Figure 4 RAPseq

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2 Sep 2024 (10:40:08)

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0.1 ENV settings

```
###
# @Description: Figure4
# @Description: Adapted from https://github.com/IonutAtanasoai1/RAPseq
# @Author: LiQun
# @Email: qun.li@ki.se
# @Date: 2 Sep 2024 ( 10:40:08 )
###
rm(list = ls())
setwd("/Users/liqun/Desktop/Projects/RAPseq/AnalysisQun/")
```

0.2 R libraries

```
library(ChIPpeakAnno)
## Loading required package: IRanges
## 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: S4Vectors
## Loading required package: stats4
## Attaching package: 'S4Vectors'
## The following object is masked from 'package:utils':
##
## The following objects are masked from 'package:base':
##
      expand.grid, I, unname
## Loading required package: GenomicRanges
## Loading required package: GenomeInfoDb
## Warning: package 'GenomeInfoDb' was built under R version 4.3.3
library(edgeR)
## Loading required package: limma
## Attaching package: 'limma'
## The following object is masked from 'package:BiocGenerics':
##
      plotMA
library(ggfortify)
## Loading required package: ggplot2
library(tidyverse)
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
## v dplyr 1.1.4 v readr 2.1.5
## v forcats 1.0.0 v stringr 1.5.1
## v lubridate 1.9.3 v tibble
                                 3.2.1
                      v tidyr
## v purrr
            1.0.2
                                  1.3.1
## -- Conflicts ----- tidyverse_conflicts() --
## x lubridate::%within%() masks IRanges::%within%()
## x dplyr::collapse() masks IRanges::collapse()
## x dplyr::filter()
                    masks stats::filter()
```

```
masks S4Vectors::first()
## x dplyr::first()
## x dplyr::lag()
                           masks stats::lag()
                           masks BiocGenerics::Position(), base::Position()
## x ggplot2::Position()
## x purrr::reduce()
                           masks GenomicRanges::reduce(), IRanges::reduce()
                           masks S4Vectors::rename()
## x dplyr::rename()
## x lubridate::second() masks S4Vectors::second()
## x lubridate::second<-() masks S4Vectors::second<-()</pre>
## x dplyr::slice()
                           masks IRanges::slice()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become error
library(scales)
##
## Attaching package: 'scales'
##
## The following object is masked from 'package:purrr':
##
##
       discard
## The following object is masked from 'package:readr':
##
##
       col_factor
library(ggplot2)
library(dendextend)
##
## -----
## Welcome to dendextend version 1.17.1
## Type citation('dendextend') for how to cite the package.
## Type browseVignettes(package = 'dendextend') for the package vignette.
## The github page is: https://github.com/talgalili/dendextend/
##
## Suggestions and bug-reports can be submitted at: https://github.com/talgalili/dendextend/issues
## You may ask questions at stackoverflow, use the r and dendextend tags:
##
    https://stackoverflow.com/questions/tagged/dendextend
##
## To suppress this message use: suppressPackageStartupMessages(library(dendextend))
##
##
## Attaching package: 'dendextend'
## The following object is masked from 'package:stats':
##
##
       cutree
library(dichromat)
library(reshape2)
## Attaching package: 'reshape2'
## The following object is masked from 'package:tidyr':
##
##
       smiths
library(dplyr)
library(stringr)
```

```
library(GenomicRanges)
library(GenomicFeatures)
## Warning: package 'GenomicFeatures' was built under R version 4.3.3
## Loading required package: AnnotationDbi
## Loading required package: Biobase
## Welcome to Bioconductor
##
##
       Vignettes contain introductory material; view with
       'browseVignettes()'. To cite Bioconductor, see
##
##
       'citation("Biobase")', and for packages 'citation("pkgname")'.
##
##
## Attaching package: 'AnnotationDbi'
##
## The following object is masked from 'package:dplyr':
##
##
       select
library(gplots)
## Attaching package: 'gplots'
## The following object is masked from 'package: IRanges':
##
##
       space
##
## The following object is masked from 'package:S4Vectors':
##
##
       space
##
## The following object is masked from 'package:stats':
##
##
       lowess
library(clusterProfiler)
## Warning: package 'clusterProfiler' was built under R version 4.3.3
##
## clusterProfiler v4.10.1 For help: https://yulab-smu.top/biomedical-knowledge-mining-book/
##
## If you use clusterProfiler in published research, please cite:
## T Wu, E Hu, S Xu, M Chen, P Guo, Z Dai, T Feng, L Zhou, W Tang, L Zhan, X Fu, S Liu, X Bo, and G Yu.
## Attaching package: 'clusterProfiler'
## The following object is masked from 'package: Annotation Dbi':
##
##
       select
##
## The following object is masked from 'package:purrr':
##
##
       simplify
##
## The following object is masked from 'package: IRanges':
##
##
       slice
```

```
## The following object is masked from 'package:S4Vectors':
##
       rename
##
## The following object is masked from 'package:stats':
##
##
       filter
library(org.Hs.eg.db)
library(org.Dr.eg.db)
##
library(tidyr)
library(gtools)
library(ggrepel)
library(ggbeeswarm)
library(ReactomePA)
## ReactomePA v1.46.0 For help: https://yulab-smu.top/biomedical-knowledge-mining-book/
##
## If you use ReactomePA in published research, please cite:
## Guangchuang Yu, Qing-Yu He. ReactomePA: an R/Bioconductor package for reactome pathway analysis and
library(VennDiagram)
## Loading required package: grid
## Loading required package: futile.logger
## Attaching package: 'futile.logger'
##
## The following object is masked from 'package:gtools':
##
##
       scat
##
##
## Attaching package: 'VennDiagram'
## The following object is masked from 'package:dendextend':
##
##
       rotate
library(LSD)
library(gridExtra)
## Attaching package: 'gridExtra'
## The following object is masked from 'package:Biobase':
##
##
       combine
##
## The following object is masked from 'package:dplyr':
##
       combine
## The following object is masked from 'package:BiocGenerics':
##
##
       combine
```

```
library(UpSetR)
library(ggpubr)
##
## Attaching package: 'ggpubr'
##
## The following object is masked from 'package: VennDiagram':
##
##
       rotate
##
## The following object is masked from 'package:dendextend':
##
       rotate
library(GenomicScores)
## Attaching package: 'GenomicScores'
## The following object is masked from 'package:utils':
##
##
       citation
library(phastCons7way.UCSC.hg38)
## Warning: replacing previous import 'utils::findMatches' by
## 'S4Vectors::findMatches' when loading 'phastCons7way.UCSC.hg38'
library(phastCons100way.UCSC.hg38)
## Warning: replacing previous import 'utils::findMatches' by
## 'S4Vectors::findMatches' when loading 'phastCons100way.UCSC.hg38'
library(corrplot)
## Warning: package 'corrplot' was built under R version 4.3.3
## corrplot 0.94 loaded
library(ggforce)
library(idr)
library(GGally)
## Registered S3 method overwritten by 'GGally':
##
    method from
   +.gg ggplot2
library(eulerr)
## Registered S3 method overwritten by 'eulerr':
## method from
   plot.venn gplots
##
## Attaching package: 'eulerr'
## The following object is masked from 'package:gplots':
##
##
       venn
library(ggseqlogo)
library(robustbase)
## Warning: package 'robustbase' was built under R version 4.3.3
## Attaching package: 'robustbase'
## The following object is masked from 'package:Biobase':
##
##
       rowMedians
```

```
library(rrvgo)
## Warning: package 'rrvgo' was built under R version 4.3.3
```

0.3 Color settings

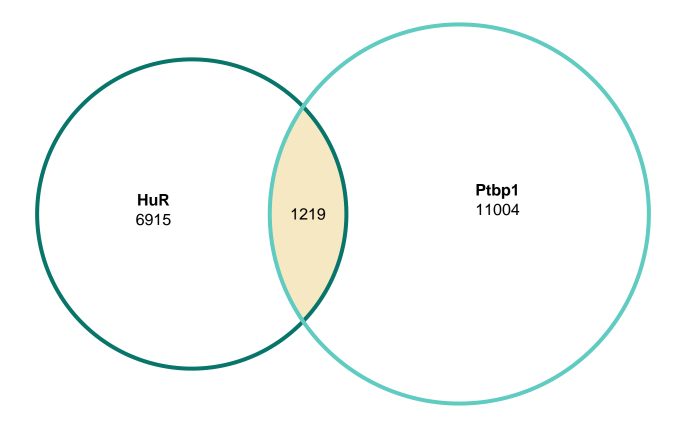
0.4 Load data

0.5 Figures

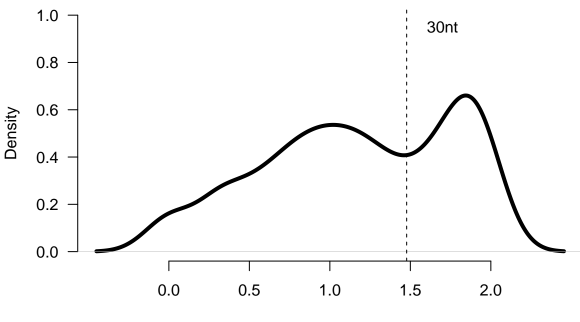
0.5.1 Figure 4A

```
# HUR
HUR_selected_forPlot <- HUR[,c(1,7,8,4:6)]
HUR_selected_forPlot$Summit_start <- HUR_selected_forPlot$Summit_start - 50
HUR_selected_forPlot$Summit_end <- HUR_selected_forPlot$Summit_end + 50
# PTBP1
PTBP1_selected_forPlot <- PTBP1[,c(1,7,8,4:6)]</pre>
```

```
PTBP1_selected_forPlot$Summit_start <- PTBP1_selected_forPlot$Summit_start - 50
PTBP1_selected_forPlot$Summit_end <- PTBP1_selected_forPlot$Summit_end + 50
HUR GRanges <- makeGRangesFromDataFrame(HUR selected forPlot)</pre>
PTBP1_GRanges <- makeGRangesFromDataFrame(PTBP1_selected_forPlot)</pre>
# uniq and common
HUR GRanges uniq <- unique(HUR[-as.data.frame(findOverlaps(HUR GRanges,PTBP1 GRanges,type = "any"))[,1]
HUR_PTBP1_GRanges_common <- as.character(seq(1,nrow( as.data.frame(findOverlaps(HUR_GRanges,PTBP1_GRang
HUR_GRanges_merge <- c(HUR_GRanges_uniq,HUR_PTBP1_GRanges_common)</pre>
PTBP1_GRanges_uniq <- unique(PTBP1[-as.data.frame(findOverlaps(HUR_GRanges,PTBP1_GRanges,type = "any"))
PTBP1_GRanges_merge <- c(PTBP1_GRanges_uniq, HUR_PTBP1_GRanges_common)
HUR_PTBP1_GRanges_list <- list(HUR_GRanges_merge, PTBP1_GRanges_merge)</pre>
names(HUR_PTBP1_GRanges_list) <- c("HuR", "Ptbp1")</pre>
# plot
HUR_PTBP1_overlaps_forPlot <- euler(HUR_PTBP1_GRanges_list, shape="circle")</pre>
HUR overlaps <- HUR[as.data.frame(</pre>
  findOverlaps(HUR_GRanges, PTBP1_GRanges, type = "any"))[,1],
  "Summit start"]
PTBP1_overlaps <- PTBP1[as.data.frame(</pre>
  findOverlaps(HUR GRanges, PTBP1 GRanges, type = "any"))[,2],
  "Summit start"]
#pdf("./Figure/Figure4/Figure4A_1.pdf", width = 8, height = 5)
plot(HUR_PTBP1_overlaps_forPlot, fills=c("white", "white", "#f6e8c3"),
     quantities=TRUE, edges=T, col=c(acqua_greens[c(3,8)]), lwd=4)
```



bw = 0.15



log10(distance between binding sites)

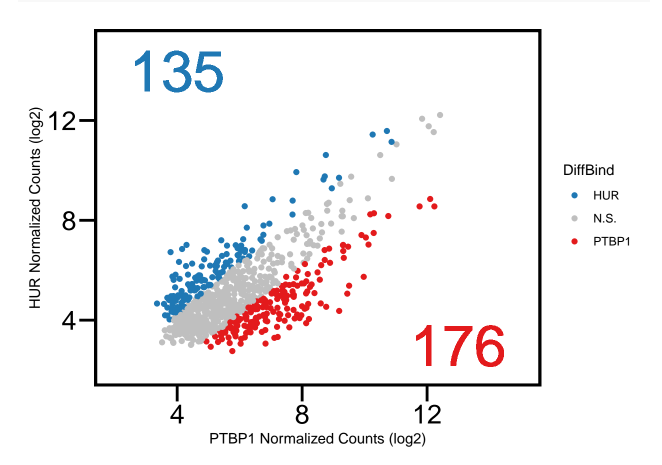
#dev.off()

0.5.2 Figure 4B

```
# extract 2 replicates
HUR_replicates <- HUR[as.data.frame(findOverlaps(HUR_GRanges, PTBP1_GRanges, type = "any"))[,1],][,c("R</pre>
PTBP1_replicates <- PTBP1[as.data.frame(findOverlaps(HUR_GRanges, PTBP1_GRanges, type = "any"))[,2],][,
HUR_comp <- HUR_replicates[log10(abs(PTBP1_overlaps-HUR_overlaps)+1)<1.492,]</pre>
PTBP1_comp <- PTBP1_replicates[log10(abs(PTBP1_overlaps-HUR_overlaps)+1)<1.492,]
HUR_rep1 <- HUR_comp$Rep1</pre>
HUR_rep2 <- HUR_comp$Rep2</pre>
PTBP1_rep1 <- PTBP1_comp$Rep1
PTBP1_rep2 <- PTBP1_comp$Rep2
# merge two data
HUR_PTBP1_tworeplicates <- data.frame(HUR_rep1, HUR_rep2, PTBP1_rep1, PTBP1_rep2)</pre>
# DEG analysis
group \leftarrow c(1,1,2,2)
DEG_data <- DGEList(counts=HUR_PTBP1_tworeplicates, group = group)</pre>
design <- model.matrix(~group)</pre>
DEG_data <- estimateDisp(DEG_data,design)</pre>
```

```
fit <- glmQLFit(DEG_data, design)</pre>
qlf <- glmQLFTest(fit)</pre>
DEG_data_table <-qlf$table</pre>
DEG_data_table$p.adjust <- -log10(p.adjust(DEG_data_table$PValue, method = "BH"))
DEG_data_table$PValue <- -log10(DEG_data_table$PValue)</pre>
# HUR PTBP1 tworeplicates
HUR_PTBP1_tworeplicates$HuR <- log2((HUR_PTBP1_tworeplicates$HUR_rep1 + HUR_PTBP1_tworeplicates$HUR_rep
HUR_PTBP1_tworeplicates$PTBP1 <- log2((HUR_PTBP1_tworeplicates$PTBP1_rep1 + HUR_PTBP1_tworeplicates$PTB</pre>
HUR_PTBP1_tworeplicates$delta_BS <- abs(HUR_PTBP1_tworeplicates$HuR - HUR_PTBP1_tworeplicates$PTBP1)</pre>
HUR_PTBP1_tworeplicates$FDR <- DEG_data_table$p.adjust</pre>
HUR_PTBP1_tworeplicates$Significant <- HUR_PTBP1_tworeplicates$FDR >= 1.30103
HUR_PTBP1_tworeplicates$logFC <- DEG_data_table$logFC</pre>
HUR_PTBP1_tworeplicates$HuR_won <- HUR_PTBP1_tworeplicates$logFC <= -1 & HUR_PTBP1_tworeplicates$FDR >=
HUR_PTBP1_tworeplicates$PTBP1_won <- HUR_PTBP1_tworeplicates$logFC >= 1 & HUR_PTBP1_tworeplicates$FDR >
HUR_PTBP1_tworeplicates$DiffBind <- paste(HUR_PTBP1_tworeplicates$HuR_won,HUR_PTBP1_tworeplicates$PTBP1
HUR_PTBP1_tworeplicates$DiffBind <- gsub("FALSE_FALSE", "N.S.", HUR_PTBP1_tworeplicates$DiffBind)
HUR_PTBP1_tworeplicates$DiffBind <- gsub("FALSE_TRUE", "PTBP1", HUR_PTBP1_tworeplicates$DiffBind)
HUR_PTBP1_tworeplicates$DiffBind <- gsub("TRUE_FALSE","HUR",HUR_PTBP1_tworeplicates$DiffBind)</pre>
#pdf("./Figure/Figure4/Figure4B.pdf", width = 8, height = 5)
HUR PTBP1 DEGPlot <- ggplot(data=HUR PTBP1 tworeplicates) +</pre>
  geom_point(aes(x=PTBP1,y=HuR,color=DiffBind)) +
  scale_color_manual(values = c("#1f78b4","grey75","#e31a1c")) +
  theme(panel.background = element_blank(),
        panel.grid = element_blank(),
        panel.border = element_rect(size=2,fill=NA,color="black"),
        axis.ticks.length = unit(4,"mm"),
        axis.ticks = element_line(color="black",size=1),
        axis.text = element_text(color="black",size=20)) +
  xlim(2,15) +
  ylim(2,15) +
  xlab("PTBP1 Normalized Counts (log2)") +
 ylab("HUR Normalized Counts (log2)")
## Warning: The `size` argument of `element_line()` is deprecated as of ggplot2 3.4.0.
## i Please use the `linewidth` argument instead.
## This warning is displayed once every 8 hours.
## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was
## generated.
## Warning: The `size` argument of `element_rect()` is deprecated as of ggplot2 3.4.0.
## i Please use the `linewidth` argument instead.
## This warning is displayed once every 8 hours.
## Call `lifecycle::last_lifecycle_warnings()` to see where this warning was
## generated.
HUR_PTBP1_DEGPlot + geom_text(aes(x=13,y=3), label = table(HUR_PTBP1_tworeplicates$DiffBind)["PTBP1"],
  geom_text(aes(x=4,y=14), label = table(HUR_PTBP1_tworeplicates$DiffBind)["HUR"], size=15, color="#1f7
## Warning in geom_text(aes(x = 13, y = 3), label = table(HUR_PTBP1_tworeplicates DiffBind)["PTBP1"], :
## i Please consider using `annotate()` or provide this layer with data containing
## a single row.
```

```
## Warning in geom_text(aes(x = 4, y = 14), label = table(HUR_PTBP1_tworeplicates$DiffBind)["HUR"], : A
## i Please consider using `annotate()` or provide this layer with data containing
## a single row.
```



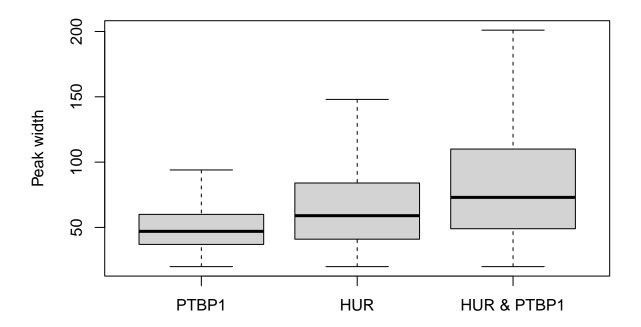
```
#dev.off()

# median sequenced fragment size HUR = 50
# median sequenced fragment size HURPTBP1_coRAP = 51
# median sequenced fragment size PTBP1 = 35

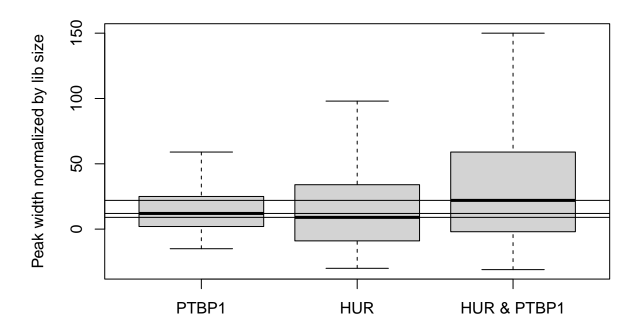
PTBP1_len <- PTBP1$end - PTBP1$start
PTBP1_len_adjust <- PTBP1_len - 35
HUR_len <- HUR$end - HUR$start
HUR_len_adjust <- HUR_len - 50
HUR_PTBP1_len <- HUR_PTBP1$end - HUR_PTBP1$start
HUR_PTBP1_len_adjust <- HUR_PTBP1_len - 51</pre>

HUR_PTBP1_len_list <- list(PTBP1_len, HUR_len, HUR_PTBP1_len)
names(HUR_PTBP1_len_list) <- c("PTBP1", "HUR", "HUR & PTBP1")
HUR_PTBP1_len_adjust_list <- list(PTBP1_len_adjust, HUR_len_adjust, HUR_PTBP1_len_adjust)
names(HUR_PTBP1_len_adjust_list) <- c("PTBP1", "HUR", "HUR & PTBP1")

boxplot(HUR_PTBP1_len_list, outline=F, ylab="Peak width")</pre>
```



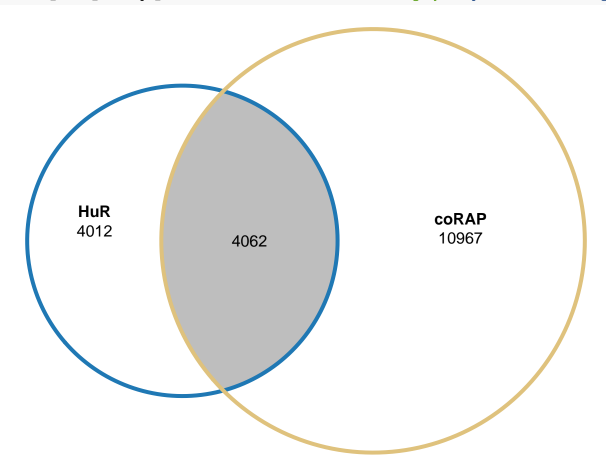
boxplot(HUR_PTBP1_len_adjust_list, outline=F, ylab="Peak width normalized by lib size")
abline(h=c(median(PTBP1_len_adjust), median(HUR_len_adjust), median(HUR_PTBP1_len_adjust)))



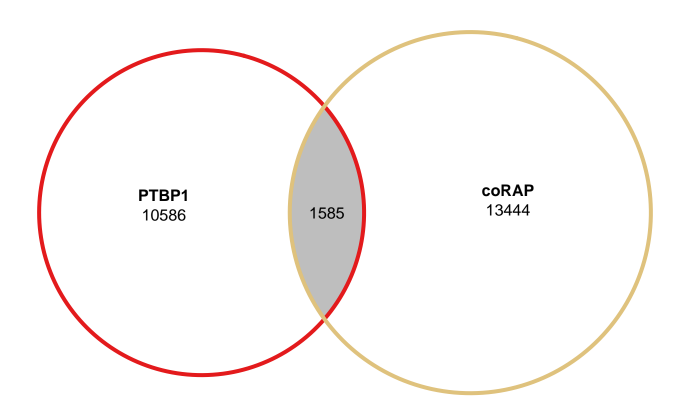
0.5.3 Figure 4E

```
\# median sequenced fragment size HUR = 50
# median sequenced fragment size HURPTBP1 = 51
\# median sequenced fragment size PTBP1 = 35
# HUR
HUR_selected_forPlot <- HUR[,c(1,7,8,4:6)]</pre>
HUR_selected_forPlot$Summit_start <- HUR_selected_forPlot$Summit_start - 50
HUR_selected_forPlot$Summit_end <- HUR_selected_forPlot$Summit_end + 50</pre>
# PTBP1
PTBP1_selected_forPlot <- PTBP1[,c(1,7,8,4:6)]</pre>
PTBP1_selected_forPlot$Summit_start <- PTBP1_selected_forPlot$Summit_start - 50
PTBP1_selected_forPlot$Summit_end <- PTBP1_selected_forPlot$Summit_end + 50
# HUR_PTBP1
HUR_PTBP1_selected_forPlot <- HUR_PTBP1[,c(1,7,8,4:6)]</pre>
HUR_PTBP1_selected_forPlot$Summit_start <- HUR_PTBP1_selected_forPlot$Summit_start - 50
HUR_PTBP1_selected_forPlot$Summit_end <- HUR_PTBP1_selected_forPlot$Summit_end + 50</pre>
HUR_GRanges <- makeGRangesFromDataFrame(HUR_selected_forPlot)</pre>
PTBP1_GRanges <- makeGRangesFromDataFrame(PTBP1_selected_forPlot)</pre>
HUR_PTBP1_GRanges <- makeGRangesFromDataFrame(HUR_PTBP1_selected_forPlot)</pre>
```

```
# uniq and common
HUR_GRanges_uniq <- unique(HUR[-as.data.frame(findOverlaps(HUR_GRanges,HUR_PTBP1_GRanges,type = "any"))</pre>
HUR_PTBP1_GRanges_uniq <- unique(HUR_PTBP1[as.data.frame(findOverlaps(HUR_GRanges,HUR_PTBP1_GRanges,typ
HUR_PTBP1_GRanges_merge <- c(HUR_GRanges_uniq,HUR_PTBP1_GRanges_uniq)</pre>
coRAP1 <- unique(HUR_PTBP1[-as.data.frame(findOverlaps(HUR_GRanges, HUR_PTBP1_GRanges, type = "any"))[,2]
coRAP1 <- c(coRAP1, HUR_PTBP1_GRanges_uniq)</pre>
HUR_coRAP <- list(HUR_PTBP1_GRanges_merge, coRAP1)</pre>
names(HUR coRAP) <- c("HuR", "coRAP")</pre>
HUR_coRAP_overlaps_forPlot <- euler(HUR_coRAP, shape="circle")</pre>
PTBP1_GRanges_uniq <- unique(PTBP1[-as.data.frame(findOverlaps(PTBP1_GRanges, HUR_PTBP1_GRanges, type = "
PTBP1_HUR_GRanges_uniq <- unique(HUR_PTBP1[as.data.frame(find0verlaps(PTBP1_GRanges,HUR_PTBP1_GRanges,t
PTBP1_HUR_GRanges_merge <- c(PTBP1_GRanges_uniq,PTBP1_HUR_GRanges_uniq)
coRAP2 <- unique(HUR_PTBP1[-as.data.frame(findOverlaps(PTBP1_GRanges, HUR_PTBP1_GRanges, type = "any"))[,
coRAP2 <- c(coRAP2,PTBP1_HUR_GRanges_uniq)</pre>
PTBP1_coRAP <- list(PTBP1_HUR_GRanges_merge, coRAP2)</pre>
names(PTBP1_coRAP) <- c("PTBP1", "coRAP")</pre>
PTBP1_coRAP_overlaps_forPlot <- euler(PTBP1_coRAP, shape="circle")
# plot
#pdf("./Figure/Figure4/Figure4E_Venn_HUR.pdf", width = 6, height = 3)
plot(HUR_coRAP_overlaps_forPlot, fills=c("white", "white", "grey"), quantities=TRUE, edges=T, col=c("#1
```

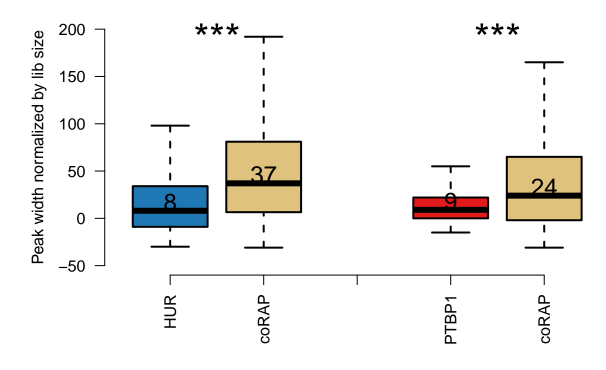


```
#dev.off()
#pdf("./Figure/Figure4/Figure4E_Venn_PTBP1.pdf", width = 6, height = 3)
plot(PTBP1_coRAP_overlaps_forPlot, fills=c("white", "white", "grey"), quantities=TRUE, edges=T, col=c("state of the color of
```



```
#dev.off()
# overlaps
HUR_PTBP1_HURPTBP1_overlaps <- HUR_PTBP1[as.data.frame(findOverlaps(HUR_GRanges,HUR_PTBP1_GRanges,type=
HUR_PTBP1_PTBP1HUR_overlaps <- HUR_PTBP1[as.data.frame(findOverlaps(PTBP1_GRanges,HUR_PTBP1_GRanges,type="any")
HUR_HURPTBP1_overlaps <- PTBP1[as.data.frame(findOverlaps(PTBP1_GRanges,HUR_PTBP1_GRanges,type="any")
HUR_HURPTBP1_overlaps <- HUR[as.data.frame(findOverlaps(HUR_GRanges,HUR_PTBP1_GRanges,type="any"))[,1],
PTBP1_HURPTBP1_overlaps_adjust <- (PTBP1_HURPTBP1_overlaps$end - PTBP1_HURPTBP1_overlaps$start) - 35
HUR_PTBP1_PTBP1HUR_overlaps_adjust <- (HUR_PTBP1_PTBP1HUR_overlaps$end-HUR_PTBP1_PTBP1HUR_overlaps$star
HUR_HURPTBP1_overlaps_adjust <- (HUR_HURPTBP1_overlaps$end - HUR_HURPTBP1_overlaps$star) - 50
HUR_PTBP1_HURPTBP1_overlaps_adjust <- (HUR_PTBP1_HURPTBP1_overlaps$end-HUR_PTBP1_HURPTBP1_overlaps$star
All_overlaps_adjust_list <- list(HUR_HURPTBP1_overlaps_adjust,HUR_PTBP1_HURPTBP1_overlaps_adjust,NULL_P")
names(All_overlaps_adjust_list) <- c("HUR", "corAP",NA,"PTBP1", "corAP")
##pdf("./Figure/Figure4/Figure4E_boxplot.pdf", width = 6, height = 7)
par(bty="n")
boxplot(All_overlaps_adjust_list, outline=F, ylab="Peak width normalized by lib size", las=2, ylim=c(-5)</pre>
```

```
text(x=1,y=median(HUR_HURPTBP1_overlaps_adjust)+9, label=median(HUR_HURPTBP1_overlaps_adjust), cex=1.5)
text(x=2,y=median(HUR_PTBP1_HURPTBP1_overlaps_adjust)+9, label=median(HUR_PTBP1_HURPTBP1_overlaps_adjust)+9
text(x=4,y=median(PTBP1_HURPTBP1_overlaps_adjust)+9, label=median(PTBP1_HURPTBP1_overlaps_adjust), cex=
text(x=5,y=median(HUR_PTBP1_PTBP1HUR_overlaps_adjust)+9, label=median(HUR_PTBP1_PTBP1HUR_overlaps_adjust)+9
pw <- wilcox.test(PTBP1_HURPTBP1_overlaps_adjust, HUR_PTBP1_PTBP1HUR_overlaps_adjust)
pwp <- pw$p.value
if ( pwp <= 0.001 ) {
 to_add <- "***"
} else if ( btp <= 0.01 ) {</pre>
 to_add <- "**"
} else if ( btp \leftarrow 0.05 ) {
  to add <- "*"
} else {
  to_add <- "n.s."
}
text(1.5,200, labels = to_add, cex=2.5)
pw <- wilcox.test(HUR_HURPTBP1_overlaps_adjust, HUR_PTBP1_HURPTBP1_overlaps_adjust)
pwp <- pw$p.value</pre>
if ( pwp <= 0.001 ) {
 to_add <- "***"
} else if ( btp <= 0.01 ) {</pre>
 to_add <- "**"
} else if ( btp <= 0.05 ) {</pre>
 to_add <- "*"
} else {
  to_add <- "n.s."
text(4.5,200,labels = to_add, cex=2.5)
```



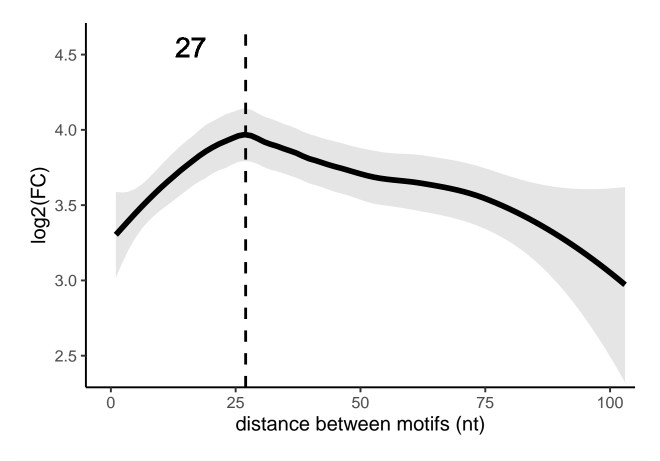
0.5.4 Figure 4F

```
HUR_PTBP1_uniq_4F <- unique(HUR_PTBP1[-as.data.frame(findOverlaps(HUR_GRanges, HUR_PTBP1_GRanges,type = HUR_PTBP1_uniq_4F_select <- HUR_PTBP1_uniq_4F[,c(1,7,8,4:6)]
HUR_PTBP1_uniq_4F_select$Summit_start <- HUR_PTBP1_uniq_4F_select$Summit_start - 50
HUR_PTBP1_uniq_4F_select$Summit_end <- HUR_PTBP1_uniq_4F_select$Summit_end + 50
HUR_PTBP1_uniq_4F_select_adjust <- makeGRangesFromDataFrame(HUR_PTBP1_uniq_4F_select)

HUR_PTBP1_uniq_4F_overlap <- unique(HUR_PTBP1_uniq_4F[-as.data.frame(findOverlaps(PTBP1_GRanges, HUR_PTBP1_uniq_4F_overlap$positive_fa <- str_sub(HUR_PTBP1_uniq_4F_overlap$positive_fa,43,157)

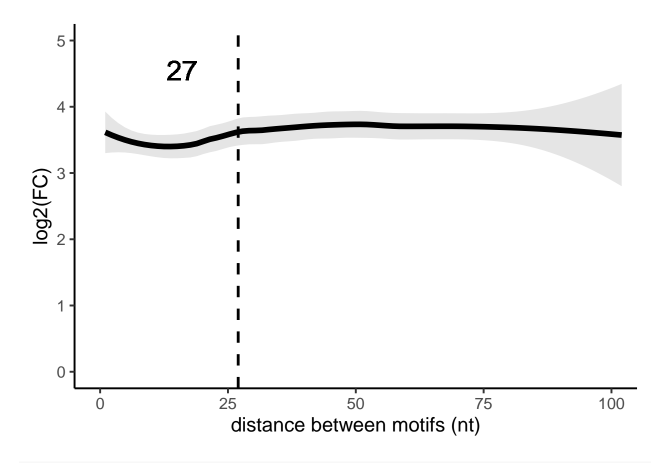
bases <- c("A", "C", "T", "G")
kmers_3 <- unite(as.data.frame(permutations(n=4,r=3,v=bases,repeats.allowed = T)),col = kmers,sep = "")
TTT_kmers_3 <- paste("TTT",kmers_3,"TTT",sep="")
kmers_2 <- unite(as.data.frame(permutations(n=4,r=2,v=bases,repeats.allowed = T)),col = kmers,sep = "")
TTT_kmers_2 <- paste("TTT",kmers_2,"TTT",sep="")
TTT_kmers_2 <- paste("TTT",kmers_2,"TTT",sep="")
TTT_kmers_2 <- paste(TTT_kmers_2,"TTT",sep="")
TTT_kmers_1 <- c("TTTTTTT","TTTGTTT","TTTGTTT","TTTGTTT")
```

```
hur_kmers <- c(TTT_kmers_1, TTT_kmers_2, TTT_kmers_3)</pre>
ptbp1_kmers <- c("TCTCT","TTTCT","CTTCT","TCTCT","CCTCT","GTTCT","GTTCT","GCTCT","GCTCT","GCTCT","GTTCT
for (i in hur kmers){
  HUR_PTBP1_uniq_4F_overlap$positive_fa <- gsub(i, "U", HUR_PTBP1_uniq_4F_overlap$positive_fa)
for (i in ptbp1 kmers){
 HUR_PTBP1_uniq_4F_overlap$positive_fa <- gsub(i,"Y",HUR_PTBP1_uniq_4F_overlap$positive_fa)
}
HUR_PTBP1_uniq_4F_overlap$\forall ys <- str_count(HUR_PTBP1_uniq_4F_overlap$\forall positive_fa,"Y")</pre>
HUR_PTBP1_uniq_4F_overlap$Us <- str_count(HUR_PTBP1_uniq_4F_overlap$positive_fa,"U")</pre>
HUR_PTBP1_uniq_4F_overlap$TTTs <- str_count(HUR_PTBP1_uniq_4F_overlap$positive_fa,"TTT")</pre>
HUR_PTBP1_uniq_4F_overlap_YsUs <- HUR_PTBP1_uniq_4F_overlap[HUR_PTBP1_uniq_4F_overlap$Ys == 1 & HUR_PTB
HUR_PTBP1_uniq_4F_overlap_YsUs$Y_loc <- unlist(str_locate_all(HUR_PTBP1_uniq_4F_overlap_YsUs$positive_f
HUR_PTBP1_uniq_4F_overlap_YsUs$U_loc <- unlist(str_locate_all(HUR_PTBP1_uniq_4F_overlap_YsUs$positive_f
HUR_PTBP1_uniq_4F_overlap_YsUs$distance <- abs(HUR_PTBP1_uniq_4F_overlap_YsUs$Y_loc - HUR_PTBP1_uniq_4F
HUR_PTBP1_uniq_4F_overlap_YsUs_plot <- ggplot2::ggplot(data=HUR_PTBP1_uniq_4F_overlap_YsUs, aes(x=dista
  stat smooth(method="loess", color="black", fill="grey", span=0.6, level=0.95, lwd=2) +
  theme_classic2(base_size = 15) +
  coord_cartesian(xlim=c(0,100), ylim=c(2.4,4.6)) +
  geom_vline(xintercept = c(27), color="black", lty=2, lwd=1) +
  geom_text(aes(x=16,y=4.55), label = 27, size=7.5, col="black") +
  xlab("distance between motifs (nt)") +
  ylab("log2(FC)")
#pdf("./Figure/Figure4/Figure4F.pdf", width = 6, height = 6)
HUR_PTBP1_uniq_4F_overlap_YsUs_plot
## Warning in geom_text(aes(x = 16, y = 4.55), label = 27, size = 7.5, col = "black"): All aesthetics h
## i Please consider using `annotate()` or provide this layer with data containing
     a single row.
## `geom_smooth()` using formula = 'y ~ x'
```



0.5.5 Figure S4A

```
hur_kmers <- c(TTT_kmers_1,TTT_kmers_2,TTT_kmers_3)</pre>
ptbp1_kmers <- c("TCTCT","TTTCT","CTTCT","CTTCT","CCTCT","GTTCT","CTTCT","GCTCT","GCTCT","GCTCT","GTTCT
for (i in hur_kmers){
 HUR_PTBP1_uniq_S4A_overlap$negative_fa <- gsub(i,"U",HUR_PTBP1_uniq_S4A_overlap$negative_fa)
for (i in ptbp1_kmers){
  HUR_PTBP1_uniq_S4A_overlap$negative_fa <- gsub(i, "Y", HUR_PTBP1_uniq_S4A_overlap$negative_fa)
}
HUR PTBP1 uniq S4A overlap$Ys <- str count(HUR PTBP1 uniq S4A overlap$negative fa, "Y")
HUR_PTBP1_uniq_S4A_overlap$Us <- str_count(HUR_PTBP1_uniq_S4A_overlap$negative_fa,"U")</pre>
HUR_PTBP1_uniq_S4A_overlap$TTTs <- str_count(HUR_PTBP1_uniq_S4A_overlap$negative_fa,"TTT")</pre>
HUR_PTBP1_uniq_S4A_overlap_YsUs <- HUR_PTBP1_uniq_S4A_overlap[HUR_PTBP1_uniq_S4A_overlap$Ys == 1 & HUR_
HUR_PTBP1_uniq_S4A_overlap_YsUs$Y_loc <- unlist(str_locate_all(HUR_PTBP1_uniq_S4A_overlap_YsUs$negative
HUR_PTBP1_uniq_S4A_overlap_YsUs$U_loc <- unlist(str_locate_all(HUR_PTBP1_uniq_S4A_overlap_YsUs$negative
HUR_PTBP1_uniq_S4A_overlap_YsUs$distance <- abs(HUR_PTBP1_uniq_S4A_overlap_YsUs$Y_loc - HUR_PTBP1_uniq_s
HUR_PTBP1_uniq_S4A_overlap_YsUs_plot <- ggplot2::ggplot(data=HUR_PTBP1_uniq_S4A_overlap_YsUs, aes(x=dis
  stat smooth(method="loess", color="black", fill="grey", span=0.6, level=0.95, lwd=2) +
  theme classic2(base size = 15) +
  coord_cartesian(xlim=c(0,100), ylim=c(0, 5)) +
  geom_vline(xintercept = c(27), color="black", lty=2, lwd=1) +
  geom_text(aes(x=16,y=4.55), label = 27, size=7.5, col="black") +
  xlab("distance between motifs (nt)") +
  ylab("log2(FC)")
#pdf("./Figure/Figure4/FigureS4A.pdf", width = 6, height = 6)
HUR_PTBP1_uniq_S4A_overlap_YsUs_plot
## Warning in geom_text(aes(x = 16, y = 4.55), label = 27, size = 7.5, col = "black"): All aesthetics h
## i Please consider using `annotate()` or provide this layer with data containing
## a single row.
## `geom_smooth()` using formula = 'y ~ x'
```



0.5.6 Figure S4B

```
HUR_uniq_S4B <- unique(HUR[-as.data.frame(findOverlaps(HUR_GRanges,HUR_PTBP1_GRanges,type = "any"))[,1]

HUR_uniq_S4B_select <- HUR_uniq_S4B[,c(1,7,8,4:6)]

HUR_uniq_S4B_select$Summit_start <- HUR_uniq_S4B_select$Summit_start - 50

HUR_uniq_S4B_select$Summit_end <- HUR_uniq_S4B_select$Summit_end + 50

HUR_uniq_S4B_select_GRanges <- makeGRangesFromDataFrame(HUR_uniq_S4B_select)

HUR_uniq_S4B_overlaps <- unique(HUR_uniq_S4B[-as.data.frame(findOverlaps(PTBP1_GRanges, HUR_uniq_S4B_seluct))

HUR_uniq_S4B_overlaps$positive_fa <- str_sub(HUR_uniq_S4B_overlaps$positive_fa,43,157)

bases <- c("A","C","T","G")

kmers_3 <- unite(as.data.frame(permutations(n=4,r=3,v=bases,repeats.allowed = T)),col = kmers,sep = "")

TTT_kmers_3 <- paste("TTT",kmers_3,"TTT",sep="")

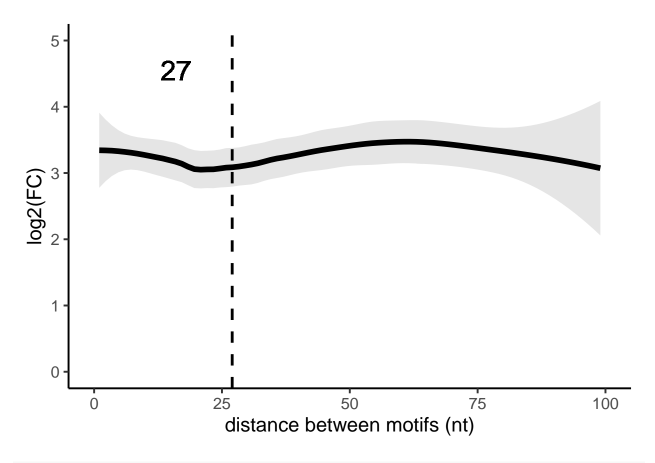
kmers_2 <- unite(as.data.frame(permutations(n=4,r=2,v=bases,repeats.allowed = T)),col = kmers,sep = "")

TTT_kmers_2 <- paste("TTT",kmers_2,sep="")

TTT_kmers_2 <- paste("TTT",kmers_2,sep="")

TTT_kmers_2 <- paste("TTT_kmers_2,"TTT",sep="")
```

```
TTT_kmers_1 <- c("TTTTTTT","TTTATTT","TTTGTTT","TTTCTTT")</pre>
hur_kmers <- c(TTT_kmers_1,TTT_kmers_2,TTT_kmers_3)</pre>
ptbp1_kmers <- c("TCTCT","TTTCT","TTTCT","CTTCT","CCTCT","GTTCT","GTTCT","GCTCT","GCTCT","GCTCT","GCTCT","GTTCT
for (i in hur kmers){
  HUR_uniq_S4B_overlaps$positive_fa <- gsub(i,"U",HUR_uniq_S4B_overlaps$positive_fa)
for (i in ptbp1 kmers){
 HUR_uniq_S4B_overlaps$positive_fa <- gsub(i,"Y",HUR_uniq_S4B_overlaps$positive_fa)
}
HUR_uniq_S4B_overlaps$Ys <- str_count(HUR_uniq_S4B_overlaps$positive_fa,"Y")</pre>
HUR_uniq_S4B_overlaps$Us <- str_count(HUR_uniq_S4B_overlaps$positive_fa,"U")
HUR_uniq_S4B_overlaps$TTTs <- str_count(HUR_uniq_S4B_overlaps$positive_fa,"TTT")</pre>
HUR_uniq_S4B_overlaps_YsUs <- HUR_uniq_S4B_overlaps[HUR_uniq_S4B_overlaps$Ys == 1 & HUR_uniq_S4B_overla
HUR_uniq_S4B_overlaps_YsUs$Y_loc <- unlist(str_locate_all(HUR_uniq_S4B_overlaps_YsUs$positive_fa,"Y"))[
HUR_uniq_S4B_overlaps_YsUs$U_loc <- unlist(str_locate_all(HUR_uniq_S4B_overlaps_YsUs$positive_fa,"U"))[
HUR_uniq_S4B_overlaps_YsUs$distance <- abs(HUR_uniq_S4B_overlaps_YsUs$Y_loc - HUR_uniq_S4B_overlaps_YsU
HUR_uniq_S4B_overlaps_YsUs_plot <- ggplot2::ggplot(data=HUR_uniq_S4B_overlaps_YsUs, aes(x=distance, y=1
  stat_smooth(method="loess", color="black", fill="grey", span=0.6, level=0.95, lwd=2) +
  theme classic2(base size = 15) +
  coord_cartesian(xlim=c(0,100), ylim=c(0,5)) +
  geom_vline(xintercept = c(27), color="black", lty=2, lwd=1) +
  geom_text(aes(x=16,y=4.55), label = 27, size=7.5, col="black") +
  xlab("distance between motifs (nt)") +
  ylab("log2(FC)")
#pdf("./Figure/Figure4/FigureS4B.pdf", width = 6, height = 6)
HUR_uniq_S4B_overlaps_YsUs_plot
## Warning in geom_text(aes(x = 16, y = 4.55), label = 27, size = 7.5, col = "black"): All aesthetics h
## i Please consider using `annotate()` or provide this layer with data containing
## a single row.
## `geom_smooth()` using formula = 'y ~ x'
```



0.5.7 Figure S4C

```
PTBP1_uniq_S4C <- unique(PTBP1[-as.data.frame(findOverlaps(PTBP1_GRanges,HUR_PTBP1_GRanges,type = "any"

PTBP1_uniq_S4C_select <- PTBP1_uniq_S4C[,c(1,7,8,4:6)]

PTBP1_uniq_S4C_select$Summit_start <- PTBP1_uniq_S4C_select$Summit_start - 50

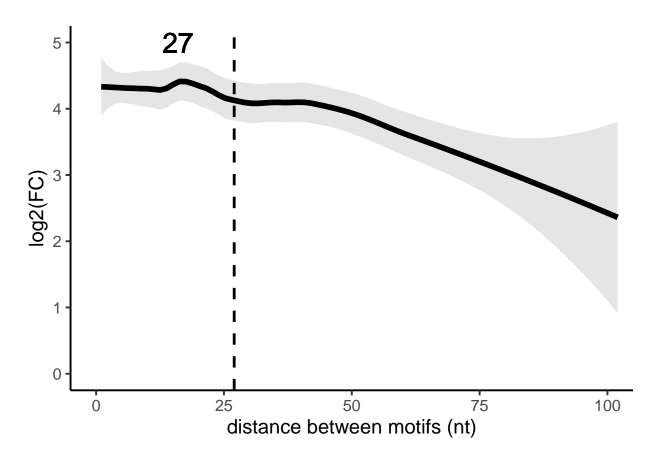
PTBP1_uniq_S4C_select$Summit_end <- PTBP1_uniq_S4C_select$Summit_end + 50

PTBP1_uniq_S4C_select_GRanges <- makeGRangesFromDataFrame(PTBP1_uniq_S4C_select)

PTBP1_uniq_S4C_select_GRanges_overlaps <- unique(PTBP1_uniq_S4C[-as.data.frame(findOverlaps(HUR_GRanges)])

PTBP1_uniq_S4C_select_GRanges_overlaps$positive_fa <- str_sub(PTBP1_uniq_S4C_select_GRanges_overlaps$positive_fa <-
```

```
TTT_kmers_1 <- c("TTTTTTT","TTTATTT","TTTGTTT","TTTCTTT")</pre>
hur_kmers <- c(TTT_kmers_1,TTT_kmers_2,TTT_kmers_3)</pre>
ptbp1_kmers <- c("TCTCT","TTTCT","CTTCT","TCTCT","CCTCT","GTTCT","GTTCT","GCTCT","GCTCT","GCTCT","GTTCT
for (i in hur kmers){
  PTBP1_uniq_S4C_select_GRanges_overlaps$positive_fa <- gsub(i, "U", PTBP1_uniq_S4C_select_GRanges_overla
for (i in ptbp1 kmers){
 PTBP1_uniq_S4C_select_GRanges_overlaps$positive_fa <- gsub(i,"Y",PTBP1_uniq_S4C_select_GRanges_overla
}
PTBP1_uniq_S4C_select_GRanges_overlaps$ys <- str_count(PTBP1_uniq_S4C_select_GRanges_overlaps$positive_
PTBP1_uniq_S4C_select_GRanges_overlaps$Us <- str_count(PTBP1_uniq_S4C_select_GRanges_overlaps$positive_
PTBP1_uniq_S4C_select_GRanges_overlaps$TTTs <- str_count(PTBP1_uniq_S4C_select_GRanges_overlaps$positiv
PTBP1_uniq_S4C_select_GRanges_overlaps_YsUs <- PTBP1_uniq_S4C_select_GRanges_overlaps[PTBP1_uniq_S4C_se
PTBP1_uniq_S4C_select_GRanges_overlaps_YsUs$Y_loc <- unlist(str_locate_all(PTBP1_uniq_S4C_select_GRange
PTBP1_uniq_S4C_select_GRanges_overlaps_YsUs$U_loc <- unlist(str_locate_all(PTBP1_uniq_S4C_select_GRange
PTBP1_uniq_S4C_select_GRanges_overlaps_YsUs$distance <- abs(PTBP1_uniq_S4C_select_GRanges_overlaps_YsUs
PTBP1_uniq_S4C_select_GRanges_overlaps_YsUs_plot <- ggplot2::ggplot(data=PTBP1_uniq_S4C_select_GRanges_
  stat smooth(method="loess", color="black", fill="grey", span=0.6, level=0.95, lwd=2) +
  theme classic2(base size = 15) +
  coord_cartesian(xlim=c(0,100), ylim=c(0,5)) +
  geom_vline(xintercept = c(27), color="black", lty=2, lwd=1) +
  geom_text(aes(x=16,y=5), label = 27, size=7.5, col="black") +
  xlab("distance between motifs (nt)") +
  ylab("log2(FC)")
#pdf("./Figure/Figure4/FigureS4C.pdf", width = 6, height = 6)
PTBP1_uniq_S4C_select_GRanges_overlaps_YsUs_plot
## Warning in geom_text(aes(x = 16, y = 5), label = 27, size = 7.5, col = "black"): All aesthetics have
## i Please consider using `annotate()` or provide this layer with data containing
##
   a single row.
## `geom_smooth()` using formula = 'y ~ x'
```



0.5.8 Figure S4D

```
PTBP1_gene <- unique(PTBP1[,"gene_name"])

HUR_gene <- unique(HUR[,"gene_name"])

coRAP_gene <- unique(HUR_PTBP1[,"gene_name"])

HUR_PTBP1_coRAP_gene <- list(HUR_gene,PTBP1_gene,coRAP_gene)

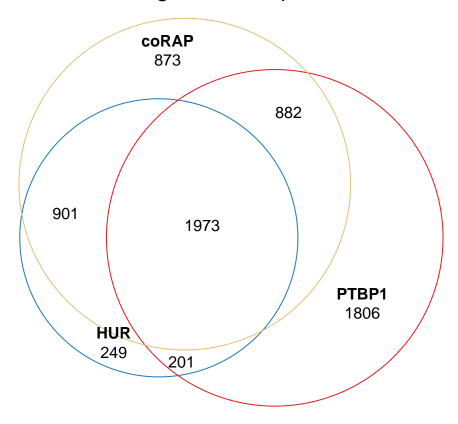
names(HUR_PTBP1_coRAP_gene) <- c("HUR","PTBP1","coRAP")

HUR_PTBP1_coRAP_gene_overlap <- euler(HUR_PTBP1_coRAP_gene, shape="circle")

#pdf("./Figure/Figure4/Figure84D.pdf", width = 6, height = 6)

plot(HUR_PTBP1_coRAP_gene_overlap, fills=c("white","white","white"), quantities=TRUE, edges=T, col=c("#
```

gene overlaps

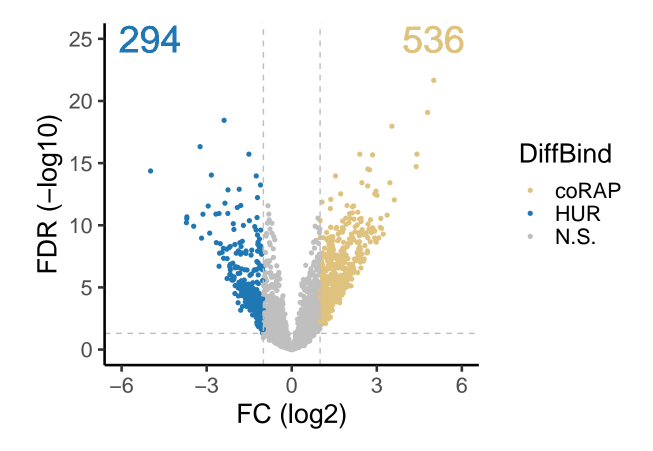


#dev.off()

###Figure S4E left

```
# load data
HUR_coRAP_gene <- as.data.frame(intersect(HUR_gene, coRAP_gene))</pre>
colnames(HUR_coRAP_gene) <- "gene_name"</pre>
HUR_coRAP_data <- merge(HUR, HUR_coRAP_gene, by="gene_name")</pre>
HUR_coRAP_data <- HUR_coRAP_data[,c("Rep1","Rep2","gene_name")]</pre>
HUR_coRAP_data_melt <- melt(HUR_coRAP_data) %>% group_by(gene_name,variable) %>% summarise(Gene_Counts =
## Using gene_name as id variables
## `summarise()` has grouped output by 'gene_name'. You can override using the
## `.groups` argument.
HUR_coRAP_data_melt <- as.data.frame(HUR_coRAP_data_melt)</pre>
HUR_coRAP_data <- reshape2::dcast(HUR_coRAP_data_melt, gene_name~variable)</pre>
## Using Gene_Counts as value column: use value.var to override.
colnames(HUR_coRAP_data) <- c("gene_name","HUR_1","HUR_2")</pre>
HURPTBP1_coRAP_data <- merge(HUR_PTBP1,HUR_coRAP_gene,by="gene_name")
HURPTBP1 coRAP data <- HURPTBP1 coRAP data[,c("Rep1","Rep2","gene name")]</pre>
HURPTBP1_coRAP_data_melt <- melt(HURPTBP1_coRAP_data) %>% group_by(gene_name,variable) %>% summarise(Ge
## Using gene_name as id variables
## `summarise()` has grouped output by 'gene_name'. You can override using the `.groups` argument.
HURPTBP1_coRAP_data_melt <- as.data.frame(HURPTBP1_coRAP_data_melt)</pre>
```

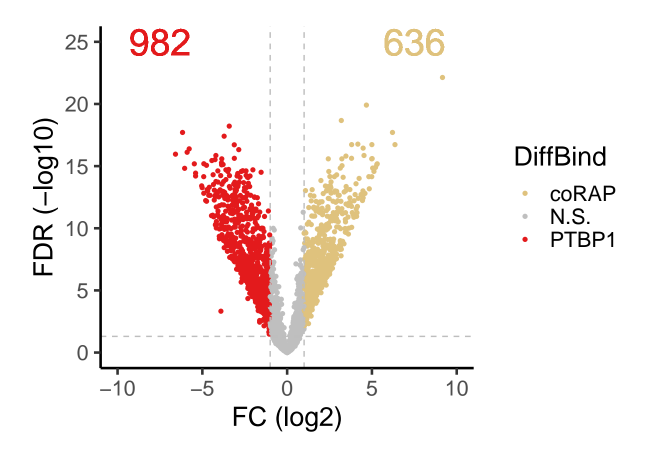
```
HURPTBP1_coRAP_data <- reshape2::dcast(HURPTBP1_coRAP_data_melt, gene_name~variable)</pre>
## Using Gene_Counts as value column: use value.var to override.
colnames(HURPTBP1_coRAP_data) <- c("gene_name","coRAP_1","coRAP_2")</pre>
HUR_coRAP_commongenes <- merge(HUR_coRAP_data, HURPTBP1_coRAP_data, by="gene_name")
# DEG analysis
x <- HUR coRAP commongenes[,2:5]
group \leftarrow c(1,1,2,2)
y <- DGEList(counts=x, group = group)</pre>
design <- model.matrix(~group)</pre>
y <- estimateDisp(y,design)
fit <- glmQLFit(y, design)</pre>
qlf <- glmQLFTest(fit)</pre>
DEG_data_table_S4EL <-qlf$table</pre>
DEG_data_table_S4EL$p.adjust <- -log10(p.adjust(DEG_data_table_S4EL$PValue, method = "BH"))
DEG_data_table_S4EL$PValue <- -log10(DEG_data_table_S4EL$PValue)
DEG_data_table_S4EL$HUR_won <- DEG_data_table_S4EL$logFC <= -1 & DEG_data_table_S4EL$p.adjust >= 1.3010
DEG_data_table_S4EL$coRAP_won <- DEG_data_table_S4EL$logFC >= 1 & DEG_data_table_S4EL$p.adjust >= 1.301
DEG_data_table_S4EL$DiffBind <- paste(DEG_data_table_S4EL$HUR_won,DEG_data_table_S4EL$coRAP_won,sep = "
DEG_data_table_S4EL$DiffBind <- gsub("FALSE_FALSE","N.S.",DEG_data_table_S4EL$DiffBind)
DEG_data_table_S4EL$DiffBind <- gsub("FALSE_TRUE", "coRAP", DEG_data_table_S4EL$DiffBind)
DEG_data_table_S4EL$DiffBind <- gsub("TRUE_FALSE", "HUR", DEG_data_table_S4EL$DiffBind)
DEG_data_table_S4EL_plot <- ggplot(data=DEG_data_table_S4EL) +</pre>
  geom_point(aes(x=logFC,y=p.adjust,color=DiffBind), pch=16) +
  geom_hline(yintercept = -log10(0.05), linetype = "dashed", color = "grey") +
  geom_vline(xintercept = c(-1,1), linetype = "dashed", color = "grey") +
  scale_color_manual(values = c("#dfc27d","#1f78b4","grey75")) +
  theme_classic(base_size = 20) +
  ylab("FDR (-log10)") +
  xlab("FC (log2)") +
  ylim(0,25) +
 xlim(-6,6)
DEG_data_table_S4EL_plot <- DEG_data_table_S4EL_plot + geom_text(aes(x=5,y=25), label = table(DEG_data_
  geom_text(aes(x=-5,y=25), label = table(DEG_data_table_S4EL$DiffBind)["HUR"], size=10, color="#1f78b4
#pdf("./Figure/Figure4/FigureS4E_left.pdf", width = 6, height = 6)
DEG_data_table_S4EL_plot
## Warning in geom_text(aes(x = 5, y = 25), label = table(DEG_data_table_S4EL$DiffBind)["coRAP"], : All
## i Please consider using `annotate()` or provide this layer with data containing
## Warning in geom_text(aes(x = -5, y = 25), label = table(DEG_data_table_S4EL$DiffBind)["HUR"], : All
## i Please consider using `annotate()` or provide this layer with data containing
## a single row.
```



0.5.9 Figure S4E right

```
PTBP1 coRAP gene <- as.data.frame(intersect(PTBP1 gene, coRAP gene))
colnames(PTBP1_coRAP_gene) <- "gene_name"</pre>
PTBP1_coRAP_data <- merge(PTBP1,PTBP1_coRAP_gene,by="gene_name")
PTBP1_coRAP_data <- PTBP1_coRAP_data[,c("Rep1","Rep2","gene_name")]</pre>
PTBP1_coRAP_data_melt <- melt(PTBP1_coRAP_data) %>% group_by(gene_name,variable) %>% summarise(Gene_Cou
## Using gene_name as id variables
## `summarise()` has grouped output by 'gene_name'. You can override using the
## `.groups` argument.
PTBP1_coRAP_data_melt <- as.data.frame(PTBP1_coRAP_data_melt)</pre>
PTBP1_coRAP_data <- reshape2::dcast(PTBP1_coRAP_data_melt, gene_name~variable)
## Using Gene_Counts as value column: use value.var to override.
colnames(PTBP1_coRAP_data) <- c("gene_name", "PTBP1_1", "PTBP1_2")</pre>
HURPTBP1_coRAP_data <- merge(HUR_PTBP1,PTBP1_coRAP_gene,by="gene_name")</pre>
HURPTBP1_coRAP_data <- HURPTBP1_coRAP_data[,c("Rep1","Rep2","gene_name")]</pre>
HURPTBP1_coRAP_data_melt <- melt(HURPTBP1_coRAP_data) %>% group_by(gene_name,variable) %>% summarise(Gene_name)
## Using gene_name as id variables
## `summarise()` has grouped output by 'gene_name'. You can override using the `.groups` argument.
HURPTBP1_coRAP_data_melt <- as.data.frame(HURPTBP1_coRAP_data_melt)</pre>
```

```
HURPTBP1_coRAP_data <- reshape2::dcast(HURPTBP1_coRAP_data_melt, gene_name~variable)</pre>
## Using Gene_Counts as value column: use value.var to override.
colnames(HURPTBP1_coRAP_data) <- c("gene_name","coRAP_1","coRAP_2")</pre>
HUR_coRAP_commongenes <- merge(PTBP1_coRAP_data,HURPTBP1_coRAP_data,by="gene_name")</pre>
# DEG analysis
x <- HUR coRAP commongenes[,2:5]
group \leftarrow c(1,1,2,2)
y <- DGEList(counts=x, group = group)</pre>
design <- model.matrix(~group)</pre>
y <- estimateDisp(y,design)
fit <- glmQLFit(y, design)</pre>
qlf <- glmQLFTest(fit)</pre>
DEG_data_table_S4ER <-qlf$table</pre>
DEG_data_table_S4ER$p.adjust <- -log10(p.adjust(DEG_data_table_S4ER$PValue, method = "BH"))
DEG_data_table_S4ER$PValue <- -log10(DEG_data_table_S4ER$PValue)
DEG_data_table_S4ER$HUR_won <- DEG_data_table_S4ER$logFC <= -1 & DEG_data_table_S4ER$p.adjust >= 1.3010
DEG_data_table_S4ER$coRAP_won <- DEG_data_table_S4ER$logFC >= 1 & DEG_data_table_S4ER$p.adjust >= 1.301
DEG_data_table_S4ER$DiffBind <- paste(DEG_data_table_S4ER$HUR_won,DEG_data_table_S4ER$coRAP_won,sep = "
DEG_data_table_S4ER$DiffBind <- gsub("FALSE_FALSE","N.S.",DEG_data_table_S4ER$DiffBind)
DEG_data_table_S4ER$DiffBind <- gsub("FALSE_TRUE", "coRAP", DEG_data_table_S4ER$DiffBind)
DEG_data_table_S4ER$DiffBind <- gsub("TRUE_FALSE", "PTBP1", DEG_data_table_S4ER$DiffBind)
DEG_data_table_S4ER_plot <- ggplot(data=DEG_data_table_S4ER) +</pre>
  geom_point(aes(x=logFC,y=p.adjust,color=DiffBind), pch=16) +
  geom_hline(yintercept = -log10(0.05), linetype = "dashed", color = "grey") +
  geom_vline(xintercept = c(-1,1), linetype = "dashed", color = "grey") +
  scale_color_manual(values = c("#dfc27d", "grey75", "#e31a1c")) +
  theme_classic(base_size = 20) +
  ylab("FDR (-log10)") +
  xlab("FC (log2)") +
 ylim(0,25) +
  xlim(-10,10)
DEG_data_table_S4ER_plot <- DEG_data_table_S4ER_plot + geom_text(aes(x=7.5,y=25), label = table(DEG_dat
  geom_text(aes(x=-7.5,y=25), label = table(DEG_data_table_S4ER$DiffBind)["PTBP1"], size=10, color="#e3
#pdf("./Figure/Figure4/Figure54E_right.pdf", width = 6, height = 6)
DEG data table S4ER plot
## Warning in geom_text(aes(x = 7.5, y = 25), label = table(DEG_data_table_S4ER$DiffBind)["coRAP"], : A
## i Please consider using `annotate()` or provide this layer with data containing
## a single row.
## Warning in geom_text(aes(x = -7.5, y = 25), label = table(DEG_data_table_S4ER$DiffBind)["PTBP1"], :
## i Please consider using `annotate()` or provide this layer with data containing
## a single row.
```



0.5.10 Figure S4F

```
PTBP1_gene <- unique(PTBP1[,"gene_name"])

HUR_gene <- unique(HUR[,"gene_name"])

coRAP_gene <- unique(HUR_PTBP1[,"gene_name"])

coRAP_gene <- unique(HUR_PTBP1[,"gene_name"])

coRAP_only <- setdiff(coRAP_gene, HUR_gene)

coRAP_only <- setdiff(coRAP_only, PTBP1_gene)

HUR_DB <- DEG_data_table_S4EL

coRAP_HU_diffb <- HUR_coRAP_commongenes[HUR_DB$DiffBind == "coRAP", "gene_name"]

PTBP1_DB <- DEG_data_table_S4ER

coRAP_PT_diffb <- HUR_coRAP_commongenes[PTBP1_DB$DiffBind == "coRAP", "gene_name"]

gene_uniq <- unique(c(coRAP_only,coRAP_HU_diffb,coRAP_PT_diffb))

entrez_IDs <- na.omit(as.data.frame(unlist(mapIds(org.Hs.eg.db, gene_uniq, 'ENTREZID', 'SYMBOL')))[,1])

## 'select()' returned 1:1 mapping between keys and columns

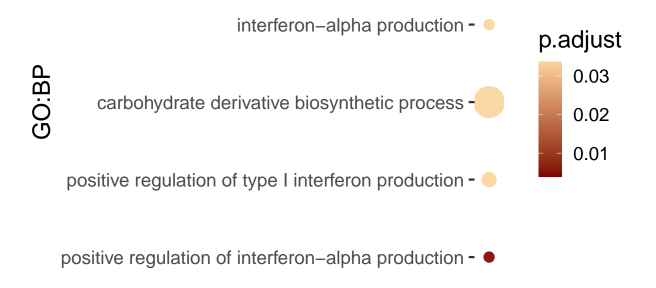
all <- unique(c(HUR_PTBP1$gene_name, HUR$gene_name, PTBP1$gene_name))

entrez_IDs_all <- na.omit(as.data.frame(unlist(mapIds(org.Hs.eg.db, all, 'ENTREZID', 'SYMBOL')))[,1])

## 'select()' returned 1:1 mapping between keys and columns
```

```
BPs <- enrichGO(
 gene = entrez_IDs,
 universe = entrez_IDs_all,
             = "ENTREZID",
 keyType
              = org.Hs.eg.db,
 OrgDb
              = "BP",
  ont
 pAdjustMethod = "BH",
 pvalueCutoff = 0.05,
 qvalueCutoff = 0.05,
 minGSSize
              = 10,
              = TRUE
 readable
)
BP <- as.data.frame(BPs)</pre>
BP$Description <- factor(BP$Description, levels = BP$Description)
CCs <- enrichGO(
 gene = entrez_IDs,
 universe = entrez_IDs_all,
              = "ENTREZID",
 keyType
              = org.Hs.eg.db,
 OrgDb
 ont
              = "CC",
 pAdjustMethod = "BH",
 pvalueCutoff = 0.05,
 qvalueCutoff = 0.05,
 minGSSize = 10,
 readable
              = TRUE
)
CC <- as.data.frame(CCs)</pre>
CC$Description <- factor(CC$Description, levels = CC$Description)</pre>
GOs <- rbind(BP,CC)</pre>
GO_plot <- ggplot2::ggplot(data=GOs, aes(x=Description, y=1)) +</pre>
  geom_point(aes(color=p.adjust), size=sqrt(GOs$Count), alpha=0.9) +
  coord flip() +
 theme classic(base size = 17.5) +
 scale_color_gradient(low="#7f0000",high="#fdd49e") +
 ylab(NULL) +
  # CC is NA
 xlab("GO:BP") +
 theme(axis.text.x = element_blank(),
       axis.line = element_blank(),
       axis.ticks.x = element_blank(),
  )
# write.csv(GOs, "./Data/GOs.csv")
#pdf("./Figure/Figure4/FigureS4F.pdf", width = 9, height = 8)
GO_plot
```

regulation of interferon-alpha production - •



```
#dev.off()
```

0.5.11 Figure S4G

```
gene_name <- unique(unlist(str_split(GOs$geneID,"\\/")))
genes <- as.data.frame(gene_name)

features <- HUR_PTBP1[,c("gene_name","feature","BS","Mean_FCI","Mean_FCH")]
features <- merge(genes,features,by="gene_name")
features$feature <- factor(features$feature, levels = c("intron","3UTR", "5UTR", "CDS"))
#pdf("./Figure/Figure4/FigureS4G.pdf", width = 8, height = 5)
par(bty="n",mfrow=c(1,2))
pie1(table(features$feature), percentage=T, col=c("#08306b", "#deebf7", "#9ecae1", "#4292c6"))
boxplot2(data=features,log2(Mean_FCI)~feature, outline=F, col=c("#08306b", "#deebf7", "#9ecae1", "#4292</pre>
```

