Supplementary Experimental Data File

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Arrangement of CLASH dataset

CLASH dataset was retrieved from PubMed (Helwak et al. 2013).

Query of Human Genome 19.

Human genome 19 information was handled through biomaRt package.

Adding miRNA and gene information

MiRNA releases are obtained from miRBase. In this step, release 21 (in Human genome 38) was downloaded.

```
read.table("mirbasehg38.txt", comment.char = "#")%>%
  filter(V3 != "miRNA_primary_transcript")%>%
  separate(V9, c("ID", "Alias", "Name", "Precusor"), sep = ";")%>%
  mutate(ID = substr(ID, 4, length(ID)),
```

CLASH dataset is published in miRBase release 15 and Human Genome 19 version.

```
read_tsv("mirna_mature.txt", col_names = FALSE)%>%
  filter(startsWith(X2, "hsa"))%>%
  dplyr::select(mirna_ID = X2, mirbase_ID = X3)%>%
  inner join(mirbasehg38%>% dplyr::select(ID, Name),
             by= c("mirbase_ID"= "ID"))%>%
  dplyr::select(mirbase_ID, Name)%>%
  distinct()%>%
  inner_join(clashelwak, by = c("mirbase_ID"= "Barcode"))%>%
  dplyr::select(Name, miRNA_seq, Ensembl_Gene_Id,
                Ensembl Transcript Id, Hugo Symbol,
                mRNA seg extended, chimeras decompressed,
                seed_type, seed_basepairs,folding_class,
                seq_ID, folding_energy,
                X5.UTR, CDS, X3.UTR)%>%
  inner_join(hg19, by = c("Ensembl_Gene_Id"= "ensembl_gene_id",
                          "Ensembl_Transcript_Id"="ensembl_transcript_id",
                          "Hugo_Symbol"="hgnc_symbol"))%>%
  mutate(region1 = ifelse(X5.UTR == "1", "5UTR", " "),
         region2= ifelse(X3.UTR == "1", "3UTR", " "),
         region3= ifelse(CDS =="1", "CDS", " "))%>%
  unite(region, c(region1, region2, region3), sep = "||")%>%
  dplyr::select(chromosome_name, start_position,
                end_position, strand, Hugo_Symbol,
                Ensembl_Gene_Id, Ensembl_Transcript_Id,
                mRNA_seq_extended, Name, miRNA_seq,
                seq_ID, seed_type,seed_basepairs,
                folding_class, folding_energy, region)%>%
  as tibble() -> clashelwakfinal
```

Converting CLASH data to human genome 38 build.

There are different liftover methods for conversion among Human Genome builds. We preffered to use UCSC liftover tool

```
#Obtaining chromosomal locations from miRNA:target interaction dataset.

lift19 <- clashelwakfinal%>%
    dplyr::select(1,2,3)%>%
    unite(start_end, c("start_position", "end_position"), sep = "-")%>%
    mutate(Chromosome = pasteO("chr",chromosome_name, ""))%>%
    unite(chromosome_name, c("Chromosome","start_end"), sep = ":")

write_tsv(lift19, "lift19.txt")
```

```
#After we searched this file in UCSC browser, the output loaded (lift19_del deleted regions on the HG38
lift19_del <- read_tsv("deleted_lift19.txt")</pre>
colnames(lift19_del)[1] <- "chromosome_loc"</pre>
lift19_del <- lift19_del%>%
  dplyr::filter(startsWith(chromosome_loc, "chr"))%>%
  separate(chromosome_loc, c("Chr", "End"), "-", remove = TRUE)%>%
  separate(Chr, c("Chr", "Start"), ":", remove = TRUE)
lift19_del$Start <- as.numeric(lift19_del$Start)</pre>
lift19_del$End <- as.numeric(lift19_del$End)</pre>
#removing deleted location from CLASH dataset
clashelwakfinal <-clashelwakfinal%>%
  mutate(Chromosome = paste0("chr",chromosome_name, ""))%>%
  dplyr::anti_join(lift19_del, by = c("Chromosome"="Chr",
                                       "start_position"="Start",
                                       "end_position"="End"))
hg38clash <- read.delim("hg38clashcomp.txt", header = FALSE, stringsAsFactors = FALSE)
# adding new location information:
clashelwakfinal <- clashelwakfinal%>%
  bind_cols(hg38clash)
colnames(clashelwakfinal)[18] <- "HG38build_loc"</pre>
clashelwakfinal <- clashelwakfinal%>%
  dplyr::mutate(Genom_build= rep("hg19"))
# Arrangement in dataset
clashelwakfinal%>% dplyr::select(cluster=seq_ID, chromosome = Chromosome,
                                  start_position, end_position,
                                  strand, hgnc_symbol = Hugo_Symbol,
                                  Ensembl_Gene_Id, Ensembl_Transcript_Id,
                                  target_seq = mRNA_seq_extended, miRNA= Name,
                                  miR_seq =miRNA_seq, seed_type,
                                  seed_type2= seed_basepairs,
                                  seed_type3= folding_class,
                                  Energy = folding_energy,
                                  HG38build_loc,
                                  Genom_build, region) -> clashelwakfinal
clashelwakfinal$strand <- as.character(clashelwakfinal$strand)</pre>
str(clashelwakfinal)
```

Interpreting the CLASH seed structures in dataset

Arrangement of CLEAR-CLiP Dataset (Moore et al. 2015)

CLASH dataset was retrieved from Nature article

```
clearclip <- read_xlsx("CLEAR-CLIP.xlsx")
#Clearclip hg18</pre>
```

Query of Human Genome 18

Adding Genome Information to dataset

Converting human genome build

```
#Obtaining chromosomal locations from miRNA:target interaction dataset.
lift18 <- clearclipfinal%>%
  unite(start end, c("start position", "end position"), sep = "-")%>%
  unite(location, c("chr", "start_end"), sep = ":")%>%
 dplyr::select(location)
write tsv(lift18, "lift18.txt")
#After we searched this file in UCSC browser, the output loaded (deleted_lift18 deleted regions on the
deleted_lift18 <- read_tsv("deleted_lift18.txt")</pre>
colnames(deleted_lift18)[1] <- "Chromosome_loc"</pre>
deleted_lift18 <- deleted_lift18%>%
  dplyr::filter(startsWith(Chromosome_loc, "chr"))%>%
  separate(Chromosome_loc, c("Chr", "End"), "-", remove = TRUE)%>%
  separate(Chr, c("Chr", "Start"), ":", remove = TRUE)
deleted_lift18$Start <- as.numeric(deleted_lift18$Start)</pre>
deleted_lift18$End <- as.numeric(deleted_lift18$End)</pre>
#removing deleted location from CLEAR-CLiP dataset
clearclipfinal <- clearclipfinal%>%
 dplyr::anti_join(deleted_lift18, by = c("chr"="Chr",
                                           "start_position"="Start",
                                           "end_position"="End"))
hg38clearclip<- read.delim("hg38clearclip.txt", header = FALSE,
                           stringsAsFactors = FALSE)
clearclipfinal <- clearclipfinal%>%
 bind_cols(hg38clearclip)
colnames(clearclipfinal)[28] <- "HG38build loc"</pre>
# adding new location information:
clearclipfinal <- clearclipfinal%>%
 dplyr::mutate(Genom_build= rep("hg18"))
# Arrangement in dataset
clearclipfinal%>%
  dplyr::select(cluster=cluster.ID, chromosome = chr, start_position,
                end_position, strand = strand.y, hgnc_symbol, Ensembl_Gene_Id =ensembl_gene_id,
                Ensembl_Transcript_Id =ensembl_transcript_id, target_seq = target.map,
                miRNA, miR_seq = miR.map, seed_type = 'seed match', Energy = MFE,
                HG38build_loc, Genom_build, region) -> clearclipfinal
```

Seed type manipulation in CLEAR-CLiP dataset

In CLEAR-CLiP dataset, seed types were shown in detail. We adjusted as canonical and non-canonical.

Integration of two experimental dataset

```
bind_rows(clashelwakfinal, clearclipfinal)%>%distinct() -> experimentalmirnagene
```

Adding Coefficients of Interaction factors

Energy values in miRNA:target pairs are represented by high-throughput studies (Helwak et al. 2013; Moore et al. 2015) which are utilized in this study. On the other hand, we have specified the other interaction factors, seed type and location of binding region on the target, as numeric values based on the previous studies. Grimson et al. (2007) have compared the effect of seed types on target repression with few miRNA had canonical seed pairing in their study. Additionally, Bartel et al. (2009) and Betel et al. (2010) have studied on functional and non-functional seed interactions. Based on results of these studies we have arranged seed types of miRNA:target interactions as numeric values. We also have redefined location of binding region on the target as numeric values, based on studies of Hausser et al. (2013) and Helwak et al. (2013). With this process, we have handled this integrated dataset in context of competitor behaviors and functionality of interactions.

In this step we added numeric intraction values at followings

Fistly, we organized these values due to the fact that the regions were defined differently in two datasets. After that, region effect was added as numeric values (shown in Table S3).

```
"0.01", ifelse(region %in% c("5UTRCDS", "CDS5UTR"),
"0.42", ifelse(region %in% c("intron",""),
"0.01", ifelse(region %in% c("exon_unclassified", ""), "0.2", NA)))))
```

Secondly, we organized seed type interactions in *Seed type manipulation* section for CLEAR-CLiP dataset to show as found in CLASH dataset. Same type formatted values added dataset as numeric values (shown in Table S2).

```
## Saving dataset
saveRDS(experimentalmirnagene, "data/experimentalmirnagene.RDS")
```

```
readRDS("data/experimentalmirnagene.RDS")->experimentalmirnagene
experimentalmirnagene
```

```
## # A tibble: 45,340 x 18
      cluster chromosome start_position end_position strand hgnc_symbol
##
                                                            <chr>>
##
      <chr> <chr>
                                  <int>
                                               <int> <chr>
## 1 0727A-... chr5
                                162864575
                                             162873157 1
                                                              CCNG1
## 2 L1HS-1... chr14
                                                              DICER1
                                 95552565
                                              95624347 -1
## 3 L2HS-8... chr6
                                109307640
                                             109416022 -1
                                                              SESN1
## 4 L2HS-1... chr5
                                              37066515 1
                                                              NIPBL
                                 36876861
## 5 L2-407... chr4
                                106603784
                                             106817143 -1
                                                              INTS12
## 6 L1HS-7... chr5
                                130977407
                                             131132710 -1
                                                              FNIP1
                                134123389
## 7 L1HS-4... chr11
                                             134135749 1
                                                              ACAD8
## 8 0727A-... chr15
                                                              CCNB2
                                59397277
                                              59417244 1
## 9 L2HS-1... chr19
                                 37001597
                                              37019562 -1
                                                              ZNF260
## 10 L2HS-9... chr11
                                              64902004 -1
                                                              SYVN1
                                 64889252
## # ... with 45,330 more rows, and 12 more variables: Ensembl_Gene_Id <chr>,
      Ensembl_Transcript_Id <chr>, target_seq <chr>, miRNA <chr>, miR_seq <chr>,
       seed_type <chr>, Energy <dbl>, HG38build_loc <chr>, Genom_build <chr>,
## #
       region <chr>, region_effect <dbl>, seed_type_effect <dbl>
```

The context of dataset is shown in Table S5 in Supplementary Tables.

REFERENCES

Bartel, David P. 2009. "MicroRNAs: Target Recognition and Regulatory Functions." Cell 136 (2): 215–33. https://doi.org/10.1016/j.cell.2009.01.002.

Betel, Doron, Anjali Koppal, Phaedra Agius, Chris Sander, and Christina Leslie. 2010. "Comprehensive Modeling of microRNA Targets Predicts Functional Non-Conserved and Non-Canonical Sites." *Genome Biology* 11 (8): R90.

Grimson, Andrew, Kyle Kai-How Farh, Wendy K. Johnston, Philip Garrett-Engele, Lee P. Lim, and David P. Bartel. 2007. "MicroRNA Targeting Specificity in Mammals: Determinants Beyond Seed Pairing." *Molecular Cell* 27 (1): 91–105. https://doi.org/10.1016/j.molcel.2007.06.017.

Hausser, J., A. P. Syed, B. Bilen, and M. Zavolan. 2013. "Analysis of CDS-Located miRNA Target Sites Suggests That They Can Effectively Inhibit Translation." *Genome Research* 23 (4): 604–15. https://doi.org/10.1101/gr.139758.112.

Helwak, Aleksandra, Grzegorz Kudla, Tatiana Dudnakova, and David Tollervey. 2013. "Mapping the Human miRNA Interactome by CLASH Reveals Frequent Noncanonical Binding." Cell 153 (3): 654–65. https://doi.org/10.1016/j.cell.2013.03.043.

Moore, Michael J., Troels K. H. Scheel, Joseph M. Luna, Christopher Y. Park, John J. Fak, Eiko Nishiuchi, Charles M. Rice, and Robert B. Darnell. 2015. "miRNA-Target Chimeras Reveal miRNA 3'-End Pairing as a Major Determinant of Argonaute Target Specificity." *Nature Communications* 6 (November): 8864. https://doi.org/10.1038/ncomms9864.