# MIGSA: Getting pbcmc datasets

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#### Abstract

In this vignette we are going to show how we got the RData pbcmcData.RData which can be loaded via the MIGSAdata package using data(pbcmcData).

Keywords: singular enrichment analysis, over representation analysis, gene set enrichment analysis, functional class scoring, big omics data.

## 1. Getting the data

Following we give the used code to download this data and their PAM50 subtypes.

```
> library(limma);
> library(pbcmc);
> # datasets included in BioConductor repository
> libNames <- c("mainz", "nki", "transbig", "unt", "upp", "vdx");</pre>
> # let's load them!
> pbcmcData <- lapply(libNames, function(actLibName) {</pre>
      print(actLibName);
      # the pbcmc package provides an easy way to download and classify them
      actLib <- loadBCDataset(Class=PAM50, libname=actLibName, verbose=FALSE);</pre>
      actLibFilt <- filtrate(actLib, verbose=FALSE);</pre>
      actLibFilt <- classify(actLibFilt, std="none", verbose=FALSE);</pre>
      actSubtypes <- classification(actLibFilt)$subtype;</pre>
      # get the expression matrix and the annotation
      actExprs <- exprs(actLib);</pre>
      actAnnot <- annotation(actLib);</pre>
```

```
# we recommend working allways with Entrez IDs, let's match them with
      # expression matrix rownames (and modify them)
      if (all(actAnnot$probe == rownames(actExprs))) {
          actExprs <- actExprs[!is.na(actAnnot$EntrezGene.ID),];</pre>
          actAnnot <- actAnnot[!is.na(actAnnot$EntrezGene.ID),];</pre>
          rownames(actExprs) <- as.character(actAnnot$EntrezGene.ID);</pre>
      } else {
          matchedEntrez <- match(rownames(actExprs), actAnnot$probe);</pre>
          # all(rownames(actExprs) %in% actAnnot$probe == !is.na(matchedEntrez));
          stopifnot(all(
               actAnnot$probe[!is.na(matchedEntrez)] ==
               rownames(actExprs)[!is.na(matchedEntrez)]));
          actExprs <- actExprs[!is.na(matchedEntrez),];</pre>
          actAnnot <- actAnnot[!is.na(matchedEntrez),];</pre>
          stopifnot(all(actAnnot$probe == rownames(actExprs)));
          actExprs <- actExprs[!is.na(actAnnot$EntrezGene.ID),];</pre>
          actAnnot <- actAnnot[!is.na(actAnnot$EntrezGene.ID),];</pre>
          rownames(actExprs) <- as.character(actAnnot$EntrezGene.ID);</pre>
      }
      # average repeated genes expression
      actExprs <- avereps(actExprs);</pre>
      stopifnot(all(colnames(actExprs) == names(actSubtypes)));
      # filtrate only these two conditions
      actExprs <- actExprs[, actSubtypes %in% c("Basal", "LumA")];</pre>
      actSubtypes <- as.character(</pre>
          actSubtypes[actSubtypes %in% c("Basal", "LumA")]);
      return(list(geneExpr=actExprs, subtypes=actSubtypes));
+ })
[1] "mainz"
[1] "nki"
[1] "transbig"
[1] "unt"
[1] "upp"
[1] "vdx"
> names(pbcmcData) <- libNames;</pre>
And let's check it is the same data.
> # save the just created pbcmcData to newPbcmcData
> newPbcmcData <- pbcmcData;</pre>
```

```
> library(MIGSAdata);
> # and load the MIGSAdata one.
> data(pbcmcData);
> all.equal(newPbcmcData, pbcmcData);
```

#### [1] TRUE

### Session Info

#### > sessionInfo()

R version 3.5.0 (2018-04-23)

Platform: x86\_64-pc-linux-gnu (64-bit) Running under: Ubuntu 16.04.4 LTS

Matrix products: default

BLAS: /home/biocbuild/bbs-3.7-bioc/R/lib/libRblas.so LAPACK: /home/biocbuild/bbs-3.7-bioc/R/lib/libRlapack.so

#### locale:

[1] LC_CTYPE=en_US.UTF-8	LC_NUMERIC=C
[3] I.C TIME=en US.UTF-8	I.C. COLLATE=C

[5] LC\_MONETARY=en\_US.UTF-8 LC\_MESSAGES=en\_US.UTF-8

[7] LC\_PAPER=en\_US.UTF-8 LC\_NAME=C
[9] LC\_ADDRESS=C LC\_TELEPHONE=C
[11] LC\_MEASUREMENT=en\_US.UTF-8 LC\_IDENTIFICATION=C

### attached base packages:

[1] stats4 parallel stats graphics grDevices utils datasets

[8] methods base

### other attached packages:

[1]	pbcmc_1.8.0	genefu_2.12.0	AIMS_1.12.0
[4]	e1071_1.6-8	iC10_1.1.3	iC10TrainingData_1.0.1
[7]	pamr_1.55	cluster_2.0.7-1	biomaRt_2.36.0
[10]	mclust_5.4	survcomp_1.30.0	prodlim_2018.04.18
[13]	survival_2.42-3	edgeR_3.22.0	MIGSAdata_1.3.0
[16]	MIGSA_1.4.0	mGSZ_1.0	ismev_1.41
[19]	mgcv_1.8-23	nlme_3.1-137	MASS_7.3-50
[22]	limma_3.36.0	GSA_1.03	BiocParallel_1.14.0
[25]	GSEABase_1.42.0	graph_1.58.0	annotate_1.58.0
[28]	XML_3.98-1.11	AnnotationDbi_1.42.0	IRanges_2.14.0
[31]	S4Vectors_0.18.0	Biobase_2.40.0	BiocGenerics_0.26.0

#### loaded via a namespace (and not attached):

[1]	survivalROC_1.0.3	Category_2.46.0
[3]	breastCancerUNT_1.17.0	bitops_1.0-6
[5]	matrixStats_0.53.1	bit64_0.9-7
[7]	progress_1.1.2	httr_1.3.1
[9]	Rgraphviz_2.24.0	tools_3.5.0
[11]	R6_2.2.2	vegan_2.5-1
[13]	KernSmooth_2.23-15	DBI_0.8
[15]	lazyeval_0.2.1	colorspace_1.3-2
[17]	rmeta_3.0	permute_0.9-4
[19]	<pre>gridExtra_2.3</pre>	prettyunits_1.0.2
[21]	bit_1.1-12	compiler_3.5.0
[23]	formatR_1.5	breastCancerNKI_1.17.0
[25]	ggdendro_0.1-20	labeling_0.3
[27]	scales_0.5.0	<pre>genefilter_1.62.0</pre>
[29]	RBGL_1.56.0	stringr_1.3.0
[31]	digest_0.6.15	${\tt breastCancerVDX\_1.17.0}$
[33]	AnnotationForge_1.22.0	pkgconfig_2.0.1
[35]	rlang_0.2.0	RSQLite_2.1.0
[37]	SuppDists_1.1-9.4	GOstats_2.46.0
[39]	RCurl_1.95-4.10	magrittr_1.5
[41]	GO.db_3.6.0	futile.logger_1.4.3
[43]	Matrix_1.2-14	Rcpp_0.12.16
[45]	munsell_0.4.3	stringi_1.1.7
	RJSONIO_1.3-0	org.Hs.eg.db_3.6.0
	plyr_1.8.4	breastCancerUPP_1.17.0
[51]	grid_3.5.0	blob_1.1.1
	breastCancerTRANSBIG_1.17.0	lattice_0.20-35
	cowplot_0.9.2	splines_3.5.0
[57]	locfit_1.5-9.1	pillar_1.2.2
	reshape2_1.4.3	<pre>futile.options_1.0.1</pre>
	lambda.r_1.2.2	data.table_1.10.4-3
	bootstrap_2017.2	gtable_0.2.0
	amap_0.8-14	assertthat_0.2.0
[67]	ggplot2_2.2.1	xtable_1.8-2
[69]	class_7.3-14	tibble_1.4.2
	memoise_1.1.0	lava_1.6.1
[73]	breastCancerMAINZ_1.17.0	

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