

iScience, Volume 25

Supplemental information

Hierarchical length and sequence preferences establish a single major piRNA 3'-end

Daniel Stoyko, Pavol Genzor, and Astrid D. Haase

Supplemental Figures (S1-3)
Stoyko et al.

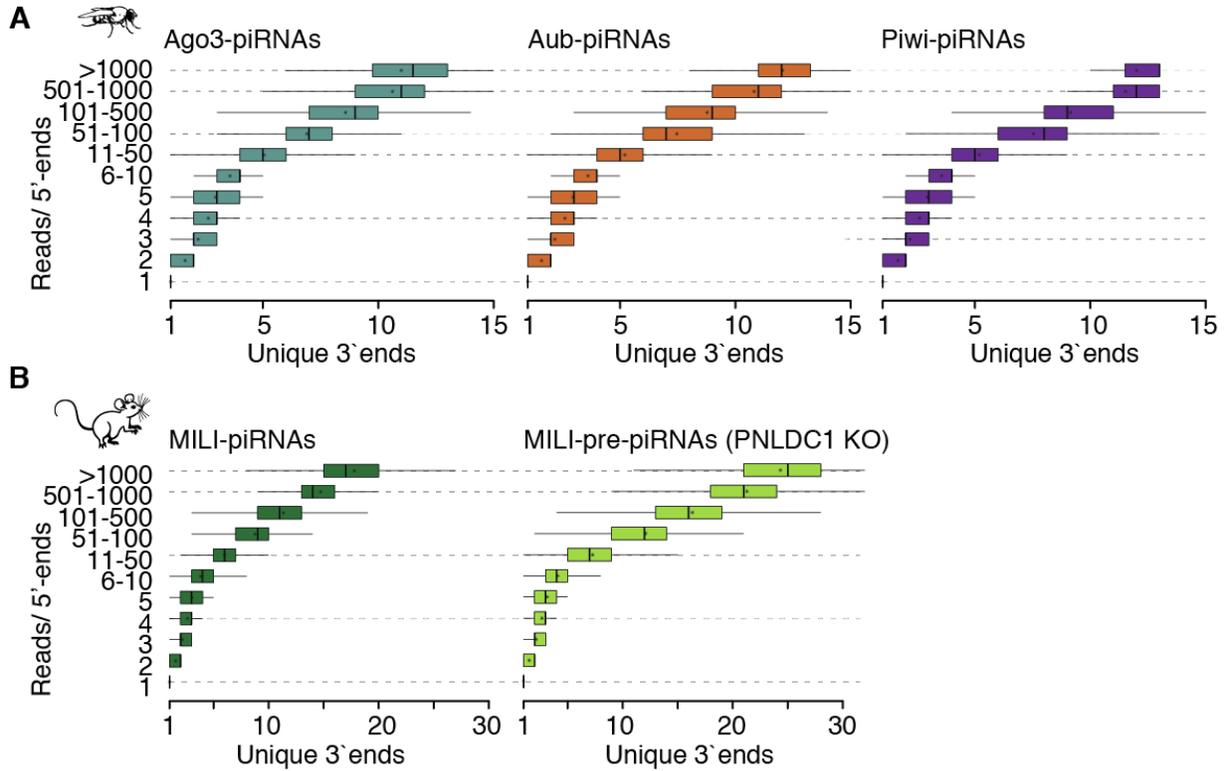


Figure S1. Fly and mouse piRNAs have a many possible 3'-ends, related to Figure 1. (A) *Drosophila* piRNAs with the same 5'-end have a number of different 3'-ends. Uniquely mapping piRNAs (18-32nt in length) were grouped by the number of reads sharing their respective 5'-ends. Number of unique 3'-ends observed per 5'-end is depicted for each group. **(B)** Both mature and untrimmed MILI-piRNAs can have several possible unique 3'-ends. (Uniquely mapping piRNAs were used.)

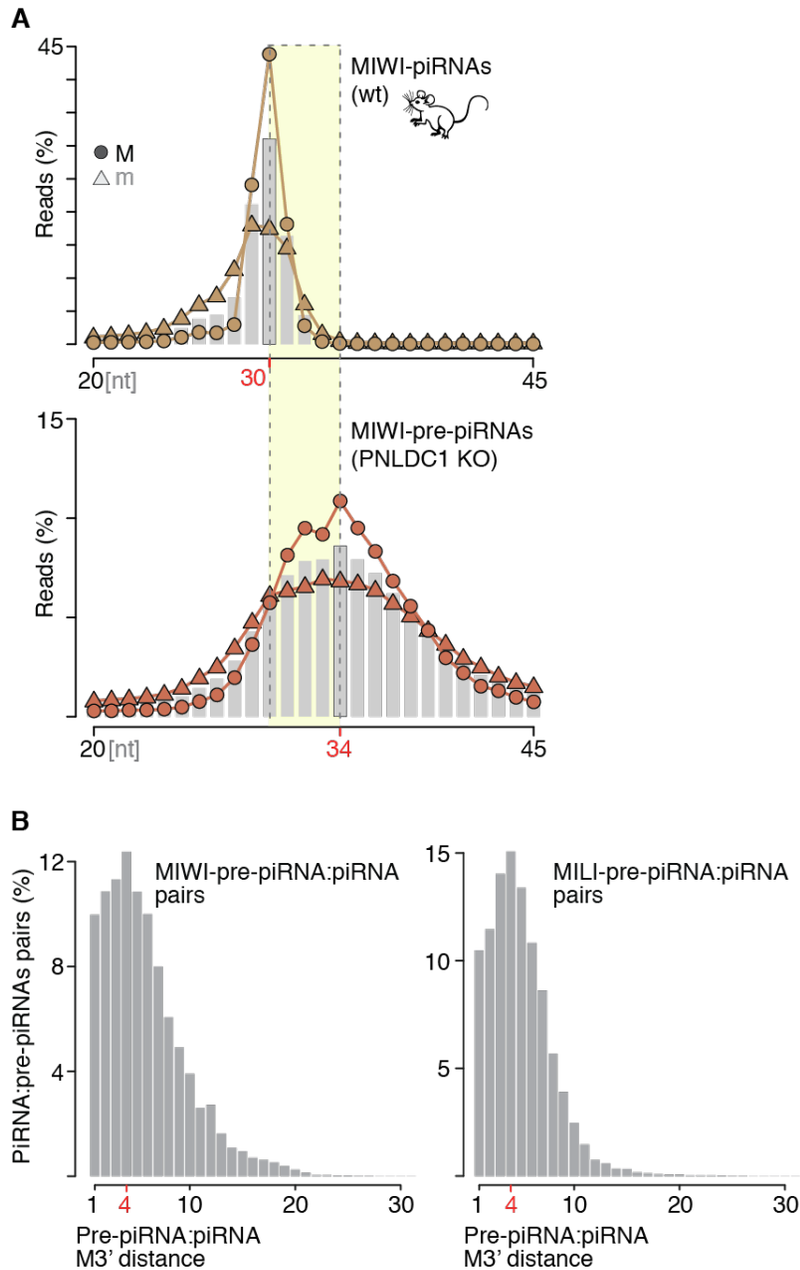


Figure S2. Major 3'-ends of murine piRNAs have specific length preferences that are four nucleotides (4nt) apart, related to Figure 2. (A) Both, trimmed and untrimmed MIWI-piRNAs with major 3'-ends have distinct length preferences at 30nt and 34nt respectively. The length distribution of major (M) and all minor (m) 3'-ends is normalized to the respective group. Bar plot depicts length distribution of the entire library. **(B)** Untrimmed murine piRNAs are preferentially 4nt longer than their mature counterparts. Untrimmed-piRNAs were paired with trimmed-piRNAs that have the same 5'-end. The distance between the major 3'-ends of the pair was calculated and scaled by the number of piRNAs in the pair.

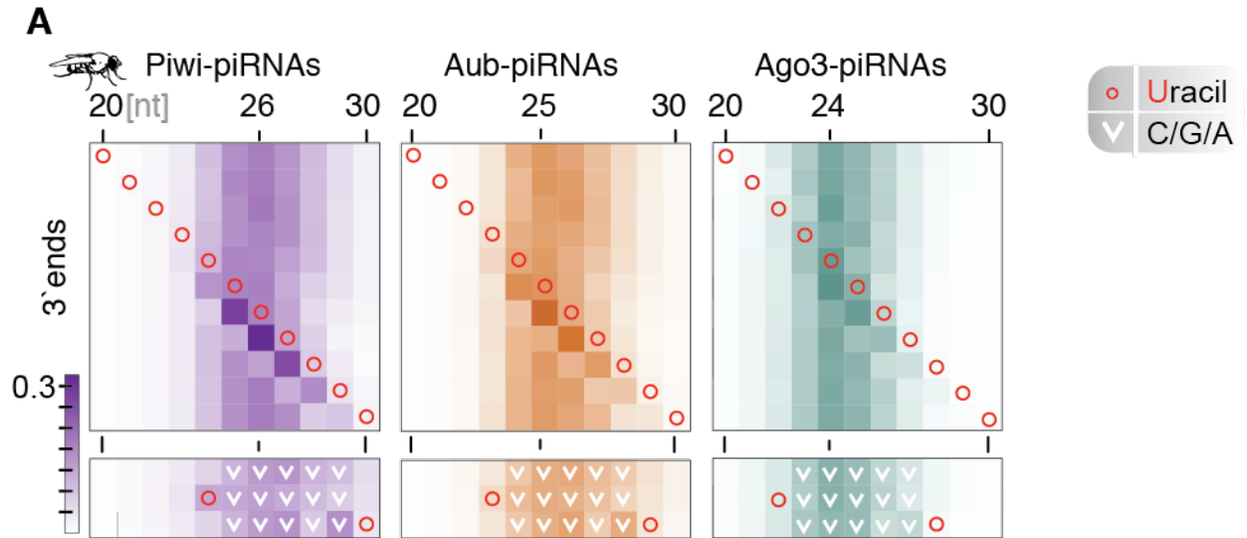


Figure S3. The presence of Uridines (U) affects the position of 3'-ends only within an optimal distance from the piRNA 5'-end, related to Figure 4. (A) In fly ovary piRNAs, the 3'-end is defined by the location of Uridine within a narrow window located at a fixed distance from the 5'-end. Rows depict distribution of 3'-ends for piRNAs where a Uridine (red circle) or absence of Uridine (V) is present at a certain distance from the 5'-end.

Data S1.
Supplemental Code
related to Figures 1-4
Stoyko et al.

(https://github.com/HaaseLab/piRNA_ThreePrimeEnds)

Diversity of piRNA 3'ends

methods by: Daniel Stoyko & Pavol Genzor

12/06/2021

Daniel Stoyko, Pavol Genzor, Astrid D. Haase

This vignette describes the computational materials & methods associated with this manuscript. Please visit [HaaseLab/piRNA_ThreePrimeEnds github repository](#) to download functions used in the various scripts and analyses. Please refer to the GEO data set **GSE156058** associated with this study for adapter sequences and raw data. The analysis in this vignette was not performed with full data sets, but only subset of the data to demonstrate the materials and methods.

About small RNA Libraries.

Small RNA libraries were prepared and pre-processed as described in Genzor et.al., 2021. Please refer to the corresponding github page for more information.

Pre-requisites & Notes

* Acquire the raw sequencing data from your facility or GEO at NCBI * NOTE: some images may not have rendered perfectly

Vignette Content

1. Figure1: Contribution of piRNA three prime ends - End Dominance
2. Figure2: Size distribution of major and minor ends of piRNAs
3. Figure3: Major and minor -dependent nucleotide frequencies of piRNAs
4. Figure4: Correlation of 3' end with nucleotide context

Environment setup & test data preparation

Prepare your R working environment by ensuring you have all necessary packages installed and loaded. Download the *.RData* object from *github* to be and load it into R before proceeding.

```
# load libraries
suppressWarnings(
  suppressPackageStartupMessages({library("data.table"); library("dplyr");
  library("ggplot2"); library("ggpubr"); library("GenomicRanges");
  library("ggseqlogo"); library("BSgenome.Dmelanogaster.UCSC.dm6");
  library("stringr"); library("parallel")}))
```

Load filtered subset of the data

```
SESSION.DIR="/Users/genzorp/Documents/GITHUB/piThreePrime/sessions/"
load(file = paste0(SESSION.DIR,"single_filtered_brep_1M.RData"))
```

Figure 1: Contribution of piRNA three prime ends

```
## Data
GR <- original.GR

## Make a table of uniquely mapping piRNAs

uniq.DT <- as.data.table(GR[mcols(GR)[["NH"]] %in% 1])

## Create a column with 5'-end coordinates
# NOTE: "start" refers to the 5'-end on the "+" strand and
# NOTE: "end" refers to 5'-end on "-" strand

uniq.DT[,FivePrime := paste0(seqnames, "_", ifelse(strand == "-", end, start), "_", strand)]

## Randomly assign each piRNA a value between 1 and N
## :- N is the number of unique sequences sharing the 5'-end without repeating
## :- This random value will be used to break ties between equally abundant 3'-ends
## :- sample() randomly assigns numbers within particular range

uniq.DT[,TieBreaker := base::sample(x = nrow(.SD), size = nrow(.SD), replace = F),
  by = FivePrime)]

## Define the Major End as the most abundant sequence per 5'-end
## :- use TieBreaker value to solve ties in abundance

uniq.DT <- uniq.DT[order(-MULT, TieBreaker)]
uniq.DT[,EndOrder := c(1:nrow(.SD)), by=FivePrime]
uniq.DT[,EndClass := ifelse(EndOrder > 1, "MinorEnds","MajorEnd")]
uniq.DT[["EndClass"]] <- factor(x = uniq.DT[["EndClass"]], levels = c("MinorEnds","MajorEnd"))

## Group 5'-ends into custom bins based on their abundance
## :- NOTE: the abundance is not normalized here

# summarize abundance
uniq.DT <- setDT(uniq.DT)
uniq.DT[,FivePrimeAbund := sum(MULT), by=FivePrime]

# find bins
aBreaks <- c(1,2,3,4,5,10,50,100,500,1000)
uniq.DT[["aBin"]] <- findInterval(x = uniq.DT[["FivePrimeAbund"]],
  vec = aBreaks, left.open = TRUE)

# add and organize bin name
uniq.DT <- setDT(uniq.DT)
uniq.DT[, "binName" := paste0(unique(c(min(FivePrimeAbund),max(FivePrimeAbund))),
  collapse = "-") , by = aBin]
uniq.DT[FivePrimeAbund > 1000][["binName"]] <- ">1000"
```

```
# order bin names
unique(unique.DT[["binName"]])
```

```
## [1] ">1000" "501-990" "101-500" "51-100" "11-50" "6-10" "5"
## [8] "4" "3" "2" "1"
```

```
binNameOrder <- c("1","2","3","4","5","6-10","11-50","51-100","101-500","501-999",>1000")
unique.DT[["aBin"]] <- factor(unique.DT[["aBin"]])
unique.DT[["binName"]] <- factor(x = unique.DT[["binName"]], levels = binNameOrder)
```

```
## Calculate contribution of the Major 3'-end to all the ends
```

```
unique.DT <- setDT(unique.DT)
unique.DT[,EndContribution := MULT/sum(MULT), by=FivePrime]
```

```
## View the generated table
```

```
unique.DT[FivePrime %in% "chr2L_3107310_-"]
```

```
##      seqnames      start      end width strand N NH MULT      FivePrime TieBreaker
## 1:   chr2L 3107284 3107310    27    - 3  1  27 chr2L_3107310_-      5
## 2:   chr2L 3107289 3107310    22    - 3  1   3 chr2L_3107310_-      1
## 3:   chr2L 3107293 3107310    18    - 2  1   2 chr2L_3107310_-      7
## 4:   chr2L 3107290 3107310    21    - 1  1   1 chr2L_3107310_-      2
## 5:   chr2L 3107285 3107310    26    - 1  1   1 chr2L_3107310_-      3
## 6:   chr2L 3107286 3107310    25    - 1  1   1 chr2L_3107310_-      4
## 7:   chr2L 3107288 3107310    23    - 1  1   1 chr2L_3107310_-      6
## 8:   chr2L 3107283 3107310    28    - 1  1   1 chr2L_3107310_-      8
##      EndOrder EndClass FivePrimeAbund aBin binName EndContribution
## 1:         1 MajorEnd          37     6 11-50      0.72972973
## 2:         2 MinorEnds         37     6 11-50      0.08108108
## 3:         3 MinorEnds         37     6 11-50      0.05405405
## 4:         4 MinorEnds         37     6 11-50      0.02702703
## 5:         5 MinorEnds         37     6 11-50      0.02702703
## 6:         6 MinorEnds         37     6 11-50      0.02702703
## 7:         7 MinorEnds         37     6 11-50      0.02702703
## 8:         8 MinorEnds         37     6 11-50      0.02702703
```

```
## Prepare data for visualization
```

```
Barplot.DT <- unique.DT[,.(Sum_Abundance = sum(MULT)), by=c("EndClass","aBin","binName")]
Barplot.DT
```

```
##      EndClass aBin binName Sum_Abundance
## 1: MajorEnd   10 >1000      381870
## 2: MinorEnds  10 >1000      247899
## 3: MajorEnd   9  <NA>        81875
## 4: MajorEnd   8 101-500     194990
## 5: MinorEnds  9  <NA>        49459
## 6: MinorEnds  8 101-500     111306
## 7: MajorEnd   7  51-100     114528
## 8: MajorEnd   6  11-50     349854
## 9: MinorEnds  7  51-100     59529
## 10: MinorEnds 6  11-50     149928
## 11: MajorEnd  5  6-10      212093
```

```
## 12: MajorEnd 4 5 78938
## 13: MinorEnds 5 6-10 61081
## 14: MajorEnd 3 4 112682
## 15: MajorEnd 2 3 247555
## 16: MajorEnd 1 2 327858
## 17: MinorEnds 4 5 15932
## 18: MinorEnds 3 4 15770
## 19: MajorEnd 0 1 177097
## 20: MinorEnds 2 3 9194
## 21: MinorEnds 1 2 4202
## EndClass aBin binName Sum_Abundance
```

```
NumberOfEnds.DT <- uniq.DT[, .N, by=c("FivePrime", "aBin", "binName")]
NumberOfEnds.DT
```

```
##           FivePrime aBin binName N
## 1: chr2L_16698758_+ 10 >1000 10
## 2: chr2L_16697936_+ 10 >1000 9
## 3: chr2L_7425853_+ 10 >1000 7
## 4: chr2L_7425979_+ 10 >1000 9
## 5: chr2L_16698418_+ 10 >1000 7
## ---
## 546175: chr2R_7522774_- 0 1 1
## 546176: chr2R_7522843_- 0 1 1
## 546177: chr2R_7522842_- 0 1 1
## 546178: chr2R_7522898_- 0 1 1
## 546179: chr2R_7523220_- 0 1 1
```

```
MeanNumberOfEnds.DT <- NumberOfEnds.DT[, lapply(.SD, mean), by=c("aBin", "binName"), .SDcols="N"]
MeanNumberOfEnds.DT
```

```
## aBin binName N
## 1: 10 >1000 8.812500
## 2: 9 <NA> 7.797872
## 3: 8 101-500 5.689873
## 4: 7 51-100 4.194179
## 5: 6 11-50 2.594930
## 6: 5 6-10 1.775432
## 7: 4 5 1.506377
## 8: 3 4 1.338025
## 9: 2 3 1.107428
## 10: 1 2 1.025309
## 11: 0 1 1.000000
```

```
## NOTE: Using the three tables generated in previous chunk
```

```
# Barplot.DT
```

```
# NumberOfEnds.DT
```

```
# MeanNumberOfEnds.DT
```

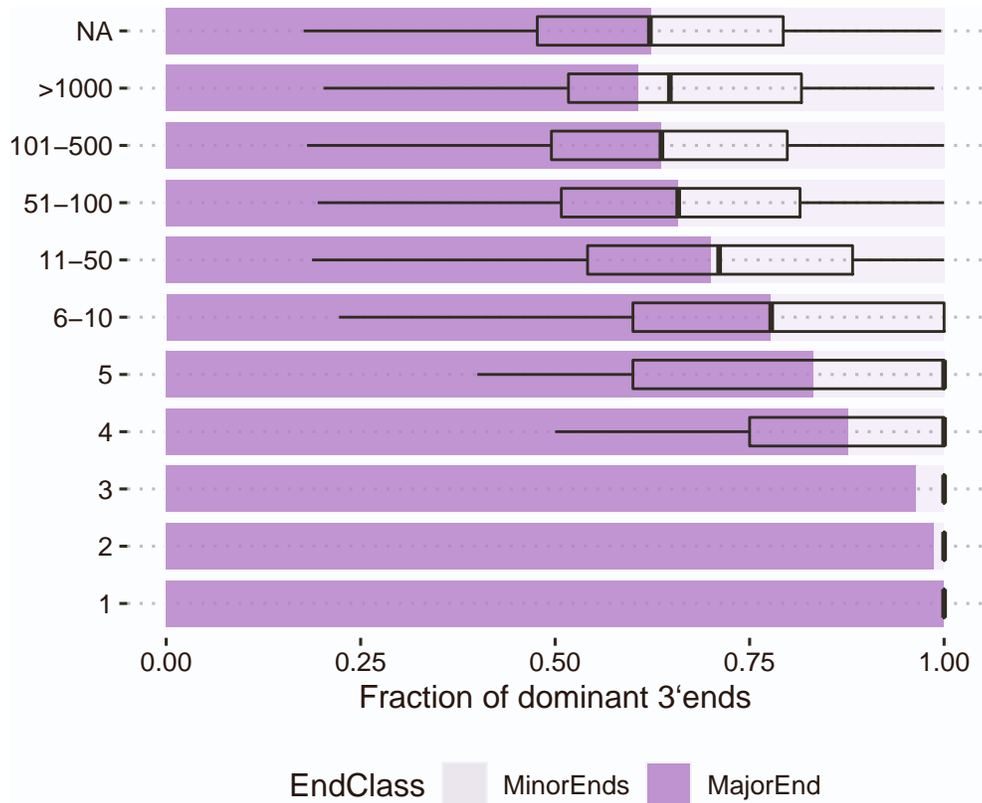
```
##
```

```
EndContributionPlot <- ggplot() + theme_pubclean() +
  geom_bar(data = Barplot.DT,
    aes(x = binName, y = Sum_Abundance, alpha = EndClass, group = EndClass),
```

```

stat = "identity", position = "fill", fill="#B37FC7",
width = 0.8) +
geom_boxplot(data = uniq.DT[EndClass %in% "MajorEnd"],
aes(x = binName, y= EndContribution),
outlier.shape = NA, width = 0.5, fill = NA) +
ylab("Fraction of dominant 3`ends") + xlab("") + coord_flip() +
scale_alpha_manual(values=c(0.1,0.8)) +
theme(legend.position="bottom",
axis.text = element_text(family = "Helvetica",colour = "black", size = 10),
aspect.ratio=0.75); EndContributionPlot

```



```

##
NumberOfEndsPlot <- ggplot() + theme_pubclean() +
geom_boxplot(data = NumberOfEnds.DT,
aes(x = binName, y = N),
color="black", fill="#C49CD3",
outlier.shape = NA) +
geom_point(data = MeanNumberOfEnds.DT,
aes(x= binName, y = N),
shape = 16, color = "firebrick3", size = 6) +
ylab("number of unique 3`ends") + xlab("") +
ggtitle(paste0("red = mean")) + coord_flip() +
theme(aspect.ratio = 0.75,
axis.text = element_text(family = "Helvetica",colour = "black", size = 10),
legend.position = "none"); NumberOfEndsPlot

```

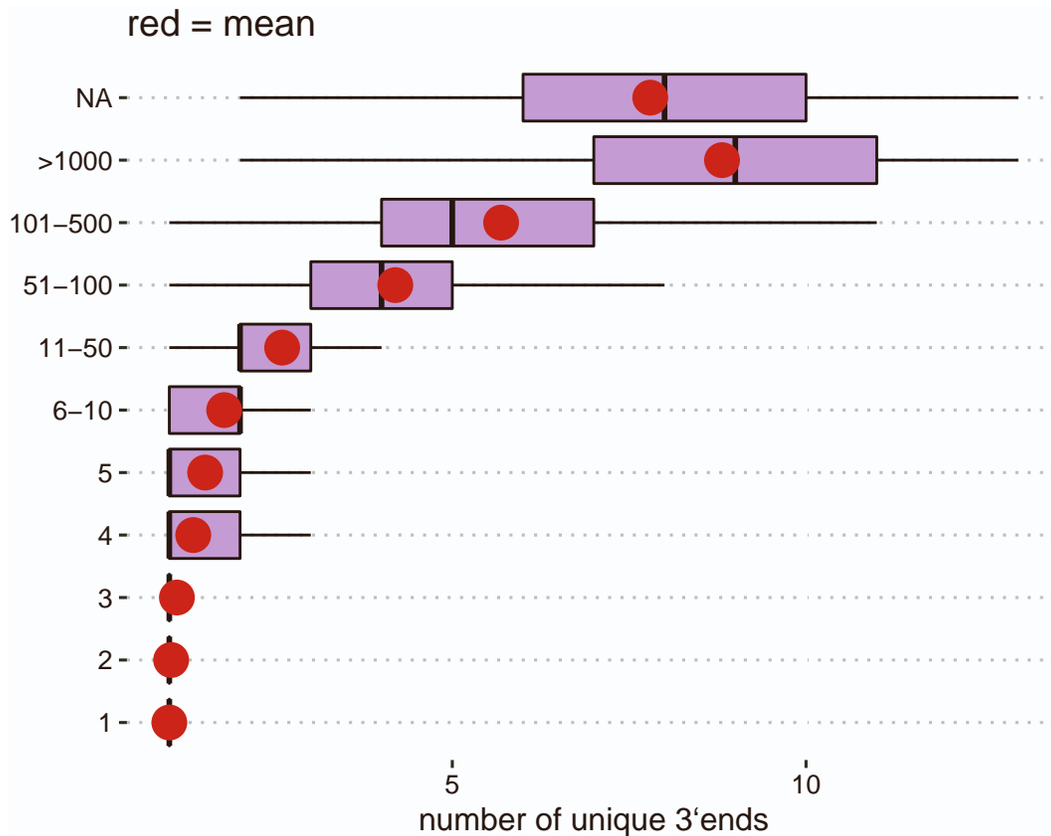


Figure 2: Size distribution of major and minor ends of piRNAs

```
## NOTE: Use that table generated in the previous chunk
##   uniq.DT

## Table by end type
MajorMinorLengths.DT <- uniq.DT[,lapply(.SD,sum), by=c("EndClass","width"), .SDcols="MULT"]
MajorMinorLengths.DT <- MajorMinorLengths.DT[order(EndClass,width)]
MajorMinorLengths.DT[,"groupReadSum" := sum(.SD),by="EndClass", .SDcols="MULT"]
MajorMinorLengths.DT[,"totalReads" := sum(.SD), .SDcols="MULT"]
MajorMinorLengths.DT[,"groupPercent" := (MULT/groupReadSum)*100]
MajorMinorLengths.DT[,"totalPercent" := (MULT/totalReads)*100]
MajorMinorLengths.DT
```

##	EndClass	width	MULT	groupReadSum	totalReads	groupPercent	totalPercent
##	1: MinorEnds	18	14295	724300	3003640	1.973630e+00	4.759225e-01
##	2: MinorEnds	19	17585	724300	3003640	2.427861e+00	5.854563e-01
##	3: MinorEnds	20	29861	724300	3003640	4.122739e+00	9.941604e-01
##	4: MinorEnds	21	58641	724300	3003640	8.096231e+00	1.952331e+00
##	5: MinorEnds	22	83027	724300	3003640	1.146307e+01	2.764213e+00
##	6: MinorEnds	23	48168	724300	3003640	6.650283e+00	1.603654e+00
##	7: MinorEnds	24	116744	724300	3003640	1.611818e+01	3.886751e+00
##	8: MinorEnds	25	123195	724300	3003640	1.700884e+01	4.101523e+00
##	9: MinorEnds	26	116621	724300	3003640	1.610120e+01	3.882656e+00
##	10: MinorEnds	27	99043	724300	3003640	1.367431e+01	3.297432e+00
##	11: MinorEnds	28	16153	724300	3003640	2.230153e+00	5.377808e-01

```

## 12: MinorEnds 29 857 724300 3003640 1.183211e-01 2.853205e-02
## 13: MinorEnds 30 90 724300 3003640 1.242579e-02 2.996364e-03
## 14: MinorEnds 31 17 724300 3003640 2.347094e-03 5.659799e-04
## 15: MinorEnds 32 3 724300 3003640 4.141930e-04 9.987881e-05
## 16: MajorEnd 18 40386 2279340 3003640 1.771829e+00 1.344569e+00
## 17: MajorEnd 19 32049 2279340 3003640 1.406065e+00 1.067005e+00
## 18: MajorEnd 20 28716 2279340 3003640 1.259838e+00 9.560400e-01
## 19: MajorEnd 21 67803 2279340 3003640 2.974677e+00 2.257361e+00
## 20: MajorEnd 22 77983 2279340 3003640 3.421297e+00 2.596283e+00
## 21: MajorEnd 23 118871 2279340 3003640 5.215150e+00 3.957565e+00
## 22: MajorEnd 24 305666 2279340 3003640 1.341029e+01 1.017652e+01
## 23: MajorEnd 25 720392 2279340 3003640 3.160529e+01 2.398397e+01
## 24: MajorEnd 26 674169 2279340 3003640 2.957738e+01 2.244507e+01
## 25: MajorEnd 27 196817 2279340 3003640 8.634824e+00 6.552616e+00
## 26: MajorEnd 28 15607 2279340 3003640 6.847158e-01 5.196029e-01
## 27: MajorEnd 29 738 2279340 3003640 3.237779e-02 2.457019e-02
## 28: MajorEnd 30 88 2279340 3003640 3.860767e-03 2.929779e-03
## 29: MajorEnd 31 47 2279340 3003640 2.062000e-03 1.564768e-03
## 30: MajorEnd 32 8 2279340 3003640 3.509788e-04 2.663435e-04
## EndClass width MULT groupReadSum totalReads groupPercent totalPercent

```

```
## All reads
```

```

AllLengths.DT <- uniq.DT[,.(MULT = sum(.SD)), by=c("width"), .SDcols="MULT"]
AllLengths.DT[,"totalReads" := sum(.SD), .SDcols = "MULT"]
AllLengths.DT[,"totalPercent" := (MULT/totalReads)*100]
AllLengths.DT

```

```

## width MULT totalReads totalPercent
## 1: 23 167039 3003640 5.561219e+00
## 2: 22 161010 3003640 5.360496e+00
## 3: 24 422410 3003640 1.406327e+01
## 4: 21 126444 3003640 4.209692e+00
## 5: 25 843587 3003640 2.808549e+01
## 6: 26 790790 3003640 2.632772e+01
## 7: 20 58577 3003640 1.950200e+00
## 8: 27 295860 3003640 9.850049e+00
## 9: 19 49634 3003640 1.652462e+00
## 10: 18 54681 3003640 1.820491e+00
## 11: 28 31760 3003640 1.057384e+00
## 12: 29 1595 3003640 5.310224e-02
## 13: 30 178 3003640 5.926143e-03
## 14: 31 64 3003640 2.130748e-03
## 15: 32 11 3003640 3.662223e-04

```

```
## Plot
```

```

LengthDistributionPlot <- ggplot() + theme_pubclean() +
  geom_bar(data = AllLengths.DT,
           aes(x=width, y=totalPercent),
           stat="identity", fill="gray80", width = 0.8) +
  geom_line(data = MajorMinorLengths.DT,
           aes(x = width, y = groupPercent, colour = EndClass),
           size = 1) +
  geom_point(data = MajorMinorLengths.DT,

```

```

aes(x = width, y = groupPercent, shape = EndClass, colour = EndClass),
size = 4) +
xlab("size (nt)") + ylab("percent of piRNAs") +
scale_x_continuous(breaks = seq(0,40,2)) +
scale_y_continuous(breaks = seq(0,40,5)) +
scale_colour_manual(values = c("#AB72C0", "#FF6D33")) +
theme(aspect.ratio = 1, legend.position = "top",
axis.text = element_text(family = "Helvetica", colour = "black", size = 10))
LengthDistributionPlot

```

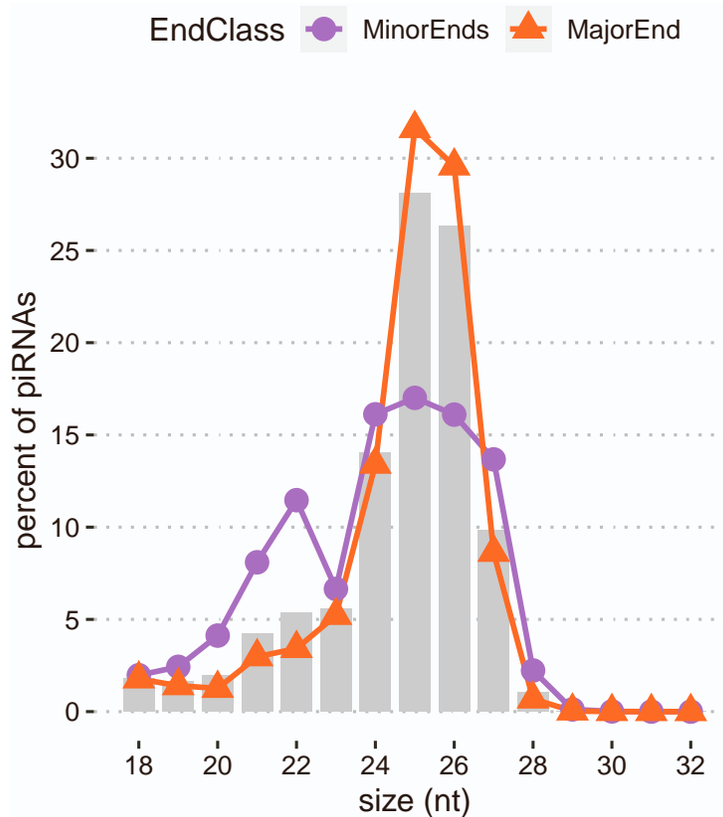


Figure S2: Distance between major ends in WT and PNLDC KO

```

## NOTE: Scripts used for comparing mouse WT to PNLDC KO piRNA major ends
## NOTE: Sample data is not provided. Data can be obtained from PRJNA421205
## See methods for details

## Set GR1 as the WT library
## Set GR2 as the PNLDC1 KO library

GR1 <- MOUSE.BAM.GR.WT
GR2 <- MOUSE.BAM.GR.KO

## For GR1, determine the major 3'-end in same manner as described previously
uniq.DT1 <- as.data.table(GR1[mcols(GR1)[["NH"]] %in% 1])
uniq.DT1[,FivePrime1 := paste0(seqnames, "_", ifelse(strand == "-", end, start), "_", strand)]

```

```

uniq.DT1[,TieBreaker := base::sample(x = nrow(.SD), size = nrow(.SD), replace = F),
      by = FivePrime1]
uniq.DT1 <- uniq.DT1[order(-MULT, TieBreaker)]
uniq.DT1[,EndOrder := c(1:nrow(.SD)), by=FivePrime1]
uniq.DT1[,EndClass := ifelse(EndOrder > 1, "MinorEnds","MajorEnd")]

## keep only major 3'-ends
uniq.DT1.maj <- uniq.DT1[EndClass %in% c("MajorEnd")]

## Determine 3'-end coordinate and discard unnecessary columns
uniq.DT1.maj[,ThreePrime1 := ifelse(strand == "-", start, end)]
uniq.DT1.maj <- uniq.DT1.maj[,c(5, 7, 8, 12)]
colnames(uniq.DT1.maj) <- c("strand1", "MULT1", "FivePrime1", "ThreePrime1")

uniq.DT1.maj

## Repeat for GR2
uniq.DT2 <- as.data.table(GR2[mcols(GR2)[["NH"]] %in% 1])
uniq.DT2[,FivePrime2 := paste0(seqnames, "_", ifelse(strand == "-", end, start), "_", strand)]
uniq.DT2[,TieBreaker := base::sample(x = nrow(.SD), size = nrow(.SD), replace = F),
      by = FivePrime2]
uniq.DT2 <- uniq.DT2[order(-MULT, TieBreaker)]
uniq.DT2[,EndOrder := c(1:nrow(.SD)), by=FivePrime2]
uniq.DT2[,EndClass := ifelse(EndOrder > 1, "MinorEnds","MajorEnd")]
uniq.DT2.maj <- uniq.DT2[EndClass %in% c("MajorEnd")]
uniq.DT2.maj[,ThreePrime2 := ifelse(strand == "-", start, end)]
uniq.DT2.maj <- uniq.DT2.maj[,c(5, 7, 8, 12)]
colnames(uniq.DT2.maj) <- c("strand2", "MULT2", "FivePrime2", "ThreePrime2")

## Merge the two files by 5'-end coordinate
## Discard sequences with 5'-ends not present in both datasets
DT.combined <- merge(uniq.DT1.maj, uniq.DT2.maj, by.x="FivePrime1", by.y="FivePrime2", all.x=F, all.y =

## Calculate total abundance of piRNAs in a pair
DT.combined[, MULTtotal := MULT1 + MULT2]

## Determine distance between the two major 3'-ends
DT.combined[, Distance := ifelse(strand1 == "+", ThreePrime2-ThreePrime1, ThreePrime1-ThreePrime2)]

## Keep sequences where PNLDC KO piRNA is longer than the WT piRNA
DT.combined <- DT.combined[Distance > 0]

## Count number of reads for each distance
DT.combined.sum <- DT.combined[, lapply(.SD, sum), .SDcols="MULTtotal", by="Distance"]

## Convert to % of total
DT.combined.sum$MULTtotal <- (DT.combined.sum$MULTtotal/ sum(DT.combined.sum$MULTtotal))*100

## Plot
DistancePlot <- ggplot() + theme_bw() +
  geom_bar(data=DT.combined.sum,
    aes(x=Distance, y = MULTtotal),
    color= "black",

```

```

    stat="identity",
    width= 0.85) +
labs(title = "SampleData",
     x= "Distance between the major 3'ends in WT and PNLDC KO",
     y= "% of total") +
theme(panel.grid = element_blank(),
     aspect.ratio = 1)

```

Figure 3: Major and minor -dependent nucleotide frequencies of piRNAs

```

## Data
GR <- original.GR

## Use only uniquely mapping piRNAs
uniq.GR <- GR[mcols(GR)[["NH"]] %in% "1"]
uniq.DT <- as.data.table(uniq.GR)

## Expand the GR to include surrounding nucleotides
uniq.GRE <- uniq.GR
end(uniq.GRE) <- end(uniq.GRE) + 4
start(uniq.GRE) <- start(uniq.GRE) - 4

## Add sequence to the table
uniq.DT[["seq"]] <- as.vector(getSeq(BSgenome.Dmelanogaster.UCSC.dm6, uniq.GRE))

## Create a column with 5'-end coordinates
uniq.DT <- setDT(uniq.DT)
uniq.DT[,FivePrime := paste0(seqnames, "_", ifelse(strand == "-", end, start), "_", strand)]

## Randomly assign each piRNA a value between 1 and N
uniq.DT[,TieBreaker := sample(nrow(.SD), size = nrow(.SD), replace = F), by = FivePrime]

## Define the Major End as the most abundant sequence per 5'-end
uniq.DT <- uniq.DT[order(-MULT, TieBreaker)]
uniq.DT[,EndOrder := c(1:nrow(.SD)), by=FivePrime]
uniq.DT[,EndClass := ifelse(EndOrder > 1, "MinorEnds","MajorEnd")]
uniq.DT[["EndClass"]] <- factor(x = uniq.DT[["EndClass"]], levels = c("MinorEnds","MajorEnd"))

## Group 5'-ends into custom bins based on their abundance
uniq.DT <- setDT(uniq.DT)
uniq.DT[,FivePrimeAbund := sum(MULT), by=FivePrime]
uniq.DT

```

```

##      seqnames      start      end width strand N NH  MULT
##    1:  chr2L 16698758 16698780   23    + 3  1 28686
##    2:  chr2L 16697936 16697957   22    + 3  1 27064
##    3:  chr2L 16698758 16698779   22    + 3  1 26662
##    4:  chr2L  7425853  7425875   23    + 3  1 24812
##    5:  chr2L  7425979  7426002   24    + 3  1 23638
##    ---
## 666754:  chr2L 19467210 19467238   29    + 1  1    1
## 666755:  chr2L 19467405 19467432   28    + 1  1    1

```

```

## 666756: chr2R 5050923 5050953 31 + 1 1 1
## 666757: chr2R 5055725 5055744 20 + 1 1 1
## 666758: chr2R 5055342 5055370 29 + 1 1 1
##
## seq FivePrime TieBreaker
## 1: TTGCTCTTTGGTGATTTAGCTGTATGGTGT chr2L_16698758_+ 7
## 2: TGTTTCTTTGGTATTCTAGCTGTAGATTGT chr2L_16697936_+ 9
## 3: TTGCTCTTTGGTGATTTAGCTGTATGGTGT chr2L_16698758_+ 6
## 4: ACAGTCAGGTACCTGAAGTAGCGCGGTGGT chr2L_7425853_+ 2
## 5: GTCTATTGTACTTCATCAGGTGCTCTGGTGTG chr2L_7425979_+ 7
## ---
## 666754: ACGCTTATTTGTTGATTAGTTCTAGCCTTAGTTTCCC chr2L_19467210_+ 10
## 666755: AAGTTAAACACCCGAAGCTGGAAGAACCGATGTAT chr2L_19467405_+ 10
## 666756: CAACTTACGCATATGTGAGTGGGGAAAGGACTCGGACAG chr2R_5050923_+ 10
## 666757: AATTTGTTTCGTCACAGTATGCAATATT chr2R_5055725_+ 10
## 666758: TAAATCTTGATTTGCGGTGCTTCCACCTGCAAACCTCT chr2R_5055342_+ 11
## EndOrder EndClass FivePrimeAbund
## 1: 1 MajorEnd 67950
## 2: 1 MajorEnd 41154
## 3: 2 MinorEnds 67950
## 4: 1 MajorEnd 46882
## 5: 1 MajorEnd 53587
## ---
## 666754: 11 MinorEnds 384
## 666755: 10 MinorEnds 648
## 666756: 11 MinorEnds 840
## 666757: 10 MinorEnds 308
## 666758: 12 MinorEnds 1238

```

```

## View the generated table
uniq.DT[FivePrime %in% "chr2L_3107310_-"]

```

```

## seqnames start end width strand N NH MULT
## 1: chr2L 3107284 3107310 27 - 3 1 27
## 2: chr2L 3107289 3107310 22 - 3 1 3
## 3: chr2L 3107293 3107310 18 - 2 1 2
## 4: chr2L 3107290 3107310 21 - 1 1 1
## 5: chr2L 3107285 3107310 26 - 1 1 1
## 6: chr2L 3107283 3107310 28 - 1 1 1
## 7: chr2L 3107288 3107310 23 - 1 1 1
## 8: chr2L 3107286 3107310 25 - 1 1 1
##
## seq FivePrime TieBreaker EndOrder
## 1: CAACTCTCTCTGCGTCTCTCTATAGACCCGATTAC chr2L_3107310_- 7 1
## 2: CAACTCTCTCTGCGTCTCTCTATAGACCCG chr2L_3107310_- 5 2
## 3: CAACTCTCTCTGCGTCTCTCTATAGA chr2L_3107310_- 8 3
## 4: CAACTCTCTCTGCGTCTCTCTATAGACCC chr2L_3107310_- 1 4
## 5: CAACTCTCTCTGCGTCTCTCTATAGACCCGATTA chr2L_3107310_- 2 5
## 6: CAACTCTCTCTGCGTCTCTCTATAGACCCGATTACC chr2L_3107310_- 3 6
## 7: CAACTCTCTCTGCGTCTCTCTATAGACCCGA chr2L_3107310_- 4 7
## 8: CAACTCTCTCTGCGTCTCTCTATAGACCCGATT chr2L_3107310_- 6 8
## EndClass FivePrimeAbund
## 1: MajorEnd 37
## 2: MinorEnds 37
## 3: MinorEnds 37
## 4: MinorEnds 37

```

```
## 5: MinorEnds          37
## 6: MinorEnds          37
## 7: MinorEnds          37
## 8: MinorEnds          37
```

```
## Plotting LOGO
## 1. select type of end and its sequences
## 2. extract sub-string
## 3. expand to reads
## 4. replace nucleotides

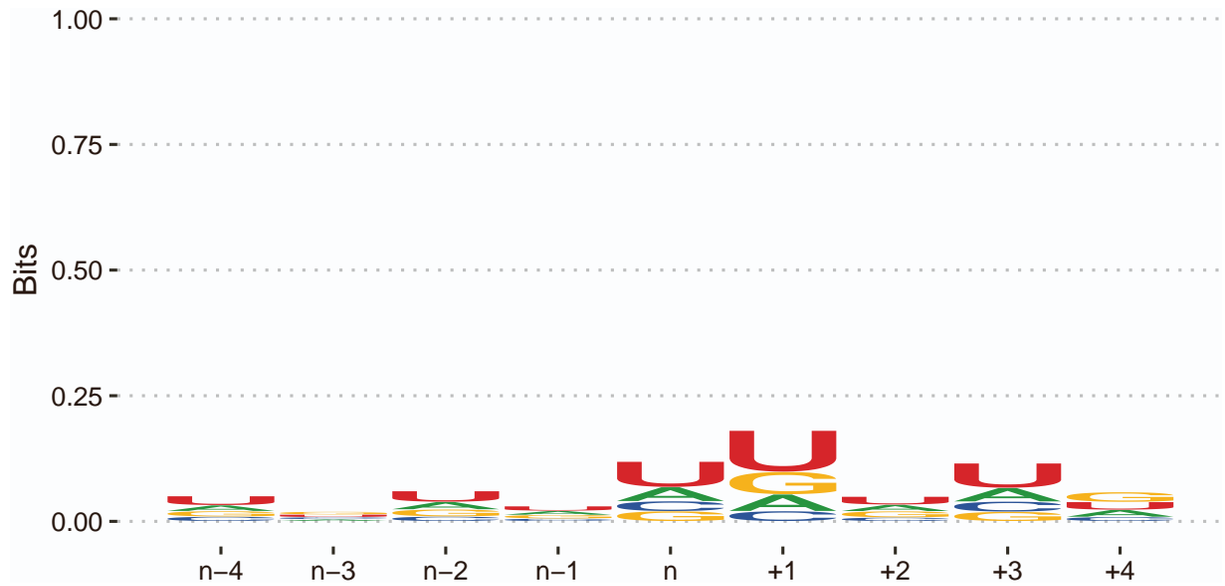
ForLogo.second.DT <- uniq.DT[EndOrder %in% 2][,c("MULT", "seq")]
ForLogo.second.DT[,Seq_Range := substring(seq, nchar(seq)-8 ,nchar(seq))]
ForLogo.second.DT <- ForLogo.second.DT[ rep( c(1:nrow(ForLogo.second.DT)),
                                           times = MULT)]
ForLogo.second.DT[["Seq_Range"]] <- gsub("T","U",ForLogo.second.DT[["Seq_Range"]])

##
## Plot LOGO
##

LogoPlot <- ggplot() + theme_pubclean() +
  geom_logo(data=ForLogo.second.DT[["Seq_Range"]],
            method="bits", seq_type="rna") +
  scale_y_continuous(limits = c(0,1)) +
  scale_x_continuous(
    breaks = 1:9,
    labels = paste(c("n-4","n-3","n-2","n-1","n","+1","+2","+3","+4")) ) +
  theme(panel.grid = element_blank(),
        axis.text = element_text(family = "Helvetica", colour = "black", size = 10),
        aspect.ratio = 0.5); LogoPlot
```

```
## Warning: 'guides(<scale> = FALSE)' is deprecated. Please use 'guides(<scale> =
## "none")' instead.
```

```
## Scale for 'x' is already present. Adding another scale for 'x', which will
## replace the existing scale.
```



```
##
## Plot +1 nucleotide frequency
##

## For barplots with +1 nucleotide composition:
## isolate the +1 nucleotide

uniq.DT[, plusOne := substring(seq, nchar(seq)-3, nchar(seq)-3)]
PlusOneComposition.DT <- uniq.DT[, .(MULT = sum(MULT)), by=c("EndClass","plusOne")]
PlusOneComposition.DT[["plusOne"]] <- factor(PlusOneComposition.DT[["plusOne"]], c("A","G","C","T"))

## set colors
nuc_colors <- c("#00AF54", "#FFD639", "#447EC5", "#DF2935")

## plot
PlusOneCompositionPlot <- ggplot() + theme_pubclean()+
  geom_bar(data = PlusOneComposition.DT,
           aes(x = EndClass, y = MULT, fill = plusOne),
           stat="identity", position="fill", width = 0.8) +
  scale_fill_manual(values=nuc_colors) +
  ylab("fraction nucleotide") +
  theme(aspect.ratio = 2,
        legend.position = "right",
        axis.text = element_text(family = "Helvetica",colour = "black", size = 10),
        panel.grid = element_blank()); PlusOneCompositionPlot
```

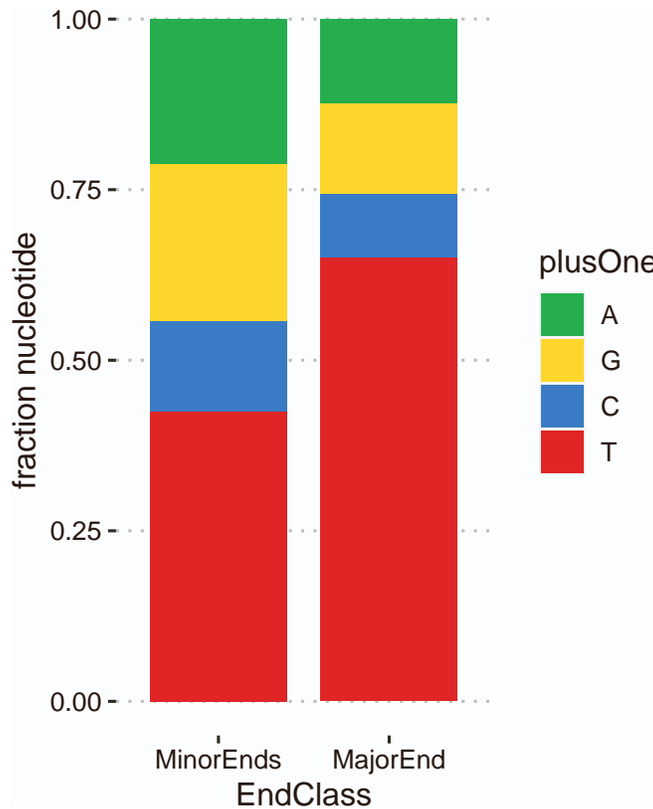


Figure 4 Heatmaps: Distribution of 3'-ends relative to 5'-ends

```
## Crate main objects

GR <- original.GR
uniq.GR <- GR[mcols(GR)[["NH"]] %in% 1]
uniq.DT <- as.data.table(uniq.GR)

## Extend the 3'-end of piRNAs so all piRNAs are 50-nt long & add sequence to the list

uniq.GRE <- uniq.GR
end(uniq.GRE[strand(uniq.GRE)=="+"]) <- start(uniq.GRE[strand(uniq.GRE)=="+"])+50
start(uniq.GRE[strand(uniq.GRE)=="-"]) <- end(uniq.GRE[strand(uniq.GRE)=="-"])-50
uniq.DT[["seq"]] <- as.vector(getSeq(BSgenome.Dmelanogaster.UCSC.dm6, uniq.GRE))

## Set analysis range, trim reads to this range & replace T's for U's
aRange <- c(18:32)
uniq.DT[,seq := substring(seq, min(aRange), max(aRange))]
uniq.DT[["seq"]] <- gsub("T", "U", uniq.DT[["seq"]])

## Create a vector of sequence contexts you wish to analyze. Must be same length as range above.
## Add any custom contexts you would like to use
## - shorthand symbols:
## N: U, C, G, or A
## V: C, G, or A (no U)
## i.e.: "NUNNNNNNNNNNNNNNN"
## This sequence will be used to select have a U 19-nt away from the 5'-end
```

```

aContexts <- unlist(lapply(c(0: (length(aRange)-1)), function(i){
  paste(c(rep("N", i), "U", rep("N", (length(aRange)-1)-i)), collapse=""))))
customContexts <- c("NNNNNNNNNNNNNNNN", "NNNNNNVVVVVVNNN", "NNNNNNVVVVVVUNN", "NNNNNUVVVVVVNNN")
aContexts <- c(aContexts, customContexts)

## Calculate size distribution for all piRNAs
## Make sure all lengths in the window of analysis are represented in the Distribution
## Normalize by converting to fraction / percentage of total

DistributionOfAll <- uniq.DT[, .(MULT = sum(MULT)), by="width"][order(width)]
if(any(!aRange %in% DistributionOfAll[["width"]])){
  DistributionOfAll <- bind_rows(DistributionOfAll,
    data.table(width = setdiff(aRange, DistributionOfAll[["width"]]),
      MULT = 0))}
DistributionOfAll[, "Percent" := (MULT/sum(MULT))*100]

## Determine the piRNA 3'-end distribution for each context

a_context <- aContexts[1]
Regex <- gsub("N", "[UCGA]", a_context)

## Loop through contexts
EndDistribution <- rbindlist(lapply(seq_along(aContexts), function(a){
  # Start and report progress
  message(paste0("a context: ",a))
  a_context <- aContexts[a]

  # Convert context to regex format and simplify
  a_regex_format <- gsub("N", "[UCGA]", a_context)
  a_regex_format <- gsub("V", "[CGA]", a_regex_format)

  # Find sequences which match the context and determine size distribution
  subset.DT <- uniq.DT[str_detect(seq, a_regex_format)][, .(MULT = sum(MULT)), by="width"]

  # Make sure all lengths in the total distribution are represented in the subset
  if(any(!DistributionOfAll[["width"]] %in% subset.DT[["width"]])){
    subset.DT <- bind_rows(subset.DT,
      data.table(width = setdiff(DistributionOfAll[["width"]],
        subset.DT[["width"]]), MULT = 0))}

  # Make sure all lengths in the window of analysis are represented in the subset
  if(any(!aRange %in% subset.DT[["width"]])){
    subset.DT <- bind_rows(subset.DT,
      data.table(width = setdiff(aRange,
        subset.DT[["width"]]), MULT = 0))}

  # order and normalize, and find deviation
  subset.DT <- subset.DT[order(width)]
  subset.DT[, "Percent" := (MULT/sum(MULT))*100]
  subset.DT[, "DeviationFromNormalDistribution" := Percent - DistributionOfAll[["Percent"]]]

```

```
# Add data identifiers
subset.DT[["ContextNumber"]] <- a
subset.DT[["Seq"]] <- strsplit(a_context, "")[[1]]
return(subset.DT )})
```

a context: 1

a context: 2

a context: 3

a context: 4

a context: 5

a context: 6

a context: 7

a context: 8

a context: 9

a context: 10

a context: 11

a context: 12

a context: 13

a context: 14

a context: 15

a context: 16

a context: 17

a context: 18

a context: 19

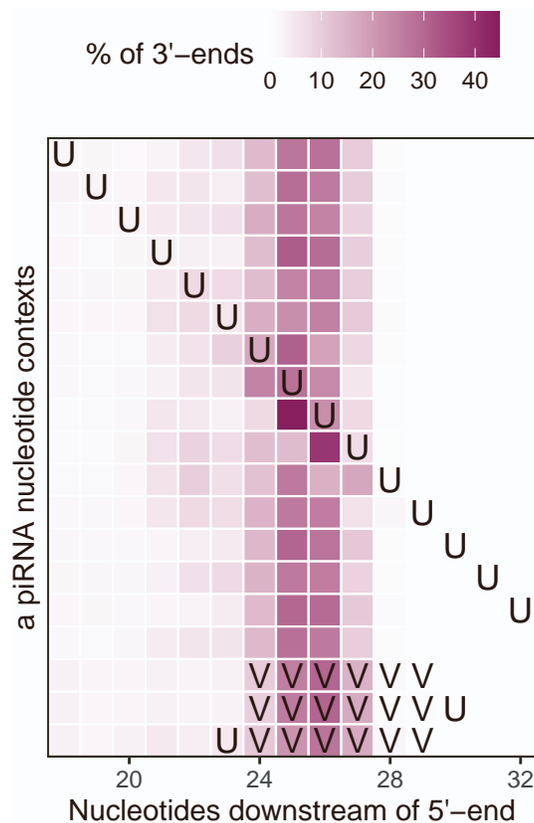
EndDistribution

```
##      width  MULT      Percent DeviationFromNormalDistribution ContextNumber Seq
##  1:    18 18915  2.059815787                0.239324650             1  U
##  2:    19 18430  2.006999998                0.354538318             1  N
##  3:    20 13728  1.494959087                -0.455241337            1  N
##  4:    21 24285  2.644600920                -1.565091321            1  N
##  5:    22 45587  4.964357510                -0.396138422            1  N
##  ---
## 281:   28  1112  1.932434311                0.875050603             19  V
## 282:   29   89  0.154664257                0.101562021             19  V
## 283:   30   10  0.017378006                0.011451863             19  N
## 284:   31    3  0.005213402                0.003082654             19  N
## 285:   32    1  0.001737801                0.001371578             19  N
```

```
## only show values within your window
EndDistribution <- EndDistribution[width %in% aRange]
```

```
##
## Plot heatmap of contexts
##

HeatmapPlot <- ggplot() + theme_bw() +
  geom_tile(data=EndDistribution,
            aes(x=width, y=ContextNumber, fill=Percent),
            color="white", size = 0.4) +
  geom_text(data=EndDistribution[Seq != "N"],
            aes(x=width, y= ContextNumber, label=Seq, size=4)) +
  scale_fill_gradient(low="white",high="maroon4",
                     limits=c(0, max(EndDistribution[["Percent"]])),
                     name = "% of 3'-ends") +
  xlab("Nucleotides downstream of 5'-end") +
  ylab("a piRNA nucleotide contexts") +
  scale_x_continuous(expand = c(0.000,0.0030)) +
  scale_size_continuous(guide="none") +
  scale_y_continuous(trans="reverse", breaks = NULL, expand = c(0,0)) +
  coord_equal() +
  theme(panel.background = element_blank(),
        panel.grid = element_blank(),
        legend.position = "top",
        axis.text.y=element_blank()); HeatmapPlot
```



```
##
## Plot Heatmap Deviation
##

HeatmapDeviationPlot <- ggplot() + theme_bw() +
  geom_tile(data=EndDistribution,
            aes(x=width, y=ContextNumber, fill=DeviationFromNormalDistribution),
            color="white", size = 0.4) +
  geom_text(data=EndDistribution[Seq != "N"],
            aes(x=width, y= ContextNumber, label=Seq, size=4))+
  scale_fill_gradient2(low="navy", mid="white", high="firebrick4",
                       limits = c(min(EndDistribution[["DeviationFromNormalDistribution"]]),
                                   max(EndDistribution[["DeviationFromNormalDistribution"]])) +
  xlab("Nucleotides downstream of 5'-end") +
  ylab("a piRNA nucleotide contexts") +
  scale_x_continuous(expand = c(0.000,0.0030))+
  scale_size_continuous(guide= "none")+
  scale_y_continuous(trans="reverse", breaks = NULL, expand = c(0,0)) +
  coord_equal() +
  theme(panel.background = element_blank(),
        panel.grid = element_blank(),
        legend.position = "top",
        axis.text.y=element_blank()); HeatmapDeviationPlot
```

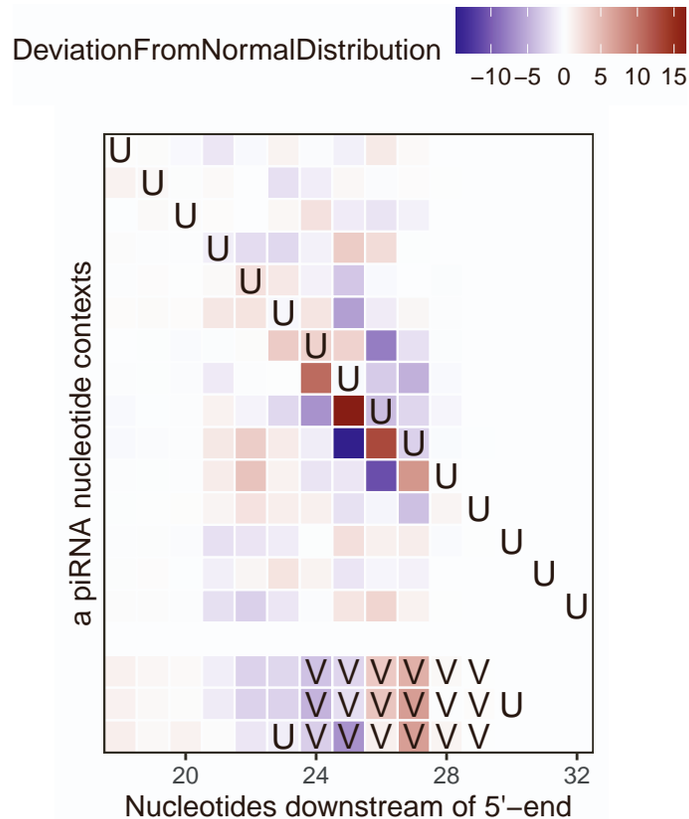


Figure 4 Z-score: Distribution of 3'-end relative to sequence context of interest

```
## Crate main objects

GR <- original.GR
uniq.GR <- GR[mcols(GR)[["NH"]] %in% 1]

## Extend both ends of piRNAs to see sequence immediately down- and up-stream

ExpandBy <- 60
end(uniq.GR) <- end(uniq.GR)+ExpandBy
start(uniq.GR) <- start(uniq.GR)-ExpandBy

## Make sure that the extended sequences are still within the boundaries of chromosomes
## Use the appropriate genome
RefGen <- BSgenome::getBSgenome("BSgenome.Dmelanogaster.UCSC.dm6")
RefGen <- GRanges(seqnames = names(RefGen),
                  ranges = IRanges(start = 1,
                                   end = GenomeInfoDb::seqlengths(RefGen)))
uniq.GR <- IRanges::subsetByOverlaps(uniq.GR, RefGen, type = "within")

## Convert to data.table and obtain sequence

uniq.DT <- as.data.table(uniq.GR)
uniq.DT[["seq"]] <- as.vector(getSeq(BSgenome.Dmelanogaster.UCSC.dm6, uniq.GR))

## Select the range of analysis, must be less than what the original sequence were expanded by (60)
```

```

Range <- 51

## Select sequence context to analyze
## Add any custom contexts you would like to use
## - use shorthands such as V = C, G, or A

Context <- "VVUVV"

## Make sequence context same length as the Range by adding N's
Context <- paste(c(paste(rep("N", floor((Range-nchar(Context))/2)), collapse=""), Context, paste(rep("N",
Context

```

```
## [1] "NNNNNNNNNNNNNNNNNNNNNNNNNNNNNNVVUVVNNNNNNNNNNNNNNNNNNNNNNNNNNNNNN"
```

```

## Convert Context to R regex format
Regex <- gsub("N", "[UCGA]", Context)
Regex <- gsub("V", "[CGA]", Regex)

## Convert to DNA
Regex <- gsub("U", "T", Regex)

## Create vector of positions for analysis
PosVector <- c(-((Range-1)/2):((Range-1)/2))

## Loop through the searching of the Context motif
## Use parallel package to speed up the Calculations
NumberOfCores <- 2

MotifSearch <- bind_rows(mclapply(seq_along(PosVector), function(a){

  Position <- PosVector[a]
  MaxWindow <- max(PosVector)

  ## Calculate total number of 3'-ends at (PosVector[a]) nucleotides away from center of context
  ThreePrimeSum <- sum(uniq.DT[str_detect(substring(seq,
                                                    (width-ExpandBy)-(a+MaxWindow),
                                                    (((width-ExpandBy)-(a+MaxWindow))+(Range-1))), Regex)]["MU

  ## prepare output
  Output <- data.table(
    Position = Position,
    Nucleotide = substring(Context, a, a),
    ThreePrime = ThreePrimeSum
  )

  return(Output)
}, mc.cores = NumberOfCores))

MotifSearch[20:35]

```

```
##      Position Nucleotide ThreePrime
```

```
## 1:      -6      N      182832
## 2:      -5      N      185349
## 3:      -4      N      148123
## 4:      -3      N      132726
## 5:      -2      V      208009
## 6:      -1      V      206033
## 7:       0      U      131668
## 8:       1      V      133841
## 9:       2      V      174869
## 10:     3      N      203119
## 11:     4      N      165821
## 12:     5      N      174248
## 13:     6      N      165076
## 14:     7      N      187432
## 15:     8      N      174311
## 16:     9      N      168634
```

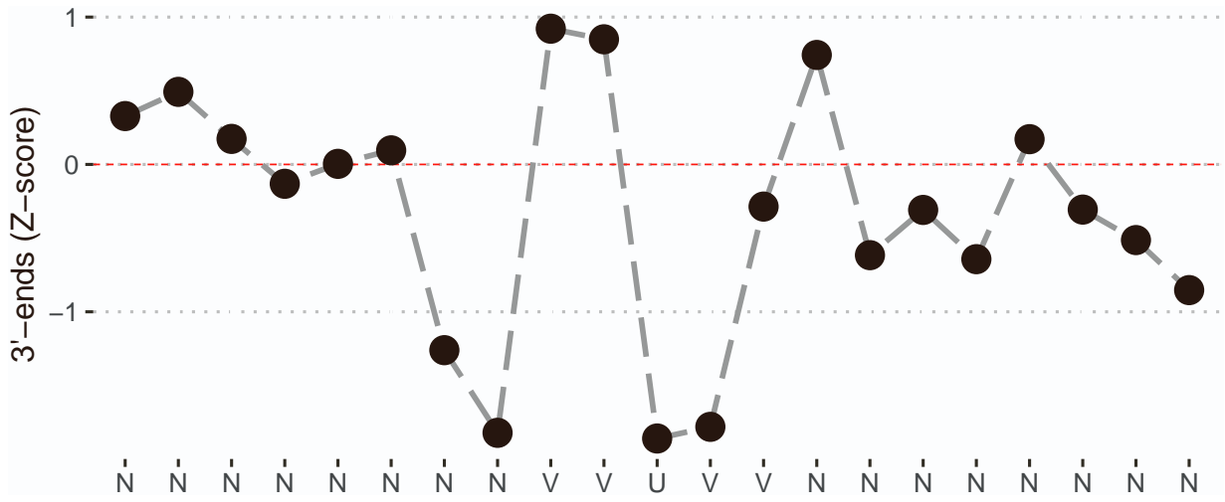
```
## Convert sum of 3'-ends at each position into a Z-score
MotifSearch[["Zscore"]] <- (MotifSearch[["ThreePrime"]] - mean(MotifSearch[["ThreePrime"]])) / sd(MotifSearch[["ThreePrime"]])

## Only depict central 21 nucleotides
MotifSearchGraph <- MotifSearch[Position %in% c(-10:10)]
```

```
## Organize data
MotifSearchGraph[["Position"]] <- factor(MotifSearchGraph[["Position"]])

## Plot the Z-score graph

ZscorePlot <- ggplot() + theme_pubclean() +
  geom_hline(yintercept = 0, linetype="dashed", color = "red", lwd = 0.3) +
  geom_line(data=MotifSearchGraph,
            aes(x=Position, y=Zscore, group = 1),
            color = "gray60", size = 1, linetype = "longdash") +
  geom_point(data=MotifSearchGraph,
             aes(x=Position, y=Zscore),
             shape=16, size=5, fill = "gray60", color = "black") +
  labs(y= "3'-ends (Z-score)", x="")+
  scale_x_discrete(labels = MotifSearchGraph[["Nucleotide"]]) +
  theme(aspect.ratio = 0.4);ZscorePlot
```



•
•
•

This concludes the methods.

THE END.