IOBR Tutorial

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Contents

IC	$_{ m BR}$	introduction 5										
	Pref	ace										
	0.1	Introduction										
	0.2	License										
	0.3	Publishment										
	0.4	Reporting bugs										
1	How to install IOBR											
	1.1	Installing Dependency Packages										
	1.2	Installing IOBR										
	1.3	The main pipeline of IOBR										
2	RN	RNA Data preprocessing 1										
	2.1	Library										
	2.2	Downloading data for example										
	2.3	Identifying outlier samples										
	2.4	PCA analysis of molecular subtypes										
	2.5	Batch effect correction										
3	Tumor ecosystem analysis 21											
	3.1	Determine TME subtype of gastric cancer using TME classifier										
	3.2	DEG analysis										
	3.3	GSEA analysis of DEGs										
	3.4	DEG analysis										
	3.5	Identifying signatures associated with TME clusters										
4	Signatures and relevant phenotypes 41											
	4.1	Loading packages										
	4.2	Downloading data for example										
	4.3	Signature score estimation										
	4.4	Identifying features associated with survival										
	4.5	Visulization using heatmap										
	4.6	Focus on target signatures										
	4.7	Survival analysis										

4 CONTENTS

	4.8	Batch correlation analysis	52									
	4.9	Visulization of correlations	55									
5	TM	E deconvolution 5	57									
	5.1	Loading packages	57									
	5.2	Downloading data for example	57									
	5.3	Available Methods to Decode TME Contexture	59									
	5.4	Method 1: CIBERSORT	60									
	5.5	Method 3: MCPcounter	62									
	5.6	Method 5: ESTIMATE	63									
	5.7	Method 6: TIMER	64									
	5.8	Method 7: quanTIseq	64									
	5.9	8) Method 8: IPS	66									
	5.10	9) Combination of above deconvolution results	67									
	5.11	Licenses of the deconvolution methods	67									
6	Cros	Cross-references 73										
	6.1	Chapters and sub-chapters	73									
	6.2		73									
7	Footnotes and citations 77											
-	7.1		77									
	7.2		77									
8	Blocks 79											
	8.1		79									
	8.2		79									
	8.3	-	80									
9	Shar	ring your book	31									
0	9.1		31									
	9.2		81									
	9.3		81									
10	Refe	erences	33									
			33									
			34									
			35									

IOBR introduction

Preface

0.1 Introduction

IOBR is design for Immuno-Oncology Biological Research. Recent advance in next-generation sequencing has triggered the rapidly accumulating publicly available multi-omics data. The application of integrated omics to exploring robust signatures for clinical translation is increasingly highlighted in immuno-oncology but raises computational and biological challenges. This vignette aims to demonstrate how to utilize the package named IOBR to perform multi-omics immuno-oncology biological research to decode tumor microenvironment and signatures for clinical translation.

This R package integrates 8 published methodologies for decoding tumor microenvironment (TME) contexture: CIBERSORT, TIMER, xCell, MCPcounter, ESITMATE, EPIC, IPS, quantIseq. Moreover, 255 published signature gene sets were collected by IOBR, involving tumor microenvironment, tumor metabolism, m6A, exosomes, microsatellite instability, and tertiary lymphoid structure. Run the function signature_collection_citation to obtain the source papers, and the function signature_collection returns the detail signature genes of all given signatures. Subsequently, IOBR adopts three computational methods to calculate the signature score, comprising PCA, z-score, and ssGSEA. To note, IOBR collected and employed multiple approaches for variable transition, visualization, batch survival analysis, feature selection, and statistical analysis. Batch analysis and visualization of corresponding results are supported. The details of how IOBR works are described below.

6 CONTENTS

IOBR R package workflow

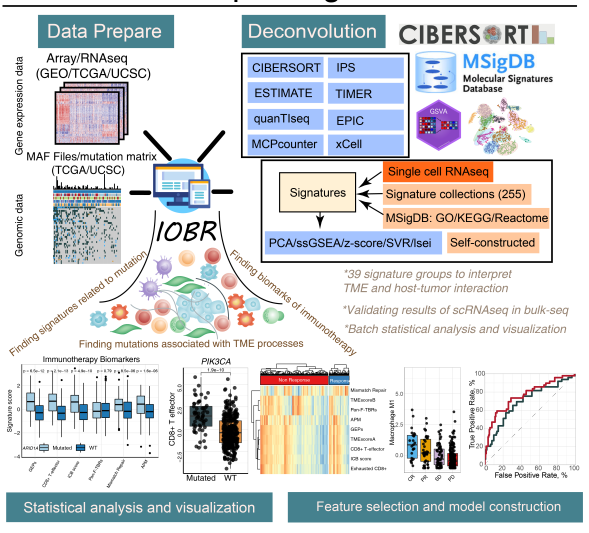


Figure 1: The workflow of IOBR

0.2. LICENSE 7

0.2 License

IOBR is released under the GPL v3.0 license. See LICENSE for details. The code contained in this book is simultaneously available under the GPL license; this means that you are free to use it in your packages, as long as you cite the source. The online version of this book is licensed under the Creative Commons Attribution-NonCommercial-ShareAlike 4.0 International License.

0.3 Publishment

Zeng D, Ye Z, Shen R, Yu G, Wu J, Xiong Y,..., Liao W (2021) **IOBR**: Multi-Omics Immuno-Oncology Biological Research to Decode Tumor Microenvironment and Signatures. *Frontiers in Immunology*. 12:687975. doi: 10.3389/fimmu.2021.687975

0.4 Reporting bugs

Please report bugs to the Github issues page

E-mail any questions to dongqiangzeng0808@gmail.com

8 CONTENTS

Chapter 1

How to install IOBR

1.1 Installing Dependency Packages

It is essential that you have R 3.6.3 or above already installed on your computer or server. IOBR is a pipeline that utilizes many other R packages that are currently available from CRAN, Bioconductor and GitHub.

1.2 Installing IOBR

Then, you can start to install IOBR from github by typing the following code into your R session:

```
if (!requireNamespace("IOBR", quietly = TRUE)) devtools::install_github("IOBR/IOBR")

## Warning: package 'tidyHeatmap' was built under R version 4.2.3

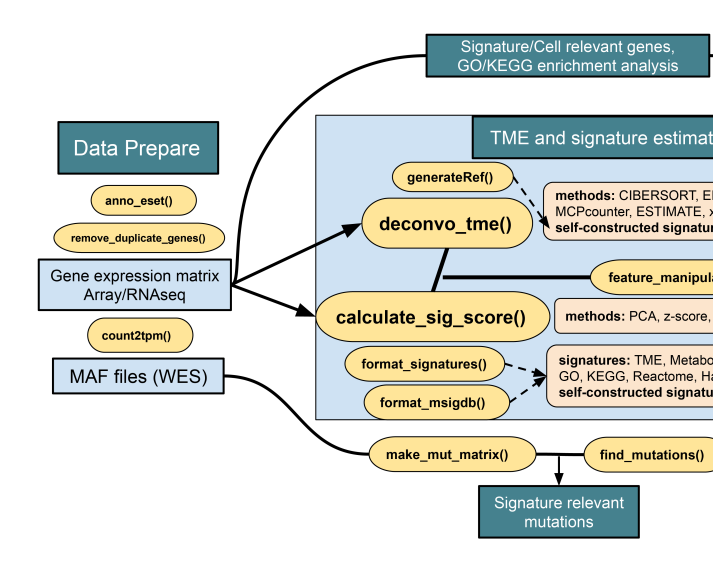
library(IOBR)

## Warning: package 'tibble' was built under R version 4.2.3

## Warning: package 'dplyr' was built under R version 4.2.3

## Warning: package 'ggplot2' was built under R version 4.2.3
```

1.3 The main pipeline of IOBR



IOBR (Immuno-Oncology B)

Figure 1.1: The main pipeline of IOBR

Chapter 2

RNA Data preprocessing

2.1 Library

Load the IOBR package in your R session after the installation is complete:

```
library(IOBR)
library(tidyverse)
library(clusterProfiler)
```

2.2 Downloading data for example

Obtaining data set from GEO Gastric cancer: GSE62254 using GEOquery R package.

```
if (!requireNamespace("GEOquery", quietly = TRUE)) BiocManager::install("GEOquery")
library("GEOquery")
# NOTE: This process may take a few minutes which depends on the internet connection s
eset_geo<-getGEO(GEO = "GSE62254", getGPL = F, destdir = "./")
eset <-eset_geo[[1]]
eset <-exprs(eset)
eset[1:5,1:5]</pre>
```

```
GSM1523727 GSM1523728 GSM1523729 GSM1523744 GSM1523745
##
## 1007 s at 3.2176645 3.0624323 3.0279131
                                            2.921683 2.8456013
## 1053_at 2.4050109 2.4394879 2.2442708 2.345916 2.4328582
## 117 at 1.4933412 1.8067380 1.5959665 1.839822 1.8326058
## 121 at
           2.1965561 2.2812181 2.1865556 2.258599 2.1874363
## 1255 g at 0.8698382 0.9502466 0.8125414 1.012860 0.9441993
Annotation of genes in the expression matrix and removal of duplicate genes.
# Load the annotation file `anno_hug133plus2` in IOBR.
head(anno hug133plus2)
## # A tibble: 6 x 2
##
   probe_id symbol
    <fct>
##
              <fct>
## 1 1007_s_at MIR4640
## 2 1053_at
              RFC2
## 3 117 at
              HSPA6
## 4 121 at PAX8
## 5 1255 g at GUCA1A
## 6 1294 at
              MIR5193
# Conduct gene annotation using `anno_hug133plus2` file; If identical gene symbols exi
eset<-anno_eset(eset = eset,</pre>
               annotation = anno_hug133plus2,
               symbol = "symbol",
               probe = "probe_id",
               method = "mean")
eset[1:5, 1:3]
```

```
## GSM1523727 GSM1523728 GSM1523729
## SH3KBP1 4.327974 4.316195 4.351425
```

```
## RPL41 4.246149 4.246808 4.257940

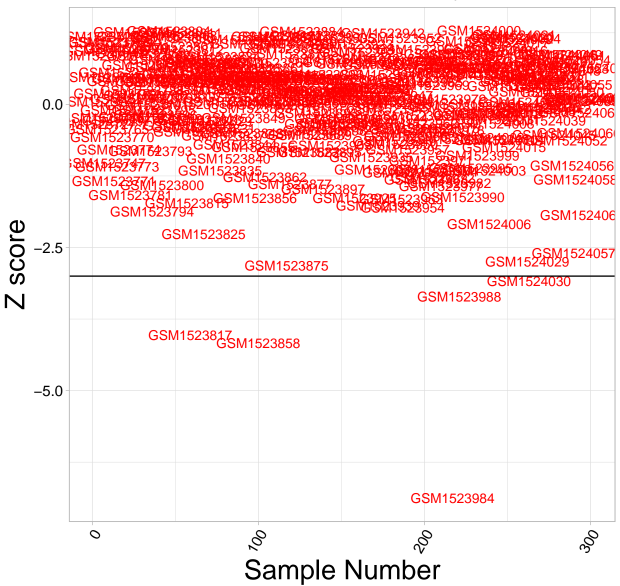
## EEF1A1 4.293762 4.291038 4.262199

## COX2 4.250288 4.283714 4.270508

## LOC101928826 4.219303 4.219670 4.213252
```

```
# source("E:/18-Github/Organization/IOBR/R/find_outlier_samples.R")
res <- find_outlier_samples(eset = eset, project = "ACRG", show_plot = TRUE)</pre>
```

Sample Connectivity



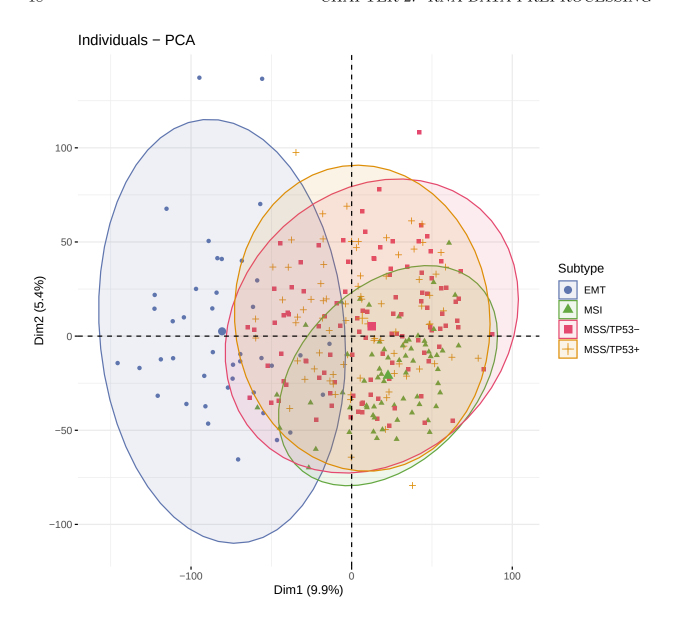
```
## [1] "GSM1523817" "GSM1523858" "GSM1523984" "GSM1523988" "GSM1524030" eset1 <- eset[, !colnames(eset)%in%res]
```

2.4 PCA analysis of molecular subtypes

```
data("pdata_acrg")
res<- iobr_pca(data = eset1,</pre>
```

```
is.matrix = TRUE,
scale = TRUE,
is.log
        = FALSE,
        = pdata_acrg,
pdata
id_pdata = "ID",
        = "Subtype",
group
geom.ind = "point",
cols
         = "normal",
palette = "jama",
repel
         = FALSE,
         = 5,
ncp
axes = c(1, 2),
addEllipses = TRUE)
```

```
##
##
        CIN
                  EBV
                            EMT
                                      GS
                                               MSI MSS/TP53- MSS/TP53+
                    0
                            42
                                       0
          0
                                                68
                                                         106
                                                                   79
##
## [1] ">>-- colors for PCA: #5f75ae" ">>-- colors for PCA: #64a841"
## [3] ">>-- colors for PCA: #e5486e" ">>-- colors for PCA: #de8e06"
res
```



2.5 Batch effect correction

Obtaining another data set from GEO Gastric cancer: GSE57303 using GEOquery R package.

```
# NOTE: This process may take a few minutes which depends on the internet connection s
eset_geo<-getGEO(GEO = "GSE57303", getGPL = F, destdir = "./")
eset2 <-eset_geo[[1]]
eset2 <-exprs(eset2)
eset2[1:5,1:5]</pre>
```

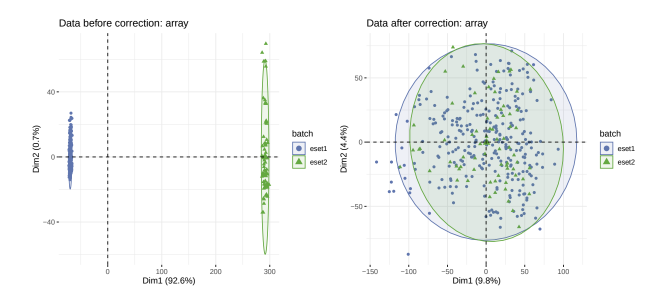
```
GSM1379261 GSM1379262 GSM1379263 GSM1379264 GSM1379265
##
## 1007 s at
             8.34746
                       9.67994
                                8.62643
                                          8.59301
                                                   8.63046
## 1053_at 5.07972 4.46377 5.29685
                                          5.78983
                                                   4.33359
## 117 at
            5.65558 4.48732 4.21615 5.47984
                                                   5.20816
## 121 at
            5.95123 7.09056 6.19903
                                          5.89872
                                                   5.91323
## 1255 g at
            1.66923 1.98758
                               1.73083
                                          1.56687
                                                   1.63332
```

Annotation of genes in the expression matrix and removal of duplicate genes.

```
GSM1379261 GSM1379262 GSM1379263 GSM1379264 GSM1379265
##
## ND4
           13.1695
                    13.1804
                             13.0600
                                       12.4544
                                               13.0457
## ATP6
          13.1433 13.0814 13.0502
                                      12.4831 13.1168
## SH3KBP1 12.9390 13.1620
                             12.9773
                                      12.8745 13.1169
## COX2
          13.0184 13.0489
                             12.8621 12.7489
                                               12.9732
## RPL41 13.0201
                    12.6034
                              12.7929
                                       13.0153
                                               12.9404
```

```
adjust_eset = TRUE,
repel = FALSE,
path = "result")
```

##
eset1 eset2
295 70
[1] ">>-- colors for PCA: #5f75ae" ">>-- colors for PCA: #64a841"
##
eset1 eset2
295 70
[1] ">>-- colors for PCA: #5f75ae" ">>-- colors for PCA: #64a841"



dim(eset_com)

[1] 21752 365

Chapter 3

Tumor ecosystem analysis

3.1 Determine TME subtype of gastric cancer using TME classifier

TME R TME classifier

##

```
library(TMEclassifier)
tme <- tme_classifier(eset = eset1, scale = TRUE)</pre>
## Step-1: Expression data preprocessing...
## Step-2: TME deconvolution...
## Step-3: Predicting TME phenotypes...
## [10:58:25] WARNING: amalgamation/../src/learner.cc:1040:
##
     If you are loading a serialized model (like pickle in Python, RDS in R) generated by
     older XGBoost, please export the model by calling `Booster.save_model` from that ve
##
##
     first, then load it back in current version. See:
##
##
       https://xgboost.readthedocs.io/en/latest/tutorials/saving model.html
##
```

for more details about differences between saving model and serializing.

```
##
## [10:58:25] WARNING: amalgamation/../src/learner.cc:749: Found JSON model saved before
## >>>--- DONE!
table(tme$TMEcluster)
##
##
  IA IE IS
## 106 94 95
head(tme)
                        ΙE
                                  IS
##
             ID
                                             IA TMEcluster
## 1 GSM1523727 0.21034596 0.1089890 0.68066503
                                                        IΑ
## 2 GSM1523728 0.01008306 0.1120120 0.87790490
                                                        ΙA
## 3 GSM1523729 0.85603729 0.1119151 0.03204757
                                                        ΙE
## 4 GSM1523744 0.05645817 0.0770771 0.86646473
                                                        ΙA
## 5 GSM1523745 0.05785239 0.8128097 0.12933787
                                                        IS
## 6 GSM1523746 0.55697476 0.3827613 0.06026396
                                                        ΙE
table(tme$TMEcluster)
##
## IA IE IS
## 106 94 95
head(tme)
##
             ID
                        ΙE
                                  IS
                                             IA TMEcluster
## 1 GSM1523727 0.21034596 0.1089890 0.68066503
                                                        ΙA
## 2 GSM1523728 0.01008306 0.1120120 0.87790490
                                                        ΙA
## 3 GSM1523729 0.85603729 0.1119151 0.03204757
                                                        ΙE
## 4 GSM1523744 0.05645817 0.0770771 0.86646473
                                                        ΙA
## 5 GSM1523745 0.05785239 0.8128097 0.12933787
                                                        IS
## 6 GSM1523746 0.55697476 0.3827613 0.06026396
                                                        ΙE
```

3.2. DEG ANALYSIS 23

3.2 DEG analysis

pdata <- tme[!tme\$TMEcluster=="IS",]</pre>

IA IE

3 HHIP

```
deg <- iobr_deg(eset</pre>
                        = eset,
                 annoation = NULL,
                 pdata = pdata,
                 group id = "TMEcluster",
                 pdata id = "ID",
                 array
                            = TRUE,
                            = "limma",
                 method
                            = c("deg_group","IA","IE"),
                 contrast
                            = NULL,
                 path
                 padj_cutoff = 0.01,
                 logfc cutoff = 0.5)
##
## Attaching package: 'limma'
## The following object is masked from 'package:BiocGenerics':
##
      plotMA
##
## group1 = IA
## group2 = IE
## # A tibble: 6 x 11
##
    symbol log2FoldChange AveExpr t pvalue
                                                  padj
                                                           B sigORnot
                                                                        label
                           <dbl> <dbl> <dbl>
                                                 <dbl> <dbl> <chr>
##
    <chr>>
                   <dbl>
                                                                        <chr>>
## 1 TMEM100
                  -0.755 1.83 -14.0 1.54e-31 3.35e-27 60.9 Down regul~ Both
## 2 ABCA8
                  -0.923 1.89 -12.9 3.18e-28 3.46e-24 53.4 Down regul~ Both
```

-0.625 1.74 -12.4 1.52e-26 1.10e-22 49.6 Down regul~ Both

3.3 GSEA analysis of DEGs

IOBR signature collection

```
head(deg)
```

```
## # A tibble: 6 x 11
     symbol log2FoldChange AveExpr t
                                                                 B sigORnot
##
                                            pvalue
                                                       padj
                                                                               label
##
     <chr>>
                      <dbl>
                              <dbl> <dbl>
                                              <dbl>
                                                       <dbl> <dbl> <chr>
                                                                               <chr>
                     -0.755
## 1 TMEM100
                               1.83 -14.0 1.54e-31 3.35e-27 60.9 Down regul~ Both
## 2 ABCA8
                     -0.923
                               1.89 -12.9 3.18e-28 3.46e-24 53.4 Down regul~ Both
## 3 HHIP
                     -0.625
                               1.74 -12.4 1.52e-26 1.10e-22 49.6 Down regul~ Both
## 4 ADH1B
                     -0.901
                               1.84 -12.2 4.71e-26 2.09e-22 48.5 Down regul~ Both
## 5 LMNB2
                                     12.2 4.80e-26 2.09e-22 48.5 NOT
                      0.285
## 6 FCER1A
                     -0.552
                               1.57 -12.0 2.34e-25 7.27e-22 46.9 Down_regul~ Both
## # i 2 more variables: IA <dbl>, IE <dbl>
sig_list <- signature_collection[c("TMEscoreB_CIR", "TMEscoreA_CIR", "DNA_replication",</pre>
                                   "Pan_F_TBRs", "TGFb.myCAF", "Ferroptosis", "TLS_Natur
sig list
## $TMEscoreB_CIR
##
     [1] "DCN"
                        "SEPP1"
                                        "ACTA2"
                                                       "SPARCL1"
                                                                      "BEX3"
     [6] "MYLK"
                        "AKR1C1"
                                        "TIMP2"
                                                       "MXRA7"
##
                                                                      "C11orf96"
                        "PDGFRA"
                                                       "MGP"
    [11] "CAV1"
                                        "FHL1"
                                                                      "EID1"
##
    [16] "LOC101930400" "DST"
                                        "GREM1"
                                                       "FERMT2"
                                                                      "TNC"
##
    [21] "CYBRD1"
                        "LTBP1"
                                        "ACTG2"
                                                       "TMEM47"
                                                                      "SERPINE2"
```

##	[26]	"ANTXR2"	"GNG11"	"TAGLN"	"GSTA4"	"PKIG"
##	[31]	"MAOA"	"PTRF"	"FAM3B"	"PBX1"	"WLS"
##	[36]	"SELM"	"SVIL"	"MYH11"	"AGT"	"SPON1"
##	[41]	"TGFB1I1"	"PDLIM3"	"PDK4"	"SYNPO2"	"MSRB3"
##	[46]	"PROS1"	"EDNRA"	"AKAP12"	"PSD3"	"TNS1"
##	[51]	"JAM3"	"PDZRN3"	"DDR2"	"HMGCS2"	"SGCE"
##	[56]	"MRVI1"	"WFDC1"	"FBLN1"	"FM05"	"MAOB"
##	[61]	"AMOTL1"	"AKT3"	"CNRIP1"	"CPE"	"MAP1B"
##	[66]	"RBP1"	"GNAI1"	"FOXF2"	"SORBS2"	"ZCCHC24"
##	[71]	"ZNF704"	"ARMCX1"	"DIXDC1"	"SSTR1"	"THRB"
##	[76]	"C3orf70"	"PKIB"	"CNN1"	"SYTL5"	"DACT1"
##	[81]	"SYNPO"	"GAS1"	"DPYSL3"	"CCDC80"	"TSPYL5"
##	[86]	"DCHS1"	"SOBP"	"AOC3"	"NDN"	"FGF7P3"
##	[91]	"SMAD9"	"MCC"	"CLMP"	"MYL9"	"RBP4"
##	[96]	"PLN"	"SPOCK1"	"COL14A1"	"CRYAB"	"SRPX"
##	[101]	"EML1"	"RERG"	"PPP1R3C"	"L0C100506718"	"CH25H"
##	[106]	"HSPB8"	"PID1"	"TTC28"	"STON1"	"ABCG2"
##	[111]	"ZSCAN18"	"SCIN"	"C14orf132"	"TMEM55A"	"WASF3"
##	[116]	"PAPLN"	"COLEC12"	"ACKR1"	"TMEM150C"	"RAI2"
##	F 7					
##	[121]	"TSPAN7"	"MRGPRF"	"ABCA8"	"CHIC1"	"NBEA"
ππ	[121]		"MRGPRF" "SETBP1"	"ABCA8" "LDOC1"	"CHIC1" "TMEM100"	"NBEA" "LOC101930349"
	[126]					
##	[126] [131]	"FAM13C"	"SETBP1"	"LDOC1"	"TMEM100"	"L0C101930349"
## ##	[126] [131] [136]	"FAM13C" "PRICKLE2"	"SETBP1" "TSPAN18"	"LDOC1" "FABP4"	"TMEM100" "ARHGEF26"	"LOC101930349" "ERICH5"
## ## ##	[126] [131] [136] [141]	"FAM13C" "PRICKLE2" "MYOCD"	"SETBP1" "TSPAN18" "BEX2"	"LDOC1" "FABP4" "PPP1R14A"	"TMEM100" "ARHGEF26" "FGF13"	"LOC101930349" "ERICH5" "RUNX1T1"
## ## ##	[126] [131] [136] [141] [146]	"FAM13C" "PRICKLE2" "MYOCD" "MAGI2-AS3"	"SETBP1" "TSPAN18" "BEX2" "LINC01279"	"LDOC1" "FABP4" "PPP1R14A" "REEP1"	"TMEM100" "ARHGEF26" "FGF13" "PLAC9"	"LOC101930349" "ERICH5" "RUNX1T1" "MYEF2"
## ## ## ##	[126] [131] [136] [141] [146] [151]	"FAM13C" "PRICKLE2" "MYOCD" "MAGI2-AS3" "PRKD1"	"SETBP1" "TSPAN18" "BEX2" "LINC01279" "RGN"	"LDOC1" "FABP4" "PPP1R14A" "REEP1" "CLDN11"	"TMEM100" "ARHGEF26" "FGF13" "PLAC9" "ANK2"	"LOC101930349" "ERICH5" "RUNX1T1" "MYEF2" "ESRRG"
## ## ## ## ##	[126] [131] [136] [141] [146] [151] [156]	"FAM13C" "PRICKLE2" "MYOCD" "MAG12-AS3" "PRKD1" "SYNC"	"SETBP1" "TSPAN18" "BEX2" "LINCO1279" "RGN" "ZNF667-AS1" "OGN"	"LDOC1" "FABP4" "PPP1R14A" "REEP1" "CLDN11" "FGF7"	"TMEM100" "ARHGEF26" "FGF13" "PLAC9" "ANK2" "SFRP1"	"LOC101930349" "ERICH5" "RUNX1T1" "MYEF2" "ESRRG" "HMCN1"
## ## ## ## ##	[126] [131] [136] [141] [146] [151] [156]	"FAM13C" "PRICKLE2" "MYOCD" "MAG12-AS3" "PRKD1" "SYNC" "TCEAL7" "LOC100507334"	"SETBP1" "TSPAN18" "BEX2" "LINCO1279" "RGN" "ZNF667-AS1" "OGN"	"LDOC1" "FABP4" "PPP1R14A" "REEP1" "CLDN11" "FGF7" "MAGI2"	"TMEM100" "ARHGEF26" "FGF13" "PLAC9" "ANK2" "SFRP1" "MIR100HG"	"LOC101930349" "ERICH5" "RUNX1T1" "MYEF2" "ESRRG" "HMCN1" "FILIP1"

##	[176]	[176] "SERP2"		"ZNF415"		"MAMDC2"		"RBM2	24"	"MEOX2"	
##	‡										
##	## \$TMEscoreA_CIR										
##	[1]	"HLA-DPB1"		"UBD"		"L0C100509457"		9457"	"WARS"		
##	[5]	"TAP1"		"HLA-DMA"		"TRIM22"		"PSAT1"			
##	[9]	"CXCL10"		"SOCS3"		"CXCL9"		"PBK"			
##	[13]	"CCL4"		"CCL5"		"BCL2A1"		"TRBC1"			
##	[17]	"ID01"		"NFE2L3"		"CCL3L3"		"DTL"			
##	[21]	"MMP9"		"SLC2A	"SLC2A3" "ZNF367"			"RCC1"			
##	[25]	"STIL"		"TRAC"	"RAC" "HELLS"			"GZMB"			
##	[29]	"RTEL1-TNFF	RSF6B"	"CXCL1	11"		"GBP5"		"CD2	"CD2"	
##	[33]	"CDCA2"		"CDT1"	1		"TNFAIP2"		"TYMP"		
##	[37]	"MICB"		"SLC2A	114"	"GZMK" "		"CD8A"			
##	[41]	"CENPH"		"MND1"	"MND1" "BATF2"			"BRIP1"			
##	[45]	"E2F7"		"KIF18A"		"AIM2"		"ETV7"			
##	[49]	"ITK"		"GNLY"			"GPR171"		"WDHD1"		
##	[53]	"GBP4"		"MB21D1" "NLF		"NLRP3" "MCE		EMP1"			
##	[57]	"POLR3G"		"NLRC3"		"KLRC2"		"CLE	EC5A"		
##	[61]	"ARHGAP11A'	ıı	"GPR84" "IFNG"			"ZBED2"				
##											
##	\$DNA_	replication	ı								
##	[1]	"RNASEH2A"	"POLD3	3" "	'DNA2"	11	FEN1"	"POLA2	•	"RNASEH1"	
##	[7]	"RPA4"	"LIG1	11	'MCM2"	11	MCM3"	"MCM4"		"MCM5"	
##	[13]	"MCM6"	"MCM7"	11	'PCNA"	"	POLE3"	"POLA1	1	"POLD1"	
##	[19]	"POLD2"	"POLE	11	'POLE2"	"	'PRIM1"	"PRIM2"	1	"POLE4"	
##	[25]	"POLD4"	"RFC1	11 11	'RFC2"	11	'RFC3"	"RFC4"		"RFC5"	
##	[31]	"RPA1"	"RPA2	11	'RPA3"	11	SSBP1"	"RNASEI	12B"	"RNASEH2C"	
##											
##	\$Base	e_excision_	repair								
##	[1]	"PARP2" "PA	ARP3" '	"POLD3"	"PARP1"	"P	PARP4" "FE	N1" "SN	'UG1	' "NEIL2" "APEX2"	

```
## [10] "POLL"
                "HMGB1" "APEX1" "LIG1" "LIG3"
                                                  "MPG"
                                                           "MUTYH" "NTHL1" "OGG1"
## [19] "PCNA"
                "POLE3" "POLB"
                                 "POLD1" "POLD2" "POLE"
                                                           "POLE2" "NEIL3" "POLE4"
## [28] "POLD4" "UNG"
                         "XRCC1" "NEIL1" "MBD4"
##
## $Pan_F_TBRs
    [1] "ACTA2"
                    "ACTG2"
                                "ADAM12"
                                           "ADAM19"
                                                       "CNN1"
                                                                   "COL4A1"
##
    [7] "CTGF"
##
                    "CTPS1"
                               "FAM101B"
                                           "FSTL3"
                                                       "HSPB1"
                                                                   "IGFBP3"
## [13] "PXDC1"
                               "SH3PXD2A" "TAGLN"
                    "SEMA7A"
                                                       "TGFBI"
                                                                   "TNS1"
## [19] "TPM1"
##
## $TGFb.myCAF
##
    [1] "CST1"
                   "LAMP5"
                             "LOXL1"
                                        "EDNRA"
                                                   "TGFB1"
                                                             "TGFB3"
                                                                        "TNN"
    [8] "CST2"
                             "COL10A1" "ELN"
##
                   "HES4"
                                                   "THBS4"
                                                             "NKD2"
                                                                        "OLFM2"
## [15] "COL6A3"
                   "LRRC17"
                             "COL3A1"
                                        "THY1"
                                                   "HTRA3"
                                                             "TMEM204" "11-Sep"
## [22] "COMP"
                   "TNFAIP6" "ID4"
                                        "GGT5"
                                                   "INAFM1"
                                                             "CILP"
                                                                        "OLFML2B"
##
## $Ferroptosis
    [1] "ACSL4"
                      "AKR1C1-3"
                                    "ALOXs"
                                                  "ATP5G3"
                                                               "CARS"
##
    [6] "CBS"
                      "CD44v"
                                    "CHAC1"
                                                  "CISD1"
                                                               "CS"
##
                      "FANCD2"
## [11] "DPP4"
                                    "GCLC/GCLM"
                                                  "GLS2"
                                                               "GPX4"
## [16] "GSS"
                                    "HSPB1/5"
                                                  "KOD"
                                                               "LPCAT3"
                      "HMGCR"
## [21] "MT1G"
                      "NCOA4"
                                    "NFE2L2"
                                                  "PTGS2"
                                                               "RPL8"
## [26] "SAT1"
                      "SLC7A11"
                                                  "TFRC"
                                                               "TP53"
                                    "SQS"
## [31] "TTC35/EMC2" "MESH1"
##
## $TLS Nature
## [1] "CD79B"
                "CD1D"
                                                                "EIF1AY" "RBP5"
                          "CCR6"
                                    "LAT"
                                             "SKAP1"
                                                       "CETP"
## [9] "PTGDS"
##
## $Glycolysis
```

```
[1] "ACSS1"
                   "ACSS2"
                             "ADH1A"
                                        "ADH1B"
                                                   "ADH1C"
                                                             "ADH4"
                                                                        "ADH5"
##
    [8] "ADH6"
                   "ADH7"
                             "ADPGK"
                                        "AKR1A1"
                                                   "ALDH1A3" "ALDH1B1" "ALDH2"
##
## [15] "ALDH3A1" "ALDH3A2"
                             "ALDH3B1" "ALDH3B2"
                                                  "ALDH7A1" "ALDH9A1" "ALDOA"
   [22] "ALDOB"
                   "ALDOC"
                             "BPGM"
                                        "DLAT"
                                                   "DLD"
                                                             "EN01"
                                                                        "EN02"
  [29] "ENO3"
                   "FBP1"
                             "FBP2"
                                        "G6PC"
                                                   "G6PC2"
                                                             "GALM"
                                                                        "GAPDH"
## [36] "GAPDHS"
                             "GPI"
                                                   "HK2"
                                                             "HK3"
                                                                        "HKDC1"
                   "GCK"
                                        "HK1"
## [43] "LDHA"
                   "LDHAL6A" "LDHAL6B" "LDHB"
                                                   "LDHC"
                                                             "PANK1"
                                                                        "PCK1"
## [50] "PCK2"
                             "PDHA2"
                   "PDHA1"
                                        "PDHB"
                                                   "PFKFB1"
                                                             "PFKFB2"
                                                                        "PFKFB3"
## [57] "PFKFB4"
                   "PFKL"
                             "PFKM"
                                        "PFKP"
                                                   "PGAM1"
                                                             "PGAM2"
                                                                        "PGAM4"
## [64] "PGK1"
                   "PGK2"
                             "PGM1"
                                        "PGM2"
                                                   "PKLR"
                                                             "PKM"
                                                                        "SLC2A2"
## [71] "TPI1"
```

3.4 DEG analysis

find_markers_in_bulk TME

```
##
## IA IE IS
## 106 94 95
## # A tibble: 56 x 7
```

3.4. DEG ANALYSIS 29

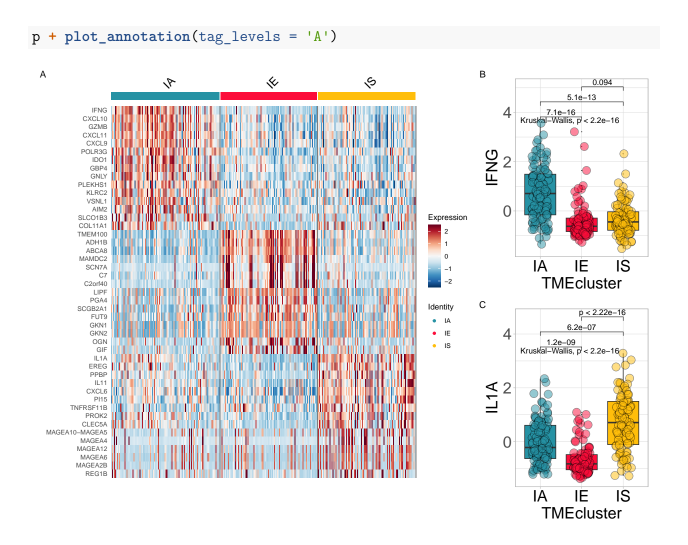
```
## # Groups:
              cluster [3]
        p_val avg_log2FC pct.1 pct.2 p_val_adj cluster gene
##
        <dbl>
                   <dbl> <dbl> <dbl> <fct>
##
                                                     <chr>>
   1 4.66e-19
                   0.208
                                  1 1.01e-14 IA
                                                     IFNG
##
                            1
##
   2 7.41e-18
                   0.170
                                  1 1.61e-13 IA
                                                     CXCL10
                            1
## 3 1.21e-15
                   0.177
                                  1 2.62e-11 IA
                                                     GZMB
                            1
## 4 1.34e-15
                   0.247
                                  1 2.91e-11 IA
                                                     CXCL11
## 5 5.33e-15
                                  1 1.16e-10 IA
                                                     CXCL9
                   0.166
## 6 6.17e-15
                                  1 1.34e-10 IA
                                                     POLR3G
                   0.158
## 7 3.06e-14
                   0.214
                                  1 6.66e-10 IA
                                                     IDO1
                            1
## 8 1.95e-13
                                  1 4.24e- 9 IA
                                                     GBP4
                   0.179
                            1
## 9 4.17e-13
                   0.150
                            1
                                  1 9.07e- 9 IA
                                                     ZBED2
## 10 3.74e-11
                   0.153
                                  1 8.13e- 7 IA
                                                     GNLY
## # i 46 more rows
```

top15 <- res\$top_markers %>% dplyr:: group_by(cluster) %>% dplyr::top_n(15, avg_log2F0
top15\$gene

##	[1]	"IFNG"	"CXCL10"	"GZMB"	"CXCL11"
##	[5]	"CXCL9"	"POLR3G"	"ID01"	"GBP4"
##	[9]	"GNLY"	"PLEKHS1"	"KLRC2"	"VSNL1"
##	[13]	"AIM2"	"SLC01B3"	"COL11A1"	"TMEM100"
##	[17]	"ADH1B"	"ABCA8"	"MAMDC2"	"SCN7A"
##	[21]	"C7"	"C2orf40"	"LIPF"	"PGA4"
##	[25]	"SCGB2A1"	"FUT9"	"GKN1"	"GKN2"
##	[29]	"OGN"	"GIF"	"IL1A"	"EREG"
##	[33]	"PPBP"	"IL11"	"CXCL6"	"PI15"
##	[37]	"TNFRSF11B"	"PROK2"	"CLEC5A"	"MAGEA10-MAGEA5"
##	[41]	"MAGEA4"	"MAGEA12"	"MAGEA6"	"MAGEA2B"
##	[45]	"REG1B"			

Seurat DoHeatmap

```
cols <- c('#2692a4','#fc0d3a','#ffbe0b')</pre>
p1 <- DoHeatmap(res$sce, top15$gene, group.colors = cols )+
  scale_fill_gradientn(colours = rev(colorRampPalette(RColorBrewer::brewer.pal(11, "RdBu
      TME
input <- combine_pd_eset(eset = eset, pdata = tme, feas = top15$gene, scale = T)
p2 <- sig_box(input, variable = "TMEcluster", signature = "IFNG", jitter = TRUE,
              cols = cols, show pvalue = TRUE, size of pvalue = 4)
## # A tibble: 3 x 8
##
               group1 group2
                                         p.adj p.format p.signif method
     .у.
                                    р
               <chr> <chr>
                                <dbl>
                                         <dbl> <chr>
##
     <chr>
                                                         <chr>
                                                                  <chr>
                             7.08e-16 2.10e-15 7.1e-16 ****
## 1 signature IA
                      IE
                                                                  Wilcoxon
## 2 signature IA
                      IS
                             5.12e-13 1 e-12 5.1e-13 ****
                                                                  Wilcoxon
## 3 signature IE
                      IS
                             9.36e- 2 9.4 e- 2 0.094
                                                                  Wilcoxon
                                                        ns
p3 <- sig_box(input, variable = "TMEcluster", signature = "IL1A",
              jitter = TRUE, cols = cols, show pvalue = TRUE, size of pvalue = 4)
## # A tibble: 3 x 8
##
               group1 group2
                                         p.adj p.format p.signif method
     .v.
                                    р
     <chr>
               <chr>
                      <chr>
                                <dbl>
                                         <dbl> <chr>
                                                         <chr>
                                                                  <chr>
##
## 1 signature IA
                      ΙE
                             1.17e- 9 2.3 e- 9 1.2e-09 ****
                                                                  Wilcoxon
                             6.19e- 7 6.20e- 7 6.2e-07 ****
## 2 signature IA
                      IS
                                                                  Wilcoxon
## 3 signature IE
                      IS
                             4.51e-19 1.4 e-18 < 2e-16 ****
                                                                  Wilcoxon
if (!requireNamespace("patchwork", quietly = TRUE)) install.packages("patchwork")
library(patchwork)
p \leftarrow (p1|p2/p3) + plot_layout(widths = c(2.3,1))
```



3.5 Identifying signatures associated with TME clusters

Calculate TME associated signatures-(through PCA method).

sig_tme[1:5, 1:3]

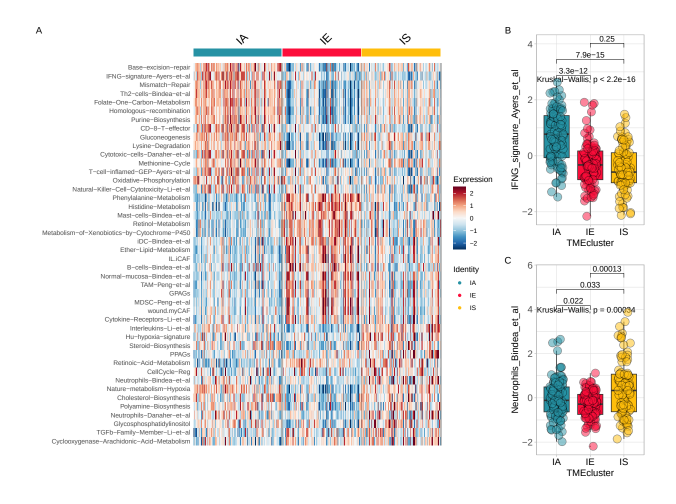
```
## CD_8_T_effector -2.5513794 0.7789141 -2.1770675
## DDR -0.8747614 0.7425162 -1.3272054
## APM 1.1098368 2.1988688 -0.9516419
## Immune_Checkpoint -2.3701787 0.9455120 -1.4844104
## CellCycle_Reg 0.1063358 0.7583302 -0.3649795
```

TMEcluster

```
res <- find_markers_in_bulk(pdata = tme, eset = sig_tme, group = "TMEcluster", nfeature
```

```
##
##
   ΙA
       IE IS
       94 95
## 106
## # A tibble: 60 x 7
## # Groups:
              cluster [3]
        p_val avg_log2FC pct.1 pct.2 p_val_adj cluster gene
##
         <dbl>
                    <dbl> <dbl> <dbl>
                                          <dbl> <fct>
##
                                                        <chr>>
    1 2.73e-18
                    17.8 0.792 0.365 6.98e-16 IA
##
                                                        Base-excision-repair
   2 8.18e-18
                    3.64 0.698 0.323 2.09e-15 IA
##
                                                        IFNG-signature-Ayers-et-al
   3 9.03e-18
                    8.94 0.84 0.381 2.31e-15 IA
##
                                                        Mismatch-Repair
   4 9.73e-18
                    4.17 0.821 0.402 2.49e-15 IA
                                                        Th2-cells-Bindea-et-al
##
   5 1.20e-16
                    3.97 0.821 0.381 3.08e-14 IA
                                                        Folate-One-Carbon-Metaboli~
##
   6 1.14e-15
                    4.36 0.811 0.354 2.91e-13 IA
                                                        Homologous-recombination
## 7 2.13e-14
                    3.09 0.83 0.397 5.45e-12 IA
                                                        Purine-Biosynthesis
## 8 3.61e-14
                    4.20 0.67 0.28
                                       9.25e-12 IA
                                                        CD-8-T-effector
## 9 1.49e-13
                    2.70 0.708 0.344 3.82e-11 IA
                                                        Th1-cells-Bindea-et-al
## 10 7.03e-13
                    2.61 0.792 0.429 1.80e-10 IA
                                                        TIP-Release-of-cancer-cell~
## # i 50 more rows
```

```
top15 <- res$top markers %>% dplyr:: group_by(cluster) %>% dplyr::top_n(15, avg log2F
p1 <- DoHeatmap(res$sce, top15$gene, group.colors = cols)+
  scale_fill_gradientn(colours = rev(colorRampPalette(RColorBrewer::brewer.pal(11, "RdBu))
top15$gene <- gsub(top15$gene, pattern = "\\-", replacement = "\\_")
input <- combine_pd_eset(eset = sig_tme, pdata = tme, feas = top15$gene, scale = T)</pre>
p2 <- sig_box(input, variable = "TMEcluster", signature = "IFNG signature Ayers et al",
              cols = cols, show pvalue = TRUE, size of pvalue = 4, size of font = 6)
## # A tibble: 3 x 8
##
     .у.
               group1 group2
                                         p.adj p.format p.signif method
                                    р
##
     <chr>
               <chr>
                      <chr>
                                <dbl>
                                         <dbl> <chr>
                                                         <chr>
                                                                  <chr>>
                             3.34e-12 6.70e-12 3.3e-12 ****
## 1 signature IA
                      ΙE
                                                                  Wilcoxon
## 2 signature IA
                      IS
                             7.93e-15 2.40e-14 7.9e-15 ****
                                                                  Wilcoxon
## 3 signature IE
                      IS
                             2.54e- 1 2.5 e- 1 0.25
                                                                  Wilcoxon
                                                         ns
p3 <- sig_box(input, variable = "TMEcluster", signature = "Neutrophils_Bindea_et_al",
              jitter = TRUE, cols = cols, show_pvalue = TRUE, size_of_pvalue = 4, size_
## # A tibble: 3 x 8
##
     .у.
               group1 group2
                                    p p.adj p.format p.signif method
     <chr>
                                <dbl> <dbl> <chr>
##
               <chr> <chr>
                                                       <chr>
                                                                <chr>
## 1 signature IA
                      ΙE
                             0.0222
                                      0.044 0.02217 *
                                                                Wilcoxon
## 2 signature IA
                      IS
                             0.0333
                                      0.044 0.03326 *
                                                                Wilcoxon
                             0.000133 0.0004 0.00013 ***
## 3 signature IE
                      IS
                                                                Wilcoxon
p \leftarrow (p1|p2/p3) + plot_layout(widths = c(2.3,1))
p + plot_annotation(tag_levels = 'A')
```



library(survminer)

##

>>> Dataset's survival follow up time is range between 1 to 105.7 months

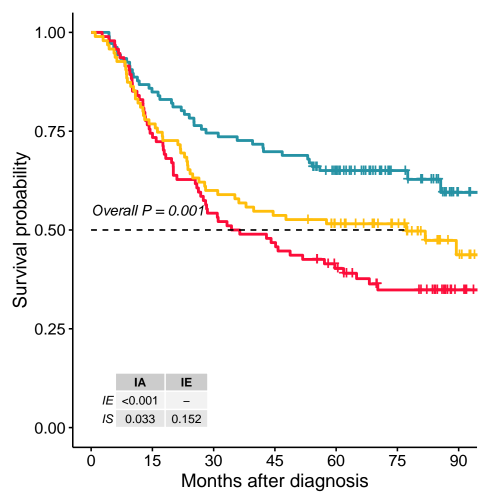
```
## IA IE IS
## 106 94 95
```

1069495

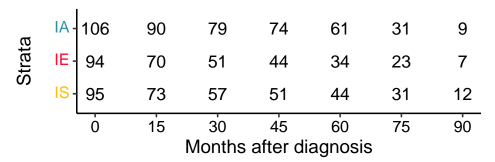
```
## Maximum of follow up time is 105.7 months; and will be divided into 6 sections;
p1
```

TMEcluster-in-ACRG





Number at risk



p1<- percent_bar_plot(input, x = "TMEcluster" , y = "Subtype", palette = "jama")</pre>

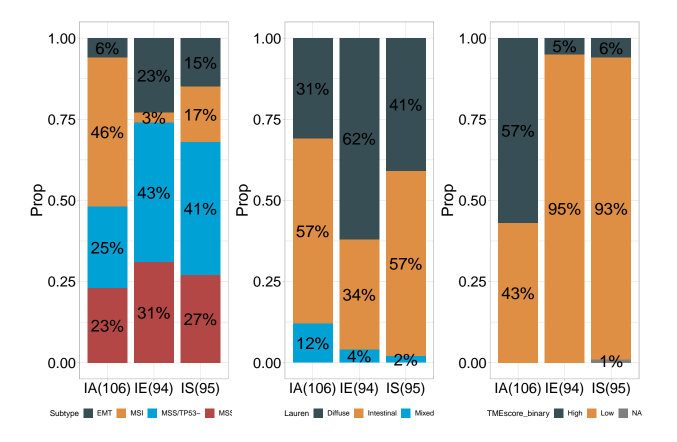
A tibble: 12 x 5

```
## # Groups:
              TMEcluster [3]
##
     TMEcluster Subtype Freq Prop count
                <fct> <dbl> <dbl> <dbl>
##
     <chr>
                              6 0.06
                                       106
##
   1 IA
                EMT
##
   2 IA
                MSI
                            49 0.46
                                       106
##
   3 IA
                MSS/TP53-
                            27 0.25
                                       106
##
   4 IA
                MSS/TP53+
                            24 0.23
                                       106
  5 IE
                EMT
                             22 0.23
##
                                        94
  6 IE
                MSI
                             3 0.03
##
                                        94
  7 IE
                MSS/TP53-
                            40 0.43
                                        94
##
  8 IE
                MSS/TP53+
                            29 0.31
                                        94
##
## 9 IS
                EMT
                            14 0.15
                                        95
## 10 IS
                MSI
                            16 0.17
                                        95
                MSS/TP53-
## 11 IS
                            39 0.41
                                        95
## 12 IS
                MSS/TP53+
                             26 0.27
                                        95
## [1] "'#374E55FF', '#DF8F44FF', '#00A1D5FF', '#B24745FF', '#79AF97FF', '#6A6599FF', '#
p2<- percent_bar_plot(input, x = "TMEcluster", y = "Lauren", palette = "jama")
## # A tibble: 9 x 5
## # Groups:
              TMEcluster [3]
##
    TMEcluster Lauren
                        Freq Prop count
```

```
##
    <chr>
              <fct> <dbl> <dbl> <dbl>
## 1 IA
                            33 0.31
                                      106
              Diffuse
## 2 IA
                            60 0.57
              Intestinal
                                      106
## 3 IA
                           13 0.12
                                      106
              Mixed
## 4 IE
              Diffuse
                           58 0.62
                                       94
## 5 IE
                            32 0.34
              Intestinal
                                       94
                           4 0.04
## 6 IE
              Mixed
                                       94
## 7 IS
              Diffuse
                            39 0.41
                                       95
## 8 IS
              Intestinal
                           54 0.57
                                       95
```

```
## 9 IS Mixed 2 0.02 95
## [1] "'#374E55FF', '#DF8F44FF', '#00A1D5FF', '#B24745FF', '#79AF97FF', '#6A6599FF', '#
p3<- percent_bar_plot(input, x = "TMEcluster", y = "TMEscore_binary", palette = "jama")</pre>
```

```
## # A tibble: 7 x 5
## # Groups: TMEcluster [3]
## TMEcluster TMEscore_binary Freq Prop count
##
    <chr> <fct> <dbl> <dbl> <dbl><</pre>
## 1 IA
          High
                              60 0.57
                                        106
## 2 IA
                                        106
                            46 0.43
           Low
## 3 IE
                            5 0.05
          High
                                       94
                           89 0.95
                                         94
## 4 IE
             Low
## 5 IS
             High
                            6 0.06
                                         95
## 6 IS
                              88 0.93
                                         95
             Low
## 7 IS
             <NA>
                              1 0.01
                                         95
## [1] "'#374E55FF', '#DF8F44FF', '#00A1D5FF', '#B24745FF', '#79AF97FF', '#6A6599FF', '#
p1|p2|p3
```



Chapter 4

Signatures and relevant phenotypes

4.1 Loading packages

Load the IOBR package in your R session after the installation is complete:

```
library(IOBR)
library(survminer)
library(tidyverse)
```

4.2 Downloading data for example

Obtaining data set from GEO Gastric cancer: GSE62254 using GEOquery R package.

```
if (!requireNamespace("GEOquery", quietly = TRUE)) BiocManager::install("GEOquery")
library("GEOquery")
# NOTE: This process may take a few minutes which depends on the internet connection s
eset_geo <- getGEO(GEO = "GSE62254", getGPL = F, destdir = "./")
eset <-eset_geo[[1]]
eset <-exprs(eset)
eset[1:5,1:5]</pre>
```

```
GSM1523727 GSM1523728 GSM1523729 GSM1523744 GSM1523745
##
## 1007 s at 3.2176645 3.0624323 3.0279131 2.921683 2.8456013
## 1053_at 2.4050109 2.4394879 2.2442708 2.345916 2.4328582
## 117 at 1.4933412 1.8067380 1.5959665 1.839822 1.8326058
## 121_at
           2.1965561 2.2812181 2.1865556 2.258599 2.1874363
## 1255_g_at 0.8698382 0.9502466 0.8125414 1.012860 0.9441993
Annotation of genes in the expression matrix and removal of duplicate genes.
# Load the annotation file `anno_hug133plus2` in IOBR.
head(anno_hug133plus2)
## # A tibble: 6 x 2
##
   probe_id symbol
    <fct>
##
              <fct>
## 1 1007_s_at MIR4640
## 2 1053_at
              RFC2
## 3 117 at
              HSPA6
## 4 121_at PAX8
## 5 1255 g at GUCA1A
## 6 1294 at
              MIR5193
# Conduct gene annotation using `anno_hug133plus2` file; If identical gene symbols exi
eset<-anno_eset(eset = eset,</pre>
               annotation = anno_hug133plus2,
               symbol = "symbol",
               probe = "probe_id",
               method = "mean")
```

```
## GSM1523727 GSM1523728 GSM1523729
## SH3KBP1 4.327974 4.316195 4.351425
```

eset[1:5, 1:3]

```
## RPL41 4.246149 4.246808 4.257940

## EEF1A1 4.293762 4.291038 4.262199

## COX2 4.250288 4.283714 4.270508

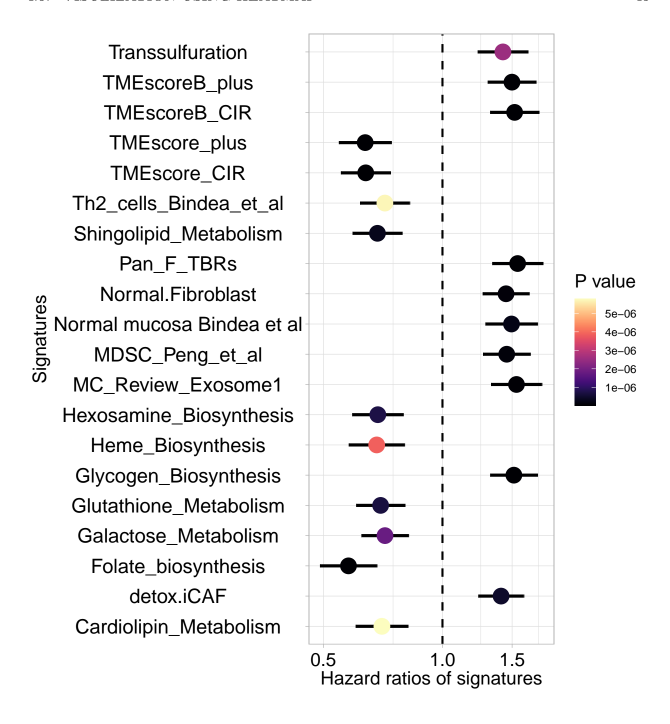
## LOC101928826 4.219303 4.219670 4.213252
```

4.3 Signature score estimation

4.4 Identifying features associated with survival

head(res)

```
## # A tibble: 6 x 5
                                    HR CI_low_0.95 CI_up_0.95
##
    ID
                                Ρ
                            <dbl> <dbl>
                                             <dbl>
                                                        <dbl>
##
    <chr>
## 1 Folate_biosynthesis 1.00e-10 0.579
                                             0.490
                                                        0.683
## 2 TMEscore CIR
                    1.32e- 9 0.640
                                             0.554
                                                        0.739
## 3 Glycogen Biosynthesis 3.24e- 9 1.52
                                            1.32
                                                       1.74
## 4 Pan_F_TBRs
                       6.33e- 9 1.55
                                            1.34
                                                        1.80
                      7.17e- 9 1.52
## 5 TMEscoreB_CIR
                                            1.32
                                                        1.75
## 6 TMEscore_plus
                        8.08e- 9 0.638
                                             0.547
                                                        0.743
res<- res[nchar(res$ID)<=28, ]
p1<- sig_forest(res, signature = "ID", n = 20)
```



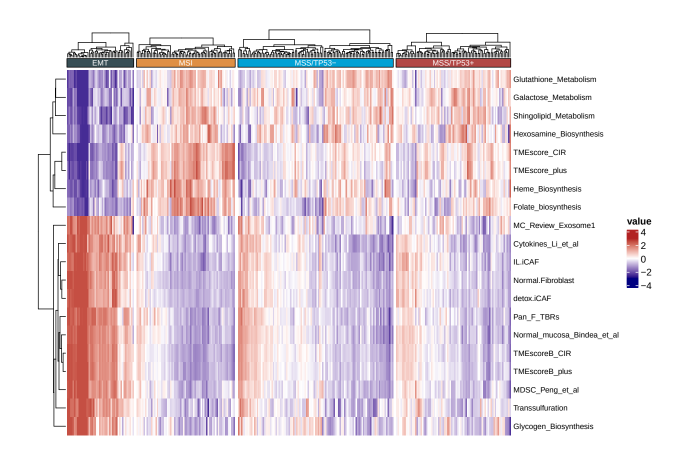
4.5 Visulization using heatmap

```
Signatures IOBR sig_heatmap
```

```
group = "Subtype",

palette_group = "jama",

palette = 6)
```



4.6 Focus on target signatures

```
size_of_pvalue = 5,
hjust = 1,
angle_x_text = 60,
size_of_font = 8)
```

```
## # A tibble: 6 x 8
##
               group1
                         group2
                                               p.adj p.format p.signif method
     .у.
                                          p
                                                              <chr>
##
     <chr>
               <chr>
                         <chr>
                                      <dbl>
                                               <dbl> <chr>
                                                                        <chr>
                                   5.39e-15 3.20e-14 5.4e-15 ****
## 1 signature EMT
                         MSI
                                                                       Wilcoxon
## 2 signature EMT
                         MSS/TP53- 5.53e-13 2.8 e-12 5.5e-13
                                                              ****
                                                                       Wilcoxon
                         MSS/TP53+ 1.90e-12 7.6 e-12 1.9e-12
## 3 signature EMT
                                                              ****
                                                                       Wilcoxon
## 4 signature MSI
                         MSS/TP53- 1.14e- 3 3.4 e- 3 0.0011
                                                                       Wilcoxon
## 5 signature MSI
                         MSS/TP53+ 7.05e- 3 1.4 e- 2 0.0071
                                                                       Wilcoxon
## 6 signature MSS/TP53- MSS/TP53+ 7.16e- 1 7.2 e- 1 0.7161
                                                                       Wilcoxon
                                                              ns
p2 <- sig_box(data
                             = input,
                             = "Pan F TBRs",
              signature
                             = "Subtype",
              variable
              jitter
                             = TRUE,
              cols
                             = NULL,
                        = "jama",
              palette
              show_pvalue
                            = TRUE,
              angle_x_text
                             = 60,
              hjust
                             = 1,
              size of pvalue = 5,
              size of font
                             = 8)
```

```
## # A tibble: 6 x 8
                                              p.adj p.format p.signif method
##
              group1
                        group2
     . V .
                                          р
                                               <dbl> <chr>
##
     <chr>
               <chr>
                         <chr>
                                      <dbl>
                                                              <chr>
                                                                       <chr>
## 1 signature EMT
                        MSI
                                  7.98e-17 3.20e-16 <2e-16
                                                              ****
                                                                       Wilcoxon
```

hjust

size_of_pvalue = 5,

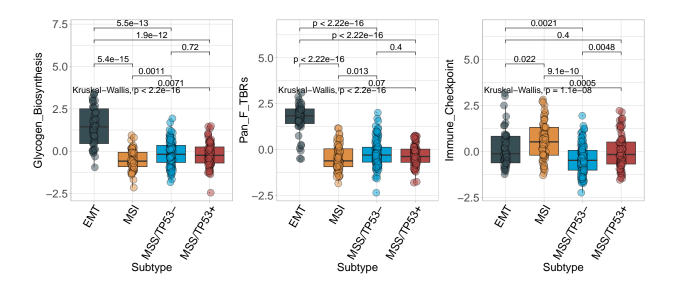
size_of_font

```
## 2 signature EMT
                         MSS/TP53- 1.70e-17 1
                                                e-16 <2e-16
                                                                        Wilcoxon
                                                               ****
## 3 signature EMT
                         MSS/TP53+ 2.57e-17 1.3 e-16 <2e-16
                                                                        Wilcoxon
                                                               ****
## 4 signature MSI
                         MSS/TP53- 1.32e- 2 4
                                                e- 2 0.013
                                                                        Wilcoxon
## 5 signature MSI
                         MSS/TP53+ 6.99e- 2 1.4 e- 1 0.070
                                                               ns
                                                                        Wilcoxon
## 6 signature MSS/TP53- MSS/TP53+ 4.02e- 1 4 e- 1 0.402
                                                                        Wilcoxon
                                                               ns
p3 <- sig_box(data
                             = input,
                             = "Immune Checkpoint",
              signature
              variable
                             = "Subtype",
              jitter
                              = TRUE,
              cols
                             = NULL,
                             = "jama",
              palette
                             = TRUE,
              show pvalue
              angle_x_text
                             = 60,
```

= 1,

= 8)

```
## # A tibble: 6 x 8
##
                                                    p.adj p.format p.signif method
               group1
                         group2
     .y.
                                           р
     <chr>
                                                    <dbl> <chr>
##
               <chr>
                         <chr>
                                       <dbl>
                                                                    <chr>
                                                                             <chr>
                                    2.20e- 2 0.044
## 1 signature EMT
                         MSI
                                                          0.0220
                                                                             Wilcoxon
## 2 signature EMT
                         MSS/TP53- 2.11e- 3 0.0085
                                                                             Wilcoxon
                                                          0.0021
                                                                    **
## 3 signature EMT
                         MSS/TP53+ 4.03e- 1 0.4
                                                          0.4026
                                                                             Wilcoxon
                                                                    ns
## 4 signature MSI
                         MSS/TP53- 9.13e-10 0.0000000055 9.1e-10
                                                                             Wilcoxon
                                                                    ****
## 5 signature MSI
                         MSS/TP53+ 5.03e- 4 0.0025
                                                          0.0005
                                                                             Wilcoxon
## 6 signature MSS/TP53- MSS/TP53+ 4.82e- 3 0.014
                                                          0.0048
                                                                             Wilcoxon
p1|p2|p3
```



4.7 Survival analysis

Signature

```
sig_surv_plot(input_pdata
                                               = input,
res <-
                                               = "Glycogen_Biosynthesis",
                            signature
                            cols
                                               = NULL,
                                               = "jco",
                            palette
                            project
                                               = "ACRG",
                            time
                                               = "OS_time",
                                               = "OS_status",
                            status
                                               = "month",
                            time_type
                                               = "result")
                            save_path
```

bestcutoff	group2	group3	${\tt Glycogen_Biosynthesis}$	status	time	ID	##
Low	Low	Middle	-0.3612213	0	88.73	1 GSM1523727	## :
Low	Low	Low	-0.6926726	0	88.23	2 GSM1523728	## 2
Low	Low	Low	-0.9388531	0	88.23	3 GSM1523729	## 3
Low	Low	Low	-1.1825136	0	105.70	4 GSM1523744	## 4
Low	Low	Middle	-0.3034304	0	105.53	5 GSM1523745	## !

6 GSM1523746 25.50

1

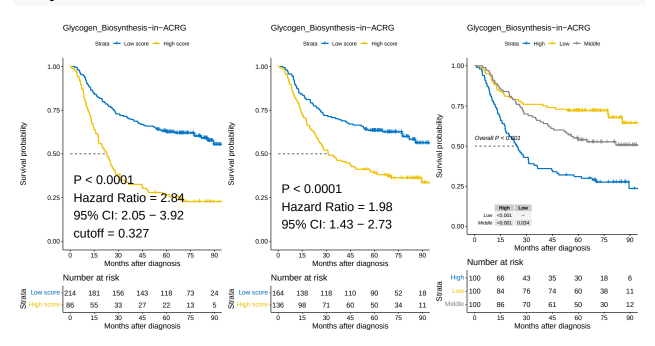
0.7517934 Hi

High High

High

[1] ">>>>>

res\$plots



Signature ROC

```
p1<- roc_time(input = input,</pre>
                        = "Glycogen Biosynthesis",
             vars
                        = "OS time",
             time
                        = "OS_status",
             status
             time_point = c(12, 24, 36),
             time_type = "month",
             palette
                        = "jama",
             cols
                        = "normal",
             seed
                        = 1234,
                        = FALSE,
             show_col
                         = "result",
             path
                         = "0S",
             main
             index
                         = 1,
```

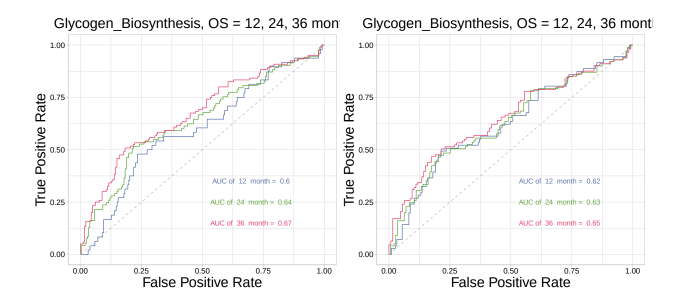
```
fig.type = "pdf",
           width = 5,
           height = 5.2)
## [1] ">>>-- Range of Time: "
## [1] 1.0 105.7
p2<- roc_time(input = input,</pre>
           vars = "Glycogen_Biosynthesis",
           time = "RFS_time",
           status = "RFS_status",
          time_point = c(12, 24, 36),
          time_type = "month",
           palette = "jama",
           cols = "normal",
           seed = 1234,
           show_col = FALSE,
           path = "result",
           main
                   = "OS",
           index
                   = 1,
           fig.type = "pdf",
```

```
## [1] ">>>-- Range of Time: "
## [1] 0.10 100.87

p1 p2
```

width = 5,

height = 5.2)



4.8 Batch correlation analysis

signatures

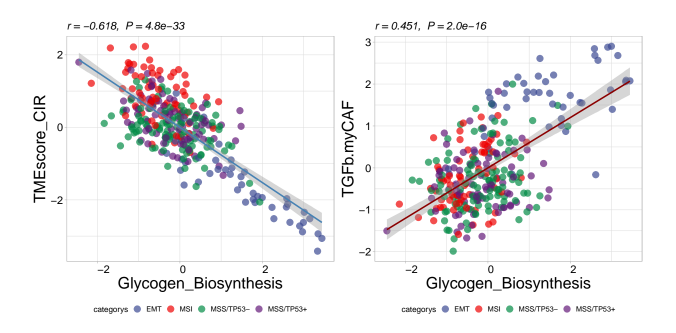
signature

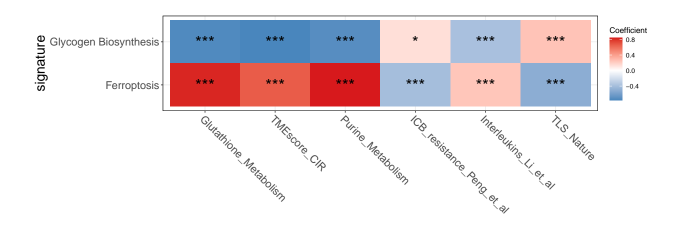
```
res <- batch_cor(data = input, target = "Glycogen_Biosynthesis", feature = colnames(input)
head(res)</pre>
```

```
## # A tibble: 6 x 6
                                                             p.adj log10pvalue stars
##
     sig names
                                       p.value statistic
                                         <dbl>
                                                    <dbl>
                                                             <dbl>
##
     <chr>
                                                                         <dbl> <fct>
                                                    0.678 2.27e-39
## 1 TMEscoreB_CIR
                                      8.89e-42
                                                                          41.1 ****
## 2 Glycine__Serine_and_Threonine_M~ 7.49e-40
                                                   -0.666 9.54e-38
                                                                          39.1 ****
## 3 Ether_Lipid_Metabolism
                                      3.84e-39
                                                    0.662 3.27e-37
                                                                          38.4 ****
## 4 MDSC_Peng_et_al
                                      1.13e-38
                                                  0.659 7.21e-37
                                                                          37.9 ****
## 5 Glycerophospholipid_Metabolism
                                      8.72e-38
                                                   -0.653 4.44e-36
                                                                          37.1 ****
## 6 TIP_Release_of_cancer_cell_anti~ 2.32e-37
                                                   -0.650 9.86e-36
                                                                          36.6 ****
p1<- get_cor(eset = sig tme, pdata = pdata acrg, var1 = "Glycogen Biosynthesis", var2 =
```

##
Spearman's rank correlation rho

```
##
## data: data[, var1] and data[, var2]
## S = 7282858, p-value < 2.2e-16
## alternative hypothesis: true rho is not equal to 0
## sample estimates:
##
          rho
## -0.6184309
##
## [1] ">>>--- The exact p value is: 4.78971420439895e-33"
                   MSI MSS/TP53- MSS/TP53+
         EMT
##
##
          46
                    68
                              107
                                         79
p2<- get_cor(eset = sig_tme, pdata = pdata_acrg, var1 = "Glycogen_Biosynthesis", var2 =
##
## Spearman's rank correlation rho
##
## data: data[, var1] and data[, var2]
## S = 2471758, p-value < 2.2e-16
## alternative hypothesis: true rho is not equal to 0
## sample estimates:
         rho
##
## 0.4507143
##
## [1] ">>>--- The exact p value is: 2.04505761057615e-16"
                   MSI MSS/TP53- MSS/TP53+
         EMT
##
          46
                    68
                              107
                                         79
##
p1|p2
```

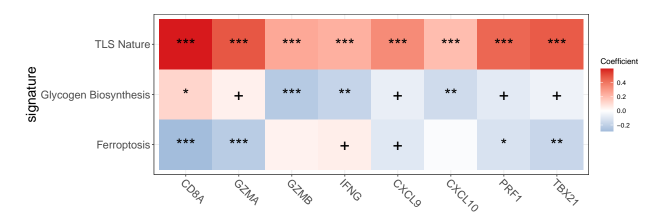


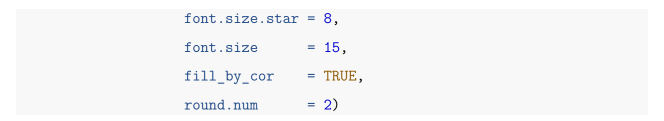


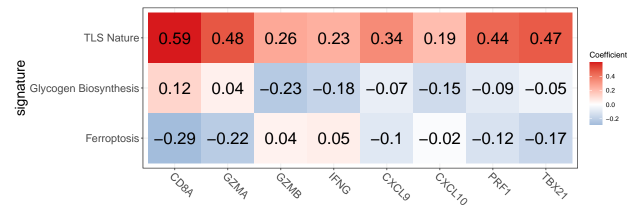
4.9 Visulization of correlations

```
input2 <- combine_pd_eset(eset = eset, pdata = input[, c("ID", "Glycogen_Biosynthesis"
feas1 <- c("Glycogen_Biosynthesis", "TLS_Nature", "Ferroptosis")
feas2 <- signature_collection$CD_8_T_effector
feas2</pre>
```

```
## [1] "CD8A"
                                        "CXCL9" "CXCL10" "PRF1"
              "GZMA"
                       "GZMB"
                                "IFNG"
                                                                 "TBX21"
p <- get_cor_matrix(data</pre>
                           = input2,
                  feas1
                             = feas2,
                  feas2
                             = feas1,
                  method
                             = "pearson",
                  scale
                               = T,
                  font.size.star = 8,
                  font.size
                                = 15,
                  fill_by_cor
                               = FALSE,
                                = 1)
                  round.num
```







Chapter 5

TME deconvolution

5.1 Loading packages

Load the IOBR package in your R session after the installation is complete:

```
library(IOBR)
library(survminer)
library(tidyverse)
```

5.2 Downloading data for example

Obtaining data set from GEO Gastric cancer: GSE62254 using GEOquery R package.

```
if (!requireNamespace("GEOquery", quietly = TRUE)) BiocManager::install("GEOquery")
library("GEOquery")
# NOTE: This process may take a few minutes which depends on the internet connection s
eset_geo<-getGEO(GEO = "GSE62254", getGPL = F, destdir = "./")
eset <-eset_geo[[1]]
eset <-exprs(eset)
eset[1:5,1:5]</pre>
```

```
## 1007_s_at 3.2176645 3.0624323 3.0279131 2.921683 2.8456013  
## 1053_at 2.4050109 2.4394879 2.2442708 2.345916 2.4328582  
## 117_at 1.4933412 1.8067380 1.5959665 1.839822 1.8326058  
## 121_at 2.1965561 2.2812181 2.1865556 2.258599 2.1874363  
## 1255_g_at 0.8698382 0.9502466 0.8125414 1.012860 0.9441993
```

```
Annotation of genes in the expression matrix and removal of duplicate genes.
library(IOBR)
# Load the annotation file `anno_hug133plus2` in IOBR.
head(anno_hug133plus2)
## # A tibble: 6 x 2
##
    probe_id symbol
    <fct>
              <fct>
##
## 1 1007_s_at MIR4640
## 2 1053 at
              RFC2
## 3 117 at
              HSPA6
## 4 121 at
              PAX8
## 5 1255 g at GUCA1A
## 6 1294 at
              MIR5193
# Conduct gene annotation using `anno_hug133plus2` file; If identical gene symbols exi
eset<-anno_eset(eset = eset,</pre>
               annotation = anno_hug133plus2,
               symbol = "symbol",
               probe = "probe_id",
               method
                        = "mean")
eset[1:5, 1:3]
```

:	##		GSM1523727	GSM1523728	GSM1523729
:	##	SH3KBP1	4.327974	4.316195	4.351425
:	##	RPL41	4.246149	4.246808	4.257940
:	##	EEF1A1	4.293762	4.291038	4.262199
:	##	COX2	4.250288	4.283714	4.270508
:	##	L0C101928826	4.219303	4.219670	4.213252

5.3 Available Methods to Decode TME Contexture

tme_d	econvolution_methods			
##	MCPcounter	EPIC	xCell	CIBERSORT
##	"mcpcounter"	"epic"	"xcell"	"cibersort"
## CI	BERSORT Absolute	IPS	ESTIMATE	SVR
##	"cibersort_abs"	"ips"	"estimate"	"svr"
##	lsei	TIMER	${\tt quanTIseq}$	
##	"lsei"	"timer"	"quantiseq"	
# Ret	urn available parameter	options of	deconvolution methods	

The input data is a matrix subseted from ESET of ACRG cohort, with genes in rows and samples in columns. The row name must be HGNC symbols and the column name must be sample names.

```
eset_acrg <- eset[, 1:50]
eset_acrg[1:5, 1:3]</pre>
```

##		GSM1523727	GSM1523728	GSM1523729
##	SH3KBP1	4.327974	4.316195	4.351425
##	RPL41	4.246149	4.246808	4.257940
##	EEF1A1	4.293762	4.291038	4.262199
##	COX2	4.250288	4.283714	4.270508
##	L0C101928826	4.219303	4.219670	4.213252

Check detail parameters of the function

```
# help(deconvo_tme)
```

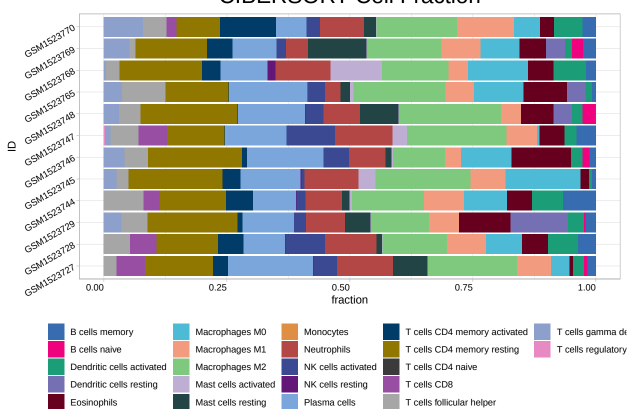
5.4 Method 1: CIBERSORT

```
cibersort<-deconvo_tme(eset = eset_acrg, method = "cibersort", arrays = TRUE, perm = 100
##
## >>> Running CIBERSORT
# head(cibersort)
res<-cell_bar_plot(input = cibersort[1:12,], title = "CIBERSORT Cell Fraction")</pre>
```

There are seven categories you can choose: box, continue2, continue, random, heatmap,

>>>=== Palette option for random: 1: palette1; 2: palette2; 3: palette3; 4: palette





5.4.1 Method 2: EPIC

```
# help(deconvo_epic)
epic<-deconvo_tme(eset = eset_acrg, method = "epic", arrays = TRUE)</pre>
##
## >>> Running EPIC
## Warning in IOBR::EPIC(bulk = eset, reference = ref, mRNA cell = NULL, scaleExprs = TF
## GSM1523744; GSM1523746; GSM1523781; GSM1523786
## - check fit.gof for the convergeCode and convergeMessage
## Warning in IOBR::EPIC(bulk = eset, reference = ref, mRNA cell = NULL, scaleExprs
## = TRUE): mRNA cell value unknown for some cell types: CAFs, Endothelial - using
## the default value of 0.4 for these but this might bias the true cell proportions
## from all cell types.
head(epic)
## # A tibble: 6 x 9
             Bcells EPIC CAFs EPIC CD4 Tcells EPIC CD8 Tcells EPIC Endothelial EPIC
##
                   <dbl>
##
     <chr>>
                              <dbl>
                                              <dbl>
                                                               <dbl>
                                                                                <dbl>
## 1 GSM152~
                  0.0292
                           0.00888
                                              0.145
                                                              0.0756
                                                                               0.0876
## 2 GSM152~
                  0.0293
                           0.0109
                                                                               0.0954
                                              0.159
                                                              0.0745
## 3 GSM152~
                  0.0308
                           0.0106
                                              0.149
                                                              0.0732
                                                                               0.0941
## 4 GSM152~
                  0.0273
                           0.0108
                                              0.145
                                                              0.0704
                                                                               0.0860
## 5 GSM152~
                  0.0280
                           0.0111
                                              0.151
                                                              0.0707
                                                                               0.0928
```

i 3 more variables: Macrophages_EPIC <dbl>, NKcells_EPIC <dbl>,

0.148

0.0716

0.0907

0.00958

otherCells EPIC <dbl>

0.0320

6 GSM152~

##

<chr>

<dbl>

5.5 Method 3: MCPcounter

```
mcp<-deconvo_tme(eset = eset acrg, method = "mcpcounter")</pre>
##
## >>> Running MCP-counter
head(mcp)
## # A tibble: 6 x 11
##
     ID
                T_cells_MCPcounter CD8_T_cells_MCPcounter Cytotoxic_lymphocytes_M~1
                              <dbl>
##
     <chr>
                                                      <dbl>
                                                                                 <dbl>
## 1 GSM1523727
                               1.47
                                                      1.11
                                                                                  1.33
## 2 GSM1523728
                              1.53
                                                      1.05
                                                                                  1.60
## 3 GSM1523729
                              1.47
                                                      1.07
                                                                                  1.37
## 4 GSM1523744
                              1.46
                                                      1.02
                                                                                  1.44
## 5 GSM1523745
                               1.51
                                                      1.10
                                                                                  1.49
## 6 GSM1523746
                               1.51
                                                      0.992
                                                                                  1.40
## # i abbreviated name: 1: Cytotoxic_lymphocytes_MCPcounter
## # i 7 more variables: B_lineage_MCPcounter <dbl>, NK_cells_MCPcounter <dbl>,
       Monocytic lineage MCPcounter <dbl>,
## #
## #
       Myeloid dendritic cells MCPcounter <dbl>, Neutrophils MCPcounter <dbl>,
## #
       Endothelial cells MCPcounter <dbl>, Fibroblasts MCPcounter <dbl>
5.5.1
        Method 4: xCELL
xcell<-deconvo_tme(eset = eset acrg, method = "xcell", arrays = TRUE)</pre>
head(xcell)
## # A tibble: 6 x 68
                aDC xCell Adipocytes xCell Astrocytes xCell `B-cells xCell`
##
```

<dbl>

<dbl>

<dbl>

```
## 1 GSM1523727 4.78e-19
                                   0.0250
                                                   0
                                                                     0
## 2 GSM1523728 9.41e- 2
                                                   7.70e- 3
                                   0.00433
                                                                     0
## 3 GSM1523729 1.02e- 1
                                   0.0789
                                                   2.04e- 2
                                                                     0
## 4 GSM1523744 7.88e- 2
                                   0.0538
                                                   4.82e-18
                                                                     0.0126
## 5 GSM1523745 9.02e- 2
                                   0.0136
                                                   1.93e- 2
                                                                     0
## 6 GSM1523746 3.40e- 2
                                                   9.22e- 2
                                   0.0331
                                                                     0
## # i 63 more variables: Basophils xCell <dbl>,
       `CD4+_memory_T-cells_xCell` <dbl>, `CD4+_naive T-cells xCell` <dbl>,
## #
       `CD4+_T-cells_xCell` <dbl>, `CD4+_Tcm_xCell` <dbl>, `CD4+_Tem_xCell` <dbl>,
## #
       `CD8+ naive T-cells xCell` <dbl>, `CD8+ T-cells xCell` <dbl>,
## #
## #
       `CD8+ Tcm xCell` <dbl>, `CD8+ Tem xCell` <dbl>, cDC xCell <dbl>,
## #
       Chondrocytes_xCell <dbl>, `Class-switched_memory_B-cells_xCell` <dbl>,
## #
       CLP_xCell <dbl>, CMP_xCell <dbl>, DC_xCell <dbl>, ...
```

5.6 Method 5: ESTIMATE

```
estimate<-deconvo_tme(eset = eset_acrg, method = "estimate")

## [1] "Merged dataset includes 9940 genes (472 mismatched)."

## [1] "1 gene set: StromalSignature overlap= 136"

## [1] "2 gene set: ImmuneSignature overlap= 138"

head(estimate)</pre>
```

A tibble: 6 x 5 StromalScore_estimate ImmuneScore_estimate ESTIMATEScore_estimate ## ID ## <chr> <dbl> <dbl> <dbl> -982. ## 1 GSM1523727 -1250.268. ## 2 GSM1523728 197. 1334. 1531. ## 3 GSM1523729 -111. 822. 711. ## 4 GSM1523744 -119. 662. 544.

```
## 5 GSM1523745 324. 1015. 1339.
## 6 GSM1523746 -594. 621. 27.0
```

i 1 more variable: TumorPurity_estimate <dbl>

5.7 Method 6: TIMER

```
timer<-deconvo_tme(eset = eset_acrg, method = "timer", group_list = rep("stad",dim(eset)</pre>
## [1] "Outlier genes: AGR2 B2M COL1A2 COL3A1 COX2 CYAT1 EEF1A1 EIF1 FTH1 GKN1 HUWE1 IGH
head(timer)
## # A tibble: 6 x 7
                B_cell_TIMER T_cell_CD4_TIMER T_cell_CD8_TIMER Neutrophil_TIMER
     <chr>
                       <dbl>
                                        <dbl>
                                                          <dbl>
                                                                           <dbl>
##
## 1 GSM1523727
                       0.104
                                       0.128
                                                          0.183
                                                                           0.108
## 2 GSM1523728
                       0.103
                                       0.130
                                                          0.192
                                                                           0.118
## 3 GSM1523729
                       0.106
                                      0.130
                                                          0.190
                                                                           0.110
## 4 GSM1523744
                       0.101
                                       0.126
                                                          0.187
                                                                           0.111
## 5 GSM1523745
                       0.104
                                        0.127
                                                          0.191
                                                                           0.116
## 6 GSM1523746
                       0.105
                                        0.129
                                                          0.192
                                                                           0.111
```

i 2 more variables: Macrophage_TIMER <dbl>, DC_TIMER <dbl>

5.8 Method 7: quanTIseq

```
quantiseq<-deconvo_tme(eset = eset_acrg, tumor = TRUE, arrays = TRUE, scale_mrna = TRUE
##
## Running quantIseq deconvolution module
## Gene expression normalization and re-annotation (arrays: TRUE)
## Removing 17 genes with high expression in tumors</pre>
```

Signature genes found in data set: 152/153 (99.35%)

Mixture deconvolution (method: lsei)

Deconvolution sucessful!

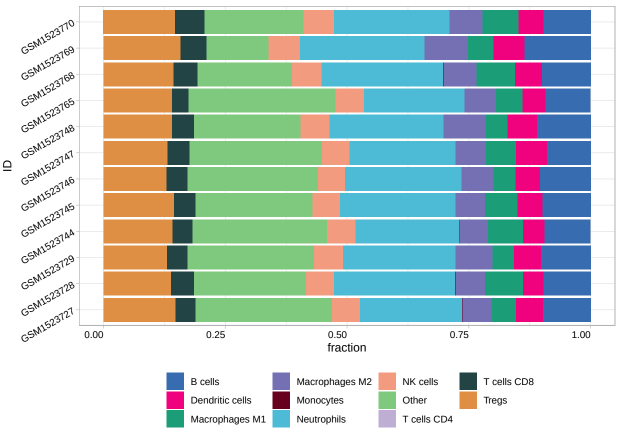
head(quantiseq)

```
## # A tibble: 6 x 12
               B cells quantiseq Macrophages M1 quantiseq Macrophages M2 quantiseq
##
    TD
##
    <chr>>
                           <dbl>
                                                   <dbl>
                                                                            <dbl>
## 1 GSM1523727
                     0.0983
                                                  0.0510
                                                                          0.0598
## 2 GSM1523728
                 0.0967
                                                  0.0795
                                                                          0.0607
## 3 GSM1523729
                 0.102
                                                  0.0450
                                                                          0.0758
                 0.0954
## 4 GSM1523744
                                                  0.0725
                                                                          0.0579
                   0.0991
## 5 GSM1523745
                                                  0.0669
                                                                          0.0613
## 6 GSM1523746
                          0.105
                                                  0.0453
                                                                           0.0662
## # i 8 more variables: Monocytes_quantiseq <dbl>, Neutrophils_quantiseq <dbl>,
## #
      NK_cells_quantiseq <dbl>, T_cells_CD4_quantiseq <dbl>,
## #
      T_cells_CD8_quantiseq <dbl>, Tregs_quantiseq <dbl>,
## #
      Dendritic_cells_quantiseq <dbl>, Other_quantiseq <dbl>
res<-cell_bar_plot(input = quantiseq[1:12, ], title = "quanTIseq Cell Fraction")</pre>
```

There are seven categories you can choose: box, continue2, continue, random, heatmap,

>>>>=== Palette option for random: 1: palette1; 2: palette2; 3: palette3; 4: palette





5.9 8) Method 8: IPS

```
ips<-deconvo_tme(eset = eset_acrg, method = "ips", plot= FALSE)
head(ips)</pre>
```

```
## # A tibble: 6 x 7
               MHC_IPS EC_IPS SC_IPS CP_IPS AZ_IPS IPS_IPS
##
    ID
                 <dbl>
                        <dbl> <dbl>
                                       <dbl>
                                              <dbl>
                                                      <dbl>
##
     <chr>
## 1 GSM1523727
                  2.25
                        0.404 - 0.192
                                     0.220
                                               2.68
                                                          9
## 2 GSM1523728 2.37
                        0.608 -0.578 -0.234
                                                          7
                                               2.17
## 3 GSM1523729 2.10
                        0.480 -0.322
                                      0.0993
                                               2.36
                                                          8
## 4 GSM1523744 2.12 0.535 -0.333
                                      0.0132
                                               2.34
                                                          8
## 5 GSM1523745 1.91 0.559 -0.479 0.0880
                                               2.08
                                                          7
```

```
## 6 GSM1523746    1.94    0.458    -0.346    0.261    2.31    8
```

5.10 9) Combination of above deconvolution results

```
tme_combine<-cibersort %>%
  inner_join(.,mcp,by = "ID") %>%
  inner_join(.,xcell,by = "ID") %>%
  inner_join(.,epic,by = "ID") %>%
  inner_join(.,estimate,by = "ID") %>%
  inner_join(.,timer,by = "ID") %>%
  inner_join(.,quantiseq,by = "ID") %>%
  inner_join(.,quantiseq,by = "ID") %>%
  inner_join(.,ips,by = "ID")
```

[1] 50 138

If you use this package in your work, please cite both our package and the method(s) you are using.

5.11 Licenses of the deconvolution methods

method	license	citation
CIBERSORT	free for non-commerical use	Newman, A. M., Liu, C. L.,
	only	Green, M. R., Gentles, A. J.,
		Feng, W., Xu, Y., Alizadeh,
		A. A. (2015). Robust
		enumeration of cell subsets
		from tissue expression profiles.
		Nature Methods, 12(5),
		453–457. https:
		//doi.org/10.1038/nmeth.3337
ESTIMATE	free (GPL2.0)	Vegesna R, Kim H,
		Torres-Garcia W,, Verhaak R
		(2013). Inferring tumour purity
		and stromal and immune cell
		admixture from expression data
		Nature Communications 4,
		2612. http:
		//doi.org/10.1038/ncomms3612
quanTIseq	free (BSD)	Finotello, F., Mayer, C.,
		Plattner, C., Laschober, G.,
		Rieder, D., Hackl, H.,,
		Sopper, S. (2019). Molecular
		and pharmacological
		modulators of the tumor
		immune contexture revealed by
		deconvolution of RNA-seq data.
		Genome medicine, 11(1), 34.
		https://doi.org/10.1186/s13073-
		019-0638-6

method	license	citation
TIMER	free (GPL 2.0)	Li, B., Severson, E., Pignon,
		JC., Zhao, H., Li, T., Novak,
		J., Liu, X. S. (2016).
		Comprehensive analyses of
		tumor immunity: implications
		for cancer immunotherapy.
		Genome Biology, 17(1), 174.
		https://doi.org/10.1186/s13059-
		016-1028-7
IPS	free (BSD)	P. Charoentong et al.,
		Pan-cancer Immunogenomic
		Analyses Reveal
		Genotype-Immunophenotype
		Relationships and Predictors of
		Response to Checkpoint
		Blockade. Cell Reports 18,
		248-262 (2017). https://doi.org/
		10.1016/j.celrep.2016.12.019

method	license	citation
MCPCounter	free (GPL 3.0)	Becht, E., Giraldo, N. A.,
		Lacroix, L., Buttard, B.,
		Elarouci, N., Petitprez, F., de
		Reyniès, A. (2016). Estimating
		the population abundance of
		tissue-infiltrating immune and
		stromal cell populations using
		gene expression. Genome
		Biology, 17(1), 218.
		https://doi.org/10.1186/s13059-
		016-1070-5
xCell	free (GPL 3.0)	Aran, D., Hu, Z., & Butte, A. J.
		(2017). xCell: digitally
		portraying the tissue cellular
		heterogeneity landscape.
		Genome Biology, $18(1)$, 220 .
		https://doi.org/10.1186/s13059-
		017-1349-1
EPIC	free for non-commercial use	Racle, J., de Jonge, K.,
	only (Academic License)	Baumgaertner, P., Speiser, D.
		E., & Gfeller, D. (2017).
		Simultaneous enumeration of
		cancer and immune cell types
		from bulk tumor gene
		expression data. ELife, 6,
		e26476. https:
		$//{\rm doi.org}/10.7554/{\rm eLife.26476}$

5.11.1 Licenses of the signature-esitmation method

method	license	citation
GSVA	free (GPL (>= 2))	Hänzelmann S, Castelo R, Guinney J (2013). "GSVA: gene set variation analysis for microarray and RNA-Seq data." BMC Bioinformatics, 14, 7. doi: 10.1186/1471-2105-14-7, http://www.biomedcentral.co m/1471-2105/14/7

Cross-references

Cross-references make it easier for your readers to find and link to elements in your book.

6.1 Chapters and sub-chapters

There are two steps to cross-reference any heading:

- 1. Label the heading: # Hello world {#nice-label}.
 - Leave the label off if you like the automated heading generated based on your heading title: for example, # Hello world = # Hello world {#hello-world}.
 - To label an un-numbered heading, use: # Hello world {-#nice-label} or {# Hello world .unnumbered}.
- 2. Next, reference the labeled heading anywhere in the text using \@ref(nice-label); for example, please see Chapter 6.
 - If you prefer text as the link instead of a numbered reference use: any text you want can go here.

6.2 Captioned figures and tables

Figures and tables with captions can also be cross-referenced from elsewhere in your book using \@ref(fig:chunk-label) and \@ref(tab:chunk-label), respectively.

See Figure 6.1.

```
par(mar = c(4, 4, .1, .1))
plot(pressure, type = 'b', pch = 19)
```

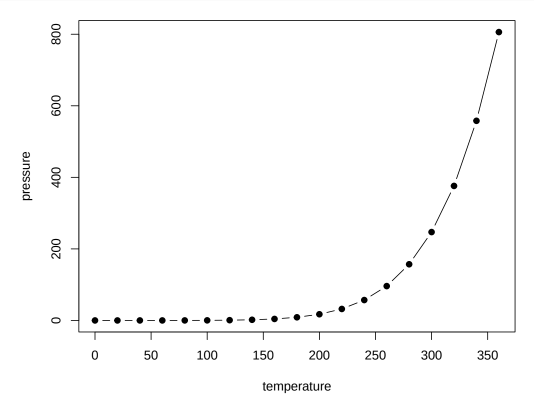


Figure 6.1: Here is a nice figure!

Don't miss Table 6.1.

```
knitr::kable(
  head(pressure, 10), caption = 'Here is a nice table!',
  booktabs = TRUE
)
```

Table 6.1: Here is a nice table!

temperature	pressure
0	0.0002
20	0.0012
40	0.0060
60	0.0300
80	0.0900
100	0.2700
120	0.7500
140	1.8500
160	4.2000
180	8.8000

Footnotes and citations

7.1 Footnotes

Footnotes are put inside the square brackets after a caret ^[]. Like this one ¹.

7.2 Citations

Reference items in your bibliography file(s) using Okey.

For example, we are using the **bookdown** package (Xie, 2023) (check out the last code chunk in index.Rmd to see how this citation key was added) in this sample book, which was built on top of R Markdown and **knitr** (Xie, 2015) (this citation was added manually in an external file book.bib). Note that the .bib files need to be listed in the index.Rmd with the YAML bibliography key.

The RStudio Visual Markdown Editor can also make it easier to insert citations: https://rstudio.github.io/visual-markdown-editing/#/citations

¹This is a footnote.

Blocks

8.1 Equations

Here is an equation.

$$f(k) = \binom{n}{k} p^k \left(1 - p\right)^{n - k} \tag{8.1}$$

You may refer to using \@ref(eq:binom), like see Equation (8.1).

8.2 Theorems and proofs

Labeled theorems can be referenced in text using \@ref(thm:tri), for example, check out this smart theorem 8.1.

Theorem 8.1. For a right triangle, if c denotes the length of the hypotenuse and a and b denote the lengths of the **other** two sides, we have

$$a^2 + b^2 = c^2$$

Read more here https://bookdown.org/yihui/bookdown/markdown-extensions-by-bookdo

wn.html.

8.3 Callout blocks

The R Markdown Cookbook provides more help on how to use custom blocks to design your own callouts: https://bookdown.org/yihui/rmarkdown-cookbook/custom-blocks.html

Sharing your book

9.1 Publishing

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9.2 404 pages

By default, users will be directed to a 404 page if they try to access a webpage that cannot be found. If you'd like to customize your 404 page instead of using the default, you may add either a _404.Rmd or _404.md file to your project root and use code and/or Markdown syntax.

9.3 Metadata for sharing

Bookdown HTML books will provide HTML metadata for social sharing on platforms like Twitter, Facebook, and LinkedIn, using information you provide in the index.Rmd YAML. To setup, set the url for your book and the path to your cover-image file. Your book's title and description are also used.

This gitbook uses the same social sharing data across all chapters in your book- all links

CHAPTER 9. SHARING YOUR BOOK

82

shared will look the same.

Specify your book's source repository on GitHub using the edit key under the configuration

options in the _output.yml file, which allows users to suggest an edit by linking to a chapter's

source file.

Read more about the features of this output format here:

https://pkgs.rstudio.com/bookdown/reference/gitbook.html

Or use:

?bookdown::gitbook

References

If IOBR R package is utilized in your published research, please cite:

Zeng D, Ye Z, Shen R, Yu G, Wu J, Xiong Y,..., Liao W (2021) **IOBR**: Multi-Omics Immuno-Oncology Biological Research to Decode Tumor Microenvironment and Signatures. *Frontiers in Immunology*. 12:687975. doi: 10.3389/fimmu.2021.687975

10.1 TME deconvolution

Please cite the following papers appropriately for TME deconvolution algorithm if used:

CIBERSORT: Newman, A. M., Liu, C. L., Green, M. R., Gentles, A. J., Feng, W., Xu, Y., ... Alizadeh, A. A. (2015). Robust enumeration of cell subsets from tissue expression profiles. Nature Methods, 12(5), 453–457. https://doi.org/10.1038/nmeth.3337

ESTIMATE: Vegesna R, Kim H, Torres-Garcia W, ..., Verhaak R.*(2013). Inferring tumour purity and stromal and immune cell admixture from expression data. Nature Communications 4, 2612. http://doi.org/10.1038/ncomms3612

quanTIseq: Finotello, F., Mayer, C., Plattner, C., Laschober, G., Rieder, D., Hackl, H., ..., Sopper, S.* (2019). Molecular and pharmacological modulators of the tumor immune contexture revealed by deconvolution of RNA-seq data. Genome medicine, 11(1), 34. https:

//doi.org/10.1186/s13073-019-0638-6

TIMER: Li, B., Severson, E., Pignon, J.-C., Zhao, H., Li, T., Novak, J., ... Liu, X. S.* (2016). Comprehensive analyses of tumor immunity: implications for cancer immunotherapy. Genome Biology, 17(1), 174.

IPS: P. Charoentong et al.*, Pan-cancer Immunogenomic Analyses Reveal Genotype-Immunophenotype Relationships and Predictors of Response to Checkpoint Blockade. Cell Reports 18, 248-262 (2017). https://doi.org/10.1016/j.celrep.2016.12.019

MCPCounter: Becht, E., Giraldo, N. A., Lacroix, L., Buttard, B., Elarouci, N., Petitprez, F., ... de Reyniès, A*. (2016). Estimating the population abundance of tissue-infiltrating immune and stromal cell populations using gene expression. Genome Biology, 17(1), 218. https://doi.org/10.1186/s13059-016-1070-5

xCell: Aran, D., Hu, Z., & Butte, A. J.* (2017). xCell: digitally portraying the tissue cellular heterogeneity landscape. Genome Biology, 18(1), 220. https://doi.org/10.1186/s13059-017-1349-1

EPIC: Racle, J., de Jonge, K., Baumgaertner, P., Speiser, D. E., & Gfeller, D*. (2017). Simultaneous enumeration of cancer and immune cell types from bulk tumor gene expression data. ELife, 6, e26476. https://doi.org/10.7554/eLife.26476

10.2 TME Signatures

For signature score estimation, please cite corresponding literature below:

ssgsea: Barbie, D.A. et al (2009). Systematic RNA interference reveals that oncogenic KRAS-driven cancers require TBK1. Nature, 462(5):108-112.

gsva: Hänzelmann, S., Castelo, R. and Guinney, J. (2013). GSVA: Gene set variation analysis for microarray and RNA-Seq data. BMC Bioinformatics, 14(1):7.

zscore: Lee, E. et al (2008). Inferring pathway activity toward precise disease classification. PLoS Comp Biol, 4(11):e1000217.

10.3. DATA SETS 85

10.3 Data sets

For the datasets enrolled in IOBR, please cite the data sources:

UCSCXena: Wang et al., et al (2019). The UCSCXenaTools R package: a toolkit for accessing genomics data from UCSC Xena platform, from cancer multi-omics to single-cell RNA-seq. Journal of Open Source Software, 4(40), 1627

TLSscore: Helmink BA, Reddy SM, Gao J, et al. B cells and tertiary lymphoid structures promote immunotherapy response. Nature. 2020 Jan;577(7791):549-555.

IMvigor210 immuntherapy cohort: Mariathasan S, Turley SJ, Nickles D, et al. TGF attenuates tumour response to PD-L1 blockade by contributing to exclusion of T cells. Nature. 2018 Feb 22;554(7693):544-548. HCP5: Kulski, J.K. Long Noncoding RNA HCP5, a Hybrid HLA Class I Endogenous Retroviral Gene: Structure, Expression, and Disease Associations. Cells 2019, 8, 480.

HCP5: Li, Y., Jiang, T., Zhou, W. et al. Pan-cancer characterization of immune-related lncRNAs identifies potential oncogenic biomarkers. Nat Commun 11, 1000 (2020). HCP5: Sun J, Zhang Z, Bao S, et alIdentification of tumor immune infiltration-associated lncRNAs for improving prognosis and immunotherapy response of patients with non-small cell lung cancer Journal for ImmunoTherapy of Cancer 2020;8:e000110.

LINC00657: Feng Q, Zhang H, Yao D, Chen WD, Wang YD. Emerging Role of Non-Coding RNAs in Esophageal Squamous Cell Carcinoma. Int J Mol Sci. 2019 Dec 30;21(1):258. doi: 10.3390/ijms21010258.

LINC00657: Qin X, Zhou M, Lv H, Mao X, Li X, Guo H, Li L, Xing H. Long noncoding RNA LINC00657 inhibits cervical cancer development by sponging miR-20a-5p and targeting RUNX3. Cancer Lett. 2020 Oct 28:S0304-3835(20)30578-4. doi: 10.1016/j.canlet.2020.10.044. LINC00657: Zhang XM, Wang J, Liu ZL, Liu H, Cheng YF, Wang T. LINC00657/miR-26a-5p/CKS2 ceRNA network promotes the growth of esophageal cancer cells via the MDM2/p53/Bcl2/Bax pathway. Biosci Rep. 2020;40(6):BSR20200525.

TCGA-STAD: Cancer Genome Atlas Research Network. Comprehensive molecular charac-

terization of gastric adenocarcinoma. Nature. 2014 Sep 11;513(7517):202-9. doi: 10.1038/nature13480. TCGA.STAD MAF data: https://api.gdc.cancer.gov/data/c06465a3-50e7-46f7-b2dd-7bd654ca206b

10.4 Others

- Newman, A. M., Liu, C. L., Green, M. R., Gentles, A. J., Feng, W., Xu, Y., ... Alizadeh, A. A. (2015). Robust enumeration of cell subsets from tissue expression profiles. Nature Methods, 12(5), 453–457.
- 2. Vegesna R, Kim H, Torres-Garcia W, ..., Verhaak R.*(2013). Inferring tumour purity and stromal and immune cell admixture from expression data. Nature Communications 4, 2612.
- 3. Rieder, D., Hackl, H., ..., Sopper, S.* (2019). Molecular and pharmacological modulators of the tumor immune contexture revealed by deconvolution of RNA-seq data. Genome medicine, 11(1), 34.
- 4. Li, B., Severson, E., Pignon, J.-C., Zhao, H., Li, T., Novak, J., ... Liu, X. S.* (2016). Comprehensive analyses of tumor immunity: implications for cancer immunotherapy. Genome Biology, 17(1), 174.
- 5. P. Charoentong et al.*, Pan-cancer Immunogenomic Analyses Reveal Genotype-Immunophenotype Relationships and Predictors of Response to Checkpoint Blockade. Cell Reports 18, 248-262 (2017).
- 6. Becht, E., Giraldo, N. A., Lacroix, L., Buttard, B., Elarouci, N., Petitprez, F., ... de Reyniès, A*. (2016). Estimating the population abundance of tissue-infiltrating immune and stromal cell populations using gene expression. Genome Biology, 17(1), 218.
- 7. Aran, D., Hu, Z., & Butte, A. J.* (2017). xCell: digitally portraying the tissue cellular heterogeneity landscape. Genome Biology, 18(1), 220.
- 8. Racle, J., de Jonge, K., Baumgaertner, P., Speiser, D. E., & Gfeller, D*. (2017).

10.4. OTHERS 87

Simultaneous enumeration of cancer and immune cell types from bulk tumor gene expression data. ELife, 6, e26476.

- 9. Barbie, D.A. et al (2009). Systematic RNA interference reveals that oncogenic KRAS-driven cancers require TBK1. Nature, 462(5):108-112.
- 10. Hänzelmann, S., Castelo, R. and Guinney, J. (2013). GSVA: Gene set variation analysis for microarray and RNA-Seq data. BMC Bioinformatics, 14(1):7.
- 11. Lee, E. et al (2008). Inferring pathway activity toward precise disease classification. PLoS Comp Biol, 4(11):e1000217.
- 12. Wang et al., et al (2019). The UCSCXenaTools R package: a toolkit for accessing genomics data from UCSC Xena platform, from cancer multi-omics to single-cell RNA-seq. Journal of Open Source Software, 4(40), 1627
- 13. Helmink BA, Reddy SM, Gao J, et al. B cells and tertiary lymphoid structures promote immunotherapy response. Nature. 2020 Jan;577(7791):549-555.
- 14. Mariathasan S, Turley SJ, Nickles D, et al. TGF attenuates tumour response to PD-L1 blockade by contributing to exclusion of T cells. Nature. 2018 Feb 22;554(7693):544-548.
- 15. Kulski, J.K. Long Noncoding RNA HCP5, a Hybrid HLA Class I Endogenous Retroviral Gene: Structure, Expression, and Disease Associations. Cells 2019, 8, 480.
- 16. Li, Y., Jiang, T., Zhou, W. et al. Pan-cancer characterization of immune-related lncR-NAs identifies potential oncogenic biomarkers. Nat Commun 11, 1000 (2020).
- 17. Sun J, Zhang Z, Bao S, et alldentification of tumor immune infiltration-associated lncRNAs for improving prognosis and immunotherapy response of patients with non-small cell lung cancerJournal for ImmunoTherapy of Cancer 2020;8:e000110.
- 18. Feng Q, Zhang H, Yao D, Chen WD, Wang YD. Emerging Role of Non-Coding RNAs in Esophageal Squamous Cell Carcinoma. Int J Mol Sci. 2019 Dec 30;21(1):258. doi: 10.3390/ijms21010258.

- 19. Qin X, Zhou M, Lv H, Mao X, Li X, Guo H, Li L, Xing H. Long noncoding RNA LINC00657 inhibits cervical cancer development by sponging miR-20a-5p and targeting RUNX3. Cancer Lett. 2020 Oct
- 20. Zhang XM, Wang J, Liu ZL, Liu H, Cheng YF, Wang T. LINC00657/miR-26a-5p/CKS2 ceRNA network promotes the growth of esophageal cancer cells via the MDM2/p53/Bcl2/Bax pathway. Biosci Rep. 2020;40(6):BSR20200525.
- 21. Cancer Genome Atlas Research Network. Comprehensive molecular characterization of gastric adenocarcinoma. Nature. 2014 Sep 11;513(7517):202-9. doi: 10.1038/nature13480.

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Xie, Y. (2015). Dynamic Documents with R and knitr. Chapman and Hall/CRC, Boca Raton, Florida, 2nd edition. ISBN 978-1498716963.

Xie, Y. (2023). bookdown: Authoring Books and Technical Documents with R Markdown. R package version 0.35.