# SCEPTRE Vs Seurat CHIP-seq Analysis

#### 2023-03-20

```
#load required packages. Sometimes need to redownload ondisc
library(biomaRt)
library(plyranges)
## Loading required package: BiocGenerics
## Attaching package: 'BiocGenerics'
## The following objects are masked from 'package:stats':
##
##
       IQR, mad, sd, var, xtabs
## The following objects are masked from 'package:base':
##
##
       anyDuplicated, aperm, append, as.data.frame, basename, cbind,
##
       colnames, dirname, do.call, duplicated, eval, evalq, Filter, Find,
##
       get, grep, grepl, intersect, is.unsorted, lapply, Map, mapply,
##
       match, mget, order, paste, pmax, pmax.int, pmin, pmin.int,
##
       Position, rank, rbind, Reduce, rownames, sapply, setdiff, sort,
##
       table, tapply, union, unique, unsplit, which.max, which.min
## Loading required package: IRanges
## Loading required package: S4Vectors
## Loading required package: stats4
## Attaching package: 'S4Vectors'
## The following objects are masked from 'package:base':
##
       expand.grid, I, unname
## Loading required package: GenomicRanges
## Loading required package: GenomeInfoDb
## Attaching package: 'plyranges'
```

```
## The following object is masked from 'package: IRanges':
##
       slice
##
## The following object is masked from 'package:biomaRt':
##
##
       select
## The following object is masked from 'package:stats':
##
##
       filter
library(GenomicRanges)
library(genomation)
## Loading required package: grid
## Warning: replacing previous import 'Biostrings::pattern' by 'grid::pattern'
## when loading 'genomation'
#devtools::install_qithub('timothy-barry/ondisc')
library(ondisc)
library(sceptre3)
library(BH)
library(varhandle)
##
## Attaching package: 'varhandle'
## The following object is masked from 'package:S4Vectors':
##
##
       unfactor
library(kableExtra)
library(rjson)
library(ggplot2)
```

#### Goal

The goal of this report is to compare the genes that are found to be significantly affected by IRF1 perturbation via Seurat and SCEPTRE to CHIPseq data.

#### Read in Promoter Data

```
#read in papalexi data
LOCAL_SCEPTRE2_DATA_DIR <-.get_config_path("LOCAL_SCEPTRE2_DATA_DIR")</pre>
papalexi dir <- pasteO(LOCAL SCEPTRE2 DATA DIR, "data/papalexi/eccite screen/")</pre>
# gene info
gene_odm_fp <- paste0(papalexi_dir, "gene/matrix.odm")</pre>
gene_metadata_fp <- paste0(papalexi_dir, "gene/metadata_qc.rds")</pre>
gene_odm <- read_odm(odm_fp = gene_odm_fp, metadata_fp = gene_metadata_fp)</pre>
#get TSS for each gene
ensembl <- useEnsembl(host = 'https://grch37.ensembl.org',biomart = "genes",</pre>
                       dataset = "hsapiens_gene_ensembl")
A = getBM(attributes=c("hgnc_symbol", "chromosome_name", "start_position",
                        "end_position", "strand"),
          filters=c('hgnc_symbol'),
          value = gene_odm |> get_feature_ids(), mart=ensembl) |>
  filter(chromosome_name %in% c(1:22, "X", "Y"))
#qet start and end site depending on whether the strand is postive or negative
TSS_start = rep(NA,nrow(A))
TSS_{end} = rep(NA, nrow(A))
for(j in c(1:nrow(A))){
  #if strand positive, use [start-500, start]
  if(A$strand[j]==1){
    TSS_end[j] = A$start_position[j]
    TSS_start[j] = A$start_position[j]-500
  }else{
    #if strand negative use [end, end + 500]
    TSS_start[j] = A$end_position[j]
    TSS_{end}[j] = A\end_{position}[j] + 500
 }
}
#add to A matrix
A$TSS start = TSS start
A$TSS_end = TSS_end
#add chr to chromosome name
A$chromosome_name = paste0("chr", A$chromosome_name)
#use A to make a promoter granges object
promoters <- GRanges(</pre>
 seqnames = A$chromosome_name,
 ranges = IRanges(start = A$TSS start, end = A$TSS end),
 TF = A$hgnc symbol)
```

#### Read in CHIPseq Data and Join the Datasets

## [1] "The number of promoters that overlap with Chipseq peaks is 1644"

#### Read in Seurat and SCEPTRE Results

```
#read in sceptre and seurat analysis data
sceptre_path = pasteO(data.dir,
                      'sceptre_full_mrna_results_with_effect_size.rds')
seurat_path = pasteO(data.dir,
                     'seurat_all_perturbations_results.rds')
#read in sceptre and seurat results
sceptre = readRDS(sceptre_path)
seurat = readRDS(seurat_path)
#adjust pvalues according to BH
sceptre$p_value_adj = sceptre$p_value
for(val in unique(sceptre$grna group)){
  ind = which(sceptre$grna group == val)
  sceptre$p_value_adj[ind] = p.adjust(sceptre$p_value[ind], method = "BH")
for(j in c(1:length(seurat))){
  seurat[[j]]$p_val_adj = p.adjust(seurat[[j]]$p_val,method = "BH")
}
```

#### Filter to Get IRF1 Results

```
#get sceptre and seurat IRF1 results
PRTB = 'IRF1'
sceptre_prtb = subset(sceptre,grna_group == PRTB)
seurat_prtb = seurat[[PRTB]]
```

### Compare Significant Genes to CHIPseq Results

```
alpha = 0.05
#get all significant genes from each method
sceptre_sig = unfactor(sceptre_prtb$response_id[sceptre_prtb$p_value_adj < alpha])
seurat_sig = rownames(seurat_prtb)[seurat_prtb$p_val_adj < alpha]
#get true positive sets
sceptre_true = sceptre_sig[sceptre_sig %in% overlap_genes]
seurat_true = seurat_sig[seurat_sig %in% overlap_genes]</pre>
```

Table 1: SCEPTRE Vs Seurat:IRF1 Perturbation True Positives

	Total Positive Genes	Unique to SCEPTRE	Unique to Seurat	Shared
True Positives	1644	66	44	160

```
#get number of shared and unique true postives
shared true = sceptre true[sceptre true %in% seurat true]
sceptre true unique = sceptre true[(sceptre true %in% seurat true) == F]
seurat_true_unique = seurat_true[(seurat_true %in% sceptre_true) == F]
#qet sensitivity
sceptre_sensitivity = length(sceptre_true)/length(overlap_genes)
seurat_sensitivity = length(seurat_true)/length(overlap_genes)
#qet false positive sets
sceptre_false = sceptre_sig[sceptre_sig %in% null_genes]
seurat_false = seurat_sig[seurat_sig %in% null_genes]
#get number of shared and unique false postives
shared_false = sceptre_false[sceptre_false %in% seurat_false]
sceptre_false_unique = sceptre_false[(sceptre_false %in% seurat_false) == F]
seurat_false_unique = seurat_false[(seurat_false %in% sceptre_false) == F]
#qet specificity
sceptre_specificity = 1-length(sceptre_false)/length(null_genes)
seurat_specificity = 1-length(seurat_false)/length(null_genes)
```

#### True Positive Table

Table 2: SCEPTRE Vs Seurat:IRF1 Perturbation False Positives

	Total Null Genes	Unique to SCEPTRE	Unique to Seurat	Shared
False Positives	11483	433	258	1341

#### True Positive Table

Table 3: SCEPTRE Vs Seurat:IRF1 Perturbation Sensitivity and Specificity

	Sensitivity	Specificity
SCEPTRE Seurat	0.1374696 0.1240876	$0.8455108 \\ 0.8607507$

## Sensitivity Specificity Table

Table 4: SCEPTRE Vs Seurat:IRF1 Perturbation CHIPseq and Reference Agreement

	Unique to CHIPseq	Unique to Reference	Shared
IRF1 Targets	491	345	9

#### Get Top Genes by Score

```
TF = direct_effects$TF
scores = direct_effects$score
top_genes = sort(TF[order(scores, decreasing = T)][1:500])
```

### Read in Reference Dataset For IRF1 TF Targets

```
data.dir = paste0(code_dir,"/sceptre2-manuscript/writeups/papalexi_analysis/")
targets.dir = paste0(data.dir,'IRF1_targets.json')
prtb_targets = fromJSON(file = targets.dir)
targets = c()
for(j in c(1:length(prtb_targets$associations))){
   targets = c(targets,prtb_targets$associations[[j]]$gene$symbol)
}
targets = targets[(targets%in%sceptre_prtb$response_id)]
null_targets =sceptre_prtb$response_id[(sceptre_prtb$response_id%in%targets)==F]
```

### Table For Overlap Of Gene Sets

## Compare Significant Gene Sets to Reference Data

```
alpha = 0.05
#qet all significant genes from each method
sceptre_sig = unfactor(sceptre_prtb$response_id[sceptre_prtb$p_value_adj < alpha])</pre>
seurat sig = rownames(seurat prtb)[seurat prtb$p val adj < alpha]</pre>
#get true positive sets
sceptre_true = sceptre_sig[sceptre_sig %in% targets]
seurat_true = seurat_sig[seurat_sig %in% targets]
#get number of shared and unique true postives
shared_true = sceptre_true[sceptre_true %in% seurat_true]
sceptre_true_unique = sceptre_true[(sceptre_true %in% seurat_true) == F]
seurat_true_unique = seurat_true[(seurat_true %in% sceptre_true) == F]
#qet sensitivity
sceptre_sensitivity = length(sceptre_true)/length(targets)
seurat_sensitivity = length(seurat_true)/length(targets)
#qet false positive sets
sceptre_false = sceptre_sig[sceptre_sig %in% null_targets]
seurat_false = seurat_sig[seurat_sig %in% null_targets]
#get number of shared and unique false postives
shared false = sceptre false[sceptre false %in% seurat false]
sceptre_false_unique = sceptre_false[(sceptre_false %in% seurat_false) == F]
seurat false unique = seurat false[(seurat false %in% sceptre false) == F]
#get specificity
sceptre specificity = 1-length(sceptre false)/length(null targets)
seurat_specificity = 1-length(seurat_false)/length(null_targets)
```

## True Positive Table

Table 5: SCEPTRE Vs Seurat:IRF1 Perturbation Reference Target True Positives

	Total Positive Genes	Unique to SCEPTRE	Unique to Seurat	Shared
True Positives	354	11	7	98

Table 6: SCEPTRE Vs Seurat:IRF1 Perturbation Reference Target False Positives

	Total Null Genes	Unique to SCEPTRE	Unique to Seurat	Shared
False Positives	13658	507	301	1465

#### False Positive Table

Table 7: SCEPTRE Vs Seurat:IRF1 Perturbation Reference Target Sensitivity and Specificity

	Sensitivity	Specificity
SCEPTRE Seurat	$\begin{array}{c} 0.3079096 \\ 0.2966102 \end{array}$	$\begin{array}{c} 0.8556158 \\ 0.8706985 \end{array}$

## Sensitivity and Specificity Table

## Bonus: Pvalue Scatterplot

```
sceptre_genes = unfactor(sceptre_prtb$response_id)
seurat_pvals = log(seurat_prtb[sceptre_genes,]$p_val,10)
sceptre_pvals = log(sceptre_prtb$p_value,10)
```

#### ## Warning: NaNs produced

```
#remove those with sceptre pval = 0
sceptre_0 = which(is.na(sceptre_pvals) == T)
sceptre_pvals = sceptre_pvals[-sceptre_0]
seurat_pvals = seurat_pvals[-sceptre_0]
```

```
#volcano plot
ggplot(Pval,aes(x = seuratP,y = sceptreP)) + geom_point() +
    ggtitle('Identity Plot of CUL3 Pvalues: SCEPTRE vs Unfiltered Seurat') +
    labs(y = 'Sceptre Pvalues',x = 'Seurat Pvalues')+
    geom_abline(slope=1, intercept = 0,color = 'red')+
    geom_vline(xintercept = log(0.05,10),color = 'red')+
    geom_hline(yintercept = log(0.05,10), color = 'red')
```

# Identity Plot of CUL3 Pvalues: SCEPTRE vs Unfiltered Seurat

