# The SDAMS package

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#### **Abstract**

This vignette introduces the use of the Bioconductor package SDAMS, which is designed for differential abundance analysis for metabolomics and proteomics data from mass spectrometry. These data may not be normally distributed and contain a large fraction of zero values. SDAMS considers a two-part semi-parametric mdoel, a logistic regression for the zero proportion and a semi-parametric log-linear model for the non-zero values. A kernel-smoothed likelihood method is proposed to estimate regression coefficients in the two-part model and a likelihood ratio test is constructed for differential abundant analysis.

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#### 1 Citation

The package SDAMS implements statistical methods from the following publication. If you use SDAMS in the published research, please cite:

Yuntong Li, Teresa W.M. Fan, Andrew N. Lane, Woo-Young Kang, Susanne M. Arnold, Arnold J. Stromberg, Chi Wang and Li Chen: A Two-Part Semi-Parametric Model for Metabolomics and Proteomics Data. (Manuscript)

### 2 Quick Start

This section show the most basic SDAMS work flow for a differential abundance analysis for metabolomics and proteomics data:

- 1. Create a MSset object using function createMSsetFromEnvir or createMSsetFromCSV. In this section we use an example MSset directly, which is an object of MSset class named exampleMSset contained in this package.
- 2. Perform a differential abundance analysis using SDA.
- > library("SDAMS")
- > data("exampleMSset")
- > results=SDA(exampleMSset)

Here, the MSset class exampleMSset contained in the package is the proteomics dataset, which has two categories, a matrix for proteomics features and a single column phenotype data for grouping information. There are 560 features for 202 experimental subjects (0 for control and 1 for patient). This is a 10% subsample of the original dataset. Each row in this matrix repressents a proteomics feature. See Reference [1] for detailed information regarding this dataset.

### 3 Data Input

#### 3.1 Create MSset from csv.files

The proteomics or metabolomics data is stroed as a matrix with each row being a feature and each column corresponding to a subject. All data in this matrix are non-negative. Another information

required is the phenotype covariates. Here we focus on the binary grouping information, such as numeric 1 for control group and 0 for case group. But it can also be characters, such as "healthy" and "disease". To utilize SDAMS package, we should have two separate csv.files (for example 'feature.csv' and 'group.csv') as inputs for createMSsetfromCSV to creat a MSset object.

#### Note:

- 1. The  $1^{st}$  column in 'feature.csv' represents feature names and the  $1^{st}$  row represents subject codes.
- 2. The  $1^{st}$  column in 'group.csv' represents subject codes, for example, Subject1, Subject2....

The format for "csv.files" should looks like as Figure 1 and Figure 2:

	9308	9324	9447	9449	9457	9459	9464	9466	9467	9471
54	0	0	116.9700012	0	0	0	0	19.01000023	0	0
94	52.40000153	0	0	139.1199951	82.37000275	0	58.22999954	0	22.32999992	25.09000015
97	0	0	53.45000076	62.72999954	47.15999985	0	0	24.44000053	0	22.77000046
191	0	0	22.18000031	170.6499939	0	0	0	0	0	26.52000046
219	58.47000122	0	0	64.5	28.11000061	0	39.77999878	7.429999828	0	5.230000019
286	0	0	0	19.26000023	89.55999756	0	162.8200073	109.8600006	75.48999786	115.6999969
366	0	0	0	0	28.97999954	0	0	0	0	0
388	29.43000031	0	0	0	0	0	0	32.20000076	0	59.97000122
420	65.79000092	0	87.44000244	314.0100098	160.4400024	0	0	0	0	26.85000038
463	0	0	0	12.15999985	38.52999878	0	0	0	0	0
499	0	0	0	26.12999916	14.22000027	0	0	0	0	1.919999957
583	43.74000168	0	0	59.29000092	7.099999905	0	0	0	0	3.809999943
605	0	0	0	657.8400269	661.9500122	0	0	32.22999954	0	0
624	0	96.94999695	0	97.40000153	167.0599976	232.9499969	89.54000092	55.5	35.15999985	0
652	0	0	9.68999958	0	0	0	0	0	0	3.599999905
709	0	0	38.45000076	0	0	0	0	8.229999542	0	6.380000114
889	0	0	0	0	0	0	0	0	0	0
912	569.3200073	0	0	0	0	0	23.57999992	120.6800003	73.69000244	100.4100037

Figure 1: Example of 'feature.csv' pattern

After creating the two csv.files, we need the paths for the two csv.files:

- > path1 <- "/path/to/your/feature.csv/"</pre>
- > path2 <- "/path/to/your/group.csv/"</pre>

Here for demonstration purposes, we use the data in the SDA package

- > directory1 <- system.file("extdata", package="SDAMS", mustWork=TRUE)</pre>
- > path1<-paste(directory1, "ProstateFeature.csv", sep="/")</pre>
- > directory2 <- system.file("extdata", package="SDAMS", mustWork=TRUE)</pre>
- > path2<-paste(directory2, "ProstateGroup.csv", sep="/")</pre>

then use the function getMSsetfromCSV after loading the SDA package

87209

29633

0.00

0.00

	grouping
9308	0
9324	0
9447	0
9449	0
9457	0
9459	0
9464	0
9466	0
9467	0
9471	0
9495	0
9497	0
9507	0
9512	0
9959	1
9960	1

Figure 2: Example of 'group.csv' pattern

0.00 0.00 0.00

0.00 0.00 0.00 57.21

0.00 0.00

0.00 43.87

0.00 0.00

0.00

```
> library("SDAMS")
> exampleMSset1 = createMSsetfromCSV(path1,path2)
> exampleMSset1
MSset (storageMode: lockedEnvironment)
assayData: 560 features, 202 samples
  element names: exprs
protocolData: none
phenoData
  sampleNames: 9512 9963 ... 49586 (202 total)
  varLabels: grouping
  varMetadata: labelDescription
featureData: none
experimentData: use 'experimentData(object)'
Annotation:
> head(featuredata(exampleMSset1)[,1:10])
        9512
               9963 9965 9975
                                  9979 9997 10015 10034 10044 10047
               0.00 0.00 0.00
93922
        0.00
                                  0.00 0.00 68.97
                                                     0.00
```

0.00 0.00 0.00

0.00 0.00 0.00

```
40225 0.00 226.40 0.00 19.65 0.00 0.00 0.00 0.00 0.00 0.00 126342 0.00 0.00 0.00 0.00 0.00 20.43 0.00 0.00 109.93 0.00 42832 52.32 137.76 70.25 0.00 453.23 92.20 0.00 352.55 496.71 0.00
```

> head(phenotypedata(exampleMSset1))

	grouping
9512	0
9963	1
9965	1
9975	1
9979	1
9997	1

### 3.2 Create MSset from R global environment

If the two datasets have been already claeaned and loaded into the R global environment, we can use createMSsetFromEnvir to create a MSset object.

```
> set.seed(100)
> featureInfo = matrix(runif(800,-2,5),ncol = 40)
> featureInfo[featureInfo<0] = 0</pre>
> rownames(featureInfo) = paste("feature",1:20,sep = '')
> colnames(featureInfo) = paste('subject',1:40,sep = '')
> groupInfo = data.frame(grouping=matrix(sample(0:1,40,replace = TRUE),ncol = 1))
> rownames(groupInfo)=colnames(featureInfo)
> exampleMSset2 = createMSsetFromEnvir(feature = featureInfo,group = groupInfo)
> exampleMSset2
MSset (storageMode: lockedEnvironment)
assayData: 20 features, 40 samples
  element names: exprs
protocolData: none
phenoData
  sampleNames: subject1 subject2 ... subject40 (40 total)
 varLabels: grouping
 varMetadata: labelDescription
```

```
featureData: none
experimentData: use 'experimentData(object)'
Annotation:
> head(featuredata(exampleMSset2)[,1:10])
          subject1 subject2 subject3 subject4 subject5 subject6 subject7
feature1 0.1543628 1.7506781 0.3146237 1.2459082 1.216678 0.2919056 0.7766364
feature2 0.0000000 2.9756269 4.0558438 2.5297084 2.195787 0.7263509 0.7489158
feature3 1.8662570 1.7684409 3.4430911 4.7240116 4.438053 0.0000000 1.3078982
feature4 0.0000000 3.2428056 3.7911241 2.7347872 4.879769 0.5297764 2.0855984
feature5 1.2798450 0.9407102 2.2232705 1.1160362 0.000000 1.9968466 0.4667102
feature6 1.3863951 0.0000000 1.4386228 0.5044165 2.045562 2.7941617 0.0000000
         subject8 subject9 subject10
feature1 2.712745 3.705652 0.02105356
feature2 0.000000 1.978800 3.02481591
feature3 0.000000 1.360440 4.46378210
feature4 4.359349 0.000000 2.72457641
feature5 0.000000 0.000000 0.00000000
feature6 3.102040 0.000000 0.43647014
> head(phenotypedata(exampleMSset2))
         grouping
subject1
subject2
                0
subject3
                0
subject4
                1
subject5
                1
subject6
                0
>
```

### 4 Data Analysis

Finally, we perform differential abundance analysis using MSset created in the last section. This can be done by using function SDA. And a list with point estimates, p-values, q-values and corresponding

feature names is returned. Below is results generated by using the MSset exampleMSset1.

- [1,] 0.4622205
- [2,] 0.1380966
- [3,] 0.1192246
- [4,] 0.4323376
- [5,] 0.1806273
- [6,] 0.1345612
- > head(results\$feat.names)
- [1] "93922" "29633" "40225" "126342" "42832" "127351"

### 5 Session Info

- > toLatex(sessionInfo())
  - R version 3.4.3 (2017-11-30), x86\_64-apple-darwin15.6.0
  - Locale: en\_US.UTF-8/en\_US.UTF-8/en\_US.UTF-8/C/en\_US.UTF-8/en\_US.UTF-8
  - Running under: macOS High Sierra 10.13.1
  - Matrix products: default

- BLAS:
  - /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRblas.0.dylib
- LAPACK:
  - /Library/Frameworks/R.framework/Versions/3.4/Resources/lib/libRlapack.dylib
- Base packages: base, datasets, graphics, grDevices, methods, stats, utils
- Other packages: SDAMS 0.99.0
- Loaded via a namespace (and not attached): Biobase 2.36.2, BiocGenerics 0.22.1, colorspace 1.3-2, compiler 3.4.3, ggplot2 2.2.1, grid 3.4.3, gtable 0.2.0, lazyeval 0.2.0, magrittr 1.5, munsell 0.4.3, parallel 3.4.3, plyr 1.8.4, qvalue 2.8.0, Rcpp 0.12.11, reshape2 1.4.2, rlang 0.1.1, scales 0.4.1, splines 3.4.3, stringi 1.1.5, stringr 1.2.0, tibble 1.3.3, tools 3.4.3, trust 0.1-7

### References

[1] Justyna Siwy, William Mullen, Igor Golovko, Julia Franke, and Petra Zürbig. Human urinary peptide database for multiple disease biomarker discovery. *PROTEOMICS-Clinical Applications*, 5(5-6):367–374, 2011.