Figure S2

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Aim

The aim of this analysis is to define promoter types in Drosophila kc167 cells based on the presence of specific core promoter elements

Environment setup

Set working directory and load required libraries and lab functions

```
setwd("/n/projects/sga/analysis/SAGA/saga_publication/")
options(knitr.figure_dir = "plots/figure_s2/")
# Standard packages
library(GenomicAlignments)
library(GenomicRanges)
library(Biostrings)
library(BSgenome.Dmelanogaster.UCSC.dm6)
library(TxDb.Dmelanogaster.UCSC.dm6.ensGene)
library(dplyr)
library(reshape2)
library(plyranges)
library(CAGEr)
library(magrittr)
library(ggplot2)
library(cowplot)
library(ggseqlogo)
library(gridExtra)
library(ggpubr)
# Lab sources Lab sources
source("./shared_code/granges_common.r")
source("./shared_code/metapeak_common.r")
source("./shared_code/knitr_common.r")
```

Analysis

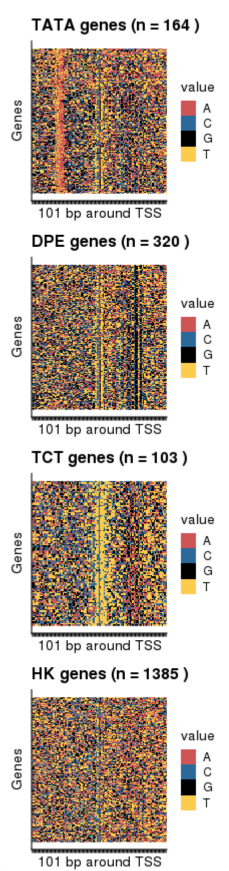
1. Define promoter types

```
# Load TSS
tss <- get(load("./rdata/cage_kc167_tss_pLaw_2tpm.RData"))</pre>
## Define function to find promoter element (motifs) in each active tss
find_motif <- function(motif_name, fb_t_id, mismatch = 0) {</pre>
    motif_info <- subset(promoter_table, name == motif_name)</pre>
    motif <- DNAString(motif_info$motif)</pre>
    up_dis <- motif_info$window_start</pre>
    down_dis <- motif_info$window_end</pre>
    gene_tss <- tss[tss$fb_t_id %in% fb_t_id]</pre>
    if (up_dis >= 0 & down_dis >= 0) {
        tss_r <- resize(gene_tss, down_dis, "start") %>%
             resize(., down_dis - up_dis, "end")
    if (up_dis < 0 & down_dis >= 0) {
        tss_r <- resize(gene_tss, down_dis, "start") %>%
             resize(., abs(up_dis) + down_dis, "end")
    if (up_dis < 0 & down_dis < 0) {</pre>
        tss_r <- resize(gene_tss, abs(up_dis), "end") %>%
             resize(., abs(up_dis) - abs(down_dis), "start")
    promoter_seq <- getSeq(Dmelanogaster, tss_r)</pre>
    names(promoter_seq) <- tss_r$fb_t_id</pre>
    count_df <- vcountPattern(motif, promoter_seq, fixed = FALSE, min.mismatch = 0,</pre>
        max.mismatch = mismatch) %>%
        data.frame(fb_t_id = fb_t_id, count = .)
    count_df$count <- ifelse(count_df$count > 0, T, F)
    colnames(count_df)[2] <- motif_name</pre>
    count_df
}
## Provide promoter element (motif) search information (motif sequence
## composition and search window relative to the TSS)
promoter_table <- read.table("./promoter_elements_sga.txt", header = T)</pre>
motifs <- promoter_table$name</pre>
## Find motifs across TSSs allowing 0 and 1 mismatch
motif_list_1mm <- mclapply(as.character(motifs), function(x) {</pre>
    motif <- find_motif(motif_name = x, tss$fb_t_id, 1)</pre>
    motif
\}, mc.cores = 3)
```

```
motif_list_0mm <- mclapply(as.character(motifs), function(x) {</pre>
    motif <- find_motif(motif_name = x, tss$fb_t_id, 0)</pre>
    motif
\}, mc.cores = 3)
motif_df_1mm <- reshape::merge_recurse(motif_list_1mm)</pre>
motif_df_0mm <- reshape::merge_recurse(motif_list_0mm)</pre>
save(motif df 1mm, file = "./rdata/motif df kc167 1mm.RData")
save(motif_df_0mm, file = "./rdata/motif_df_kc167_0mm.RData")
tss_info <- as.data.frame(tss)[c(1:16)]
motif_info_df_0 <- merge(tss_info, motif_df_0mm)</pre>
motif_info_df_1 <- merge(tss_info, motif_df_1mm)</pre>
## Define promoter groups
tata_tss <- tss[tss$fb_t_id %in% subset(motif_df_1mm, TATA)$fb_t_id]
dpe_tss <- tss[tss$fb_t_id %in% subset(motif_df_1mm, !(TATA) & DPE_0 | PB)$fb_t_id]</pre>
tct tss <- tss[tss$fb t id %in% subset(motif df 0mm, TCT)$fb t id]
hk_tss <- tss[tss$fb_t_id %in% subset(motif_df_0mm, !(TATA | TCT | MTE | DPE | DPE_K |
    DPE_0 | PB | Inr) & (DRE | Motif1 | Motif6 | Motif7))$fb_t_id]
motif_list_kc167 <- list(tata = tata_tss, dpe = dpe_tss, tct = tct_tss, hk = hk_tss)</pre>
save(motif_list_kc167, file = "./rdata/motif_list_kc167.RData")
```

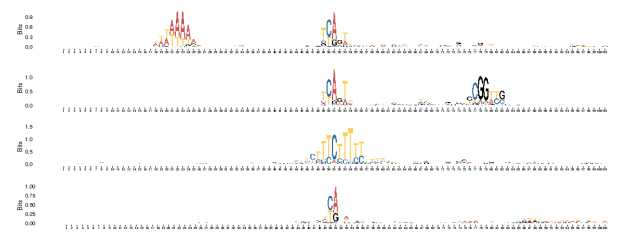
2. Plot a DNA-sequence heatmap of the different promoters types

```
## Define function
get_heatmap <- function(tss, window, direction, name) {</pre>
    seq <- getSeq(Dmelanogaster, resize(tss, window, direction))</pre>
    seq_df <- as.character(seq) %>%
        lapply(., function(x) strsplit(x, "")) %>%
        unlist(., recursive = F) %>%
        do.call(rbind, .) %>%
        as.data.frame()
    seq_df$id <- 1:nrow(seq_df)</pre>
    seq_df_m <- reshape2::melt(seq_df, id.vars = "id")</pre>
    ATGC_plot <- ggplot(seq_df_m, aes(x = variable, y = id, fill = value)) + geom_raster() +
        scale_fill_manual(values = c("indianred3", "#2C699B", "black", "#FDCC4E")) +
        xlab(paste(window, "bp around TSS")) + ylab("Genes") + ggtitle(name) + theme_cowplot() +
        theme(axis.ticks.y = element_blank(), axis.text.y = element_blank(), axis.text.x = element_blank
}
## Generate heatmaps
tata_hm <- get_heatmap(tata_tss, 101, "center", paste("TATA genes", "(n =", length(tata_tss),</pre>
dpe_hm <- get_heatmap(dpe_tss, 101, "center", paste("DPE genes", "(n =", length(dpe_tss),</pre>
```



heatmap-1.png

3. Plot a position weight matrix (PWM) across the promoter types TSSs



Session Info

sessionInfo()

```
## [9] LC ADDRESS=C
                                   LC TELEPHONE=C
## [11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
## attached base packages:
## [1] stats4
                 parallel stats
                                     graphics grDevices utils
                                                                   datasets
## [8] methods
                 base
## other attached packages:
## [1] digest 0.6.27
  [2] pander_0.6.3
## [3] data.table_1.14.0
## [4] lattice_0.20-44
## [5] ggpubr_0.4.0
## [6] gridExtra_2.3
## [7] ggseqlogo_0.1
##
   [8] cowplot_1.1.1
## [9] ggplot2_3.3.3
## [10] magrittr 2.0.1
## [11] CAGEr_1.34.0
## [12] MultiAssayExperiment 1.18.0
## [13] plyranges_1.12.0
## [14] reshape2 1.4.4
## [15] dplyr_1.0.6
## [16] TxDb.Dmelanogaster.UCSC.dm6.ensGene 3.12.0
## [17] GenomicFeatures 1.44.0
## [18] AnnotationDbi 1.54.0
## [19] BSgenome.Dmelanogaster.UCSC.dm6_1.4.1
## [20] BSgenome_1.60.0
## [21] rtracklayer_1.52.0
## [22] GenomicAlignments_1.28.0
## [23] Rsamtools_2.8.0
## [24] Biostrings_2.60.1
## [25] XVector_0.32.0
## [26] SummarizedExperiment_1.22.0
## [27] Biobase 2.52.0
## [28] MatrixGenerics_1.4.0
## [29] matrixStats 0.59.0
## [30] GenomicRanges_1.44.0
## [31] GenomeInfoDb 1.28.0
## [32] IRanges_2.26.0
## [33] S4Vectors 0.30.0
##
  [34] BiocGenerics_0.38.0
## loaded via a namespace (and not attached):
     [1] VGAM_1.1-5
                                colorspace_2.0-1
                                                       ggsignif_0.6.1
##
     [4] rjson_0.2.20
                                rio_0.5.26
                                                       ellipsis_0.3.2
##
     [7] som_0.3-5.1
                                farver_2.1.0
                                                       bit64_4.0.5
##
  [10] fansi_0.5.0
                                splines_4.1.0
                                                       cachem_1.0.5
## [13] knitr_1.33
                                broom_0.7.6
                                                       cluster_2.1.2
## [16] dbplyr_2.1.1
                                png_0.1-7
                                                       compiler_4.1.0
## [19] httr_1.4.2
                                backports_1.2.1
                                                       assertthat_0.2.1
## [22] Matrix_1.3-4
                                fastmap_1.1.0
                                                       formatR_1.11
                                                       tools_4.1.0
## [25] htmltools_0.5.1.1
                                prettyunits_1.1.1
## [28] gtable_0.3.0
                                glue_1.4.2
                                                       GenomeInfoDbData 1.2.6
```

	F047	1: 0.0.0	D 4 0 0	D . O O 1
##		rappdirs_0.3.3	Rcpp_1.0.6	carData_3.0-4
##	[34]	cellranger_1.1.0	vctrs_0.3.8	$nlme_3.1-152$
##	[37]	xfun_0.23	stringr_1.4.0	openxlsx_4.2.3
##	[40]	lifecycle_1.0.0	restfulr_0.0.13	<pre>formula.tools_1.7.1</pre>
##	[43]	gtools_3.9.2	rstatix_0.7.0	XML_3.99-0.6
##	[46]	beanplot_1.2	stringdist_0.9.6.3	zlibbioc_1.38.0
##	[49]	MASS_7.3-54	scales_1.1.1	hms_1.1.0
##	[52]	yaml_2.2.1	curl_4.3.1	memoise_2.0.0
##	[55]	biomaRt_2.48.0	reshape_0.8.8	stringi_1.6.2
##	[58]	RSQLite_2.2.7	highr_0.9	BiocIO_1.2.0
##	[61]	permute_0.9-5	filelock_1.0.2	zip_2.2.0
##	[64]	BiocParallel_1.26.0	operator.tools_1.6.3	rlang_0.4.11
##	[67]	pkgconfig_2.0.3	bitops_1.0-7	evaluate_0.14
##	[70]	purrr_0.3.4	labeling_0.4.2	bit_4.0.4
##	[73]	tidyselect_1.1.1	plyr_1.8.6	R6_2.5.0
##	[76]	generics_0.1.0	DelayedArray_0.18.0	DBI_1.1.1
##	[79]	haven_2.4.1	foreign_0.8-81	pillar_1.6.1
##	[82]	withr_2.4.2	mgcv_1.8-36	abind_1.4-5
##	[85]	KEGGREST_1.32.0	RCurl_1.98-1.3	tibble_3.1.2
##	[88]	crayon_1.4.1	car_3.0-10	KernSmooth_2.23-20
##	[91]	utf8_1.2.1	BiocFileCache_2.0.0	rmarkdown_2.8
##	[94]	progress_1.2.2	readxl_1.3.1	grid_4.1.0
##	[97]	blob_1.2.1	vegan_2.5-7	forcats_0.5.1
##	[100]	tidyr_1.1.3	munsell_0.5.0	
		•		