## Patient FLC and FLC Sphereoid miR correlation

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1/27/2021

```
library(dplyr)
library(ggpubr)
library(tibble)
```

#### Load CSV

```
flc.counts <- read.csv("../NML-PRI-MET_noDups_avg-normalized-counts-by-group.csv")
flc.sphereoids <- read.csv("FLC-PDX_DESeq_miRs.csv")
flc.pri <- flc.counts %>%
    select(miR, avg_PRI)
flc.mets <- flc.counts %>%
    select(miR, avg_MET)
```

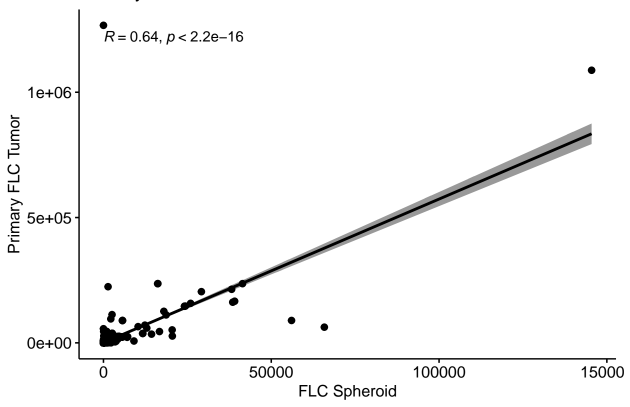
#### Workflow with primary patient FLC primary tumors

```
#Join intracellular sphereoids miRs with primary tumor miRs
flc.combined.deseq <- full_join(flc.sphereoids, flc.pri, by = "miR")
#change is.na rows to 0
flc.combined.deseq[is.na(flc.combined.deseq)] <- 0
#Move the column to rowname. this is necessary for the final clean up step of removing rows that contai
flc.combined.deseq <- column_to_rownames(flc.combined.deseq, var = "miR")
flc.clean.100 <- flc.combined.deseq %>%
    filter_all(any_vars(. > 100))
#this is done again with reads at least 50 reads in either column
flc.clean.50 <- flc.combined.deseq %>%
    filter_all(any_vars(. > 50))
```

Plots for primary FLC primary tumors with and without log10 transformation

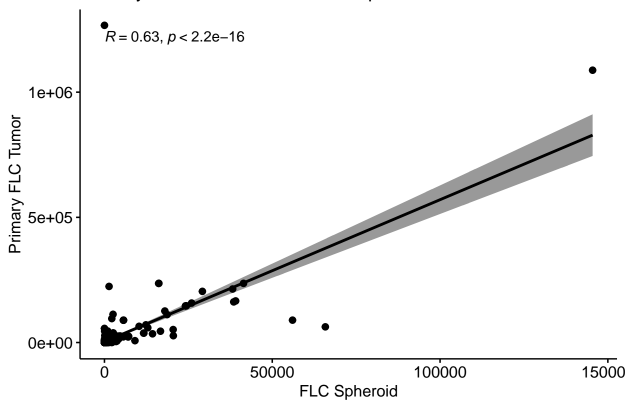
```
add = "reg.line", conf.int = TRUE,
          cor.coef = TRUE, cor.method = "pearson",
          xlab = "FLC Spheroid", ylab = "Primary FLC Tumor", title = "Primary FLC tumor miRs vs FLC Sph
p.pri.100.log10 <- ggscatter(flc.clean.100, x = "avg_Intra_FLC", y = "avg_PRI",
          add = "reg.line", conf.int = TRUE,
          cor.coef = TRUE, cor.method = "pearson",
          xlab = "log10 FLC Spheroid", ylab = "log 10 Primary FLC", title = "log10(Primary FLC tumor mi)
  scale_x_log10() +
  scale_y_log10()
p.pri.50 <- ggscatter(flc.clean.50, x = "avg_Intra_FLC", y = "avg_PRI",</pre>
          add = "reg.line", conf.int = TRUE,
          cor.coef = TRUE, cor.method = "pearson",
          xlab = "FLC Spheroid", ylab = "Primary FLC Tumor", title = "Primary FLC tumor miRs vs FLC Sph
p.pri.50.log10 <- ggscatter(flc.clean.50, x = "avg_Intra_FLC", y = "avg_PRI",
          add = "reg.line", conf.int = TRUE,
          cor.coef = TRUE, cor.method = "pearson",
          xlab = "log10 FLC Spheroid", ylab = "log 10 Primary FLC Tumor", title = "log10(Primary FLC tumor")
  scale_x_log10() +
  scale_y_log10()
p.pri.noclean
```

# Primary FLC tumor miRS vs FLC-PDX miRs



p.pri.100

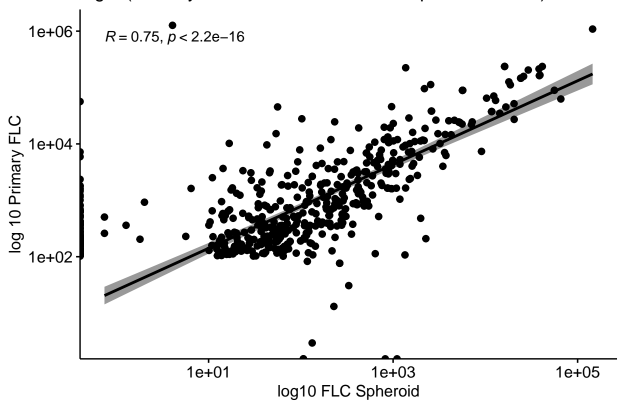
### Primary FLC tumor miRs vs FLC Sphereoid miRs - 100+ reads in



### p.pri.100.log10

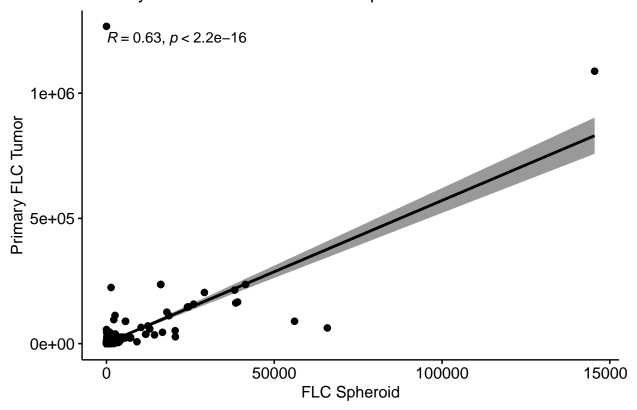
## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## 'geom\_smooth()' using formula 'y ~ x'
## Warning: Removed 119 rows containing non-finite values (stat\_smooth).
## Warning: Removed 119 rows containing non-finite values (stat\_cor).

log10(Primary FLC tumor miRs vs FLC Sphereoid miRs) 100+ rea



p.pri.50

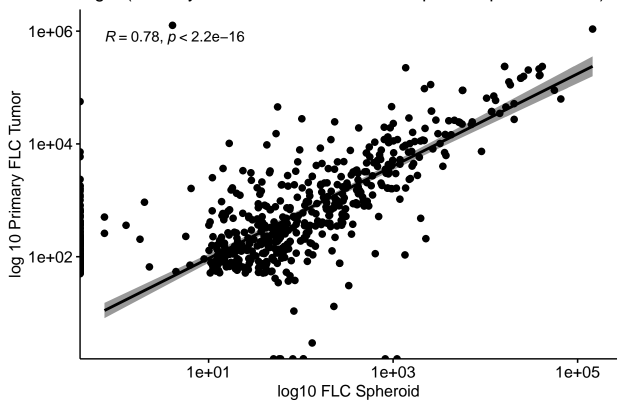
### Primary FLC tumor miRs vs FLC Sphereoid miRs - 50+ reads in a



### p.pri.50.log10

## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## 'geom\_smooth()' using formula 'y ~ x'
## Warning: Removed 213 rows containing non-finite values (stat\_smooth).
## Warning: Removed 213 rows containing non-finite values (stat\_cor).

### log10(Primary FLC tumor miRs vs DESeq\_FLC Spheroid miRs) 50

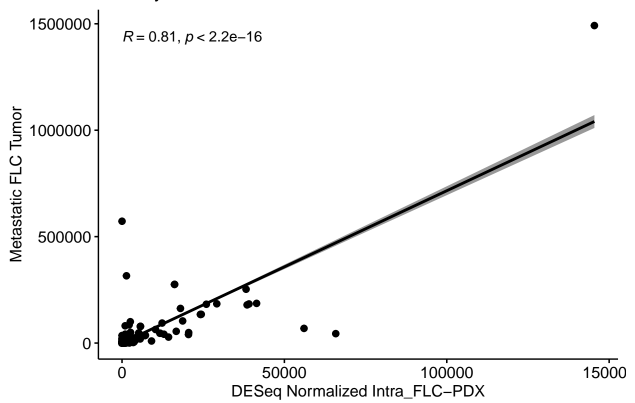


Workflow with primary patient FLC metastatic tumors

```
#The steps here are similar to that seen in code chunk 4
flc.combined.mets <- full_join(flc.mets, flc.sphereoids, by = "miR")</pre>
flc.combined.mets[is.na(flc.combined.mets)] <- 0</pre>
flc.combined.mets <- column_to_rownames(flc.combined.mets, var = "miR")
flc.clean.mets.100 <- flc.combined.mets %>%
  filter_all(any_vars(. > 100))
flc.clean.mets.50 <- flc.combined.mets %>%
  filter_all(any_vars(. > 50))
p.met.noclean <- ggscatter(flc.combined.mets, x = "avg_Intra_FLC", y = "avg_MET",</pre>
          add = "reg.line", conf.int = TRUE,
          cor.coef = TRUE, cor.method = "pearson",
          xlab = "DESeq Normalized Intra_FLC-PDX", ylab = "Metastatic FLC Tumor", title = "Primary FLC
p.met.100 <- ggscatter(flc.clean.mets.100, x = "avg_Intra_FLC", y = "avg_MET",
          add = "reg.line", conf.int = TRUE,
          cor.coef = TRUE, cor.method = "pearson",
          xlab = "FLC Spheroid", ylab = "Metastatic FLC Tumor", title = "Metastatic FLC tumor miRs vs F
p.met.100.log10 <- ggscatter(flc.clean.mets.100, x = "avg_Intra_FLC", y = "avg_MET",
          add = "reg.line", conf.int = TRUE,
          cor.coef = TRUE, cor.method = "pearson",
```

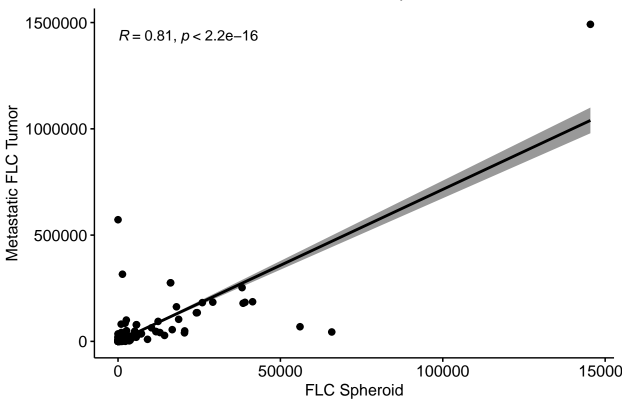
## 'geom\_smooth()' using formula 'y ~ x'

### Primary FLC tumor miRS vs FLC miRs



```
p.met.100
```

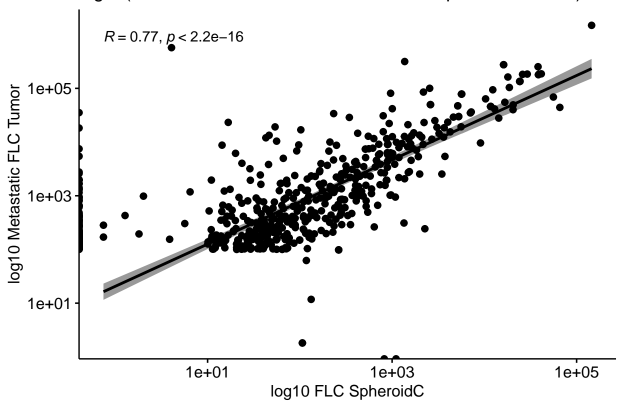




### p.met.100.log10

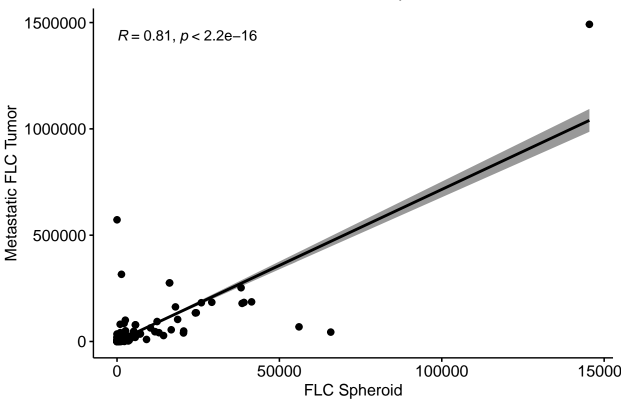
## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## 'geom\_smooth()' using formula 'y ~ x'
## Warning: Removed 120 rows containing non-finite values (stat\_smooth).
## Warning: Removed 120 rows containing non-finite values (stat\_cor).

log10(Metastatic FLC tumor miRs vs \_FLC sphereoids miRs) 100+



p.met.50





### p.met.50.log10

## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## Warning: Transformation introduced infinite values in continuous x-axis
## Warning: Transformation introduced infinite values in continuous y-axis
## 'geom\_smooth()' using formula 'y ~ x'
## Warning: Removed 203 rows containing non-finite values (stat\_smooth).
## Warning: Removed 203 rows containing non-finite values (stat\_cor).



