RWC23_ELT2_ChIP_Boeck_Time_Resolved_RNA

```
Install libraries
# fill this in
Note: you must load biomaRt before loading tidyverse
Load libraries
library(biomaRt)
library(tidyverse)
## -- Attaching packages -----
## v ggplot2 3.3.0
                      v purrr
                                0.3.4
## v tibble 3.0.1
                                0.8.5
                      v dplyr
## v tidyr
            1.0.3
                      v stringr 1.4.0
## v readr
            1.3.1
                      v forcats 0.5.0
## -- Conflicts ------
## x dplyr::filter() masks stats::filter()
## x dplyr::lag()
                    masks stats::lag()
## x dplyr::select() masks biomaRt::select()
library(ComplexHeatmap)
## Loading required package: grid
## ComplexHeatmap version 2.2.0
## Bioconductor page: http://bioconductor.org/packages/ComplexHeatmap/
## Github page: https://github.com/jokergoo/ComplexHeatmap
## Documentation: http://jokergoo.github.io/ComplexHeatmap-reference
```

Load custom functions

```
source("../RWC23_Functions.R")
```

If you use it in published research, please cite:

genomic data. Bioinformatics 2016.

Note: Ensure BioConductor is version 3.10 or above

Pseudocode: - Bring in Boeck Data - Translate to WBGeneID - Filter for ELT-2 ChIP bound genes, make heatmap - Filter for intestine expressed genes (spencer data), make heatmap, add row annotation for binding cluster

Gu, Z. Complex heatmaps reveal patterns and correlations in multidimensional

Import Time-resolved RNA

```
time_resolved_rna <-
  read.delim(
    "../02 Public Intesine RNA/01 input/9 Boeck et al 2016 time-resolved transcriptome/Unified dcpm per
   quote = "",
    stringsAsFactors = FALSE
  )
paramart <-
  useMart("parasite_mart",
          dataset = "wbps_gene",
          host = "https://parasite.wormbase.org",
          port = 443)
time_resolved_rna <- getBM(</pre>
 mart = paramart,
 filter = c("wormbase_gseqname"),
 value = time_resolved_rna$WormbaseName,
  attributes = c("wormbase_gseq", "wbps_gene_id", "wikigene_name")
) %>% right_join(time_resolved_rna, by = c("wormbase_gseq" = "WormbaseName"))
## Cache found
time_resolved_rna <- time_resolved_rna %>% drop_na(wbps_gene_id)
intestine gene list <-
 read_csv("../02_Public_Intesine_RNA/02_output/RWC23_Public_Intestine_RNA_Data.csv")
## Parsed with column specification:
## cols(
##
     WBGeneID = col_character()
## )
```

Import wTF3.0 worm transcription factor database

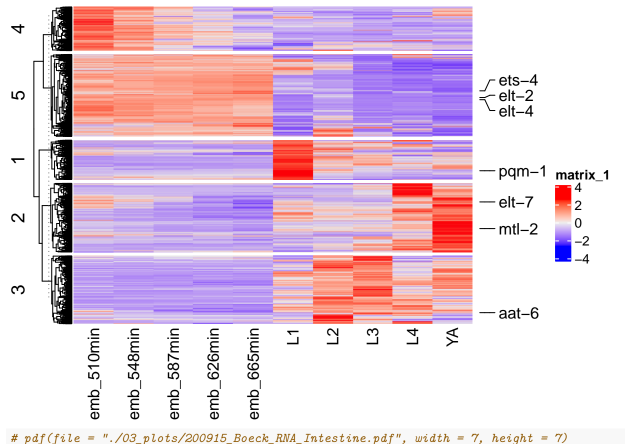
```
wTF3.0 <-
  read.csv(
    "../01_ChIPseq_RNAseq_Integration/01_input/TF3-0_namesonly.txt",
    sep = "\t",
    header = TRUE
) %>% select(WBGeneID)
```

Filter time-resolved RNA-seq based on intestine expression

```
time_resolved_rna_intestine_df <- time_resolved_rna %>%
  remove_rownames() %>%
  arrange(wbps_gene_id) %>%
  filter(wbps_gene_id %in% intestine_gene_list$WBGeneID) %>%
  select(-(emb_4cell:emb_471min),-DE,-D,-DX,-Soma,-Male,-AdultSPE9,-gonad,-LENGTH)
head(time_resolved_rna_intestine_df)
                     wbps_gene_id wikigene_name emb_510min emb_548min emb_587min
     wormbase_gseq
## 1
         T13A10.10 WBGene00000005
                                          aat-4
                                                    0.1841
                                                               0.1632
                                                                          0.1776
## 2
          T11F9.4 WBGene00000007
                                          aat-6
                                                    0.1513
                                                               0.1482
                                                                          0.1586
```

```
## 3
           ZK455.1 WBGene00000040
                                                     2.3243
                                                                 1.9498
                                                                            1.8170
                                           aco-1
## 4
           T25C8.2 WBGene00000067
                                                    15.2874
                                                                16.6729
                                                                           16.8900
                                           act-5
## 5
           F57F5.4 WBGene00000073
                                           add-2
                                                     0.7871
                                                                 0.7277
                                                                            0.6445
## 6
          D2030.10 WBGene00000084
                                                     0.1429
                                                                 0.1878
                                                                            0.1805
                                           aex-1
##
     emb 626min emb 665min
                                   L1
                                               L2
                                                          1.3
                                                                     T.4
                                                                               YΑ
         0.1677
                    0.1630 0.0436931
## 1
                                       0.2184170
                                                   0.265660
                                                              0.3224440
                                                                         0.408817
## 2
         0.1554
                    0.1584 0.1681510
                                        0.2751570
                                                   0.349014
                                                              0.3264440
                                                                         0.271406
## 3
         1.8299
                    1.9978 5.0249900
                                        5.9824800
                                                   8.917410
                                                              2.3600200
                                                                         4.554760
## 4
        16.6729
                   18.0843 29.1548000 49.1039000 71.569300 29.3725000 34.417200
## 5
         0.4962
                    0.3919 0.5606450
                                        0.3947570
                                                   0.335628
                                                              0.1979400
                                                                         0.387552
## 6
         0.1832
                    0.1716 0.1049800
                                        0.0941584
                                                   0.122310
                                                              0.0752852
                                                                         0.155656
time_resolved_rna_intestine_matrix <-</pre>
  time_resolved_rna_intestine_df %>%
  select(-wormbase_gseq, -wikigene_name) %>%
  remove_rownames() %>%
  arrange(wbps_gene_id) %>%
  column_to_rownames(var = "wbps_gene_id") %>%
  as.matrix()
head(time_resolved_rna_intestine_matrix)
##
                  emb_510min emb_548min emb_587min emb_626min emb_665min
                                  0.1632
## WBGene0000005
                      0.1841
                                             0.1776
                                                        0.1677
                                                                    0.1630
## WBGene0000007
                      0.1513
                                  0.1482
                                             0.1586
                                                         0.1554
                                                                    0.1584
## WBGene00000040
                      2.3243
                                  1.9498
                                             1.8170
                                                        1.8299
                                                                    1.9978
## WBGene00000067
                     15.2874
                                 16.6729
                                            16.8900
                                                        16.6729
                                                                   18.0843
## WBGene00000073
                      0.7871
                                  0.7277
                                             0.6445
                                                        0.4962
                                                                    0.3919
## WBGene00000084
                      0.1429
                                  0.1878
                                             0.1805
                                                         0.1832
                                                                    0.1716
##
                          L1
                                      L2
                                                L3
                                                            L4
                                                                      YA
## WBGene0000005
                   0.0436931
                              0.2184170 0.265660
                                                    0.3224440
                                                               0.408817
## WBGene00000007
                   0.1681510
                               0.2751570
                                         0.349014
                                                    0.3264440
                                                                0.271406
## WBGene00000040
                   5.0249900
                              5.9824800 8.917410
                                                    2.3600200
                                                                4.554760
## WBGene00000067 29.1548000 49.1039000 71.569300 29.3725000 34.417200
## WBGene00000073
                   0.5606450
                               0.3947570
                                         0.335628
                                                    0.1979400
                                                               0.387552
## WBGene00000084
                   0.1049800
                              0.0941584
                                         0.122310 0.0752852 0.155656
Perform row normalization
time_resolved_rna_intestine_matrix_scaled <-</pre>
  t(apply(unlist(time_resolved_rna_intestine_matrix), 1, scale))
colnames(time_resolved_rna_intestine_matrix_scaled) <-</pre>
  colnames(time_resolved_rna_intestine_matrix)
Store index of relavent genes for row annotations. Use custom function
gene names <-
  c("elt-2", "elt-7", "elt-4", "pqm-1", "mtl-2", "ets-4", "aat-6")
GOI_df <-
  GOI_annotate_heatmap(gene_names, time_resolved_rna_intestine_df$wikigene_name)
GOI_df
##
      name index
             159
## 1 elt-2
## 2 elt-7
            2115
## 3 elt-4
             161
## 4 pqm-1
             457
## 5 mtl-2
             350
```

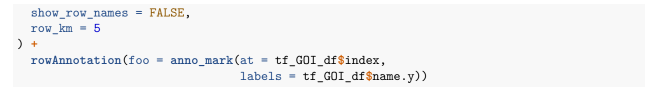
```
## 6 ets-4 2459
## 7 aat-6
time_resolved_rna_intestine_df %>% filter(wikigene_name %in% GOI_df$name)
     wormbase_gseq
                     wbps_gene_id wikigene_name emb_510min emb_548min emb_587min
## 1
           T11F9.4 WBGene00000007
                                           aat-6
                                                     0.1513
                                                                0.1482
                                                                            0.1586
## 2
           C33D3.1 WBGene00001250
                                           elt-2
                                                     1.0771
                                                                0.7690
                                                                            0.7890
## 3
          C39B10.6 WBGene00001252
                                                     0.4558
                                                                            0.5394
                                           elt-4
                                                                0.5370
## 4
          T08G5.10 WBGene00003474
                                                     0.0855
                                                                0.0869
                                                                            0.0960
                                          mtl-2
## 5
           F40F8.7 WBGene00004096
                                           pqm-1
                                                     1.1130
                                                                0.9787
                                                                            0.9506
## 6
           C18G1.2 WBGene00015981
                                           elt-7
                                                     0.3221
                                                                0.3533
                                                                            0.3047
## 7
           F22A3.1 WBGene00017687
                                           ets-4
                                                     1.6876
                                                                1.9241
                                                                            2.0529
##
     emb_626min emb_665min
                                            L2
                                                      L3
                                  L1
                                                                L4
                    0.1584 0.1681510 0.275157 0.3490140 0.3264440
## 1
         0.1554
                                                                    0.2714060
## 2
         0.8991
                    0.8958 0.2715040 0.531793 0.5176460 0.3497100
                                                                    0.3836130
## 3
         0.5519
                    0.5908 0.0731321 0.093782 0.0656069 0.0929745 0.0279745
## 4
         0.1147
                    0.1162 2.6928700 3.669790 6.0893500 6.8993700 11.9476000
## 5
         0.9540
                    1.0173 2.4094400 1.813160 1.0956700 1.0476000 0.8723370
## 6
                    0.0675 0.3254970 0.393028 0.2046660 0.4542380 0.2526850
         0.1257
## 7
         2.0319
                    2.0283 0.4815530 1.165750 1.3026800 0.4953370 0.5412020
Boeck_intestine_RNA <-</pre>
 Heatmap(
   time resolved rna intestine matrix scaled,
    cluster_columns = FALSE,
    show row names = FALSE,
   row_km = 5
  ) +
  rowAnnotation(foo = anno_mark(GOI_df$index, labels = GOI_df$name))
Boeck intestine RNA
```

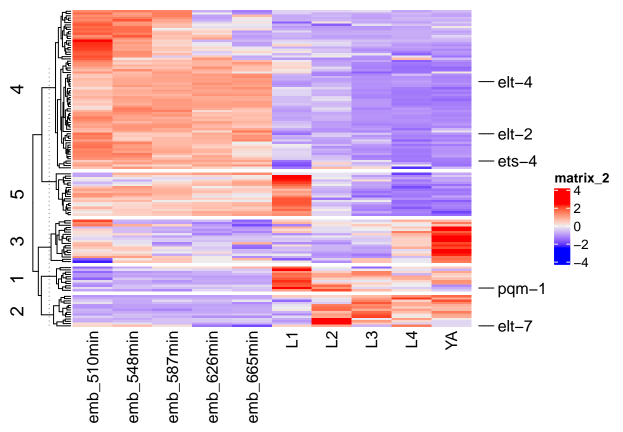


```
# Boeck_intestine_RNA
# dev.off()
```

Filter heatmap for only transcription factors. This is very ugly, fix later.

```
time_resolved_rna_intestine_matrix_scaled_TFONLY <-</pre>
  matrix_select(time_resolved_rna_intestine_matrix_scaled, wTF3.0$WBGeneID)
tf_GOI_df <-
  GOI_df %>% left_join(time_resolved_rna_intestine_df, by = c("name" = "wikigene_name")) %>% select(nam
tf_GOI_df
##
      name wormbase_gseq
                           wbps_gene_id
## 1 elt-2
                 C33D3.1 WBGene00001250
## 2 elt-7
                 C18G1.2 WBGene00015981
## 3 elt-4
                C39B10.6 WBGene00001252
                 F40F8.7 WBGene00004096
## 4 pqm-1
## 5 ets-4
                 F22A3.1 WBGene00017687
tf_GOI_df <-
  GOI_annotate_heatmap(
    tf_GOI_df$wbps_gene_id,
    rownames(time_resolved_rna_intestine_matrix_scaled_TFONLY)
  ) %>% full_join(tf_GOI_df, by = c("name" = "wbps_gene_id"))
Heatmap(
  time_resolved_rna_intestine_matrix_scaled_TFONLY,
  cluster_columns = FALSE,
```





Import ELT-2 ChIP-seq binding data

```
chip_df <-
  read_csv(file = "../01_ChIPseq_RNAseq_Integration/01_input/200719_annotatedPeaks.csv")
## Parsed with column specification:
## cols(
     .default = col_double(),
##
     name = col_character(),
##
     cluster.description = col_character(),
##
##
     peak = col_character(),
     WBGeneID = col_character(),
##
##
     feature_strand = col_character(),
     insideFeature = col_character(),
##
     fromOverlappingOrNearest = col_character()
##
## See spec(...) for full column specifications.
head(chip_df)
```

```
## # A tibble: 6 x 32
     LE_1 LE_2 L1_1 L1_2 L3_1 L3_2 LE_IDR L1_IDR L3_IDR summit_agreement
##
    <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
                                         <dbl>
                                                <dbl>
                                                       <dbl>
     1.93
           1.60 4.25
                       3.77 4.88
                                   5.01
                                                                         27.4
## 1
     2.11
           1.94
                 4.05
                       4.46
                             4.95
                                   5.94
                                             0
                                                                         12.4
                                             0
                                                    0
## 3 1.22 1.53 2.61
                       2.85 2.45 2.86
                                                           1
                                                                        137
          1.42 2.74
    1.81
                       3.28
                            4.18 4.49
                                             0
                                                                          2.5
     2.22 2.17
                 2.24
                       2.13 4.02 4.10
                                             1
                                                           1
                                                                         10
     1.89 2.10 3.43 2.85 3.42 3.53
                                             Λ
                                                                        124.
## # ... with 22 more variables: k4cluster <dbl>, k11cluster <dbl>,
      k4weights <dbl>, k11weights <dbl>, LE_nonNormed <dbl>, L1_nonNormed <dbl>,
      L3_nonNormed <dbl>, LE_std <dbl>, L1_std <dbl>, L3_std <dbl>, name <chr>>,
## #
## #
      cluster.description <chr>, variance <dbl>, peak <chr>, WBGeneID <chr>,
## #
      start_position <dbl>, end_position <dbl>, feature_strand <chr>,
## #
      insideFeature <chr>, distancetoFeature <dbl>, shortestDistance <dbl>,
## #
      fromOverlappingOrNearest <chr>
```

Subset ELT-2 ChIP with literature Intestine Expression

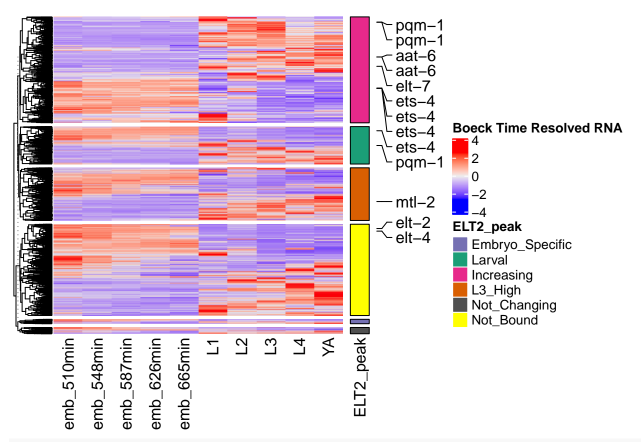
Do this earlier in the code to have k4labels stored in the time_resolved_rna dataframe and subsequent subsetting

```
chip_rna_df <- chip_df %>%
  select(name, cluster.description, WBGeneID) %>%
  right_join(time_resolved_rna_intestine_df,
             by = c("WBGeneID" = "wbps_gene_id")) %>%
  replace_na(list("cluster.description" = "Not_Bound", "name" = "Not_Bound"))
chip_rna_df$cluster.description <-</pre>
  factor(
    chip_rna_df$cluster.description,
    levels = c(
      "Embryo_Specific",
      "Larval",
      "Increasing",
      "L3_High",
      "Not_Changing",
      "Not_Bound"
    )
```

Subset heatmap based on ELT-2 binding pattern

```
## aco-1 -0.5644348 -0.7208340 -0.7762942 -0.7709069 -0.7007882 0.5634316
## aco-1 -0.5644348 -0.7208340 -0.7762942 -0.7709069 -0.7007882 0.5634316
## aco-1 -0.5644348 -0.7208340 -0.7762942 -0.7709069 -0.7007882 0.5634316
##
                 L2
                           L3
                                      L4
## aat-4 0.06919896 0.5393195 1.1043837 1.9638914
## aat-6 0.73548911 1.6569684 1.3753731 0.6886896
## aat-6 0.73548911 1.6569684 1.3753731 0.6886896
## aco-1 0.96330009 2.1889901 -0.5495173 0.3670534
## aco-1 0.96330009 2.1889901 -0.5495173 0.3670534
## aco-1 0.96330009 2.1889901 -0.5495173 0.3670534
for (name in gene_names) {
  index <- which(rownames(chip_rna_matrix_scaled) == name)</pre>
  for (i in 1:length(index)) {
   print(c(name, index[i]))
  }
}
## [1] "elt-2" "241"
## [1] "elt-7" "2877"
## [1] "elt-4" "244"
## [1] "pqm-1" "701"
## [1] "pqm-1" "702"
## [1] "pqm-1" "703"
## [1] "mtl-2" "513"
## [1] "ets-4" "3337"
## [1] "ets-4" "3338"
## [1] "ets-4" "3339"
## [1] "ets-4" "3340"
## [1] "aat-6" "2"
## [1] "aat-6" "3"
chip GOI df <- data.frame(name = NULL, index = NULL)</pre>
for (name in gene_names) {
  index <- which(rownames(chip_rna_matrix_scaled) == name)</pre>
 for (i in 1:length(index)) {
   toappend <-
      data.frame(name = name,
                 index = index[i],
                 stringsAsFactors = FALSE)
    chip_GOI_df <- bind_rows(chip_GOI_df, toappend)</pre>
 }
}
chip_GOI_df
##
       name index
## 1 elt-2
              241
## 2 elt-7 2877
## 3 elt-4
## 4 pqm-1
             701
## 5 pqm-1
              702
## 6
              703
     pqm-1
## 7 mtl-2
            513
## 8 ets-4 3337
## 9 ets-4 3338
```

```
## 10 ets-4 3339
## 11 ets-4 3340
## 12 aat-6 2
## 13 aat-6
BoeckRNA_ELT2_chip <- Heatmap(</pre>
  chip_rna_matrix_scaled,
 name = "Boeck Time Resolved RNA",
 row_split = chip_rna_df$cluster.description,
 row_title = NULL,
 cluster_columns = FALSE
 rowAnnotation(
   ELT2_peak = chip_rna_df$cluster.description,
   col = list(
     ELT2_peak = c(
        "Embryo_Specific" = "#7570B3",
        "Larval" = "#1B9E77",
        "Increasing" = "#E7298A",
        L3_{High} = \#D95F02,
        "Not_Changing" = "#505050",
        "Not_Bound" = "yellow"
     )
    ),
    border = TRUE
  rowAnnotation(foo = anno_mark(at = chip_GOI_df$index,
                                labels = chip_GOI_df$name))
BoeckRNA_ELT2_chip
```



 $\begin{tabular}{ll} \# pdf(file = "./03_plots/200915_Boeck_RNA_ELT2ChIP_Patterns.pdf", width = 7, height = 7) \\ \# Boeck_RNA_ELT2_chip \end{tabular}$

dev.off()