Figure 7 YBX1

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2025-05-13

This markdown show how to generate Figure S5

Loading packages and required data: YBX1 metafile with binding sites of RAPseq Input and T7 input, generated in the script "YBX1_metafile_annotation

```
library(eulerr)
## Warning: package 'eulerr' was built under R version 4.3.3
library(ggplot2)
## Warning: package 'ggplot2' was built under R version 4.3.3
library(reshape2)
library(gplots)
## Warning: package 'gplots' was built under R version 4.3.3
## Registered S3 method overwritten by 'gplots':
     method
##
               from
##
     plot.venn eulerr
##
## Attaching package: 'gplots'
## The following object is masked from 'package:eulerr':
##
##
       venn
## The following object is masked from 'package:stats':
##
##
       lowess
library(tidyverse)
## Warning: package 'purrr' was built under R version 4.3.3
## Warning: package 'lubridate' was built under R version 4.3.3
## — Attaching core tidyverse packages —
                                                                 - tidyverse
2.0.0 --
## √ dplyr
               1.1.4
                         ✓ readr
                                      2.1.5
## √ forcats
               1.0.0

√ stringr

                                      1.5.1
## ✓ lubridate 1.9.4
                          √ tibble
                                      3.2.1
## √ purrr
               1.0.4
                         √ tidyr
                                      1.3.1
```

```
## — Conflicts ——
tidyverse_conflicts() ---
## X dplyr::filter() masks stats::filter()
## X dplyr::lag()
                    masks stats::lag()
## i Use the conflicted package (<http://conflicted.r-lib.org/>) to force all
conflicts to become errors
library(ggpubr)
library(dendextend)
## Warning: package 'dendextend' was built under R version 4.3.3
##
## -----
## Welcome to dendextend version 1.19.0
## 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:ggpubr':
##
##
      rotate
## The following object is masked from 'package:stats':
##
##
      cutree
library(dplyr)
library(gtools)
library(corrplot)
## Warning: package 'corrplot' was built under R version 4.3.3
## corrplot 0.95 loaded
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. clusterProfiler 4.0: A universal enrichment tool
for interpreting omics data. The Innovation. 2021, 2(3):100141
## Attaching package: 'clusterProfiler'
## The following object is masked from 'package:purrr':
##
##
       simplify
##
## The following object is masked from 'package:stats':
##
       filter
library(org.Hs.eg.db)
## Loading required package: AnnotationDbi
## Loading required package: stats4
## Loading required package: BiocGenerics
## Attaching package: 'BiocGenerics'
##
## The following objects are masked from 'package:lubridate':
##
##
       intersect, setdiff, union
##
## The following objects are masked from 'package:dplyr':
##
##
       combine, intersect, setdiff, union
##
## 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: Biobase
## Welcome to Bioconductor
```

```
##
##
       Vignettes contain introductory material; view with
       'browseVignettes()'. To cite Bioconductor, see
##
       'citation("Biobase")', and for packages 'citation("pkgname")'.
##
##
## Loading required package: IRanges
## Loading required package: S4Vectors
## Attaching package: 'S4Vectors'
##
## The following object is masked from 'package:clusterProfiler':
##
##
       rename
##
## The following objects are masked from 'package:lubridate':
##
##
       second, second<-
##
## The following objects are masked from 'package:dplyr':
##
##
       first, rename
##
## The following object is masked from 'package:tidyr':
##
##
       expand
## The following object is masked from 'package:gplots':
##
##
       space
##
## The following object is masked from 'package:utils':
##
##
       findMatches
##
## The following objects are masked from 'package:base':
##
       expand.grid, I, unname
##
##
##
## Attaching package: 'IRanges'
##
## The following object is masked from 'package:clusterProfiler':
##
       slice
##
##
## The following object is masked from 'package:lubridate':
##
##
       %within%
##
## The following objects are masked from 'package:dplyr':
```

```
##
       collapse, desc, slice
##
##
## The following object is masked from 'package:purrr':
##
##
       reduce
##
##
## Attaching package: 'AnnotationDbi'
##
## The following object is masked from 'package:clusterProfiler':
##
##
       select
##
## The following object is masked from 'package:dplyr':
##
##
       select
library(rrvgo)
## Warning: package 'rrvgo' was built under R version 4.3.3
YBX1<-
read.table(file="/Users/riccardomosca/Desktop/RAPseq PAPER/PEAKs/ANNOTATED/T7
_Fig5/Ybx1_T7_scored_annotated.txt", header=T, stringsAsFactors = F)
head(YBX1)
##
      chr
            start
                      end
                                          peak_ID
                                                         RBPs strand
## 1 chr1 1043290 1043298 chr1_1043277_1043341_+ Ybx1_final
## 2 chr1 1046002 1046010 chr1 1045989 1046039 + Ybx1 final
                                                                   +
## 3 chr1 1046657 1046665 chr1_1046652_1046673_+ Ybx1_final
## 4 chr1 1049916 1049924 chr1 1049902 1049932 + Ybx1 final
## 5 chr1 1217314 1217322 chr1 1217305 1217329 - Ybx1 final
## 6 chr1 1228604 1228612 chr1_1228566_1228658_- Ybx1_final
     Ybx1_rep1_signal Ybx1_rep2_signal halo_signal input_signal
T7Ybx1 rep1 signal
## 1
                    34
                                     41
                                                 16
                                                               17
4
                   43
## 2
                                     24
                                                 12
                                                               10
10
## 3
                   27
                                     26
                                                 17
                                                               18
11
## 4
                   23
                                     29
                                                 10
                                                                7
6
## 5
                   21
                                     21
                                                   4
                                                               11
5
## 6
                   27
                                     30
                                                 12
                                                               14
2
##
     T7Ybx1_rep2_signal T7halo_signal T7input_signal
## 1
                      4
                                     9
                                             10.00000
## 2
                       3
                                     4
                                              3.00000
```

##	3	29	8	12.66667
##	4	6	6	9.00000
##	5	2	2	0.50000
##	6	3	3	13.50000
##				

pos_fa

1

GAGGGGGGGCTTGTGGGACCACTGAGCCCCTGTGTCCTTCCCAGACCAGGCCCCGTCCCCATGCCTCGGGGTGCAGT GTGCATTTGGGGCGACGTGTGCTGAAGAACGGGCAGCAGCGTGTGAATGCCTGCAGGCGTGCTCGAGCCTCTAC GATCCTGTGTGCGGCAGCGACGGCGTCACATACGGCAGCGCGTGCG

2

TCATGGGGTGGGGTCACCCGAGCCACAGAGGTTTCCCATGCCCGTGCCCCAGACGCTTCTGCGCCTGCGACC TGTGCGGAGATGCGCTGTGAGTTCGGTGCGCGGTGCGTGGAGGAGTCTGGCTCAGCCCACTGTGTCTGCCCGATGCT CACCTGTCCAGAGGCCAACGCTACCAAGGTGAGGGGTGTGGGATGT

3

CCACTGCCAGCGTCCCCAGGACCACCGTGTGGCCCGTGCTGACGGTGCCCCCCACGGCACCCTCCCCTGCACCCAGC CTGGTGGCGTCCGCCTTTGGTGAATCTGGCAGCACTGATGGAAGCAGCGATGAGGAACTGAGCGGGGACCAGGAGGC CAGTGGGGGTGGCTCTGGGGGTGAGCAGGGATCAAGGACTTGGGGT

4

GAGTGGGACCCCGGGGCCTGTGGGCGGTACCCAACCGACGCCTCCTGGGACCTCGGTCCCGGTCCCGTCTTCCTCCA
TCCAGGACCAACCTGTGCCGATGAGAAGAGCCCCTGCCAGCCCAACCCCTGCCATGGGGCGCGCCCTGCCGTGTGC
TGCCCGAGGGTGGTGCTCAGTGCGAGTGCCCCCTGGGGCGTGAGGG

5

GAAGCGAGATTTCGTTCTCAGAAGTGAGATGCCACGATTTCGAGGGACCAGGGAGGAGGCGGCGCCGCGGGGCCACA GCCCAGCCCCGCCCCGACCGCGTCACAGCCAAAGCCCCGCCGGGGTGCGCGCTGCAGAGGGGACACGCCGCTCAT CAACTGGGGCAGCCGCGGTCGGTCGGCCGGTGGCGGCGGCAGGGA

6

CTCCGCGTCCTCATCAAAGCCACCCAGGTCCTTGCCTAGGAAGACCTCCTGGTGGAAGCCGCGATTGAGGTGCCCGT CCATCTCCAGCTTCACCCCGTTCAGGTGGTCTGGGGGCAGGATCTCATTCTCCCCTGTTGGCTACTCTCTCGA GTGGACGAGTGGTTGGCAGGCCGTGCAGACGCGTCCATCAGAAGGA

##

neg_fa

1

2

GCTCTGATGGGAAGACGCCCTCGCTGGACGCAGAGGGCTCCAACTGCCCCGGTGAGTGGACGGCTGGGCGAGGGGAG TGTGAGGATAGCCTGGGCTCGGCCGAGGTGCTGCCCCCTCGCCTGGCAGCAGGTCAGTGCCGGGGGTTATGGTCTT GGGACTCGGCCCCCTCAAACATGTGCGTGCCGGGGACCCCACGCCT

3

CCCTGCCACAGCTGGACGACCTCTTCCGGAATTCAGACGTCAAGAAGGATTTTCGGAGTGTCCGCTTGCGGGACCTG GGGCCCGGCAAATCCGTCCGCGCCATTGTGGATGTGCACTTTGACCCCAGTGAGACCTGCACCCTGGACCCTTCCTG GGAGGCAATGGGTGGGGGATGCCTGGGGCTCTGCCATGCTCAGAGC

4

5

AAGGGGCTGAGGATGGAGCCCGGCCAGGCTCGCCTGTCTCTGATCCGTCTGAACCTAAACGCCAACAACGGCCATGC

AGCCTGGTGGACACCGCTGAGCAAACGCCCCAGTGACCAGCCCAGATGGAGTCTCAGGCCAGACACACCAGCACACCTT CCTGCCTCAGGCCCAGATCCACCAGCCCCTCCCGCCACAGCACCTC ## 6 CAGGGGCTAACCTGTCCCCATCGCCCAACACCTGCAGCACAGCTTTCCTGTGGGGCCCGGCTCTGTCCCTCCTGGCA TCATGCCACTGACCCATGGGGTTAGGGTCACTCTCCAGACCCTGAC FCH rep1 FCH rep2 FCI rep1 FCI rep2 FCH T7rep1 FCH T7rep2 FCI T7rep1 ## 1 2.125000 2.562500 2.000000 2.411765 0.4444444 0.4444444 0.4000000 ## 2 3.583333 2.000000 4.300000 2.400000 2.5000000 0.7500000 3.3333333 ## 3 1.588235 1.529412 1.500000 1.444444 1.3750000 3.6250000 0.8684211 ## 4 2.300000 2.900000 3.285714 4.142857 1.0000000 1.0000000 0.6666667 ## 5 5.250000 5.250000 1.909091 1.909091 2.5000000 1.0000000 10.0000000 ## 6 2.250000 2.500000 1.928571 2.142857 0.6666667 1.0000000 0.1481481 FCI_T7rep2 Mean_FCs_rep1 Mean_FCs_rep2 Mean_FCs_T7rep1 Mean_FCs_T7rep2 ## 1 0.400000 2.062500 2.487132 0.4222222 0.4222222 ## 2 1.0000000 3.941667 2.200000 2.9166667 0.8750000 ## 3 2.2894737 1.544118 1.486928 1.1217105 2.9572368 ## 4 0.6666667 2.792857 3.521429 0.8333333 0.8333333 ## 5 4.0000000 3.579545 6.2500000 3.579545 2.5000000 ## 6 0.2222222 2.089286 2.321429 0.4074074 0.6111111 ## BS rep2 BS T7rep1 BS T7rep2 BS T7 Mean FCH BS rep1 BS ## 1 8.600470 11.008706 0.6692064 0.6692064 9.804588 0.6692064 2.343750 7.5394740 1.1566871 12.923178 4.3480805 2.791667 ## 2 17.705388 8.140967 ## 3 5.957177 5.661263 3.0291117 11.6934950 5.809220 7.3613033 1.558824 ## 4 10.176770 13.924420 1.6666667 1.6666667 12.050595 1.6666667 2.600000 ## 5 12.612750 12.612750 11.2959683 2.5000000 12.612750 6.8979841 5.250000 ## 6 8.060425 9.285714 0.4074074 0.8078449 8.673069 0.6076262 2.375000 MeanFC MeanFC T7 ## Mean FCI Mean FCH T7 Mean FCI T7 gene_ID 0.4000000 2.274816 0.4222222 ENSG00000188157.15 ## 1 2.205882 0.444444 ## 2 3.350000 1.6250000 2.1666667 3.070833 1.8958333 ENSG00000188157.15 ## 3 1.472222 2.5000000 1.5789474 1.515523 2.0394737 ENSG00000188157.15 ## 4 3.714286 0.6666667 3.157143 0.8333333 ENSG00000188157.15 1.0000000 ## 5 1.909091 1.7500000 7.0000000 3.579545 4.3750000 ENSG00000078808.19 0.1851852 2.205357 0.5092593 ENSG00000078808.19 ## 6 2.035714 0.8333333 feature gene_name ## gene type ## 1 CDS AGRN protein coding ## 2 CDS AGRN protein_coding ## 3 CDS AGRN protein_coding ## 4 CDS AGRN protein_coding ## 5 3UTR SDF4 protein coding ## 6 CDS SDF4 protein_coding

Figure S7A

```
RAP_enrichments <- YBX1$peak_ID
T7RAP_enrichments <- YBX1[YBX1$T7Ybx1_rep1_signal/YBX1$T7input_signal > 1 &
YBX1$T7Ybx1_rep2_signal/YBX1$T7input_signal > 1 &
YBX1$T7Ybx1_rep2_signal/YBX1$T7halo_signal > 1 &
YBX1$T7Ybx1_rep2_signal/YBX1$T7halo_signal > 1 &
YBX1$T7Ybx1_rep2_signal/YBX1$T7halo_signal > 1, "peak_ID"]

AB <- list(RAP_enrichments, T7RAP_enrichments)
names(AB) <- c("with modif", "without modif")
FigS7A <- euler(AB, shape = "circle")

plot(FigS7A, fills = c("#93ADD0", "#8CBDA0"), quantities = TRUE, edges = F)</pre>
```

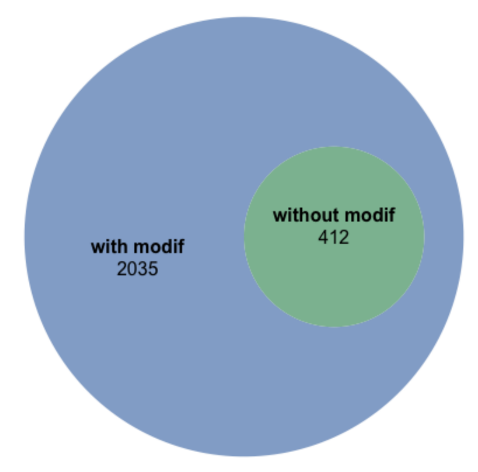


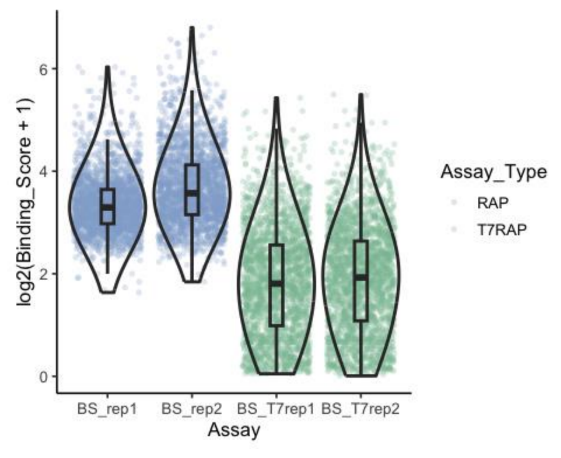
Figure S7B

```
FCs <- YBX1[, c("BS_rep1", "BS_rep2", "BS_T7rep1", "BS_T7rep2")]
FCs <- reshape2::melt(FCs, value.name = "Binding_Score", variable.name =</pre>
```

```
"Assay")
RAP <- rep("RAP", nrow(FCs)/2)
T7RAP <- rep("T7RAP", nrow(FCs)/2)
FCs$Assay_Type <- c(RAP, T7RAP)

plot <- ggplot(data = FCs, aes(x = Assay, y = log2(Binding_Score + 1))) +
geom_jitter(aes(color = Assay_Type),
    pch = 16, alpha = 0.25) + scale_color_manual(values = c("#93ADD0",
"#8ABEA0")) +
    theme_classic(base_size = 12.5)

FigS7B <- plot + geom_violin(trim = T, bw = 0.75, scale = "width", lwd = 1,
fill = NA) +
    geom_boxplot(outlier.shape = NA, width = 0.15, lwd = 1, fill = NA)</pre>
FigS7B
```



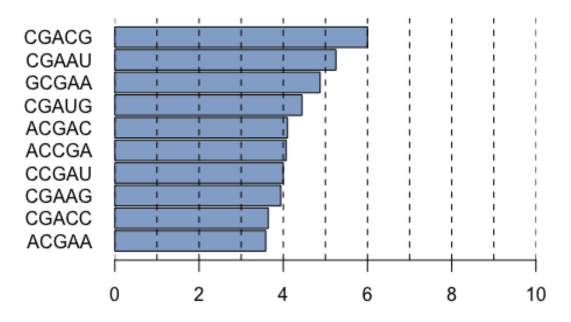
```
list <- list(BS_T71 = YBX1$BS_T7rep1, BS_T72 = YBX1$BS_T7rep2)
# run wilcox.test() for each, comparing to the overlap vector
wt <- lapply(names(list), function(nm) {
    y <- list[[nm]]</pre>
```

```
test <- wilcox.test(YBX1$BS rep1, y)
    data.frame(comparison = paste0("BS1 vs ", nm), W =
as.numeric(test$statistic),
         p.value = test$p.value)
})
# combine into one df
wt df <- do.call(rbind, wt)
Figure S7C
# Looking for the top 10 motifs k=5
pos_fa <- str_sub(YBX1$pos_fa, 75, 125)</pre>
kmer_table <- as.data.frame("YBX1")</pre>
STRINGS <- c((paste(pos_fa, collapse = "NN")))</pre>
kmer table <- cbind(kmer table, STRINGS)</pre>
kmer_table[] <- lapply(kmer_table, as.character)</pre>
bases <- c("A", "C", "T", "G")
kmers <- unite(as.data.frame(permutations(n = 4, r = k, v = bases,</pre>
repeats.allowed = T)),
    col = kmers, sep = "")
kmers <- cbind(kmers, str_count(kmer_table[1, 2], kmers[,</pre>
1])/length(kmer_table[1,
    2]))
rownames(kmers) <- kmers$kmers</pre>
pos_kmers_5 <- as.data.frame(kmers[, -1], row.names = rownames(kmers))</pre>
colnames(pos kmers 5) <- "YBX1"</pre>
neg_fa <- str_sub(YBX1$neg_fa, 75, 125)</pre>
kmer_table <- as.data.frame("YBX1")</pre>
STRINGS <- c(as.character(paste(neg_fa, collapse = "NN")))</pre>
kmer_table <- cbind(kmer_table, STRINGS)</pre>
kmer_table[] <- lapply(kmer_table, as.character)</pre>
bases <- c("A", "C", "T", "G")
kmers <- unite(as.data.frame(permutations(n = 4, r = k, v = bases,</pre>
repeats.allowed = T)),
    col = kmers, sep = "")
kmers <- cbind(kmers, str_count(kmer_table[1, 2], kmers[,</pre>
1])/length(kmer table[1,
    2]))
```

neg kmers 5 <- as.data.frame(kmers[, -1], row.names = rownames(kmers))</pre>

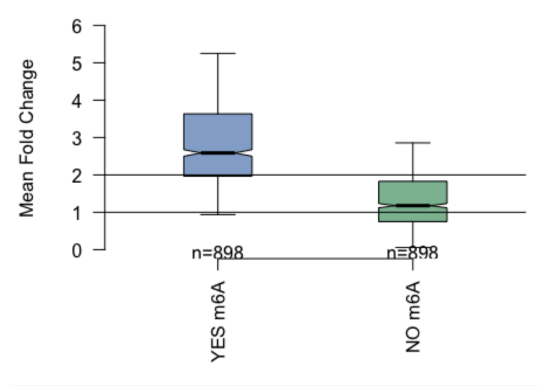
rownames(kmers) <- kmers\$kmers</pre>

```
colnames(neg kmers 5) <- "YBX1neg"</pre>
pos_kmers_5$KMERS <- rownames(pos_kmers_5)</pre>
neg kmers 5$KMERS <- rownames(neg kmers 5)</pre>
K5s_Ybx1 <- merge(pos_kmers_5, neg_kmers_5, by = "KMERS")</pre>
K5s_Ybx1 \leftarrow K5s_Ybx1[K5s_Ybx1$YBX1 >= 1, ]
K5s_Ybx1 \leftarrow K5s_Ybx1[K5s_Ybx1$YBX1neg >= 1, ]
K5s_Ybx1$Fold_Change <- K5s_Ybx1$YBX1/K5s_Ybx1$YBX1neg
# top10
K5s_Ybx1 <- K5s_Ybx1 %>%
    top_n(10, Fold_Change)
K5s_Ybx1 <- K5s_Ybx1[order(K5s_Ybx1$Fold_Change), ]</pre>
K5s_Ybx1$KMERS <- gsub("T", "U", K5s_Ybx1$KMERS)</pre>
barplot(K5s_Ybx1$Fold_Change, names.arg = K5s_Ybx1$KMERS, horiz = T, las = 1,
xlab = "Counts: Bound sites / Control sites",
    col = "#93ADD0", xlim = c(0, 10), space = 0.1)
abline(v = c(0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10), lty = 2, lwd = 1)
```



Counts: Bound sites / Control sites

m6A dependency in top10 binding motif peaks



```
wt <- wilcox.test(AB$`YES m6A`, AB$`NO m6A`, alternative = "greater")
wt <- wt$p.value</pre>
```

Figure S7E

```
# Loading the YBX1_19 metafile and the annotated m5C generated in the
markdown
# 'YBX1 liftover'
m5C annotated <-
read.table("/Users/riccardomosca/Desktop/RAPseq_PAPER/FIGURES/FIGURE5/m5C_ann
otated.txt",
    sep = "\t", header = T)
YBX1 19 <-
read.table("/Users/riccardomosca/Desktop/RAPseq_PAPER/FIGURES/Ybx1_me
tafile_19.txt",
    sep = "\t", header = T)
# Considering only the genes dependent on the modifications
YBX1 19 <- YBX1 19 %>%
    filter(!peak ID %in% T7RAP enrichments)
gene IDs <- YBX1 19 %>%
    mutate(gene_ID = strsplit(gene_ID, "\\.") %>%
        lapply(., function(x) x[1]) %>%
        unlist())
gene_IDs_Ybx1 <- gene_IDs %>%
    distinct(gene_ID, .keep_all = FALSE)
gene IDs m5C <- m5C annotated %>%
    mutate(gene_ID = strsplit(gene_ID, "\\.") %>%
        lapply(., function(x) x[1]) %>%
        unlist())
gene_IDs_m5C <- gene_IDs_m5C %>%
    distinct(gene_ID, .keep_all = FALSE)
YBX1_RAP <- gene_IDs_Ybx1$gene_ID
m5C <- gene IDs m5C$gene ID
AB <- list(YBX1 RAP, m5C)
names(AB) <- c("YBX1_RAP", "m5C")</pre>
v <- euler(AB, shape = "circle")</pre>
plot(v, fills = c("#93ADD0", "#9ECAE1"), quantities = TRUE, edges = F)
```

