CRISPRcleanR: An R package for unsupervised identification and correction of gene independent cell responses to CRISPR-cas9 targeting

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1 Quick start

1.1 Installation

First, you need to install and load the devtools package. You can do this from CRAN. Invoke R and then type:

```
install.packages("devtools")
library(devtools)
```

Secondly, install CRISPRcleanR with the following command:

```
install_github("francescojm/CRISPRcleanR")
```

1.2 Raw sgRNA count normalisation and computation of sgRNAs' log fold-changes

Load CRISPRcleanR.

```
library(CRISPRcleanR)

## Loading required package: stringr

## Loading required package: DNAcopy

## Loading required package: pROC

## Type 'citation("pROC")' for a citation.

##

## Attaching package: 'pROC'

## The following objects are masked from 'package:stats':

##

## cov, smooth, var

## Loading required package: pracma

## Loading required package: PRROC
```

Step 1: Load your sgRNA library annotation. In this example we will use a built in data frame containing the annotation of the SANGER v1.0 library, introduced in [1]:

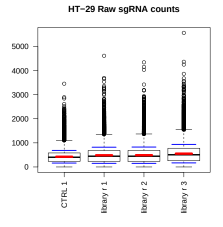
```
data(KY_Library_v1.0)
```

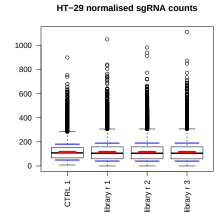
To use your own library annotation you will have to store it into a data frame with the same format of the KY_Library_v1.0 (detailed in the corresponding entry of the reference manual of the CRISPRcleanR package).

Step 2: Store the path of the tsv file containing your sgRNAs' raw counts in a temporary variable. In this example we will use counts generated upon a CRISPR-Cas9 pooled drop-out screen (described in [2]) built in this package, for an example immortalised human cancer cell line (HT-29)

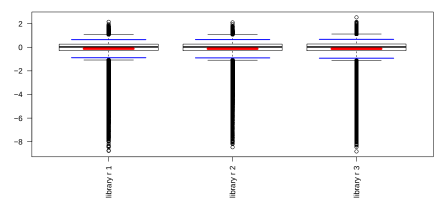
The tsv file with the sgRNAs' raw counts must be formatted as specified in the reference manual entry of the ccr.NormfoldChanges function.

Step 3: Perform normalisation of raw counts and compute sgRNAs' log fold-changes (in this example we will exclude sgRNAs with less than 30 reads in the plasmid sample).





HT-29 log Fold Changes



This function returns a list of two data frames containing, normalised counts and log fold-changes, respectively, and it saves them as Robjects in a user defined directory (specified by the parameter outdir, which is set to './' by default).

```
head(normANDfcs$norm_counts)
##
                                              sgRNA gene ERS717283.plasmid
## 1 A1BG_CCDS12976.1_ex3_19:58862927-58862950:-_5-1 A1BG
                                                                 292.14621
## 2 A1BG_CCDS12976.1_ex4_19:58863655-58863678:+_5-2 A1BG
                                                                 151.02032
## 3 A1BG_CCDS12976.1_ex4_19:58863697-58863720:-_5-3 A1BG
                                                                 209.08503
## 4 A1BG_CCDS12976.1_ex4_19:58863866-58863889:+_5-4 A1BG
                                                                 110.40106
## 5 A1BG_CCDS12976.1_ex5_19:58864367-58864390:-_5-5 A1BG
                                                                  95.81979
## 6 A1CF_CCDS7241.1_ex6_10:52588014-52588037:-_5-1 A1CF
                                                                  60.92889
##
    HT29_c904R1 HT29_c904R2 HT29_c904R3
## 1
      308.05192 354.89835
                             305.56806
## 2
      145.38048 113.16912
                             166.47364
## 3
      280.97793 203.70441
                             223.03071
## 4
       80.31191
                  64.05372
                              78.74023
## 5
       78.71932
                 123.80701
                             103.32157
## 6
       47.77762
                   76.50232
                              59.75464
head(normANDfcs$logFCs)
##
                                              sgRNA gene HT29_c904R1 HT29_c904R2
## 1 A1BG_CCDS12976.1_ex3_19:58862927-58862950:-_5-1 A1BG 0.07635566 0.28027938
## 2 A1BG_CCDS12976.1_ex4_19:58863655-58863678:+_5-2 A1BG -0.05472442 -0.41467098
## 3 A1BG_CCDS12976.1_ex4_19:58863697-58863720:-_5-3 A1BG 0.42548611 -0.03752168
## 4 A1BG_CCDS12976.1_ex4_19:58863866-58863889:+_5-4 A1BG -0.45663336 -0.78070106
## 5 A1BG_CCDS12976.1_ex5_19:58864367-58864390:-_5-5 A1BG -0.28197989
## 6 A1CF_CCDS7241.1_ex6_10:52588014-52588037:-_5-1 A1CF -0.34756262 0.32598467
##
    HT29_c904R3
## 1 0.06469488
## 2 0.14010902
## 3 0.09293733
```

```
## 4 -0.48496821
## 5 0.10820208
## 6 -0.02784489
```

IMPORTANT: if there are control replicates in your sgRNAs count file their number must be specified by in the parameter ncontrols (equal to 1 by default) of the ccr.NormfoldChanges function.

1.3 Genome sorting of sgRNAs' log fold-changes and their correction for gene independent responses to CRISPR-Cas9 targeting

Step 1: Map genome-wide sgRNAs' log fold changes (averaged across replicates) on the genome, sorted according to the of their targeted gene on the chromosomes.

```
head(gwSortedFCs)
##
                                   CHR startp
                                              endp genes
## SAMD11_CCDS2.2_ex3_1:871254-871277:+_5-1 1 871254 871277 SAMD11 -0.12965287
## SAMD11_CCDS2.2_ex4_1:874451-874474:-_5-2 1 874451 874474 SAMD11 0.09329615
## SAMD11_CCDS2.2_ex6_1:876601-876624:-_5-5 1 876601 876624 SAMD11 -0.02110076
                                    1 887388 887411 NOC2L -1.27571756
## NOC2L_CCDS3.1_ex8_1:887388-887411:+_5-1
##
                                        BP
## SAMD11_CCDS2.2_ex3_1:871254-871277:+_5-1 871265.5
## SAMD11_CCDS2.2_ex4_1:874451-874474:-_5-2 874462.5
## SAMD11_CCDS2.2_ex4_1:874487-874510:+_5-3 874498.5
## SAMD11_CCDS2.2_ex5_1:874693-874716:+_5-4 874704.5
## SAMD11_CCDS2.2_ex6_1:876601-876624:-_5-5 876612.5
## NOC2L_CCDS3.1_ex8_1:887388-887411:+_5-1 887399.5
```

Step 2: Identify and correct biased sgRNAs' log fold-changes putatively due to gene independent responses to CRISPR-Cas9 targeting, using the ccr.GWclean function. This function calls iteratively the ccr.cleanChrm function, which performs the correction on an individual chromosome). In this example we are using a completely unsuerpvised approach and correcting chromosomal segments of equal sgRNA log fold-changes if they include sgRNAs targeting at least 3 different genes, and without making any assumption on gene essentiality, nor knowing a priori the copy number status of the included genes [2].

```
correctedFCs<-ccr.GWclean(gwSortedFCs,display=TRUE,label='HT-29')</pre>
```

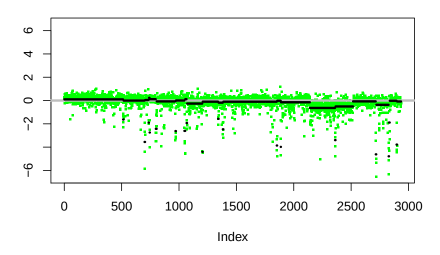
The corrected sgRNAs fold-changes are returned in a list (as a data frame), together with the annotation of the identified segments (in another data frame) and a vector of strings containing all the gemome-sorted sgRNAs' identifiers.

```
head(correctedFCs$corrected_logFCs)
##
                                                         endp genes
                                                                           avgFC
                                            CHR startp
## SAMD11_CCDS2.2_ex3_1:871254-871277:+_5-1 1 871254 871277 SAMD11 -0.12965287
                                             1 874451 874474 SAMD11 0.09329615
## SAMD11_CCDS2.2_ex4_1:874451-874474:-_5-2
## SAMD11_CCDS2.2_ex4_1:874487-874510:+_5-3
                                            1 874487 874510 SAMD11 0.25286616
## SAMD11_CCDS2.2_ex5_1:874693-874716:+_5-4
                                             1 874693 874716 SAMD11 -0.05128489
## SAMD11 CCDS2.2 ex6 1:876601-876624:- 5-5
                                             1 876601 876624 SAMD11 -0.02110076
## NOC2L_CCDS3.1_ex8_1:887388-887411:+_5-1
                                             1 887388 887411 NOC2L -1.27571756
##
                                                 BP correction correctedFC
## SAMD11_CCDS2.2_ex3_1:871254-871277:+_5-1 871265.5
                                                             0 -0.12965287
## SAMD11_CCDS2.2_ex4_1:874451-874474:-_5-2 874462.5
                                                             0 0.09329615
## SAMD11_CCDS2.2_ex4_1:874487-874510:+_5-3 874498.5
                                                             0 0.25286616
## SAMD11_CCDS2.2_ex5_1:874693-874716:+_5-4 874704.5
                                                             0 -0.05128489
## SAMD11_CCDS2.2_ex6_1:876601-876624:-_5-5 876612.5
                                                             0 -0.02110076
## NOC2L_CCDS3.1_ex8_1:887388-887411:+_5-1 887399.5
                                                             0 -1.27571756
```

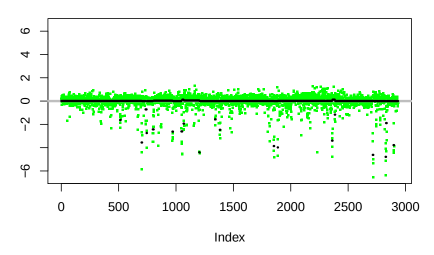
Details on how the data frame with the corrected sgRNAs' log fold-changes should be interpreted can be found in the entry of the ccr.GWclean function, of the package reference manual.

This function also produces one plot per chromosome, with segments of sgRNAs' equal log fold-change before and after the correction. An example of these plot is reported below: chromosome 8, in HT-29, showing a region containing MYC, which is highly biased toward consistent negative fold-changes.

HT.29.Chr.8.sgRNA.FCs



HT.29.Chr.8.sgRNA.FCs...post.CRISPRcleanR



$\begin{array}{cc} {\bf 1.4} & {\bf Correcting~sgRNAs'~treatment~counts~for~mean-variance} \\ & {\bf modeling} \end{array}$

In order to apply the inverse transformation described in [2], deriving corrected normalised sgRNAs' treatment counts from CRISPRcleanR corrected log fold-changes, it is sufficient to run the function ccr.correctCounts as follows:

Together with the plasmid counts, the corrected treatment counts can be used to compute depletion singificance scores with a mean-variance modeling approach (such that implemented in MAGeCK[3]).

```
head(correctedCounts)
##
                                                sgRNA gene ERS717283.plasmid
## 1 A1BG_CCDS12976.1_ex3_19:58862927-58862950:-_5-1 A1BG
                                                                    292.14621
## 2 A1BG_CCDS12976.1_ex4_19:58863655-58863678:+_5-2 A1BG
                                                                    151.02032
## 3 A1BG_CCDS12976.1_ex4_19:58863697-58863720:-_5-3 A1BG
                                                                    209.08503
## 4 A1BG_CCDS12976.1_ex4_19:58863866-58863889:+_5-4 A1BG
                                                                    110.40106
## 5 A1BG_CCDS12976.1_ex5_19:58864367-58864390:-_5-5 A1BG
                                                                     95.81979
  6 A1CF_CCDS7241.1_ex6_10:52588014-52588037:-_5-1 A1CF
                                                                     60.92889
##
     HT29_c904R1 HT29_c904R2 HT29_c904R3
## 1
       309.77863
                   356.73522
                                307.28892
## 2
       144.74469
                   112.89328
                                165.60212
## 3
       280.34458
                   203.51866
                                222.73301
## 4
        80.64680
                    64.52159
                                 79.08797
## 5
        78.22697
                   122.47062
                                102.36866
## 6
        45.21299
                    71.83850
                                 56.31473
```

This function also saves the correctedCounts as Rdata object at the location specified by the parameter OutDir. To run MAGeCK, using these corrected sgRNAs' counts you will need to save them as a tsv file, which will be ued as input for MAGeCK. This can be done by using the following function, which also returns the path of the saved file.

```
ccr.PlainTsvFile(correctedCounts,fprefix = 'HT-29')
## [1] "./HT-29_sgRNA_count.tsv"
```

IMPORTANT: the corrected sgRNAs' counts are already normalised, therefore, when executing MAGeCK, the parameter --norm-method should be set to none.

2 Visualisation and assessment of Results

2.1 Classification performances of reference sets of genes (or sgRNAs) based on depletion log fold-changes

To perform a basic quality control assessment of your data it is possible to test the genome-wide profile of sgRNAs' depletion log fold-changes (logFCs) (or gene depletion logFCs, averaged across targeting sgRNAs) as a classifier of reference sets of core-fitness essential (CFE) and non-essential genes.

What you need for this is a named vector of sgRNAs (or gene) logFCs and two reference gene sets (respectively for positive and negative cases). In this example we make use of a precomputed essentiality profile from the builtin data object EPLC.272HcorrectedFCs. This is a list containing corrected sgRNAs log fold-changes and segment annotations for an example cell line (EPLC-272H), obtained using the ccr.GWclean function, as detailed in its reference manual entry. However the data frame containing the corrected log fold-changes, included in this list, reports also the original sgRNAs logFC (column avgFC), which will be used in this example.

```
data(EPLC.272HcorrectedFCs)
```

```
head(EPLC.272HcorrectedFCs$corrected_logFCs)
##
                                                         endp genes
## SAMD11_CCDS2.2_ex3_1:871254-871277:+_5-1
                                             1 871254 871277 SAMD11 -0.20295496
## SAMD11_CCDS2.2_ex4_1:874451-874474:-_5-2
                                              1 874451 874474 SAMD11 -0.08917153
## SAMD11_CCDS2.2_ex4_1:874487-874510:+_5-3
                                             1 874487 874510 SAMD11 -0.04417670
## SAMD11_CCDS2.2_ex5_1:874693-874716:+_5-4
                                             1 874693 874716 SAMD11 0.30441537
## SAMD11_CCDS2.2_ex6_1:876601-876624:-_5-5
                                             1 876601 876624 SAMD11 -0.11240079
## NOC2L_CCDS3.1_ex8_1:887388-887411:+_5-1
                                              1 887388 887411 NOC2L -1.61370746
##
                                                  BP correction correctedFC
## SAMD11_CCDS2.2_ex3_1:871254-871277:+_5-1 871265.5
                                                              0 -0.20295496
## SAMD11_CCDS2.2_ex4_1:874451-874474:-_5-2 874462.5
                                                              0 -0.08917153
## SAMD11_CCDS2.2_ex4_1:874487-874510:+_5-3 874498.5
                                                              0 -0.04417670
## SAMD11_CCDS2.2_ex5_1:874693-874716:+_5-4 874704.5
                                                              0 0.30441537
## SAMD11_CCDS2.2_ex6_1:876601-876624:-_5-5 876612.5
                                                              0 -0.11240079
## NOC2L_CCDS3.1_ex8_1:887388-887411:+_5-1 887399.5
                                                              0 -1.61370746
```

As reference gene sets we will use lists of CFE and non-essential genes assembled from multiple RNAi studies. These are used as classification template by the BAGEL algorithm to call gene depletion significance [4], and are included in the builtin data objects BAGEL_essential and BAGEL_nonEssential.

```
data(BAGEL_essential)
data(BAGEL_nonEssential)
```

```
head(BAGEL_essential)
## [1] "ACTL6A" "ACTR6" "ALYREF" "ANAPC4" "ANAPC5" "AP2S1"
head(BAGEL_nonEssential)
## [1] "ABCG8" "ACCSL" "ACTL7A" "ACTL7B" "ACTL9" "ACTR11"
```

Finally, you will need the sgRNAs library annotation. In this case we will use the builtin object KY_KY_Library_v1.0 (introduced in the previous section) [1]. As for the previous examples, to use a different library annotation you will have to store it in a data frame with the same format of the KY_Library_v1.0 data frame (detailed in the corresponding entry of the reference manual of the CRISPRcleanR package).

```
data(KY_Library_v1.0)
```

We will start with an evualuation at the sgRNA level. As mentioned, the logFCs need to be stored a named vector:

```
FCs<-EPLC.272HcorrectedFCs$corrected_logFCs$avgFC
names(FCs)<-rownames(EPLC.272HcorrectedFCs$corrected_logFCs)
```

The ccr.genes2sgRNAs function can be used, as follows, to convert the reference sets of CFE and non-essential genes into sets of sgRNAs:

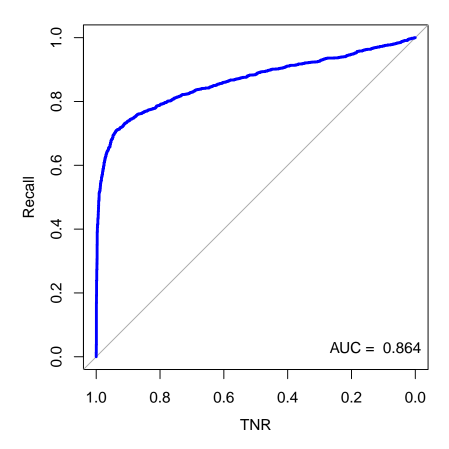
```
BAGEL_essential_sgRNAs<-
ccr.genes2sgRNAs(KY_Library_v1.0,BAGEL_essential)

BAGEL_nonEssential_sgRNAs<-
ccr.genes2sgRNAs(KY_Library_v1.0,BAGEL_nonEssential)
```

Following these calls, possible warning messages could appear informing you that some of the reference genes are not targeted by any sgRNAs in the considered library. This has no impact on the following steps and results.

Finally, to visualise the ROC curve quantifying the performances in classifying the considered reference sets based on their logFCs, it is sufficient to call:

```
sgRNA_level_ROC<-ccr.ROC_Curve(FCs,BAGEL_essential_sgRNAs,
BAGEL_nonEssential_sgRNAs)
```

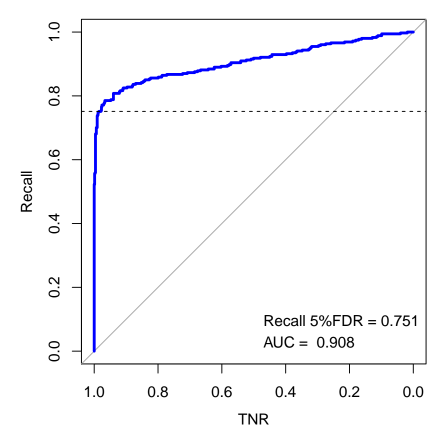


To reperform the analysis at the gene level, you should first convert the profile of sgRNAs' logFCs into gene level summaries. The function ccr.geneMeanFCs performs this conversion by considering for each gene the average logFC across its targeting sgRNAs.

```
geneFCs<-ccr.geneMeanFCs(FCs,KY_Library_v1.0)
head(geneFCs)

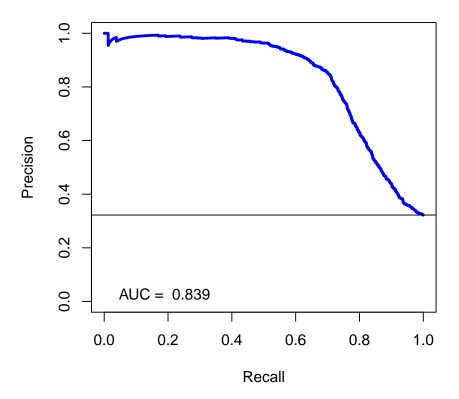
## A1BG A1CF A2M A2ML1 A3GALT2 A4GALT
## -0.2474235 -0.1550534 0.2190111 0.3736683 -0.5151889 -0.1698269</pre>
```

The following call reperforms the ROC analysis at the gene level and it also computes and shows Recall values at fixed False Discovery Rate (which is defined by the user and in this case is equal to 5%).

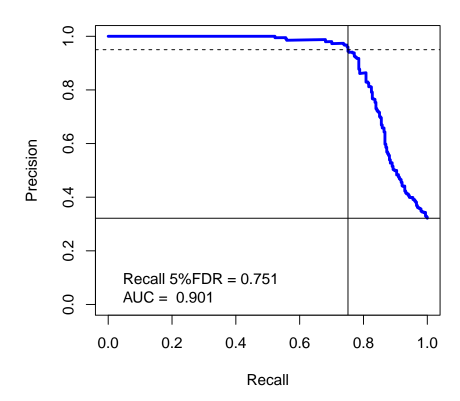


The same assessment can be performed at the level of Precision/Recall (PrRc) evaluation: to visualise the PrRc curve quantifying the performances in classifying the considered reference sets based on their logFCs, it is sufficient to call:

sgRNA_level_PrRc<-ccr.PrRc_Curve(FCs,BAGEL_essential_sgRNAs, BAGEL_nonEssential_sgRNAs)



As before, to reperform the analysis at the gene level, it is sufficient to call the following function, which also computes and shows Recall values at fixed False Discovery Rate (which is defined by the user and in this case is equal to 5%).



As can be seen above, when setting the parameter FRDth to a value different from NULL (its default value), this function also returns the log fold change threshold at which a classification FDR equal to the inputted value is achieved.

```
gene_level_PrRc$sigthreshold
## threshold
## -0.7683409
```

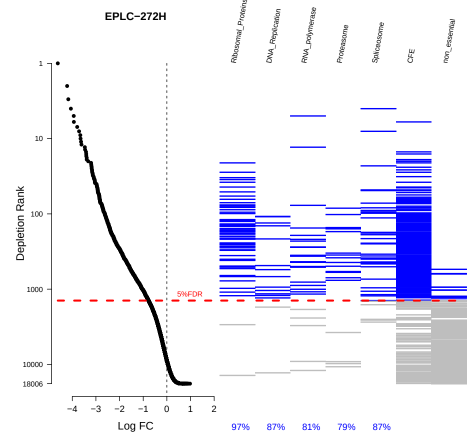
2.2 Depletion profile visualisation with genes signatures superimposed and recall computation

For another quick assessment of your data, it is possible to visually inspect enrichments of predefined sets of core-fitness essential genes near the top of the genome wide essentiality profiles (composed of sgRNA or gene depletion logFCs ranked in increasing order), and to compute their classification recall at a fixed

FDR (determined as deatiled in the previous subsection).

To this aim, in this example we will load additional sets of CFE genes assembled from MsigDB [5] as detailed in [2], and we will store all of them into a named list, as follows:

Finally a visualisation of the gene essentiality profile with superimposed these signatures, can be created as follows:



IMPORTANT: When calling ccr.VisDepAndSig it is important to correctly specify the index position of the reference gene sets that are used as classification template to derive the FDR threshold, within the list of signatures. In this case the template sets are BAGEL_essential and BAGEL_nonEssential, which in the SIGNATURE list are in position 6 and 7, respectively (this must be specified in the pIs and nIs parameters of the ccr.VisDepAndSig function). This function also returns recall values at 5% FDR for all the inputted signatures.

Recall_scores				
##	Ribosomal_Proteins	DNA_Replication	RNA_polymerase	Proteasome
##	0.96721311	0.8666667	0.80769231	0.78947368
##	Spliceosome	CFE	non_essential	
##	0.86842105	0.75141243	0.01874163	

2.3 CRISPRcleanR correction assessment: Statistical tests

To evaluate the effect of the CRISPRcleanR correction on your data it is possible to inspect the logFCs' variations for sgRNAs targeting predefined sets of genes for statistically significant differences with respect to background pre/post CRISPRcleanR correction.

To this aim, in this example we will use the builtin data object HT.29correctedFCs containing corrected sgRNAs' logFCs and segment annotations for an example cell line (HT-29), obtained using the ccr.GWclean function, as detailed in its reference manual entry.

data(HT.29correctedFCs)

The function ccr.perf_statTests performs this analysis, saving pdf figures in a user defined location ('./' by default).

Particularly, this functions assess statistical difference respect to background population pre/post CRISPRcleanR correction of logFCs for sgRNAs targeting respectively:

- copy number (CN) deleted genes according to the GDSC1000 repository
- CN deleted genes (gistic score = -2) according to the CCLE repository
- non expressed genes (FPKM lower than 0.05)
- \bullet genes with gistic score = 1
- genes with gistic score = 2
- non espressed genes (FPKM lower than 0.05) with gistic score = 1
- non espressed genes (FPKM lower than 0.05) with gistic score = 2
- genes with minimal CN = 2, according to the GDSC1000
- genes with minimal CN = 4, according to the GDSC1000
- genes with minimal CN = 8, according to the GDSC1000
- genes with minimal CN = 10, according to the GDSC1000
- \bullet non expressed genes (FPKM lower than 0.05) with minimal CN = 2, according to the GDSC1000
- \bullet non expressed genes (FPKM lower than 0.05) with minimal CN = 4, according to the GDSC1000
- \bullet non expressed genes (FPKM lower than 0.05) with minimal CN = 8, according to the GDSC1000

 \bullet non expressed genes (FPKM lower than 0.05) with minimal CN = 10, according to the GDSC1000

The call should be as follows:

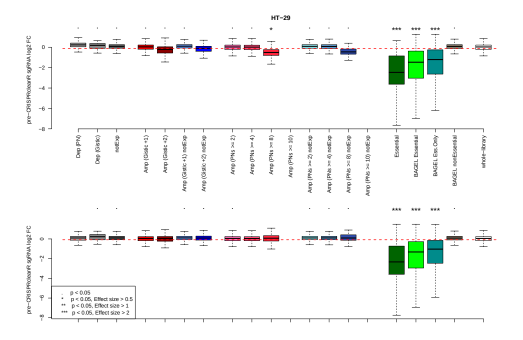
```
RES<-ccr.perf_statTests('HT-29',libraryAnnotation = KY_Library_v1.0,
                   correctedFCs = HT.29correctedFCs$corrected_logFCs,
                   GDSC.geneLevCNA = NULL,
                   CCLE.gisticCNA = NULL,
                   RNAseq.fpkms = NULL)
## [1] "Testing sgRNAs targeting: Dep (PN) genes"
  [1] "Testing sgRNAs targeting: Dep (Gistic) genes"
  [1] "Testing sgRNAs targeting: notExp genes"
  [1] "Testing sgRNAs targeting: Amp (Gistic +1) genes"
  [1] "Testing sgRNAs targeting: Amp (Gistic +2) genes"
  [1] "Testing sgRNAs targeting: Amp (Gistic +1) notExp genes"
  [1] "Testing sgRNAs targeting: Amp (Gistic +2) notExp genes"
  [1]
       "Testing sgRNAs targeting: Amp (PNs >= 2) genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 4) genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 8) genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 10) genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 2) notExp genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 4) notExp genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 8) notExp genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 10) notExp genes"
  [1] "Testing sgRNAs targeting: Essential genes"
  [1] "Testing sgRNAs targeting: BAGEL Essential genes"
  [1] "Testing sgRNAs targeting: BAGEL Ess.Only genes"
  [1] "Testing sgRNAs targeting: BAGEL nonEssential genes"
```

This saves a pdf with the following figure in a user defined location ('./' by default).

Leaving the parameters GDSC.geneLevCNA, CCLE.gisticCNA, and RNAseq.fpkms to their default NULL value will force this function to use the respective builtin data objects encompassing data only for 15 cell lines, used in [2] and in this package to assess the performances of CRISPRcleanR.

IMPORTANT: To analyse data from screening a different cell line ad-hoc GDSC.geneLevCNA, CCLE.gisticCNA, and RNAseq.fpkms data objects should be assembled and passed to this function. These should have the same format of the respective builtin counterparts, detailaed in their user reference manual entries, which contains also additional infos on how to derive this datasets for 1,000 human cance cell lines from the GDSC1000 data portal [6]).

Comprehensive statistical scores (detailed in the user reference manual) are



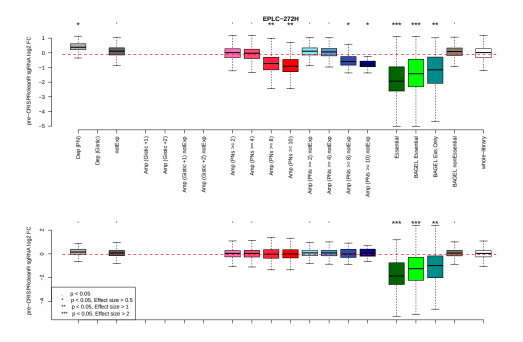
also returned in output by this function.

Another example, analysing in the same way the essentiality profile of the EPLC-272H cell line (includede in a corresponding builtin R object) is reported below,

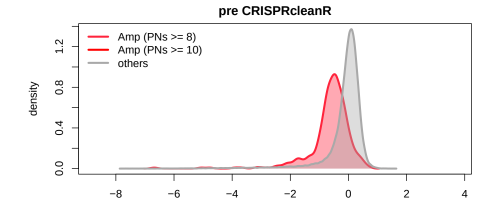
```
RES<-ccr.perf_statTests('EPLC-272H',libraryAnnotation = KY_Library_v1.0,
                    correctedFCs = EPLC.272HcorrectedFCs$corrected_logFCs)
## [1] "No gistic CNA scores available for this cell line"
  [1] "Testing sgRNAs targeting: Dep (PN) genes"
  [1] "Testing sgRNAs targeting: Dep (Gistic) genes"
      "Testing sgRNAs targeting: notExp genes"
  [1] "Testing sgRNAs targeting: Amp (Gistic +1) genes"
  [1] "Testing sgRNAs targeting: Amp (Gistic +2) genes"
  [1] "Testing sgRNAs targeting: Amp (Gistic +1) notExp genes"
   [1] "Testing sgRNAs targeting: Amp (Gistic +2) notExp genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 2) genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 4) genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 8) genes"
## [1] "Testing sgRNAs targeting: Amp (PNs >= 10) genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 2) notExp genes"
  [1] "Testing sgRNAs targeting: Amp (PNs >= 4) notExp genes"
## [1] "Testing sgRNAs targeting: Amp (PNs >= 8) notExp genes"
```

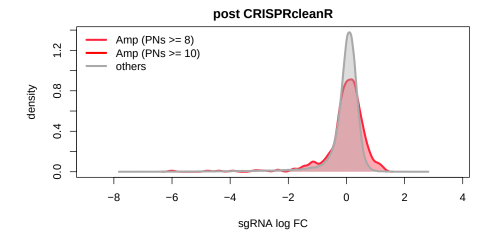
```
## [1] "Testing sgRNAs targeting: Amp (PNs >= 10) notExp genes"
## [1] "Testing sgRNAs targeting: Essential genes"
## [1] "Testing sgRNAs targeting: BAGEL Essential genes"
## [1] "Testing sgRNAs targeting: BAGEL Ess.Only genes"
## [1] "Testing sgRNAs targeting: BAGEL nonEssential genes"
```

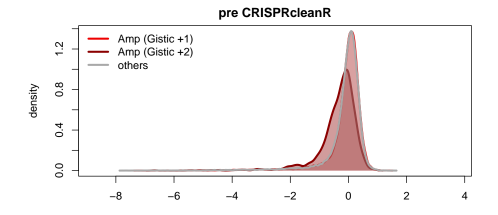
```
RES$PVALS
                       Dep (PN) Dep (Gistic) notExp Amp (Gistic +1)
## pre-CRISPRcleanR 5.145194e-27 NA 0.00000e+00 NA ## post-CRISPRcleanR 3.219730e-06 NA 5.69841e-173 NA
       Amp (Gistic +2) Amp (Gistic +1) notExp Amp (Gistic +2) notExp
## pre-CRISPRcleanR NA
## post-CRISPRcleanR NA
                                         NA
                                                     NA
## Amp (PNs >= 2) Amp (PNs >= 4) Amp (PNs >= 8) Amp (PNs >= 10)
## pre-CRISPRcleanR 1.580669e-18 1.780173e-57 1.991668e-137 1.476429e-42 ## post-CRISPRcleanR 4.951065e-05 1.593962e-02 2.036767e-01 9.122396e-02
## Amp (PNs >= 2) notExp Amp (PNs >= 4) notExp
## pre-CRISPRcleanR 2.268570e-311 2.387775e-52
## post-CRISPRcleanR 7.683083e-168 4.710481e-49
## Amp (PNs >= 8) notExp Amp (PNs >= 10) notExp Essential
## pre-CRISPRcleanR 3.575550e-07 4.600584e-05 2.736956e-175
## post-CRISPRcleanR 1.871438e-01
                                                 4.089818e-01 3.550035e-159
## BAGEL Essential BAGEL Ess.Only BAGEL nonEssential
## pre-CRISPRcleanR 2.898286e-297 1.591525e-205 8.174798e-81
## post-CRISPRcleanR 5.759565e-270 2.745117e-183
                                                     7.856856e-48
RES$EFFsizes
                   Dep (PN) Dep (Gistic) notExp Amp (Gistic +1)
## pre-CRISPRcleanR 0.7269806 NA 0.3141696 NA ## post-CRISPRcleanR 0.2874528 NA 0.2168474 NA
         Amp (Gistic +2) Amp (Gistic +1) notExp Amp (Gistic +2) notExp
## pre-CRISPRcleanR NA
## post-CRISPRcleanR NA
                                       NA
NA
                                                      NA
## Amp (PNs >= 2) Amp (PNs >= 4) Amp (PNs >= 8) Amp (PNs >= 10)
## pre-CRISPRcleanR 0.10987879 0.1142627 1.0609873 1.34598769
## post-CRISPRcleanR 0.05019186 0.0170022 0.0426818 0.09939445
## Amp (PNs >= 2) notExp Amp (PNs >= 4) notExp
## pre-CRISPRcleanR 0.3054594 0.2039852
## post-CRISPRcleanR
                             0.2183217
                                                   0.1916572
## Amp (PNs >= 8) notExp Amp (PNs >= 10) notExp Essential
## pre-CRISPRcleanR 0.6678038 0.9848744 2.859216
## post-CRISPRcleanR 0.1506673 0.1617632 2.945718
## BAGEL Essential BAGEL Ess.Only BAGEL nonEssential
## pre-CRISPRcleanR 2.223645 1.915641 0.2455203
## post-CRISPRcleanR 2.244647 1.900852 0.1799946
```

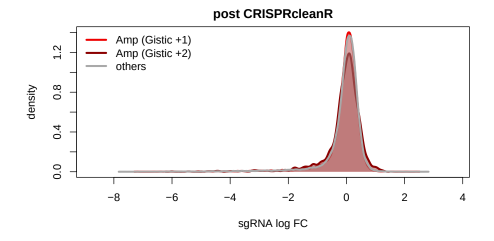


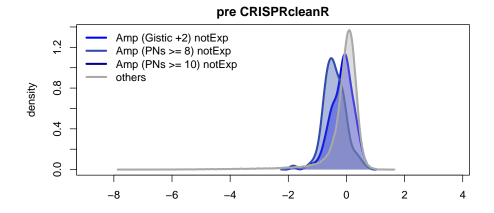
To inspect the variation induced by the CRISPRcleanR correction on the logFCs' distributions of sgRNAs targeting defined sets of genes prior/post CRISPRcleanR correction, the following function can be also used (and will produce the following density plots):

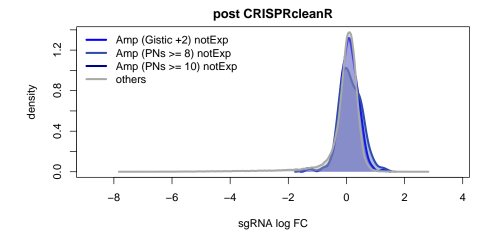


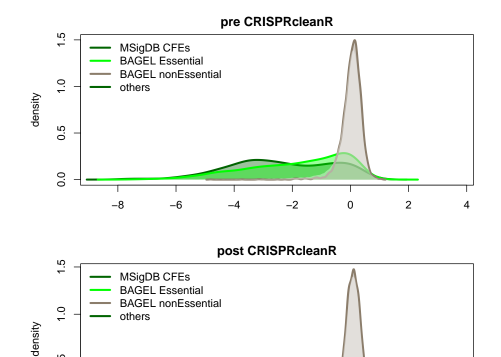












IMPORTANT: The instructions provided regarding what CN/transcriptional data object to pass to the ccr.perf_statTests apply also to this function.

-2

sgRNA log FC

0

2

0.5

-8

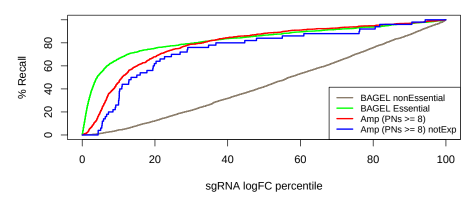
-6

Additional infos on how to use this function can be found in the user reference manual.

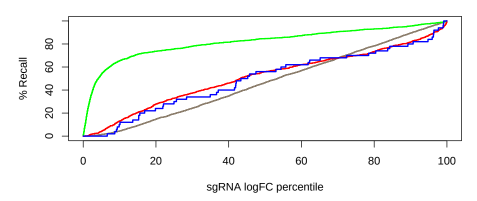
2.4 Recall variations following CRISPRcleanR correction for reference, copy number amplified, and non expressed genes

A final analysis that can be done with the CRISPRcleanR package in order to evalute the effect of its correction on the classification recall for predefined gene sets can be performed by calling the function, which can work at the sgRNA

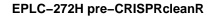
EPLC-272H pre-CRISPRcleanR

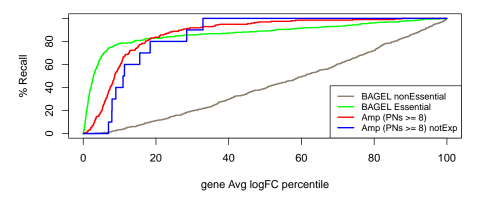


EPLC-272H post-CRISPRcleanR

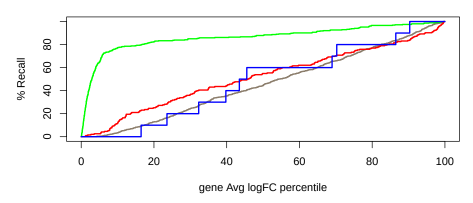


as well as the gene level:





EPLC-272H post-CRISPRcleanR



 ${\bf IMPORTANT:} \ {\bf The\ instructions\ provided\ regarding\ what\ CN/transcriptional\ data\ object\ to\ pass\ to\ the\ {\tt ccr.perf_statTests}\ apply\ also\ to\ this\ function.$

Additional infos on how to use this function can be found in the user reference manual.

2.5 CRISPRcleanR correction assessment: impact on phenotype and possible distortion

To evaluate the effect of the CRISPRcleanR correction on the genes showing a significant loss/gain-of-fitness effect (fitness genes) in the uncorrected data, a comparison of fitness gene sets (computed with MAGeCK [3]) before/after CRISPRcleanR correction can be performed as follows

Before running the below example let us clean the R environment with the following command:

```
rm(list=ls())
```

Subsequently, as detailed above, we load the reference sgRNA library annotation, we load and normalise sgRNA counts for the EPLC-272H example cell line.

Then, we save the normalised counts as plain tsv file (suitable for MAGeCK):

At this point we execute MAGeCK on the uncorrected normalised counts. **IMPORTANT:** This requires python and MAGeCK (v0.5.3). Additionally, as these counts are already normalised, we should specify that the normalisation shouldn't be reperformed by MAGeCK.

Run-time information displayed by MAGeCK (not reported in this document) will be visualised and all the outputted files will be saved in the working directory (by default), unless different specified in the corresponding argument (see reference manual entry for this function for further details).

This function returns in output the path of the gene summary file outputted by MAGeCK.

```
uncorrected_gs_fn
## [1] "./EPLC-272H.gene_summary.txt"
```

As detailed in the previous sections of this document, we now correct the sgRNA counts of the example cell lines with CRISPRcleanR and save them in a plain tsv file (suitable to be used by MAGeCK)

Then we execute MAGeCK on the corrected sgRNA counts.

Finally, we assess the impact of CRISP cleanR correction on gain/loss-of-fitness genes with the following call, which will produce bar/pie charts reported below

References

- [1] Konstantinos Tzelepis et al. "A CRISPR Dropout Screen Identifies Genetic Vulnerabilities and Therapeutic Targets in Acute Myeloid Leukemia." In: *Cell reports* 17.4 (Oct. 2016), pp. 1193–1205.
- [2] Francesco Iorio et al. "Unsupervised correction of gene-independent cell responses to CRISPR-Cas9 targeting". In: revision 0.0 (), pp. 0–0.
- [3] Wei Li et al. "MAGeCK enables robust identification of essential genes from genome-scale CRISPR/Cas9 knockout screens." In: *Genome Biology* 15.12 (2014), p. 554.
- [4] Traver Hart and Jason Moffat. "BAGEL: a computational framework for identifying essential genes from pooled library screens." In: *BMC bioinformatics* 17 (Apr. 2016), p. 164.
- [5] A Subramanian et al. "Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles". In: *Proceedings of the National Academy of Sciences of the United States of America* 102.43 (2005), p. 15545.
- [6] Francesco Iorio et al. "A Landscape of Pharmacogenomic Interactions in Cancer." In: Cell (July 2016).