

Projekt zaliczeniowy

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1. Ładowanie bibliotek

```
library(DESeq2)
library(ggplot2)
library(ComplexHeatmap)
```

2. Informacje o danych bioprojektu (PRJNA313294)

```
SRAruns <- c("SRR3191542", "SRR3191543", "SRR3191544", "SRR3191545", "SRR3194428", "SRR3194429", "SRR3194430", "SRR3194431")
SRAitems <- c("Mock1-1", "Mock2-1", "ZIKV1-1", "ZIKV2-1", "Mock1-2", "Mock2-2", "ZIKV1-2", "ZIKV2-2")
SRAdetails <- c(rep("Illumina MiSeq", times = 4), rep("Illumina NextSeq 500", times = 4))
SRAsamples <- c("Mock_1_MiSeq", "Mock_2_MiSeq", "ZIKA_1_MiSeq", "ZIKA_2_MiSeq", "Mock_1_NextSeq", "Mock_2_NextSeq", "ZIKA_1_NextSeq", "ZIKA_2_NextSeq")
```

```
SRAdata <- rbind(SRAruns, SRAitems, SRAdetails, SRAsamples)
print(SRAdata)
```

```
##           [,1]           [,2]           [,3]
## SRAruns      "SRR3191542"      "SRR3191543"      "SRR3191544"
## SRAitems      "Mock1-1"         "Mock2-1"         "ZIKV1-1"
## SRAdetails    "Illumina MiSeq"  "Illumina MiSeq"  "Illumina MiSeq"
## SRAsamples    "Mock_1_MiSeq"    "Mock_2_MiSeq"    "ZIKA_1_MiSeq"
##           [,4]           [,5]           [,6]
## SRAruns      "SRR3191545"      "SRR3194428"      "SRR3194429"
## SRAitems      "ZIKV2-1"         "Mock1-2"         "Mock2-2"
## SRAdetails    "Illumina MiSeq"  "Illumina NextSeq 500" "Illumina NextSeq 500"
## SRAsamples    "ZIKA_2_MiSeq"    "Mock_1_NextSeq"  "Mock_2_NextSeq"
##           [,7]           [,8]
## SRAruns      "SRR3194430"      "SRR3194431"
## SRAitems      "ZIKV1-2"         "ZIKV2-2"
## SRAdetails    "Illumina NextSeq 500" "Illumina NextSeq 500"
## SRAsamples    "ZIKA_1_NextSeq"  "ZIKA_2_NextSeq"
```

3. Wczytanie danych zmapowanych do genomu hg19

```
counts_all <- read.delim("~/projekt/analizaTranskryptomu/projekt/hg19/COUNTS/counts_ALL.txt", comment.char="#")
colnames(counts_all)[7:14] <- SRAsamples

counts_paired <- read.delim("~/projekt/analizaTranskryptomu/projekt/hg19/COUNTS/counts_PE.txt", comment.char="#")
colnames(counts_paired)[7:10] <- SRAsamples[1:4]
```

```
counts_single <- read.delim("~/projekt/analizaTranskryptomu/projekt/hg19/COUNTS/counts_SE.txt", comment = "#", as.is = TRUE)
colnames(counts_single)[7:10] <- SRAShortcuts[5:8]
```

4. Analiza DE

```
dds <- function(data, runs) {
  countData <- data[,7:(7+runs-1)]
  rownames(countData) = data$Geneid
  samples <- names(countData)
  if(runs > 4) {
    condition <- factor(c("mock", "mock", "zika", "zika", "mock", "mock", "zika", "zika"))
  }
  else {
    condition <- factor(c("mock", "mock", "zika", "zika"))
  }
  colData <- data.frame(samples = samples, condition = condition)
  dds <- DESeqDataSetFromMatrix(countData = countData, colData = colData, design = ~condition)

  return(dds)
}

analyseDE = function(data) {
  dds <- DESeq(data)
  res <- results(dds)

  r = res[res$baseMean!=0,]
  r = r[r$log2FoldChange > 1 | r$log2FoldChange < -1,]
  x = !is.na(r$padj)
  r = r[x,]
  r = r[r$padj<0.05,]

  print(head(r))

  return(dds)
}
```

4.1. Zbiorecz

```
DE_all <- analyseDE(dds(counts_all, 8))
```

```
## log2 fold change (MLE): condition zika vs mock
## Wald test p-value: condition zika vs mock
## DataFrame with 6 rows and 6 columns
##           baseMean  log2FoldChange      lfcSE
##           <numeric>      <numeric>      <numeric>
## VWA1          334.426316217386 -1.32578817670352 0.0969376879419521
## MMP23B         64.2662818719584 -1.20821536435696 0.256532925580776
## CFAP74         29.9194006545816 -1.81721702770035 0.335123059831466
## LOC100129534   48.4952917103514 -1.24258583913354 0.248226437135996
## TNFRSF14-AS1  50.0132552352106  5.55556197239441 0.508976370901608
## TNFRSF14       39.5661970198706  3.71195816205217 0.369528973318885
```

	stat	pvalue	padj
	<numeric>	<numeric>	<numeric>
VWA1	-13.6767051582396	1.39889139887406e-42	1.04063200715454e-40
MMP23B	-4.70978671303745	2.47976153514423e-06	1.20388217385792e-05
CFAP74	-5.4225365112572	5.87592075449711e-08	3.60238555016946e-07
L0C100129534	-5.0058561588779	5.56142771889026e-07	2.96677517640969e-06
TNFRSF14-AS1	10.9151667739569	9.75500944865484e-28	3.30917241530221e-26
TNFRSF14	10.0451072312777	9.65426209400259e-24	2.47513273088438e-22

4.2. Dla każdego z urządzeń

Illumina MiSeq

```
DE_paired <- analyseDE(dds(counts_paired, 4))
```

```
## log2 fold change (MLE): condition zika vs mock
## Wald test p-value: condition zika vs mock
## DataFrame with 6 rows and 6 columns
```

	baseMean	log2FoldChange	lfcSE
	<numeric>	<numeric>	<numeric>
VWA1	75.5398925477733	-1.27710020252877	0.310291019365312
TNFRSF14-AS1	15.7621492406834	5.98721330457336	1.54916859724598
TNFRSF14	10.0893229839815	4.30846211355709	1.26716465965906
MEGF6	417.300756458076	-1.197162158804	0.14170808792281
LNCTAM34A	18.4667693307206	-1.60355478272455	0.635892821555479
TMEM51	76.7978830409192	-1.08678049119696	0.30771355360715

```
##
```

	stat	pvalue	padj
	<numeric>	<numeric>	<numeric>
VWA1	-4.11581426088652	3.8581491608349e-05	0.000309091607984836
TNFRSF14-AS1	3.86479129206278	0.000111184229442529	0.000783191516013005
TNFRSF14	3.4000807083085	0.000673659657991891	0.00373635297489782
MEGF6	-8.44808631851775	2.96116419348808e-17	1.55932466518511e-15
LNCTAM34A	-2.52173751356721	0.0116776810442466	0.0415245815584491
TMEM51	-3.53179272884556	0.000412752741205068	0.00243836283249716

Illumina NextSeq 500

```
DE_single <- analyseDE(dds(counts_single, 4))
```

```
## log2 fold change (MLE): condition zika vs mock
## Wald test p-value: condition zika vs mock
## DataFrame with 6 rows and 6 columns
```

	baseMean	log2FoldChange	lfcSE
	<numeric>	<numeric>	<numeric>
LINC02593	155.3179313799	-1.0667898894932	0.224527222396563
VWA1	750.709188115296	-1.35230758965782	0.110248830255318
MMP23B	114.073377704044	-1.16619139667573	0.261206019938957
CFAP74	67.2196258284694	-2.00458276889009	0.366954567515667
GABRD	26.0893918827756	-1.82675486692474	0.57611650293165
L0C100129534	111.017280389548	-1.29314098690466	0.273434690240048

```
##
```

	stat	pvalue	padj
	<numeric>	<numeric>	<numeric>
LINC02593	-4.75127193088872	2.02141068919781e-06	7.35889188230861e-06
VWA1	-12.2659586185731	1.37969709828036e-34	3.22381776525957e-33

```
## MMP23B      -4.46464211256793 8.02027001183273e-06 2.73707395838519e-05
## CFAP74      -5.46275464688011 4.68802339767858e-08 2.02722708021764e-07
## GABRD       -3.17080808765075 0.00152015537041154 0.00383060370870692
## LOC100129534 -4.7292499198599 2.25350844351798e-06 8.15950963445154e-06
```

5. Analiza PCA

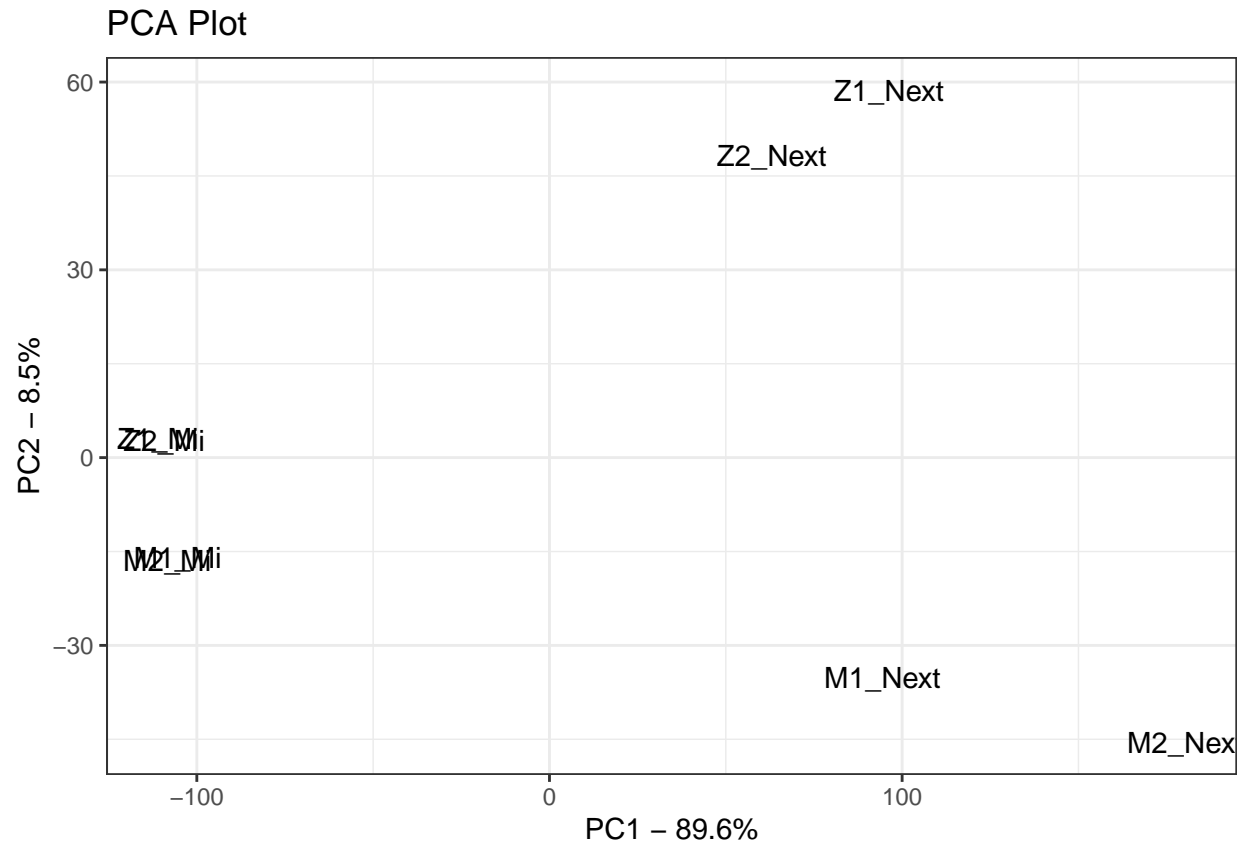
```
PCAnalysis <- function(data, runs) {
  gene_data <- data[7:14]
  rownames(gene_data) <- data$Geneid

  row_sub <- apply(gene_data, 1, function(row) all(row != 0))
  gene_data <- gene_data[row_sub,]
  gene_data_matrix <- as.matrix(gene_data)

  pca <- prcomp(t(gene_data_matrix), scale = T)
  pca.data <- data.frame(Sample = rownames(pca$x), X = pca$x[,1], Y = pca$x[,2])
  pca.var <- pca$sdev^2
  pca.var.per <- round(pca.var/sum(pca.var)*100, 1)

  ggplot(data = pca.data, aes(x = X, y = Y, label = Sample)) +
    geom_text() +
    xlab(paste("PC1 - ", pca.var.per[1], "%", sep="")) +
    ylab(paste("PC2 - ", pca.var.per[2], "%", sep="")) +
    theme_bw() +
    ggtitle("PCA Plot")
}

counts_all_display <- counts_all
colnames(counts_all_display)[7:14] <- c("M1_Mi", "M2_Mi", "Z1_Mi", "Z2_Mi", "M1_Next", "M2_Next", "Z1_N", "Z2_N")
PCAnalysis(counts_all_display)
```



6. Heatmapa

```
normalize <- function(data) {
  log_data <- rlog(data)
  normalized_data<- assay(log_data)
  normalized_data <- as.data.frame(normalized_data)

  return(normalized_data)
}

drawHeatmap <- function(data){
  threshold <- 14.5
  data <- data[
    data$Mock_1_MiSeq > threshold |
    data$Mock_2_MiSeq > threshold |
    data$ZIKA_1_MiSeq > threshold |
    data$ZIKA_2_MiSeq > threshold |
    data$Mock_1_NextSeq > threshold |
    data$Mock_2_NextSeq > threshold |
    data$ZIKA_1_NextSeq > threshold |
    data$ZIKA_2_NextSeq > threshold
  , ]

  Heatmap(data , cluster_columns = FALSE,
    row_names_side = "left",
```

```

    row_dend_sid = "left",
    row_names_gp=gpar(cex = 0.8))
}

data_for_heatmap <- normalize(dds(counts_all, 8))
data_reordered <- data_for_heatmap[c(0,1,2,5,6,3,4,7,8)]
drawHeatmap(data_reordered)

```

