

# Metabolomics Pipeline Overview

## Table of contents

<b>1</b>	<b>Introduction</b>	<b>1</b>
<b>2</b>	<b>Data Distribution</b>	<b>2</b>
<b>3</b>	<b>Sample Analysis — PCA</b>	<b>3</b>
<b>4</b>	<b>Feature Analysis — limma</b>	<b>4</b>
<b>5</b>	<b>Supervised Analysis — sparse PLS-DA</b>	<b>5</b>
<b>6</b>	<b>Putative Annotation</b>	<b>9</b>
<b>7</b>	<b>Conclusion</b>	<b>10</b>

## 1 Introduction

This document describes a basic pipeline for analyzing untargeted LC–MS/MS metabolomics data in R.

The example data consist of panels of untargeted LC–MS features from positive and negative ionization mode runs, similar to metabolites measured from tissue, stool samples, or microbial cultures. The files in `data_raw/metabolomics-pipeline-example.xlsx` and the corresponding positive- and negative-mode .csv exports come from a Thermo Fisher Compound Discoverer v3 workflow and have already undergone feature detection, basic quality control, and signal curation. The values are relative peak intensities for deconvoluted LC–MS features.

These data have been restructured into a hypothetical experimental design with samples assigned to a control group, one of two single-treatment groups (tr1 or tr2), or a dual-treatment group (tr1+tr2) to demonstrate a clear treatment-response pattern.

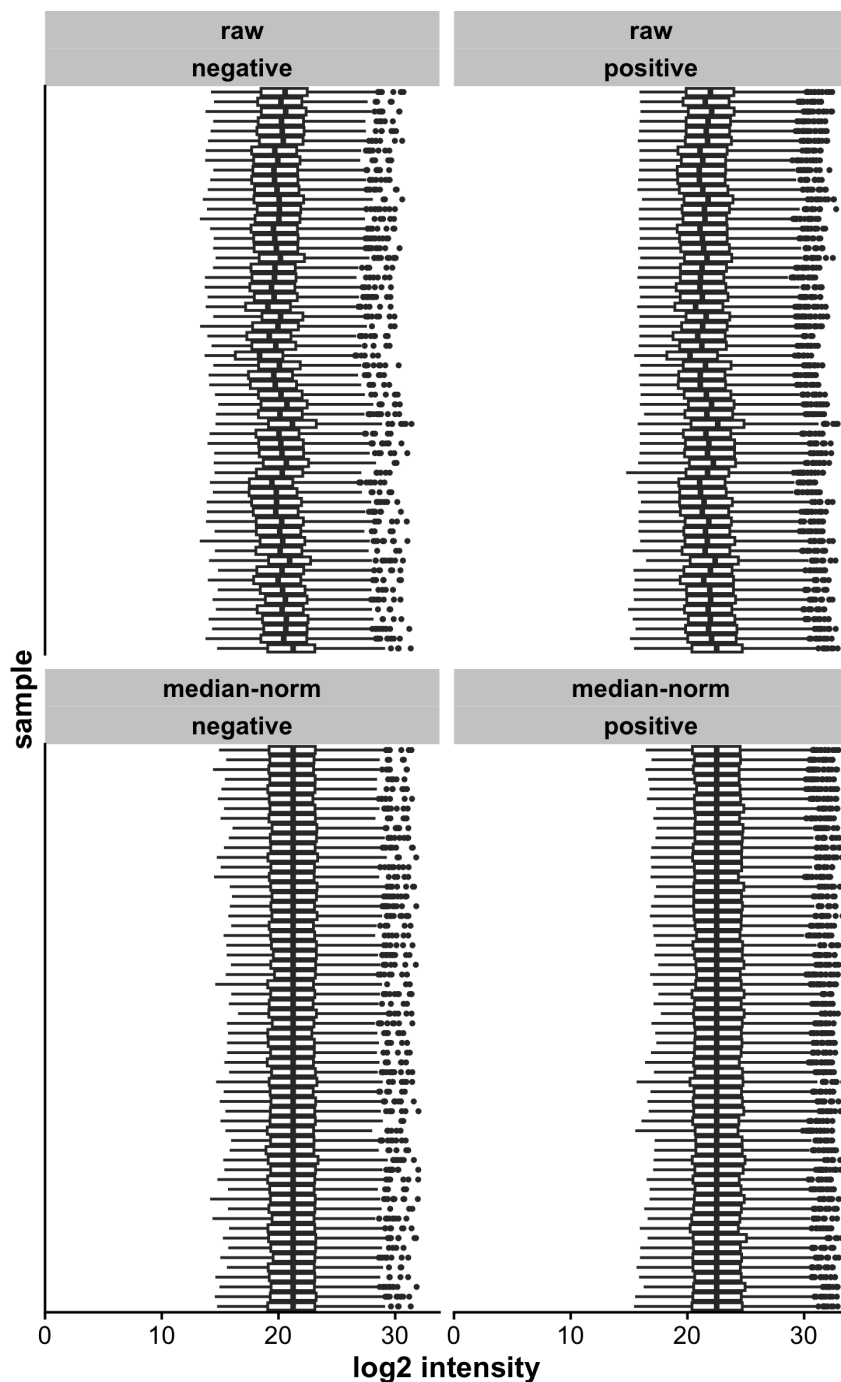
The pipeline illustrates a standard untargeted metabolomics workflow: data cleaning and normalization, unsupervised and supervised multivariate analysis, feature-level differential testing, and putative metabolite annotation using KEGG-based compound information.

All requisite R packages are managed with the renv package (see the `renv/` folder for dependencies). `R/00_setup.r` initializes the environment, loads packages, formats sample metadata, and sets global plotting aesthetics for the figures in this document.

## 2 Data Distribution

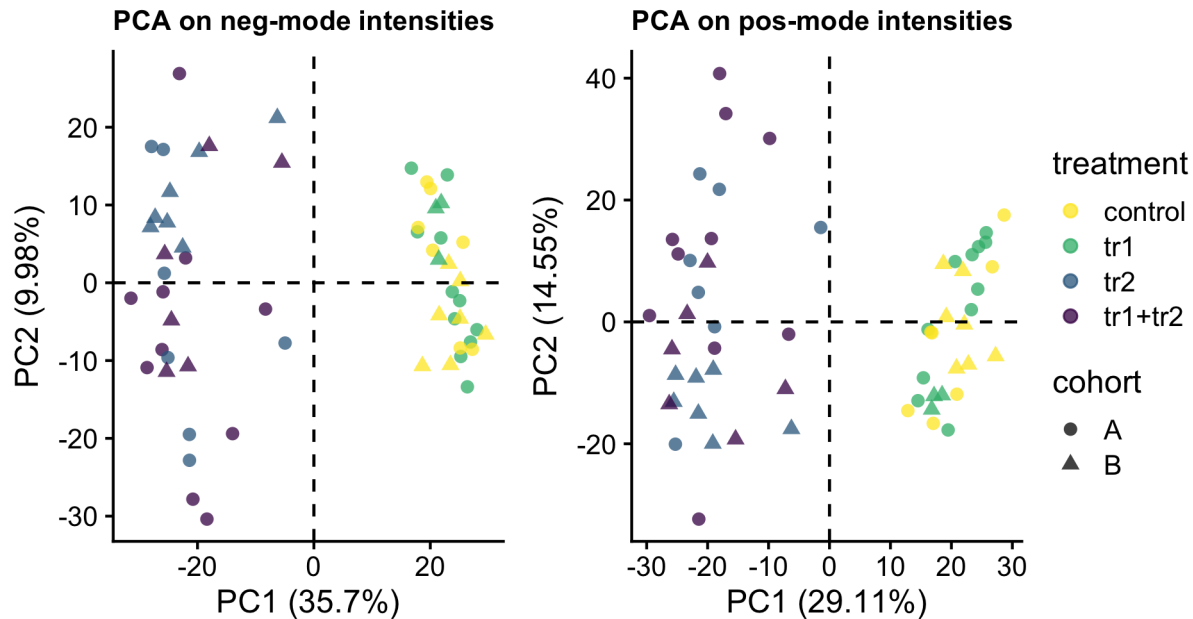
To further clean up the data, 01\_preprocess.r merges LC-MS features with identical mass-to-charge ratios and retention times. It then assigns each feature a unique ID by merging its mass-to-charge ratio and retention time (“mz\_rt\_min”).

As LC-MS intensities of metabolomics data typically skew to the right, data normalization is required to improve comparability across samples and help stabilize variance before actual analysis can begin. The figure demonstrates how median normalization reduces between-sample differences in overall intensity while preserving the underlying variability that’s of interest in the data. The data are also transformed into a log2 scale to reduce domination by extreme values and bring the data closer to normality.



### 3 Sample Analysis — PCA

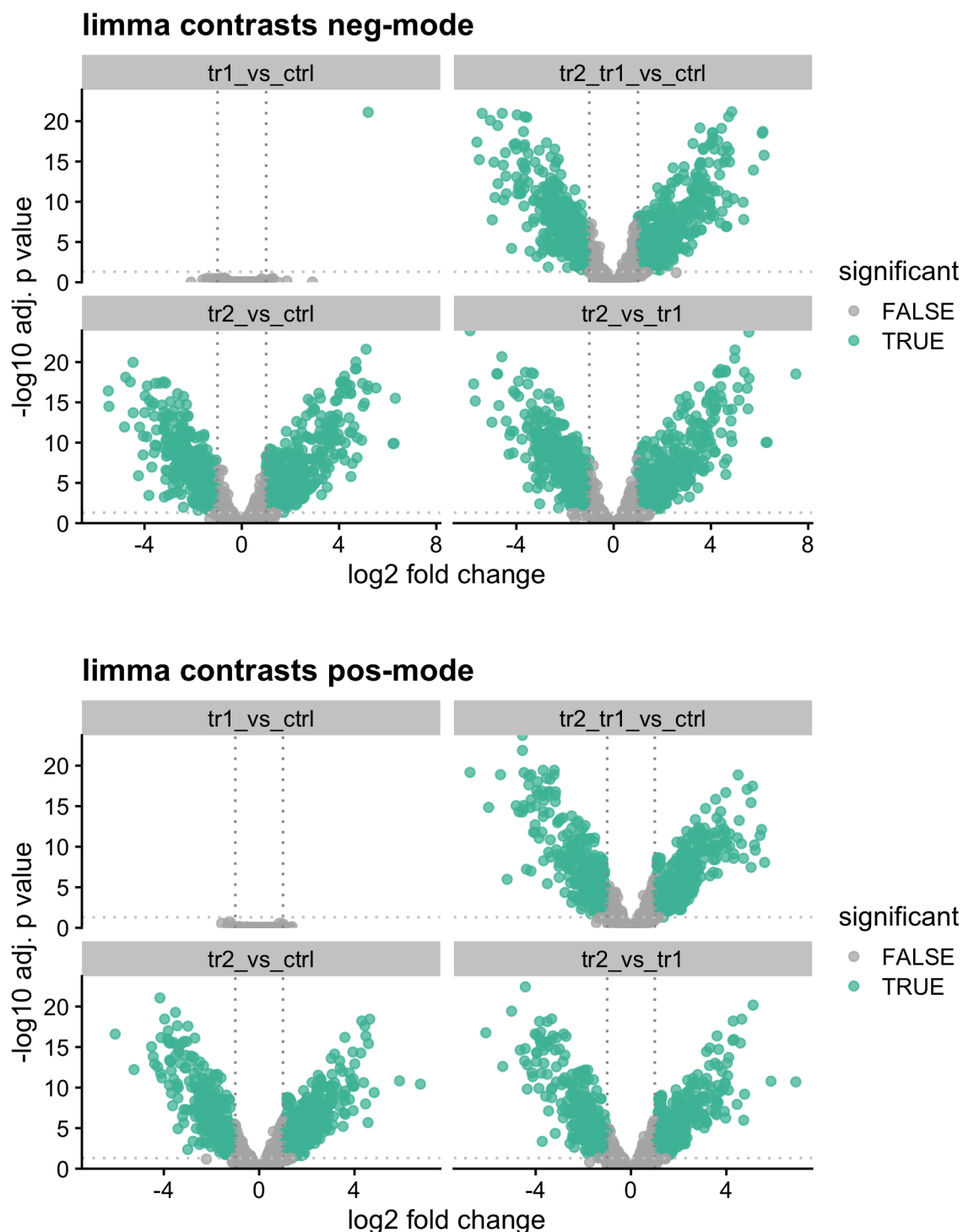
Principal Component Analysis is used to gauge the overall structure of the metabolomic profiles (samples/cases). The median-normalized, log2-transformed intensities are next centered and scaled, then decomposed into a set of orthogonal principal components that capture the dominant sources of variance in the dataset.



Principal component (PC) 1 illustrates a divide amongst the samples based on the treatment group, with controls and tr1 contrasted from tr2 and the dual-treatment (tr1+tr2). PC 1 captures 35.7% of the variance in the data for the negative-mode intensities and 29.11% for the positive-mode intensities, meaning the treatment variable has substantial influence on the overall data structure. PC 2 does not appear to highlight any meaningful secondary structure in the data.

## 4 Feature Analysis — limma

To evaluate the underlying feature fluctuations responsible for the observed sample differences, linear models for microarrays (limma) were applied to the median-normalized, log<sub>2</sub>-transformed intensities. This approach tests each LC-MS feature for differential abundance across treatment groups while controlling the false discovery rate using multiple-testing correction.



The figure depicts the differentially abundant features for each pairwise comparison, highlighting metabolites that most strongly distinguish the control, single-treatment (tr1, tr2), and dual-treatment (tr1+tr2) conditions.

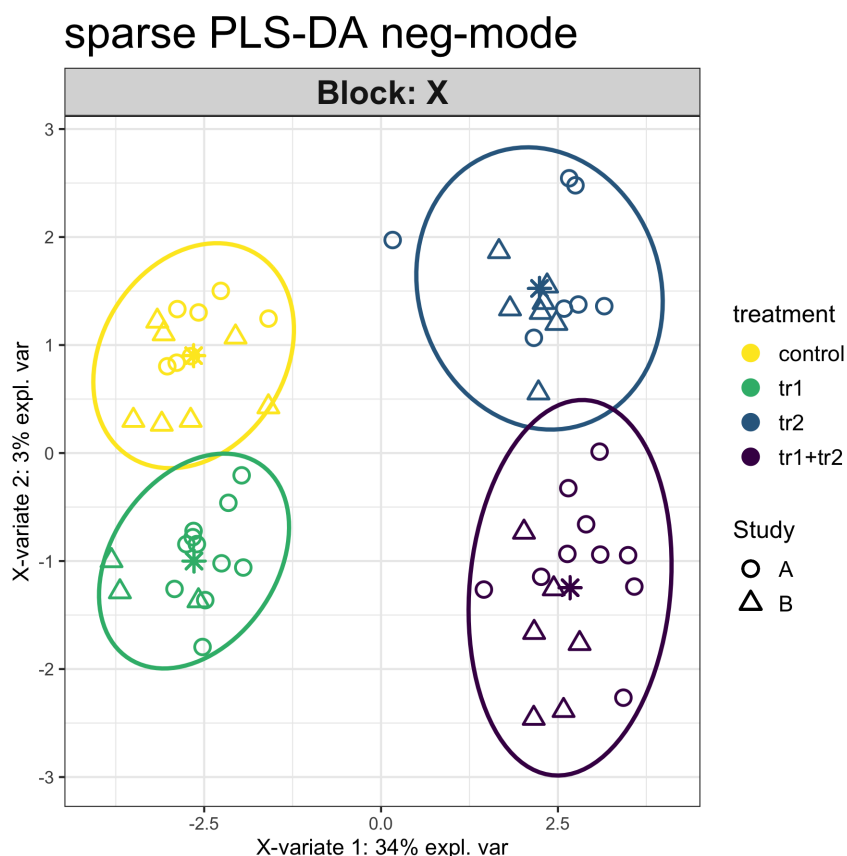
Table 1: limma differential features

Mode	tr1 vs ctrl	tr2 vs ctrl	tr1+tr2 vs ctrl	tr2 vs tr1	total features
negative	1	845	820	825	1499
positive	0	642	643	625	1498

The figure and table results show that treatment 2 induces a pronounced effect in the samples' metabolome with over half of the total features perturbed from the control baseline. Treatment 1 alone has effectively no impact on the metabolome and thus all the differential features between the dual-treatment and controls are likely a consequence of treatment 2.

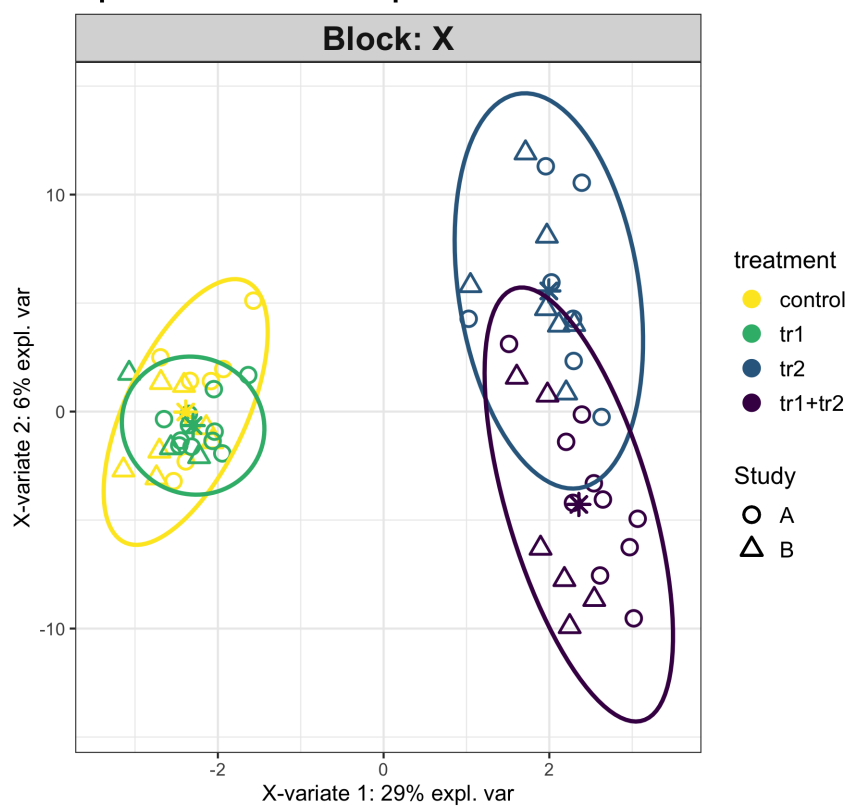
## 5 Supervised Analysis — sparse PLS-DA

Partial least squares discriminant analysis (PLS-DA) is a supervised multivariate method that models the relationship between a set of predictors (the LC-MS features) and a categorical outcome (the treatment groups). In this implementation, a sparse variant (sPLS-DA) is used so that only a subset of variables contributes non-zero loadings to the components.



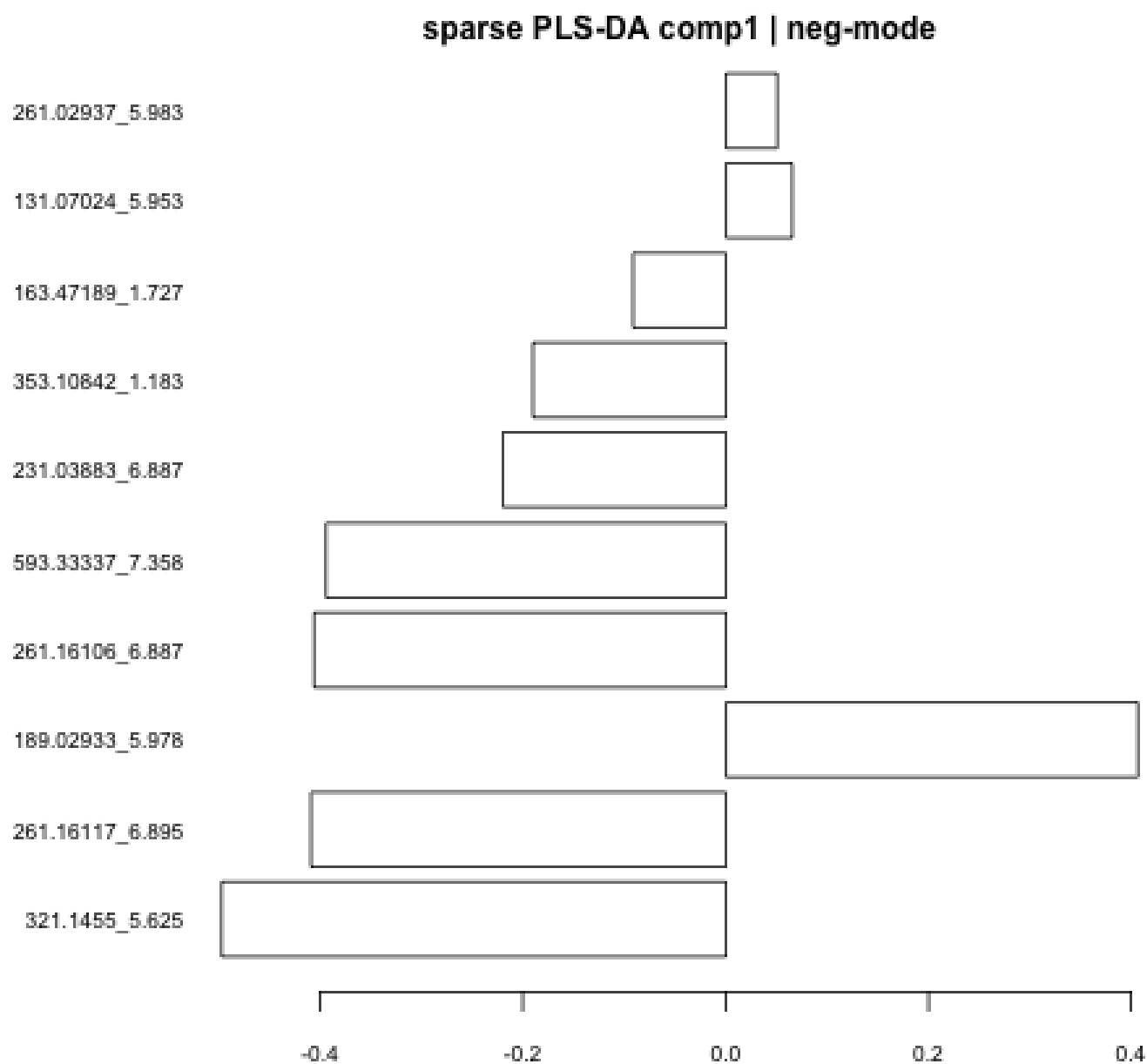
sPLS-DA is applied here to identify a minimal set of discriminative features that best separates the control, single-treatment, and dual-treatment groups in the reduced latent space. The resulting components summarize multivariate treatment effects, while the sparsity constraint highlights candidate metabolites that contribute most strongly to class separation and can be prioritized for downstream annotation.

## sparse PLS-DA pos-mode

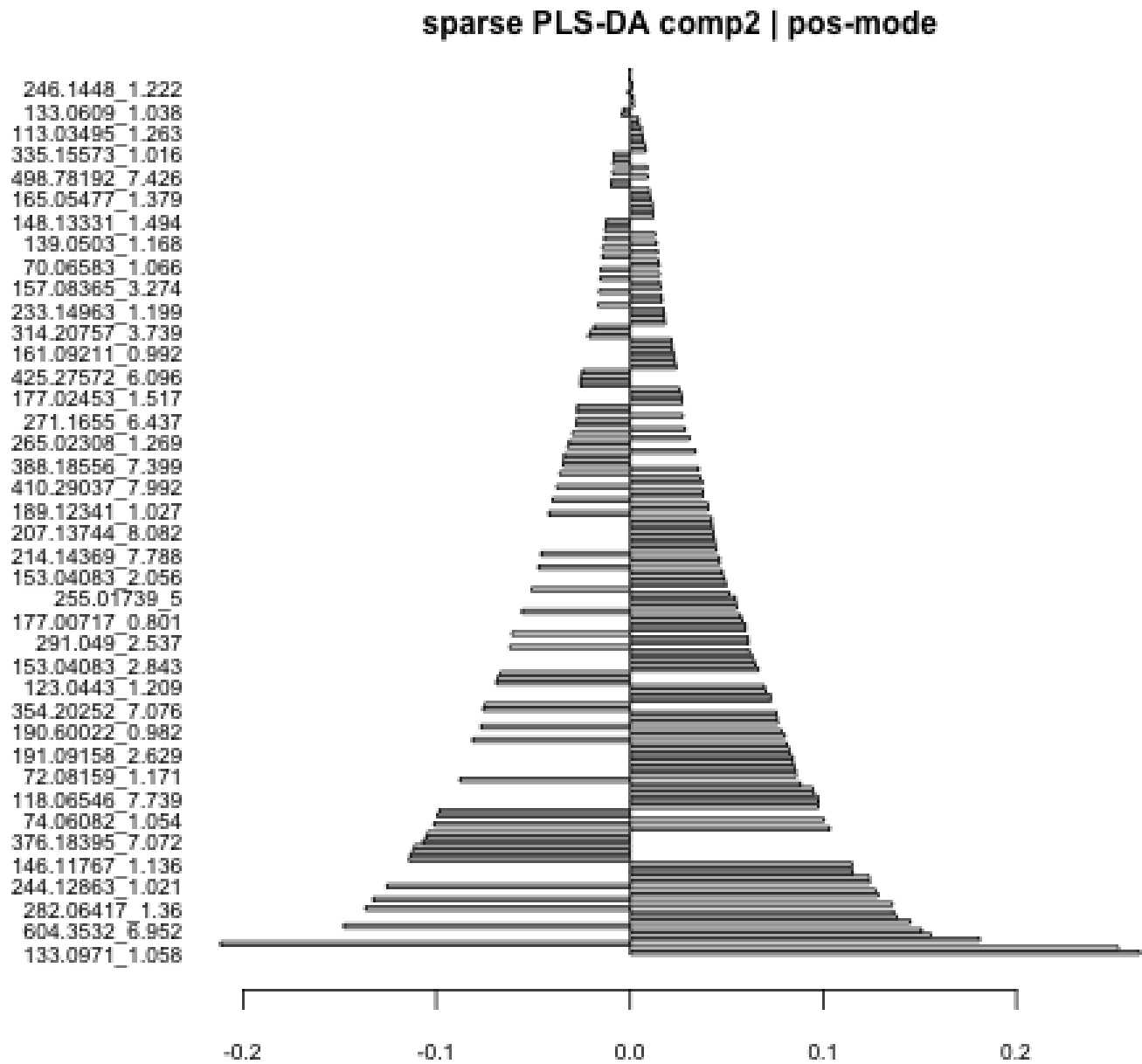


In both ionization modes, the first component emphasizes the contrasts induced by treatment 2, consistent with the unsupervised PCA results. In the negative-mode data, the second component further discriminates the groups but accounts for only 3% of the total variance, so its contribution should be interpreted cautiously.

The LC-MS features driving these separations can be examined via their loadings on each sparse component. For the negative-mode data, both components retain fewer than 50 features out of 1,499, indicating that the variation in this small subset provides an efficient summary of the treatment-related differences.



In the positive-mode data, the first component similarly selects 10 features to distinguish treatment groups. The second component (shown below), by contrast, retains 100 features and shows only weak additional separation between the treatment 2 and dual-treatment samples, consistent with the idea that a larger number of variables is required to represent a relatively subtle effect.





## 6 Putative Annotation

While sample-level analyses are useful for detecting systemic effects on the metabolome, feature-level analyses become more informative once features are placed into a biological context. For untargeted LC–MS data, this requires at least a tentative chemical assignment for each LC–MS peak. Without simultaneous runs against authentic chemical standards, absolute identification and quantification are not possible, but using the measured mass-to-charge ratios and ionization mode, the neutral molecular weight of each analyte can be calculated.

In typical LC–MS/MS workflows, this calculation is performed by vendor software during feature detection and deconvolution, producing neutral molecular weights or exact masses alongside peak intensities. These calculated molecular weights can then be compared against metabolite reference databases, such as the Kyoto Encyclopedia of Genes and Genomes (KEGG), by matching them to the exact masses of known compounds.

In this pipeline, features that were either significantly differentially abundant in the limma analysis or retained by the sparse PLS-DA models were matched to KEGG compounds using their calculated molecular weights. Each feature was compared against the locally cached KEGG compound list, and compounds with exact masses falling within a  $\pm 10$  ppm window were recorded as putative matches. The table below summarizes example features, their candidate KEGG IDs, names, formulas, and pathways. These assignments are considered putative because they are based solely on accurate-mass matching and do not incorporate confirmation against authentic standards.

```
# A tibble: 290 x 5
  mz_rt_min      calculated_mw exact_mass formula name
  <chr>          <dbl>         <dbl> <chr>    <chr>
1 104.03478_0.988      103.          103. C3H5N03 2-Aminomalonate semialdehy~
2 114.05493_1.017      115.          115. C5H9N02 Resolvin E1; 5S,12R,18R-Tr~
3 115.03951_1.818      114.          114. C5H6O3 1,4-Dichlorobenzene; p-Dic~
4 115.03955_1.121      114.          114. C5H6O3 1,4-Dichlorobenzene; p-Dic~
5 116.07098_0.876      115.          115. C5H9N02 Resolvin E1; 5S,12R,18R-Tr~
6 118.08659_1.001      117.          117. C5H11N02 Chlornaphazine; N,N-Bis(2~~
7 118.0866_1.002       100.          100. C5H8O2 5-Valerolactone; delta-Val~
8 118.0866_1.002       100.          100. C5H8O2 Tiglic acid; (E)-2,3-Dimet~
9 118.0866_1.171      117.          117. C5H11N02 Chlornaphazine; N,N-Bis(2~~
10 120.06581_1.056      87.0          87.0 C3H5N02 Galactosylglucosyldiacylg~
# i 280 more rows
```

In a real study of treatment effects on the metabolome, these candidate annotations would be manually vetted to assess their chemical plausibility and biological relevance. Multiple KEGG compounds can share similar exact masses, so a single LC–MS feature can have several potential matches, which complicates interpretation and reinforces the need for additional structural information (e.g., fragmentation spectra, retention time, or standards) before assigning a definitive identity.

## 7 Conclusion

Overall, this pipeline demonstrates a typical workflow for analyzing untargeted LC–MS/MS metabolomics data. Median normalization and log2 transformation are used to stabilize feature variances and place intensities on a comparable scale. PCA provides an overview of global metabolomic patterns and highlights the dominant effect of treatment 2. Differential analysis with limma identifies features most strongly associated with each treatment in pairwise contrasts, and sparse PLS-DA adds a supervised perspective by estimating the minimal set of features needed to classify samples into their respective treatment groups. Finally, differential features prioritized by these analyses are compared against a KEGG compound reference to assemble a set of putative chemical annotations, providing a starting point for biological interpretation and follow-up validation.