# GSE32866, Genome-scale DNA methylation profiling of lung adenocarcinoma:

#### Paulyna Magana

## Contents

0.1	Introduction	1
0.2	Importing the data	1
0.3	Extract the data	2
0.4	Exploratory analysis	2
0.5	Make sure dimensions match up	2
0.6	Look at overall distributions	9
0.7	Inspect the clinical variables	6
0.8	Sample clustering and Principal Components Analysis	7

## 0.1 Introduction

Load the following packages:

```
library(rmarkdown) # paged table
library(GEOquery) #geo query access
library(dplyr)
library(devtools)
library(ggplot2)
library(limma)
library(Glimma)
library(edgeR)
library(kableExtra)
library(ggrepel)
library(tinytex)
library(purrr)
Sys.setenv(VROOM_CONNECTION_SIZE = 25600000)
```

# 0.2 Importing the data

```
my_id <- "GSE32866"
gse <- getGEO(my_id)</pre>
```

Some datasets on GEO may be derived from different microarray platforms. Therefore the object gse is a list of different datasets. You can find out how many were used by checking the length of the gse object. Usually there will only be one platform and the dataset we want to analyse will be the first object in the list (gse[[1]]).

```
length(gse)
```

[1] 1

#### 0.3 Extract the data

```
gse <- gse[[1]]
gse
ExpressionSet (storageMode: lockedEnvironment)
assayData: 27578 features, 55 samples
  element names: exprs
protocolData: none
phenoData
  sampleNames: GSM813533 GSM813534 ... GSM813587 (55 total)
  varLabels: title geo_accession ... tissue:ch1 (58 total)
  varMetadata: labelDescription
featureData
  featureNames: cg00000292 cg00002426 ... cg27665659 (27578 total)
  fvarLabels: ID Name ... ORF (38 total)
  fvarMetadata: Column Description labelDescription
experimentData: use 'experimentData(object)'
  pubMedIds: 22613842
Annotation: GPL8490
```

#### 0.4 Exploratory analysis

The exprs function can retrieve the expression values as a data frame; with one column per-sample and one row per-gene.

```
pdata= pData(gse) #sample information
edata= exprs(gse) #expression data
fdata = fData(gse) #gene annotation
```

#### 0.5 Make sure dimensions match up

The number of rows of the feature data should match the number of rows of the genomic data (both are the number of genes). The number of rows of the phenotype data should match the number of columns of the genomic data (both are the number of samples).

```
dim(fdata)
```

```
[1] 27578 38
```

```
#Dimension of edata
dim(pdata)
```

[1] 55 58

```
#Dimension of edata
dim(edata)
```

[1] 27578 55

#### 0.6 Look at overall distributions

For visualisation and statistical analysis, we will inspect the data to discover what scale the data are presented in. The methods we will use assume the data are on a log2 scale; typically in the range of 0 to 16.

The summary function can then be used to print the distributions.

```
## exprs get the expression levels as a data frame and get the distribution
dat <- summary(exprs(gse))
m <- matrix(1:ncol(dat), 5)
for (i in 1:ncol(m)) {
  cat(kbl(dat[, m[, i]], 'latex', booktabs=TRUE), "\\newline")
}</pre>
```

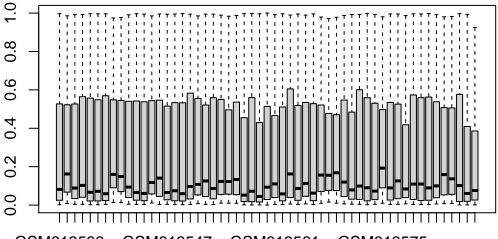
GSM813533	GSM813534	GSM813535	GSM813536	GSM813537
Min. :0.00158 1st Qu.:0.02499 Median :0.08185 Mean :0.26751 3rd Qu.:0.52701	Min. :0.0088 1st Qu.:0.0675 Median :0.1621 Mean :0.2951 3rd Qu.:0.5223	Min. :0.003852 1st Qu.:0.034014 Median :0.088613 Mean :0.270265 3rd Qu.:0.526548	Min. :0.00412 1st Qu.:0.04078 Median :0.10260 Mean :0.28832 3rd Qu.:0.56454	Min. :0.001593 1st Qu.:0.021340 Median :0.066667 Mean :0.274648 3rd Qu.:0.557164
Max. :0.99686 NA's :118		Max. :0.993288 NA's :23	Max. :0.99236 NA's :66	Max. :0.997155 NA's :1
GSM813538	GSM813539	GSM813540	GSM813541	GSM813542
Min. :0.001563 1st Qu.:0.021631 Median :0.071831 Mean :0.272946 3rd Qu.:0.548465 Max. :0.996068 NA's :4	Min. :0.002816 1st Qu.:0.023452 Median :0.05912 Mean :0.272262 3rd Qu.:0.569431 Max. :0.996306 NA's :3	0 Median :0.15982 Mean :0.30666	Mean :0.14873 Mean :0.29943	Median :0.09375 Mean :0.27430
GSM813543	GSM813544	GSM813545	GSM813546	GSM813547
Min. :0.003748 1st Qu.:0.025443 Median :0.066108 Mean :0.269724 3rd Qu.:0.542025	Min. :0.002923 1st Qu.:0.022825 Median :0.06055 Mean :0.263932 3rd Qu.:0.538074	8 Median :0.11796 Mean :0.28764	Median :0.14063 Mean :0.29780	Median :0.06540 Mean :0.25911
Max. :0.996605 NA's :1	Max. :0.994622 NA's :5	Max. :0.98549 NA's :6	Max. :0.99514 NA's :42	Max. :0.99406 NA's :2

GSM813548	GSM813549	GSM813550	GSM813551	GSM813552
Min. :0.002475 1st Qu.:0.022601 Median :0.074436 Mean :0.271670 3rd Qu.:0.533615	Min. :0.001802 1st Qu.:0.021012 Median :0.059294 Mean :0.264499 3rd Qu.:0.532461	Min. :0.00275 1st Qu.:0.03178 Median :0.09664 Mean :0.29360 3rd Qu.:0.58296	Min. :0.007388 1st Qu.:0.053120 Median :0.107693 Mean :0.286343 3rd Qu.:0.555106	Mean $:0.28056$
Max. :0.995359 NA	Max. :0.997065 NA's :6	Max. :0.99442 NA's :96	Max. :0.985281 NA's :3	Max. :0.99471 NA's :208
GSM813553	GSM813554	GSM813555	GSM813556	GSM813557
Min. :0.003055 1st Qu.:0.032660 Median :0.086232 Mean :0.280567 3rd Qu.:0.559251 Max. :0.994143 NA	Min. :0.01007 1st Qu.:0.05484 Median :0.12322 Mean :0.28903 3rd Qu.:0.55005 Max. :0.98923 NA's :51	Min. :0.0099 1st Qu.:0.0568 Median :0.1227 Mean :0.2733 3rd Qu.:0.4960 Max. :0.9836 NA's :453	1st Qu.:0.05408 Median :0.13335 Mean :0.28789 3rd Qu.:0.53638 Max. :0.98753	Min. :0.001198 1st Qu.:0.018511 Median :0.051788 Mean :0.239002 3rd Qu.:0.455191 Max. :0.996707 NA's :16
GSM813558	GSM813559	GSM813560	GSM813561	GSM813562
Min. :0.000755 1st Qu.:0.017597 Median :0.072014 Mean :0.277990 3rd Qu.:0.559703 Max. :0.997774 NA's :8	Min. :0.000893 1st Qu.:0.015929 Median :0.044893 Mean :0.230163 3rd Qu.:0.429716 Max. :0.996079 NA's :19	Min. :0.00000 1st Qu.:0.02894 Median :0.09338 Mean :0.26941 3rd Qu.:0.51380 Max. :1.00000 NA's :11	Min. :0.00329 1st Qu.:0.03900 Median :0.11016 Mean :0.26246 3rd Qu.:0.46627 Max. :0.99187 NA's :102	Min. :0.00288 1st Qu.:0.02200 Median :0.05827 Mean :0.25957 3rd Qu.:0.51087 Max. :0.99605 NA's :49
GSM813563	GSM813564	GSM813565	GSM813566	GSM813567
Min. :0.005434 1st Qu.:0.039251 Median :0.162052 Mean :0.322312 3rd Qu.:0.604959 Max. :0.989720	Min. :0.003316 1st Qu.:0.024736 Median :0.086374 Mean :0.271873 3rd Qu.:0.518953 Max. :0.995040	Min. :0.00608 1st Qu.:0.04581	Min. :0.003185 1st Qu.:0.023066	Min. :0.009481 1st Qu.:0.070661 Median :0.156346 Mean :0.297500
NA's :1	NA's :1	NA's :39	NA's :2	NA's :3
GSM813568	GSM813569	GSM813570	GSM813571	GSM813572
Min. :0.01552 1st Qu.:0.07648 Median :0.15532 Mean :0.28663 3rd Qu.:0.47719	Min. :0.00985 1st Qu.:0.07542 Median :0.16813 Mean :0.28440 3rd Qu.:0.47053	Min. :0.006009 1st Qu.:0.044466 Median :0.119853 Mean :0.290716 3rd Qu.:0.547162	Min. :0.004533 1st Qu.:0.030680 Median :0.078872 Mean :0.256968 3rd Qu.:0.484131	Min. :0.003998 1st Qu.:0.029502 Median :0.098945 Mean :0.302563 3rd Qu.:0.600460
Max. :0.97192 NA's :4	Max. :0.97799 NA's :51	Max. :0.987253 NA's :1	Max. :0.992672 NA's :3	Max. :0.992380 NA's :1
·				

GSM813573	GSM813574	GSM813575	GSM813576	GSM813577
Min. :0.001719	Min. :0.003869	Min. :0.00984	Min. :0.002911	Min. :0.003975
1st Qu.:0.022253	1st Qu.:0.028332	1st Qu.:0.08979	1st Qu.:0.026097	1st Qu.:0.033384
Median :0.090038	Median :0.072875	Median :0.19254	Median :0.089301	Median :0.125818
Mean :0.283277	Mean :0.270009	Mean :0.30278	Mean :0.274118	Mean :0.286136
3rd Qu.:0.559240	3rd Qu.:0.530568	3rd Qu.:0.49747	3rd Qu.:0.534446	3rd Qu.:0.525828
Max. :0.998242	Max. :0.992547	Max. :0.98056	Max. :0.995558	Max. :0.995050
NA's :6	NA's :3	NA's :124	NA's :4	NA's :4
GSM813578	GSM813579	GSM813580	GSM813581	GSM813582
Min. :0.00654	Min. :0.003284	Min. :0.002743	Min. :0.002231	Min. :0.005838
1st Qu.:0.03869	1st Qu.:0.029137	1st Qu.:0.023585	1st Qu.:0.025043	1st Qu.:0.038127
Median :0.08380	Median :0.109404	Median :0.109505	Median :0.089401	Median :0.100111
Mean :0.24641	Mean :0.294090	Mean :0.289157	Mean :0.282871	Mean :0.280244
3rd Qu.:0.41828	3rd Qu.:0.573597	3rd Qu.:0.559674	3rd Qu.:0.562494	3rd Qu.:0.537907
Max. :0.98813	Max. :0.995953	Max. :0.995473	Max. :0.997839	Max. :0.991527
NA's :6	NA	NA's :4	NA's :16	NA's :4
GSM813583	GSM813584	GSM813585	GSM813586	GSM813587
Min. :0.00773	Min. :0.00872	Min. :0.000538	Min. :0.001063	Min. :0.0012
1st Qu.:0.05315	1st Qu.:0.05574	1st Qu.:0.018411	1st Qu.:0.019389	1st Qu.:0.0262
Median :0.15874	Median :0.13669	Median :0.101503	Median :0.060440	Median :0.0759
Mean :0.28648	Mean :0.28092	Mean :0.294703	Mean :0.228942	Mean :0.2264
3rd Qu.:0.50774	3rd Qu.:0.50595	3rd Qu.:0.576664	3rd Qu.:0.409377	3rd Qu.:0.3859
Max. :0.98072	Max. :0.98240	Max. :0.996908	Max. :0.995679	Max. :0.9941
NA's :131	NA's :280	NA's :4	NA's :22	NA's :871

A boxplot can also be generated to see if the data have been normalised. If so, the distributions of each sample should be highly similar.

## boxplot(edata,outline=FALSE)



GSM813533 GSM813547 GSM813561 GSM813575

kable(table(pdata\$characteristics\_ch1.1), booktabs = T)

Var1		Freq
	Lung tumor Normal lung	28 27

kable(table(pdata\$characteristics\_ch1.6,pdata\$characteristics\_ch1.1), booktabs = T)

	tissue: Lung tumor	tissue: Normal lung
gender: Female	14	13
gender: Male	11	11
gender: NA	3	3

# 0.7 Inspect the clinical variables

Data submitted to GEO contain sample labels assigned by the experimenters, and some information about the processing protocol. All these data can be extracted by the pData function.

```
sampleInfo <- pData(gse)

## source_name_ch1 and characteristics_ch1.1 seem to contain factors we might need for the analysis
sampleInfo <- select(sampleInfo, source_name_ch1,characteristics_ch1.1)

#rename to more convenient column names
sampleInfo <- rename(sampleInfo, patient=characteristics_ch1.1, group = source_name_ch1)
kable(sampleInfo, longtable = T, booktabs = T, caption = "SampleInfo") %>%
kable_styling(latex_options = c("repeat_header"))
```

Table 1: SampleInfo

	group	patient
GSM813533	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813534	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813535	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813536	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813537	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813538	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813539	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813540	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813541	Fresh frozen macrodissected tissue	tissue: Normal lung
$\operatorname{GSM813542}$	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813543	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813544	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813545	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813546	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813547	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813548	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813549	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813550	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813551	Fresh frozen macrodissected tissue	tissue: Normal lung

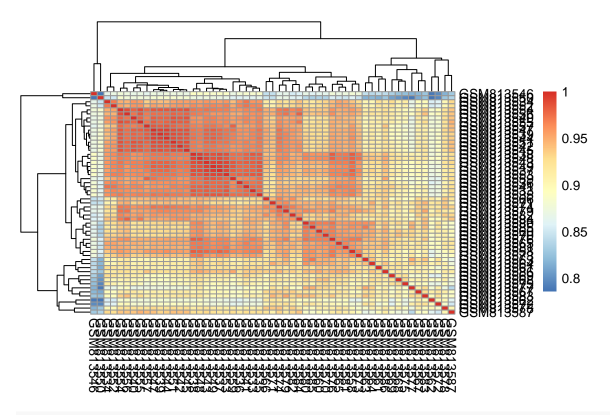
Table 1: SampleInfo (continued)

	group	patient
$\operatorname{GSM813552}$	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813553	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813554	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813555	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813556	Fresh frozen macrodissected tissue	tissue: Normal lung
$\mathbf{GSM} 813557$	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813558	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813559	Fresh frozen macrodissected tissue	tissue: Normal lung
GSM813560	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813561	Fresh frozen macrodissected tissue	tissue: Lung tumor
$\mathbf{GSM} 813562$	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813563	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813564	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813565	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813566	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813567	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813568	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813569	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813570	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813571	Fresh frozen macrodissected tissue	tissue: Lung tumor
$\mathbf{GSM} 813572$	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813573	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813574	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813575	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813576	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813577	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813578	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813579	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813580	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813581	Fresh frozen macrodissected tissue	tissue: Lung tumor
$\operatorname{GSM813582}$	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813583	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813584	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813585	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813586	Fresh frozen macrodissected tissue	tissue: Lung tumor
GSM813587	Fresh frozen macrodissected tissue	tissue: Lung tumor

# 0.8 Sample clustering and Principal Components Analysis

Unsupervised analysis is a good way to get an understanding of the sources of variation in the data. It can also identify potential outlier samples.

```
library(pheatmap)
corMatrix <- cor(exprs(gse), use="c")
pheatmap(corMatrix)</pre>
```



#### session\_info()

- Session info -----

setting value

version R version 4.0.5 (2021-03-31)

os macOS Big Sur 10.16 system x86\_64, darwin17.0

ui X11 language (EN)

collate en\_GB.UTF-8
ctype en\_GB.UTF-8
tz Europe/London
date 2022-03-09

pandoc 2.11.4 @ /Applications/RStudio.app/Contents/MacOS/pandoc/ (via rmarkdown)

- Packages ------

! pa	ckage	*	version	date	(UTC)	lib	source
an	notate		1.68.0	2020-	-10-27	[2]	Bioconductor
An	notationDbi		1.52.0	2020-	-10-27	[2]	Bioconductor
as	sertthat		0.2.1	2019-	-03-21	[2]	CRAN (R 4.0.2)
Bi	obase	*	2.50.0	2020-	-10-27	[2]	Bioconductor
Bi	ocGenerics	*	0.36.1	2021-	-04-16	[2]	Bioconductor
Bi	ocParallel		1.24.1	2020-	-11-06	[2]	Bioconductor
bi	t		4.0.4	2020-	-08-04	[2]	CRAN (R 4.0.2)
bi	t64		4.0.5	2020-	-08-30	[2]	CRAN (R 4.0.2)
bi	tops		1.0-7	2021-	-04-24	[2]	CRAN (R 4.0.2)
bl	ob		1.2.2	2021-	-07-23	[2]	CRAN (R 4.0.2)
br	io		1.1.3	2021-	-11-30	[2]	CRAN (R 4.0.2)
ca	chem		1.0.6	2021-	-08-19	[2]	CRAN (R 4.0.2)

```
3.7.0
                                     2021-04-20 [2] CRAN (R 4.0.2)
  callr
  cli
                          3.2.0
                                     2022-02-14 [2] CRAN (R 4.0.5)
                          2.0 - 3
  colorspace
                                     2022-02-21 [2] CRAN (R 4.0.5)
                                     2022-02-14 [2] CRAN (R 4.0.5)
                          1.5.0
  crayon
  curl
                          4.3.2
                                     2021-06-23 [2] CRAN (R 4.0.5)
  DBI
                                     2021-12-20 [2] CRAN (R 4.0.2)
                          1.1.2
                          0.16.3
                                     2021-03-24 [2] Bioconductor
  DelayedArray
                          1.4.0
                                     2021-09-28 [2] CRAN (R 4.0.2)
  desc
  DESeq2
                          1.30.1
                                     2021-02-19 [2] Bioconductor
                                     2021-11-30 [2] CRAN (R 4.0.2)
  devtools
                        * 2.4.3
  digest
                          0.6.29
                                     2021-12-01 [2] CRAN (R 4.0.2)
                                     2022-02-08 [2] CRAN (R 4.0.5)
                        * 1.0.8
  dplyr
                                     2021-01-14 [2] Bioconductor
                        * 3.32.1
  edgeR
  ellipsis
                                     2021-04-29 [2] CRAN (R 4.0.2)
                          0.3.2
  evaluate
                          0.15
                                     2022-02-18 [2] CRAN (R 4.0.5)
  fansi
                          1.0.2
                                     2022-01-14 [2] CRAN (R 4.0.5)
                                     2021-01-25 [2] CRAN (R 4.0.2)
  fastmap
                          1.1.0
  fs
                          1.5.2
                                     2021-12-08 [2] CRAN (R 4.0.2)
                          1.72.1
                                     2021-01-21 [2] Bioconductor
  genefilter
                                     2020-10-27 [2] Bioconductor
  geneplotter
                          1.68.0
  generics
                          0.1.2
                                     2022-01-31 [2] CRAN (R 4.0.5)
  GenomeInfoDb
                          1.26.7
                                     2021-04-08 [2] Bioconductor
  GenomeInfoDbData
                                     2021-04-30 [2] Bioconductor
                          1.2.4
                          1.42.0
                                     2020-10-27 [2] Bioconductor
  GenomicRanges
                        * 2.58.0
                                     2020-10-27 [2] Bioconductor
  GEOquery
P ggplot2
                        * 3.3.5
                                     2021-06-25 [?] CRAN (R 4.0.2)
P ggrepel
                        * 0.9.1
                                     2021-01-15 [?] CRAN (R 4.0.2)
  Glimma
                        * 2.0.0
                                     2020-10-27 [2] Bioconductor
                                     2022-02-24 [2] CRAN (R 4.0.5)
                          1.6.2
  glue
  gtable
                          0.3.0
                                     2019-03-25 [2] CRAN (R 4.0.2)
  highr
                          0.9
                                     2021-04-16 [2] CRAN (R 4.0.2)
  hms
                          1.1.1
                                     2021-09-26 [2] CRAN (R 4.0.2)
  htmltools
                          0.5.2
                                     2021-08-25 [2] CRAN (R 4.0.2)
                          1.5.4
                                     2021-09-08 [2] CRAN (R 4.0.2)
  htmlwidgets
                                     2020-07-20 [2] CRAN (R 4.0.2)
  httr
                          1.4.2
                          2.24.1
                                     2020-12-12 [2] Bioconductor
  IRanges
  jsonlite
                          1.8.0
                                     2022-02-22 [2] CRAN (R 4.0.5)
  kableExtra
                        * 1.3.4.9000 2021-09-12 [2] Github (haozhu233/kableExtra@4c93f1a)
  knitr
                          1.37
                                     2021-12-16 [2] CRAN (R 4.0.2)
                                     2021-09-22 [2] CRAN (R 4.0.2)
  lattice
                          0.20 - 45
                          1.0.1
                                     2021-09-24 [2] CRAN (R 4.0.2)
  lifecycle
  limma
                        * 3.46.0
                                     2020-10-27 [2] Bioconductor
  locfit
                                     2020-03-25 [2] CRAN (R 4.0.2)
                          1.5 - 9.4
                                     2022-01-26 [2] CRAN (R 4.0.5)
  magrittr
                          2.0.2
                                     2021-12-08 [2] CRAN (R 4.0.2)
  Matrix
                          1.4 - 0
                                     2021-01-30 [2] Bioconductor
  MatrixGenerics
                          1.2.1
                                     2021-09-17 [2] CRAN (R 4.0.2)
  matrixStats
                          0.61.0
                                     2021-11-26 [2] CRAN (R 4.0.2)
  memoise
                          2.0.1
  munsell
                          0.5.0
                                     2018-06-12 [2] CRAN (R 4.0.2)
                                     2019-01-04 [2] CRAN (R 4.0.2)
  pheatmap
                        * 1.0.12
                          1.7.0
                                     2022-02-01 [2] CRAN (R 4.0.5)
  pillar
                                     2021-12-20 [2] CRAN (R 4.0.2)
  pkgbuild
                          1.3.1
  pkgconfig
                          2.0.3
                                     2019-09-22 [2] CRAN (R 4.0.2)
                                     2021-11-30 [2] CRAN (R 4.0.2)
  pkgload
                          1.2.4
```

```
2020-01-24 [2] CRAN (R 4.0.2)
  prettyunits
                       1.1.1
                        3.5.2
                                   2021-04-30 [2] CRAN (R 4.0.2)
  processx
 ps
                       1.6.0
                                  2021-02-28 [2] CRAN (R 4.0.2)
                                   2020-04-17 [2] CRAN (R 4.0.2)
                      * 0.3.4
 purrr
                        2.5.1
                                   2021-08-19 [2] CRAN (R 4.0.2)
 RColorBrewer
                                   2014-12-07 [2] CRAN (R 4.0.2)
                        1.1-2
                                 2022-01-13 [2] CRAN (R 4.0.5)
  Rcpp
                       1.0.8
                       1.98-1.6 2022-02-08 [2] CRAN (R 4.0.5)
  RCurl
P readr
                        2.1.2
                                   2022-01-30 [?] CRAN (R 4.0.5)
                                   2021-11-30 [2] CRAN (R 4.0.2)
  remotes
                        2.4.2
                                   2022-01-24 [1] CRAN (R 4.0.5)
  renv
                        0.15.2
                                   2022-02-03 [2] CRAN (R 4.0.5)
                        1.0.1
  rlang
                                   2021-09-14 [2] CRAN (R 4.0.2)
                      * 2.11
  rmarkdown
                                   2020-11-15 [2] CRAN (R 4.0.2)
  rprojroot
                        2.0.2
 RSQLite
                        2.2.10
                                   2022-02-17 [2] CRAN (R 4.0.5)
  rstudioapi
                        0.13
                                   2020-11-12 [2] CRAN (R 4.0.2)
                        1.0.2
                                   2021-10-16 [2] CRAN (R 4.0.2)
  rvest
 S4Vectors
                       0.28.1
                                   2020-12-09 [2] Bioconductor
                                   2020-05-11 [2] CRAN (R 4.0.2)
  scales
                       1.1.1
                                   2021-12-06 [2] CRAN (R 4.0.2)
  sessioninfo
                        1.2.2
  stringi
                        1.7.6
                                   2021-11-29 [2] CRAN (R 4.0.2)
P stringr
                       1.4.0
                                  2019-02-10 [?] CRAN (R 4.0.2)
  SummarizedExperiment 1.20.0
                                2020-10-27 [2] Bioconductor
                        3.2-13
                                   2021-08-24 [2] CRAN (R 4.0.2)
  survival
                                  2022-02-03 [2] CRAN (R 4.0.5)
  svglite
                        2.1.0
  systemfonts
                        1.0.4
                                   2022-02-11 [2] CRAN (R 4.0.5)
  testthat
                        3.1.2
                                   2022-01-20 [2] CRAN (R 4.0.5)
                                   2021-11-07 [2] CRAN (R 4.0.2)
  tibble
                        3.1.6
                                   2022-02-01 [?] CRAN (R 4.0.5)
P tidyr
                       1.2.0
                                  2022-02-21 [2] CRAN (R 4.0.5)
  tidyselect
                       1.1.2
  tinytex
                      * 0.37
                                   2022-02-16 [2] CRAN (R 4.0.5)
  tzdb
                        0.2.0
                                   2021-10-27 [2] CRAN (R 4.0.2)
                                   2021-12-09 [2] CRAN (R 4.0.2)
  usethis
                      * 2.1.5
                        1.2.2
                                   2021-07-24 [2] CRAN (R 4.0.2)
  utf8
                                   2021-04-29 [2] CRAN (R 4.0.2)
                        0.3.8
  vctrs
  viridisLite
                        0.4.0
                                   2021-04-13 [2] CRAN (R 4.0.2)
  vroom
                        1.5.7
                                  2021-11-30 [2] CRAN (R 4.0.2)
  webshot
                        0.5.2
                                  2019-11-22 [2] CRAN (R 4.0.2)
                                   2021-11-30 [2] CRAN (R 4.0.2)
  withr
                        2.4.3
  xfun
                        0.29
                                   2021-12-14 [2] CRAN (R 4.0.2)
  XML
                        3.99-0.9 2022-02-24 [2] CRAN (R 4.0.5)
  xm12
                        1.3.3
                                   2021-11-30 [2] CRAN (R 4.0.2)
                                   2019-04-21 [2] CRAN (R 4.0.2)
  xtable
                        1.8 - 4
                                   2020-10-28 [2] Bioconductor
 XVector
                        0.30.0
                                   2022-02-21 [2] CRAN (R 4.0.5)
 vaml
                        2.3.5
                                   2020-10-28 [2] Bioconductor
  zlibbioc
                        1.36.0
```

P -- Loaded and on-disk path mismatch.

-----

<sup>[1] /</sup>Users/paulyna/Documents/Github/Gene Analysis/renv/library/R-4.0/x86\_64-apple-darwin17.0

<sup>[2] /</sup>Library/Frameworks/R.framework/Versions/4.0/Resources/library