361   | 0       | Fungi                 | О               | Belongs to the<br>heat shock<br>protein 70 family  | SSA2         | -                |
| k123_5131902 | _2 -3.945   | 0       | Bacteria              | N               | domain, Protein  | -            | 3.2.1.81,3.2.1.9 |

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| Query                        | Coefficient | p-Value | Taxonomic Annotation             | COG<br>Category | Description   | Gene<br>Name | EC |
|------------------------------|-------------|---------|----------------------------------|-----------------|---|--------------|----|
| k123_1897324                 | _1 -2.804   | 0       | Xanthomonadales                  | O               | Part of a stress-induced multi-chaperone system, it is involved in the recovery of the cell from heat-induced damage, in cooperation with DnaK, DnaJ and GrpE | clpB         | -  |
| k123_1007534<br>k123_1119824 |             | 0       | Actinobacteria<br>Actinobacteria | C<br>O          | Ferredoxin<br>Heat shock 70<br>kDa protein  | fdxA<br>dnaK | -  |
| k123_3696373                 | _1 -2.768   | 0       | Actinobacteria                   | O               | Prevents misfolding and promotes the refolding and proper assembly of unfolded polypeptides generated under stress conditions                                 | groL2        | -  |
| k123_3023451                 | _1 -5.388   | 0       | Fungi                            | O               | Hsp90 protein   | HSP82        | -  |

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| 10 | Divition  |  |

| Query                                     | Coefficient | p-Value | Taxonomic Annotation                  | COG<br>Category | Description  | Gene<br>Name  | EC                   |
|---|-------------|---------|---------------------------------------|-----------------|--|---------------|----------------------|
| k123_101824<br>k123_414234<br>k123_391488 | 4_1 -3.376  | 0 0 0   | Xanthomonadales  Fungi Actinobacteria | O<br>O<br>G     | Part of a stress-induced multi-chaperone system, it is involved in the recovery of the cell from heat-induced damage, in cooperation with DnaK, DnaJ and GrpE Hsp90 protein Phosphoketolase    | clpB<br>HSP82 | -<br>4.1.2.22,4.1.2. |
| k123_31221                                | 0_1 -4.387  | 0       | Fungi                                 | О               | Heat shock<br>protein  | HSP82         | -                    |
| k123_402738                               | 1_1 -1.934  | 0       | Actinobacteria                        | O               | Part of a<br>stress-induced<br>multi-chaperone<br>system, it is<br>involved in the<br>recovery of the<br>cell from<br>heat-induced<br>damage, in<br>cooperation with<br>DnaK, DnaJ and<br>GrpE | clpB          | -                    |

| Query         | Coefficient | p-Value | Taxonomic Annotation | COG<br>Category | Description                                       | Gene<br>Name | EC |
|---------------|-------------|---------|----------------------|-----------------|---|--------------|----|
| k123_572232_  | 1 -2.504    | 0       | Actinobacteria       | С               | 4Fe-4S dicluster domain                           | fadF         | -  |
| k123_559972_  | 1 -2.527    | 0       | Actinobacteria       | О               | Heat shock 70<br>kDa protein                      | dnaK         | -  |
| k123_2243380_ | _1 -4.837   | 0       | Fungi                | O               | Belongs to the<br>heat shock<br>protein 70 family | SSA2         | -  |

Supplementary Table 3. Top 10 most up- and down-regulated genes, determined by log2 fold change and adjusted p-values, for each sequential timepoint comparison. Positive log2 fold change values represent genes whose expression was higher in the later decomposition timepoint soils, while negative log2 fold change values are higher in earlier decomposition timepoint soils. Taxonomic annotation, COG categories, gene names, and EC were assigned via eggNOG-mapper. The comparison column distinguishes each timepoint comparison.

| Query          | Comparison Coeffici | ent p-Value | Taxonomic<br>Annotation | COG<br>Category | Description   | Gene<br>Name | EC        |
|----------------|---------------------|-------------|-------------------------|-----------------|---|--------------|-----------|
| k123_3351426_1 | 0-12 11.4           | 10 < 0.001  | Clostridia              | S               | Flavin reductase like domain  | -            | -         |
| k123 3689959 6 | 0-12 12.8           | 63 < 0.001  | Bacilli                 | GM              | NmrA-like family  | -            | -         |
| k123_1682409_1 | 0-12 12.8           | 14 < 0.001  | Bacilli                 | C               | Oxidoreductase required for the transfer of electrons from pyruvate to flavodoxin | nifJ         | 1.2.7.1   |
| k123_4131385_2 | 0-12 11.7           | 68 < 0.001  | Bacilli                 | M               | Domain of<br>unknown function<br>(DUF5011)  | -            | -         |
| k123_3474748_6 | 0-12 11.5           | 48 < 0.001  | Bacilli                 | E               | ABC transporter substrate-binding protein   | oppA         | -         |
| k123_2018420_1 | 0-12 11.3           | 26 < 0.001  | Bacilli                 | G               | PTS fructose<br>transporter<br>subunit IIC  | fruA         | 2.7.1.202 |
| k123_4134106_1 | 0-12 10.7           | 76 < 0.001  | Bacilli                 | С               | e3 binding<br>domain  | pdhC         | 2.3.1.12  |
| k123_2451258_1 | 0-12 12.5           | 72 < 0.001  | Clostridia              | M               | S-layer homology<br>domain  | -            | -         |
| k123_3683359_3 | 0-12 12.8           | 58 < 0.001  | Bacilli                 | P               | P-type ATPase   | copA         | 3.6.3.54  |
| k123_2018420_3 | 0-12 9.9            | 98 < 0.001  | Bacilli                 | GKT             | Transcriptional antiterminator  | manR         | -         |

| Query          | Comparison Coefficient | p-Value | Taxonomic<br>Annotation | COG<br>Category | Description   | Gene<br>Name | EC         |
|----------------|------------------------|---------|-------------------------|-----------------|---|--------------|------------|
| k123_4030449_2 | 0-12 -7.201            | < 0.001 | Gammaproteobacteria     | С               | PFAM blue (type 1) copper domain protein              | -            | -          |
| k123_1903136_2 | 0-12 -6.769            | < 0.001 | Actinobacteria          | Q               | Taurine catabolism dioxygenase TauD, TfdA family      | -            | 1.14.11.17 |
| k123_4017556_1 | 0-12 -6.740            | < 0.001 | Betaproteobacteria      | Q               | Taurine<br>dioxygenase                                | -            | 1.14.11.17 |
| k123_2470867_1 | 0-12 -6.354            | < 0.001 | Actinobacteria          | Q               | taurine<br>catabolism<br>dioxygenase                  | tauD         | 1.14.11.17 |
| k123_2231885_1 | 0-12 -5.426            | < 0.001 | Actinobacteria          | Е               | ABC-type<br>branched-chain<br>amino acid<br>transport | -            | -          |
| k123_1903136_1 | 0-12 -5.513            | < 0.001 | Actinobacteria          | K               | Bacterial regulatory proteins, tetR family            | -            | -          |
| k123_3697194_1 | 0-12 -5.446            | < 0.001 | Actinobacteria          | Q               | Taurine catabolism dioxygenase TauD, TfdA family      | tauD         | 1.14.11.17 |

| Query          | Comparison Coefficie | nt p-Value | Taxonomic<br>Annotation | COG<br>Category | Description  | Gene<br>Name | EC       |
|----------------|----------------------|------------|-------------------------|-----------------|--|--------------|----------|
| k123_4238273_1 | 0-12 -6.10           | 6 < 0.001  | Deltaproteobacteria     | S               | Part of the outer<br>membrane protein<br>assembly complex,<br>which is involved<br>in assembly and<br>insertion of<br>beta-barrel<br>proteins into the<br>outer membrane | -            | -        |
| k123_2680211_1 | 0-12 -5.15           | 5 < 0.001  | Proteobacteria          | Q               | Multicopper oxidase  | -            | 1.16.3.3 |
| k123_4688925_2 | 0-12 -6.01           | 4 < 0.001  | Betaproteobacteria      | С               | D-arabinono-1,4-<br>lactone oxidase  | -            | -        |
| k123_785677_1  | 12-58 12.10          | 6 < 0.001  | Betaproteobacteria      | J               | One of the primary rRNA binding proteins, it binds directly near the 3'-end of the 23S rRNA, where it nucleates assembly of the 50S subunit                              | rplC         | -        |
| k123_2456034_2 | 12-58 11.22          | 6 < 0.001  | Betaproteobacteria      | J               | Forms part of the polypeptide exit tunnel  | rplD         | -        |
| k123_3024550_1 | 12-58 10.81          | 5 < 0.001  | Betaproteobacteria      | J               | This is one of the proteins that binds to the 5S RNA in the ribosome where it forms part of the central protuberance   | ctc          | -        |

| Query          | Comparison Coefficient | p-Value | Taxonomic<br>Annotation | COG<br>Category | Description   | Gene<br>Name          | EC |
|----------------|------------------------|---------|-------------------------|-----------------|---|-----------------------|----|
| k123_1916443_2 | 12-58 11.317           | < 0.001 | Betaproteobacteria      | J               | Binds directly to 23S ribosomal RNA and is necessary for the in vitro assembly process of the 50S ribosomal subunit. It is not involved in the protein synthesizing functions of that subunit   | rplT                  | -  |
| k123_794397_5  | 12-58 12.046           | < 0.001 | Betaproteobacteria      | J               | Ribosomal<br>protein L17  | $\operatorname{rpl}Q$ | -  |
| k123_1010408_1 | 12-58 11.827           | < 0.001 | Betaproteobacteria      | J               | This is 1 of the proteins that binds and probably mediates the attachment of the 5S RNA into the large ribosomal subunit, where it forms part of the central protuberance. In the 70S ribosome it contacts protein S13 of the 30S subunit (bridge B1b), connecting the 2 subunits | rplE                  | -  |

| Query          | Comparison Coeffic | cient p-Value | Taxonomic<br>Annotation | COG<br>Category | Description                             | Gene<br>Name | EC |
|----------------|--------------------|---------------|-------------------------|-----------------|---|--------------|----|
| k123_1115623_7 | 12-58 10           | .795 < 0.001  | Betaproteobacteria      | JKL             | Belongs to the DEAD box helicase family | rhlE1        | -  |
| k123_3802697_1 | 12-58 10           | .705 < 0.001  | Betaproteobacteria      | JKL             | Belongs to the DEAD box helicase family | rhlE1        | -  |
| k123_5130820_1 | 12-58 11           | .586 < 0.001  | Betaproteobacteria      | JKL             | Belongs to the DEAD box helicase family | rhlE1        | -  |

| Query          | Comparison | Coefficient | p-Value | Taxonomic<br>Annotation | COG<br>Category | Description  | Gene<br>Name | EC |
|----------------|------------|-------------|---------|-------------------------|-----------------|--|--------------|----|
| k123_2572977_1 | 12-58      | 11.374      | < 0.001 | Betaproteobacteria      | U               | The central subunit of the protein translocation channel SecYEG. Consists of two halves formed by TMs 1-5 and 6-10. These two domains form a lateral gate at the front which open onto the bilayer between TMs 2 and 7, and are clamped together by SecE at the back. The channel is closed by both a pore ring composed of hydrophobic SecY resides and a short helix (helix 2A) on the extracellular side of the membrane which forms a plug. The plug probably moves laterally to allow the channel to open. The ring and the pore may move independently | secY         |    |

| Query          | Comparison C | coefficient | p-Value | Taxonomic<br>Annotation | COG<br>Category | Description   | Gene<br>Name | EC              |
|----------------|--------------|-------------|---------|-------------------------|-----------------|---|--------------|-----------------|
| k123_3351426_1 | 12-58        | -10.845     | < 0.001 | Clostridia              | S               | Flavin reductase like domain  | -            | -               |
| k123_3689959_6 | 12-58        | -12.298     | < 0.001 | Bacilli                 | GM              | NmrA-like family  | _            | -               |
| k123_4131385_2 | 12-58        | -11.203     | < 0.001 | Bacilli                 | M               | Domain of<br>unknown function<br>(DUF5011)  | -            | -               |
| k123_1682409_1 | 12-58        | -11.434     | < 0.001 | Bacilli                 | C               | Oxidoreductase required for the transfer of electrons from pyruvate to flavodoxin | nifJ         | 1.2.7.1         |
| k123_3028891_1 | 12-58        | -10.738     | < 0.001 | Clostridia              | K               | PFAM flavin reductase   | -            | -               |
| k123_2018420_3 | 12-58        | -9.433      | < 0.001 | Bacilli                 | GKT             | Transcriptional antiterminator  | manR         | -               |
| k123_3469192_2 | 12-58        | -9.634      | < 0.001 | Clostridia              | Т               | S-layer homology<br>domain  | -            | -               |
| k123_3683359_3 | 12-58        | -11.778     | < 0.001 | Bacilli                 | P               | P-type ATPase   | copA         | 3.6.3.54        |
| k123_335030_2  | 12-58        | -11.417     | < 0.001 | Bacilli                 | С               | Iron-containing<br>alcohol<br>dehydrogenase                                       | adhB         | 1.1.1.1,1.1.1.2 |
| k123_3012292_1 | 12-58        | -9.296      | < 0.001 | Bacilli                 | S               | Putative neutral zinc metallopeptidase  | yugP         | -               |
| k123 3581297 1 | 58-86        | 8.049       | < 0.001 | Fungi                   | -               | -   | _            | -               |
| k123_133381_1  | 58-86        | 8.362       | < 0.001 | Betaproteobacteria      | $\mathbf{C}$    | anaerobic<br>respiration  | -            | 1.7.2.6         |
| k123_3681927_1 | 58-86        | 8.474       | < 0.001 | Betaproteobacteria      | С               | Cytochrome C<br>oxidase subunit II,<br>periplasmic<br>domain                      | -            | 1.9.3.1         |

| Query          | Comparison | Coefficient | p-Value | Taxonomic<br>Annotation | COG<br>Category | Description   | Gene<br>Name | EC      |
|----------------|------------|-------------|---------|-------------------------|-----------------|---|--------------|---------|
| k123_4461993_6 | 58-86      | 10.379      | < 0.001 | Bacteroidetes           | S               | Tetratricopeptide repeat protein  | -            | -       |
| k123_1691746_1 | 58-86      | 8.748       | < 0.001 | Rhodospirillales        | C               | NDH-1 shuttles electrons from NADH, via FMN and iron- sulfur (Fe-S) centers, to quinones in the respiratory chain. The immediate electron acceptor for the enzyme in this species is believed to be ubiquinone. Couples the redox reaction to proton translocation (for every two electrons transferred, four hydrogen ions are translocated across the cytoplasmic membrane), and thus conserves the redox energy in a proton gradient. This subunit may bind ubiquinone | nuoH         | 1.6.5.3 |

| Query          | Comparison Coefficient | p-Value | Taxonomic<br>Annotation | COG<br>Category | Description  | Gene<br>Name | EC       |
|----------------|------------------------|---------|-------------------------|-----------------|--|--------------|----------|
| k123_1674520_1 | 58-86 8.655            | < 0.001 | Alphaproteobacteria     | С               | Produces ATP from ADP in the presence of a proton gradient across the membrane. The alpha chain is a regulatory subunit  | atpA         | 3.6.3.14 |
| k123_4907595_1 | 58-86 8.469            | < 0.001 | Oceanospirillales       | J               | Belongs to the<br>universal<br>ribosomal protein<br>uS2 family   | rpsB         | -        |
| k123_788629_1  | 58-86 8.018            | < 0.001 | Bacteroidetes           | PT              | COGs COG3712<br>Fe2 -dicitrate<br>sensor membrane<br>component   | -            | -        |
| k123_784141_1  | 58-86 6.602            |         | Bacteroidetes           | -               | -  | -            | -        |
| k123_2678388_2 | 58-86 7.966            | < 0.001 | Betaproteobacteria      | C               | Subunits I and II form the functional core of the enzyme complex. Electrons originating in cytochrome c are transferred via heme a and Cu(A) to the binuclear center formed by heme a3 and Cu(B) | $\cos B$     | 1.9.3.1  |

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| Query          | Comparison C | Coefficient | p-Value | Taxonomic<br>Annotation | COG<br>Category | Description   | Gene<br>Name             | EC |
|----------------|--------------|-------------|---------|-------------------------|-----------------|---|--------------------------|----|
| k123 903075 1  | 58-86        | -7.906      | < 0.001 | Bacteroidetes           | _               | -   | -                        | _  |
| k123_2460567_2 | 58-86        | -7.060      | < 0.001 | Bacteroidetes           | S               | Functions as a peptidoglycan terminase that cleaves nascent peptidoglycan strands endolytically to terminate their elongation | $\mathrm{mlt}\mathrm{G}$ | -  |
| k123 680086 2  | 58-86        | -6.939      | < 0.001 | Bacteria                | _               | -   | _                        | -  |
| k123_3353720_2 | 58-86        | -7.036      | < 0.001 | Bacteroidetes           | U               | Preprotein<br>translocase SecG<br>subunit   | $\sec G$                 | -  |
| k123_4342821_1 | 58-86        | -6.571      | < 0.001 | Bacteroidetes           | CO              | Redoxin   | -                        | -  |
| k123_680086_6  | 58-86        | -6.695      | < 0.001 | Bacteroidetes           | G               | Sad1 / UNC-like<br>C-terminal   | -                        | -  |
| k123_3687116_4 | 58-86        | -6.677      | < 0.001 | Bacteroidetes           | S               | Belongs to the pirin family   | yhhW                     | -  |

| Query          | Comparison | Coefficient | p-Value | Taxonomic<br>Annotation | COG<br>Category | Description  | Gene<br>Name | EC        |
|----------------|------------|-------------|---------|-------------------------|-----------------|--|--------------|-----------|
| k123_349053_1  | 58-86      | -6.992      | < 0.001 | Bacteroidetes           | F               | Catalyzes the conversion of inosine 5'-phosphate (IMP) to xanthosine 5'-phosphate (XMP), the first committed and rate- limiting step in the de novo synthesis of guanine nucleotides, and therefore plays an important role in the regulation of cell growth | guaB         | 1.1.1.205 |
| k123_2348950_2 | 58-86      | -6.358      | < 0.001 | Bacteroidetes           | О               | Heat shock<br>protein  | -            | -         |
| k123_1004837_2 | 58-86      | -7.162      | < 0.001 | Bacteroidetes           | S               | Fimbrillin-like  | -            | -         |
| k123_465331_1  | 86-168     | 6.832       | < 0.001 | Bacteroidetes           | M               | Pkd domain containing protein  | -            | -         |
| k123_1247591_2 | 86-168     | 6.662       | < 0.001 | Xanthomonadales         | -               | -  | -            | -         |
| k123_356907_3  | 86-168     | 7.008       | < 0.001 | Actinobacteria          | I               | Fatty acid<br>desaturase   | $desA3\_2$   | -         |
| k123_1802620_3 | 86-168     | 6.775       | < 0.001 | Actinobacteria          | S               | Protein of<br>unknown function<br>(DUF1360)  | -            | -         |
| k123_4137171_1 | 86-168     | 7.254       | < 0.001 | Actinobacteria          | U               | WD40-like Beta<br>Propeller Repeat   | -            | -         |
| k123_1449660_1 | 86-168     | 7.112       | < 0.001 | Actinobacteria          | $\mathbf{S}$    | LVIVD repeat   | -            | -         |

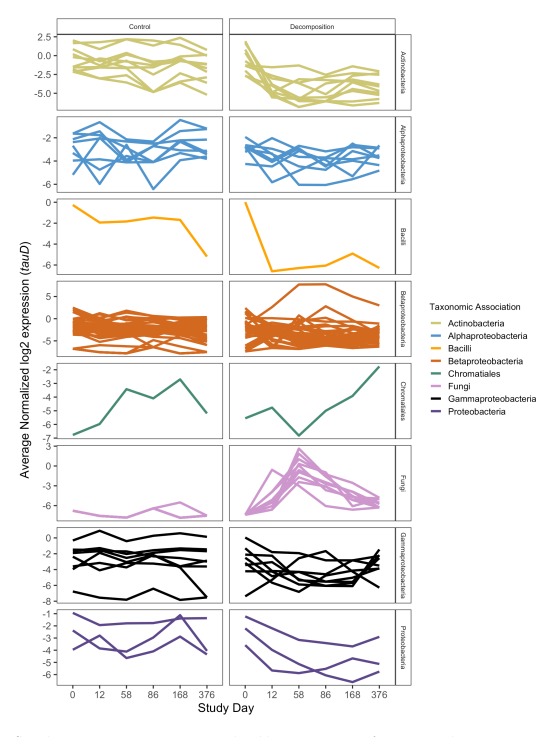
| Query                            | Comparison       | Coefficient    | p-Value | Taxonomic<br>Annotation | COG<br>Category | Description  | Gene<br>Name | EC      |
|----------------------------------|------------------|----------------|---------|-------------------------|-----------------|--|--------------|---------|
| k123_3015796_1                   | 86-168           | 6.131          | < 0.001 | Actinobacteria          | S               | Tetratricopeptide repeat   | -            | -       |
| k123_4469568_1                   | 86-168           | 8.516          | < 0.001 | Gammaproteobacteria     | NU              | Pilin (bacterial filament)   | pilA         | -       |
| k123_4013561_1<br>k123_4136401_2 | 86-168<br>86-168 | 6.723<br>6.097 | < 0.001 | Xanthomonadales         | N<br>O          | Required for morphogenesis and for the elongation of the flagellar filament by facilitating polymerization of the flagellin monomers at the tip of growing filament. Forms a capping structure, which prevents flagellin subunits (transported through the central channel of the flagellum) from leaking out without polymerization at the distal end FKBP-type peptidyl-prolyl | fliD         | 5.2.1.8 |
|                                  |                  |                |         |                         |                 | cis-trans<br>isomerase   |              |         |
| k123_2576445_1                   | 86-168           | -7.806         | < 0.001 | Xanthomonadales         | M               | Belongs to the ompA family   | mopB         | -       |

| Query          | Comparison Coefficie | nt p-Value | Taxonomic<br>Annotation | COG<br>Category | Description   | Gene<br>Name           | EC             |
|----------------|----------------------|------------|-------------------------|-----------------|---|------------------------|----------------|
| k123_2013370_2 | 86-168 -7.4          | 09 < 0.001 | Gammaproteobacteria     | S               | Protein of<br>unknown function<br>(DUF1471)   | -                      | -              |
| k123_1784040_1 | 86-168 -7.5          | 39 < 0.001 | Clostridia              | L               | PFAM Reverse<br>transcriptase<br>(RNA-dependent<br>DNA polymerase)  | -                      | -              |
| k123_2784492_8 | 86-168 -7.10         | 0.001      | Actinobacteria          | Q               | Amidohydrolase family   | -                      | -              |
| k123_1671881_1 | 86-168 -7.7          | .2 < 0.001 | Bacteroidetes           | P               | von Willebrand factor, type A   | -                      | -              |
| k123_115662_2  | 86-168 -8.99         | 27 < 0.001 | Bacilli                 | С               | succinate<br>dehydrogenase  | $\operatorname{sdh} A$ | 1.3.5.1,1.3.5. |
| k123_2013542_1 | 86-168 -7.26         | 36 < 0.001 | Betaproteobacteria      | S               | Domain of<br>unknown function<br>(DUF4148)  | -                      | -              |
| k123_784141_1  | 86-168 -6.4          | 0.001      | Bacteroidetes           | _               | -   | _                      | -              |
| k123_126934_1  | 86-168 -8.8          | 76 < 0.001 | Bacteroidetes           | M               | Belongs to the<br>membrane fusion<br>protein (MFP)<br>(TC 8.A.1) family   | -                      | -              |
| k123_3912958_2 | 86-168 -7.09         | 0.001      | Bacteroidetes           | С               | cytochrome c<br>oxidase   | ccoG                   | -              |
| k123 4347668 1 | 168-376 7.99         | 35 < 0.001 | Archaea                 | _               | -   | _                      | -              |
| k123_2010423_2 | 168-376 8.4          | 57 < 0.001 | Acidobacteria           | J               | Forms part of the ribosomal stalk, playing a central role in the interaction of the ribosome with GTP-bound translation factors | rplJ                   | -              |

| Query                            | Comparison Coeffic | cient p-Val                | ne Taxonomic Annotation | $\begin{array}{c} \operatorname{COG} \\ \operatorname{Category} \end{array}$ | Description  | Gene<br>Name          | EC      |
|----------------------------------|--------------------|----------------------------|-------------------------|--|--|-----------------------|---------|
| k123_4021403_1                   | 168-376 6          | 6.597 < 0.0                | 01 Acidobacteria        | O  | Prevents misfolding and promotes the refolding and proper assembly of unfolded polypeptides generated under stress conditions                              | groL                  | -       |
| k123_2899693_1                   | 168-376 6          | 6.546 < 0.0                | 01 Acidobacteria        | О  | Belongs to the<br>ClpA ClpB family   | -                     | -       |
| $k123\_461276\_1$                | 168-376 9          | 0.975 < 0.0                | 01 Bacteria             | N  | domain, Protein  | -                     | -       |
| k123_13363_1                     | 168-376 6          | 5.773 < 0.0                | 01 Acidobacteria        | O  | Prevents misfolding and promotes the refolding and proper assembly of unfolded polypeptides generated under stress conditions                              | $\operatorname{groL}$ | -       |
| k123_3585893_1                   | 168-376 6          | 3.944 < 0.0                | 01 Acidobacteria        | K  | DNA-dependent<br>RNA polymerase<br>catalyzes the<br>transcription of<br>DNA into RNA<br>using the four<br>ribonucleoside<br>triphosphates as<br>substrates | rpoC                  | 2.7.7.6 |
| k123_3020221_1<br>k123_4015238_2 |                    | 7.640 < 0.0<br>8.717 < 0.0 |                         | -<br>U   | -<br>MotA TolQ ExbB<br>proton channel  | -                     | -       |

| Query              | Comparison Coefficient | p-Value | Taxonomic<br>Annotation | COG<br>Category | Description  | Gene<br>Name | EC        |
|--------------------|------------------------|---------|-------------------------|-----------------|--|--------------|-----------|
| k123_4017681_1     | 168-376 8.220          | < 0.001 | Acidobacteria           | J               | Binds the lower<br>part of the 30S<br>subunit head.<br>Binds mRNA in<br>the 70S ribosome,<br>positioning it for<br>translation | rpsC         | -         |
| k123_465331_1      | 168-376 -7.066         | < 0.001 | Bacteroidetes           | M               | Pkd domain containing protein  | -            | -         |
| k123_4458935_1     | 168-376 -7.201         | < 0.001 | Epsilonproteobacteria   | Q               | calcium- and calmodulin- responsive adenylate cyclase activity   | -            | -         |
| k123_5025476_3     | 168-376 -8.470         | < 0.001 | Actinobacteria          | G               | phosphocarrier<br>protein hpr  | ptsH         | -         |
| k123_1114482_1     | 168-376 -6.397         | < 0.001 | Bacteroidetes           | MU              | Outer membrane efflux protein  | -            | -         |
| $k123\_4360823\_3$ | 168-376 -6.013         | < 0.001 | Gamma proteobacteria    | IQ              | KR domain  | -            | 1.1.1.159 |
| k123_3135574_1     | 168-376 -7.251         | < 0.001 | Actinobacteria          | K               | transcriptional<br>regulator (DeoR<br>family)  | fruR         | -         |
| k123_3354064_4     | 168-376 -6.628         | < 0.001 | Actinobacteria          | G               | Bacterial extracellular solute-binding protein   | -            | -         |
| k123_2683111_1     | 168-376 -6.487         | < 0.001 | Actinobacteria          | G               | L-arabinose<br>isomerase   | -            | 5.3.1.4   |
| k123_4463835_2     | 168-376 -6.188         | < 0.001 | Actinobacteria          | K               | DNA-templated transcription, initiation  | -            | -         |

| Query          | Comparison Coefficient | p-Value | Taxonomic<br>Annotation | COG<br>Category | Description                    | Gene<br>Name | EC |
|----------------|------------------------|---------|-------------------------|-----------------|--------------------------------|--------------|----|
| k123_2915236_1 | 168-376 -7.060         | < 0.001 | Actinobacteria          | M               | probably involved in cell wall | -            | -  |



Supplementary Fig 4. 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.