Figure-S1C-SLC1A1.R

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```
# This Script Generates Figure S1C
# Script By: Eishani Kumar Sokolowski

# Empty the environment & suppress warnings
rm(list = ls())
options(warn=-1)

# Loading libraries
library(tidyverse)
```

```
## — Attaching core tidyverse packages —
                                                             — tidyverse 2.0.0 —
             1.1.4
## ✓ dplyr
                        ✓ readr
                                    2.1.5
## ✓ forcats 1.0.0
                        ✓ stringr
                                    1.5.1
                                    3.2.1
## ✓ ggplot2 3.5.1

✓ tibble

## ✓ lubridate 1.9.3

✓ tidyr

                                    1.3.1
## ✓ purrr
              1.0.2
## — Conflicts —
                                                       — tidyverse_conflicts() —
## * dplyr::filter() masks stats::filter()
                    masks stats::lag()
## x dplyr::lag()
## i Use the conflicted package (<http://conflicted.r-lib.org/>) to force all conflicts
to become errors
```

```
library(ggplot2)
library(pheatmap)
library(ggpubr)
library(data.table)
```

```
##
## Attaching package: 'data.table'
##
## The following objects are masked from 'package:lubridate':
##
##
       hour, isoweek, mday, minute, month, quarter, second, wday, week,
##
       yday, year
##
## The following objects are masked from 'package:dplyr':
##
##
       between, first, last
##
## The following object is masked from 'package:purrr':
##
##
       transpose
```

```
library(dplyr)
library(ggpubr)
library(rstatix)
```

```
##
## Attaching package: 'rstatix'
##
## The following object is masked from 'package:stats':
##
## filter
```

```
# Load the gene of interest
DEG.of.interest <- c("SLC1A1")</pre>
# Loading the normalized expression matrix
normalized.counts.ERS <- read.csv("./normalized.with.ERCC.counts.for.heatmap.ER.Stress.c
sv")
normalized.counts.INF <- read.csv("./normalized.with.ERCC.counts.for.heatmap.Inflammatio
n.csv")
final.df <- merge(normalized.counts.ERS,normalized.counts.INF,by="Genes", all = TRUE)</pre>
final.df <- final.df[final.df$Genes %in% DEG.of.interest,]</pre>
rownames(final.df) <- final.df$Genes
final.df <- final.df[-c(1)]</pre>
# ER stress
normalized.counts.ERS.Thaps <- final.df[ , grep("Thaps", colnames(final.df))]</pre>
normalized.counts.ERS.DMSO <- final.df[ , grep("DMSO", colnames(final.df))]</pre>
# Thapsigargin expression
normalized.counts.ERS.Thaps.Islet_144_Thapsigargin<- as.data.frame(normalized.counts.ER
S. Thaps [, 1])
normalized.counts.ERS.Thaps.Islet_146_Thapsigargin<- as.data.frame(normalized.counts.ER
S.Thaps[,2])
normalized.counts.ERS.Thaps.Islet 147 Thapsigargin<- as.data.frame(normalized.counts.ER
S. Thaps [.3])
normalized.counts.ERS.Thaps.Islet_148_Thapsigargin<- as.data.frame(normalized.counts.ER
S. Thaps [, 4])
normalized.counts.ERS.Thaps.Islet 149 Thapsigargin<- as.data.frame(normalized.counts.ER
S. Thaps [,5])
normalized.counts.ERS.Thaps.Islet_75_Thapsigargin<- as.data.frame(normalized.counts.ER
S. Thaps [, 6])
normalized.counts.ERS.Thaps.Islet_78_Thapsigargin<- as.data.frame(normalized.counts.ER
S. Thaps [, 7])
normalized.counts.ERS.Thaps.Islet_81_Thapsigargin<- as.data.frame(normalized.counts.ER
S. Thaps [, 8])
normalized.counts.ERS.Thaps.Islet_82_Thapsigargin<- as.data.frame(normalized.counts.ER
S. Thaps [, 9])
normalized.counts.ERS.Thaps.Islet 86 Thapsigargin<- as.data.frame(normalized.counts.ER
S.Thaps[,10])
colnames(normalized.counts.ERS.Thaps.Islet 144 Thapsigargin) <- c("Expression")
colnames(normalized.counts.ERS.Thaps.Islet 146 Thapsigargin) <- c("Expression")
```

```
colnames(normalized.counts.ERS.Thaps.Islet 147 Thapsigargin) <- c("Expression")
colnames(normalized.counts.ERS.Thaps.Islet_148_Thapsigargin) <- c("Expression")</pre>
colnames(normalized.counts.ERS.Thaps.Islet_149_Thapsigargin) <- c("Expression")
colnames(normalized.counts.ERS.Thaps.Islet_75_Thapsigargin) <- c("Expression")</pre>
colnames(normalized.counts.ERS.Thaps.Islet_78_Thapsigargin) <- c("Expression")</pre>
colnames(normalized.counts.ERS.Thaps.Islet_81_Thapsigargin) <- c("Expression")</pre>
colnames(normalized.counts.ERS.Thaps.Islet_82_Thapsigargin) <- c("Expression")</pre>
colnames(normalized.counts.ERS.Thaps.Islet_86_Thapsigargin) <- c("Expression")</pre>
normalized.counts.ERS.Thaps.Islet_144_Thapsigargin$Islet<- c(colnames(normalized.counts.
ERS.Thaps)[1])
normalized.counts.ERS.Thaps.Islet_146_Thapsigargin$Islet<- c(colnames(normalized.counts.
ERS.Thaps)[2])
normalized.counts.ERS.Thaps.Islet_147_Thapsigargin$Islet<- c(colnames(normalized.counts.
ERS.Thaps)[3])
normalized.counts.ERS.Thaps.Islet_148_Thapsigargin$Islet<- c(colnames(normalized.counts.
ERS.Thaps)[4])
normalized.counts.ERS.Thaps.Islet_149_Thapsigargin$Islet<- c(colnames(normalized.counts.
ERS.Thaps)[5])
normalized.counts.ERS.Thaps.Islet_75_Thapsigargin$Islet<- c(colnames(normalized.counts.E
RS.Thaps)[6])
normalized.counts.ERS.Thaps.Islet_78_Thapsigargin$Islet<- c(colnames(normalized.counts.E
RS.Thaps)[7])
normalized.counts.ERS.Thaps.Islet_81_Thapsigargin$Islet<- c(colnames(normalized.counts.E
RS.Thaps)[8])
normalized.counts.ERS.Thaps.Islet_82_Thapsigargin$Islet<- c(colnames(normalized.counts.E
RS.Thaps)[9])
normalized.counts.ERS.Thaps.Islet_86_Thapsigargin$Islet<- c(colnames(normalized.counts.E
RS.Thaps)[10])
normalized.counts.ERS.Thaps.Islet_144_Thapsigargin$Genes<- rownames(normalized.counts.ER
S.Thaps)
normalized.counts.ERS.Thaps.Islet_146_Thapsigargin$Genes<- rownames(normalized.counts.ER
S.Thaps)
normalized.counts.ERS.Thaps.Islet_147_Thapsigargin$Genes<- rownames(normalized.counts.ER
S.Thaps)
normalized.counts.ERS.Thaps.Islet_148_Thapsigargin$Genes<- rownames(normalized.counts.ER
S.Thaps)
normalized.counts.ERS.Thaps.Islet_149_Thapsigargin$Genes<- rownames(normalized.counts.ER
S.Thaps)
normalized.counts.ERS.Thaps.Islet_75_Thapsigargin$Genes<- rownames(normalized.counts.ER
S.Thaps)
normalized.counts.ERS.Thaps.Islet_78_Thapsigargin$Genes<- rownames(normalized.counts.ER
S.Thaps)
normalized.counts.ERS.Thaps.Islet_81_Thapsigargin$Genes<- rownames(normalized.counts.ER
S.Thaps)
normalized.counts.ERS.Thaps.Islet_82_Thapsigargin$Genes<- rownames(normalized.counts.ER
S.Thaps)
normalized.counts.ERS.Thaps.Islet_86_Thapsigargin$Genes<- rownames(normalized.counts.ER
S.Thaps)
final.df.Thaps <- rbind(normalized.counts.ERS.Thaps.Islet_144_Thapsigargin,</pre>
```

```
normalized.counts.ERS.Thaps.Islet_146_Thapsigargin, normalized.counts.ERS.Thaps.Islet_147_Thapsigargin, normalized.counts.ERS.Thaps.Islet_148_Thapsigargin, normalized.counts.ERS.Thaps.Islet_149_Thapsigargin, normalized.counts.ERS.Thaps.Islet_75_Thapsigargin, normalized.counts.ERS.Thaps.Islet_78_Thapsigargin, normalized.counts.ERS.Thaps.Islet_81_Thapsigargin, normalized.counts.ERS.Thaps.Islet_82_Thapsigargin, normalized.counts.ERS.Thaps.Islet_86_Thapsigargin)
```

final.df.Thaps\$Treatment <- c("B_Thapsigargin")</pre>

```
normalized.counts.ERS.DMS0.Islet_144_DMS0<- as.data.frame(normalized.counts.ERS.DMS0[,
11)
normalized.counts.ERS.DMS0.Islet_146_DMS0<- as.data.frame(normalized.counts.ERS.DMS0[,
21)
normalized.counts.ERS.DMS0.Islet_147_DMS0<- as.data.frame(normalized.counts.ERS.DMS0[,
31)
normalized.counts.ERS.DMS0.Islet_148_DMS0<- as.data.frame(normalized.counts.ERS.DMS0[,
4])
normalized.counts.ERS.DMS0.Islet_149_DMS0<- as.data.frame(normalized.counts.ERS.DMS0[,
5])
normalized.counts.ERS.DMSO.Islet_75_DMSO<- as.data.frame(normalized.counts.ERS.DMSO[,
6])
normalized.counts.ERS.DMSO.Islet_78_DMSO<- as.data.frame(normalized.counts.ERS.DMSO[,
7])
normalized.counts.ERS.DMSO.Islet 81 DMSO<- as.data.frame(normalized.counts.ERS.DMSO[,
8])
normalized.counts.ERS.DMSO.Islet 82 DMSO<- as.data.frame(normalized.counts.ERS.DMSO[,
91)
normalized.counts.ERS.DMS0.Islet_86_DMS0<- as.data.frame(normalized.counts.ERS.DMS0[,1
0])
colnames(normalized.counts.ERS.DMSO.Islet_144_DMSO) <- c("Expression")</pre>
colnames(normalized.counts.ERS.DMSO.Islet_146_DMSO) <- c("Expression")</pre>
colnames(normalized.counts.ERS.DMSO.Islet_147_DMSO) <- c("Expression")</pre>
colnames(normalized.counts.ERS.DMSO.Islet 148 DMSO) <- c("Expression")
colnames(normalized.counts.ERS.DMSO.Islet_149_DMSO) <- c("Expression")</pre>
colnames(normalized.counts.ERS.DMS0.Islet 75 DMS0) <- c("Expression")</pre>
colnames(normalized.counts.ERS.DMS0.Islet_78_DMS0) <- c("Expression")</pre>
colnames(normalized.counts.ERS.DMSO.Islet 81 DMSO) <- c("Expression")</pre>
colnames(normalized.counts.ERS.DMS0.Islet_82_DMS0) <- c("Expression")</pre>
colnames(normalized.counts.ERS.DMS0.Islet_86_DMS0) <- c("Expression")</pre>
normalized.counts.ERS.DMS0.Islet_144_DMS0$Islet<- c(colnames(normalized.counts.ERS.DMS0)
[1])
normalized.counts.ERS.DMS0.Islet_146_DMS0$Islet<- c(colnames(normalized.counts.ERS.DMS0)
[2])
```

```
normalized.counts.ERS.DMSO.Islet 147 DMSO$Islet<- c(colnames(normalized.counts.ERS.DMSO)
[3])
normalized.counts.ERS.DMSO.Islet_148_DMSO$Islet<- c(colnames(normalized.counts.ERS.DMSO)</pre>
normalized.counts.ERS.DMSO.Islet 149 DMSO$Islet<- c(colnames(normalized.counts.ERS.DMSO)
[5])
normalized.counts.ERS.DMSO.Islet_75_DMSO$Islet<- c(colnames(normalized.counts.ERS.DMSO)</pre>
[6])
normalized.counts.ERS.DMS0.Islet_78_DMS0$Islet<- c(colnames(normalized.counts.ERS.DMS0)</pre>
[7])
normalized.counts.ERS.DMSO.Islet_81_DMSO$Islet<- c(colnames(normalized.counts.ERS.DMSO)</pre>
[8])
normalized.counts.ERS.DMS0.Islet_82_DMS0$Islet<- c(colnames(normalized.counts.ERS.DMS0)</pre>
[9])
normalized.counts.ERS.DMS0.Islet_86_DMS0$Islet<- c(colnames(normalized.counts.ERS.DMS0)
[10])
normalized.counts.ERS.DMSO.Islet 144 DMSO$Genes<- rownames(normalized.counts.ERS.DMSO)
normalized.counts.ERS.DMSO.Islet_146_DMSO$Genes<- rownames(normalized.counts.ERS.DMSO)
normalized.counts.ERS.DMSO.Islet 147 DMSO$Genes<- rownames(normalized.counts.ERS.DMSO)
normalized.counts.ERS.DMSO.Islet_148_DMSO$Genes<- rownames(normalized.counts.ERS.DMSO)
normalized.counts.ERS.DMSO.Islet 149 DMSO$Genes<- rownames(normalized.counts.ERS.DMSO)
normalized.counts.ERS.DMS0.Islet_75_DMS0$Genes<- rownames(normalized.counts.ERS.DMS0)</pre>
normalized.counts.ERS.DMS0.Islet 78 DMS0$Genes<- rownames(normalized.counts.ERS.DMS0)
normalized.counts.ERS.DMSO.Islet_81_DMSO$Genes<- rownames(normalized.counts.ERS.DMSO)</pre>
normalized.counts.ERS.DMS0.Islet 82 DMS0$Genes<- rownames(normalized.counts.ERS.DMS0)
normalized.counts.ERS.DMS0.Islet_86_DMS0$Genes<- rownames(normalized.counts.ERS.DMS0)</pre>
final.df.DMS0 <- rbind(normalized.counts.ERS.DMS0.Islet_144_DMS0,</pre>
                     normalized.counts.ERS.DMSO.Islet 146 DMSO,
                     normalized.counts.ERS.DMSO.Islet 147 DMSO,
                     normalized.counts.ERS.DMSO.Islet_148_DMSO,
                     normalized.counts.ERS.DMSO.Islet 149 DMSO,
                     normalized.counts.ERS.DMSO.Islet 75 DMSO,
                     normalized.counts.ERS.DMSO.Islet_78_DMSO,
                     normalized.counts.ERS.DMSO.Islet 81 DMSO,
                     normalized.counts.ERS.DMSO.Islet 82 DMSO,
                     normalized.counts.ERS.DMSO.Islet_86_DMSO)
final.df.DMSO$Treatment <- c("A_DMSO")</pre>
final.df.ERS <- rbind(final.df.DMSO,final.df.Thaps)</pre>
# Cytokine-induced inflammation
normalized.counts.Cytokines <- final.df[ , grep("Cytokines", colnames(final.df))]</pre>
normalized.counts.Untreated <- final.df[ , grep("Untreated", colnames(final.df))]</pre>
# Cytokines expression
```

```
normalized.counts.Cytokines.Islet_100_Cytokines<- as.data.frame(normalized.counts.Cytok
ines[,1])
normalized.counts.Cytokines.Islet_103_Cytokines<- as.data.frame(normalized.counts.Cytok
ines[,2])
normalized.counts.Cytokines.Islet_144_Cytokines<- as.data.frame(normalized.counts.Cytok
ines[,3])
normalized.counts.Cytokines.Islet_146_Cytokines<- as.data.frame(normalized.counts.Cytok
ines[,4])
normalized.counts.Cytokines.Islet_147_Cytokines<- as.data.frame(normalized.counts.Cytok
ines[,5])
normalized.counts.Cytokines.Islet_148_Cytokines<- as.data.frame(normalized.counts.Cytok
ines[,6])
normalized.counts.Cytokines.Islet_149_Cytokines<- as.data.frame(normalized.counts.Cytok
ines[,7])
normalized.counts.Cytokines.Islet_75_Cytokines<- as.data.frame(normalized.counts.Cytoki
nes[.8])
normalized.counts.Cytokines.Islet_78_Cytokines<- as.data.frame(normalized.counts.Cytoki
nes[.9])
normalized.counts.Cytokines.Islet_81_Cytokines<- as.data.frame(normalized.counts.Cytoki
nes[.10])
normalized.counts.Cytokines.Islet_82_Cytokines<- as.data.frame(normalized.counts.Cytoki
nes[,11])
normalized.counts.Cytokines.Islet_86_Cytokines<- as.data.frame(normalized.counts.Cytoki
nes[,12])
normalized.counts.Cytokines.Islet_92_Cytokines<- as.data.frame(normalized.counts.Cytoki
nes[,13])
normalized.counts.Cytokines.Islet_93_Cytokines<- as.data.frame(normalized.counts.Cytoki
nes[,14])
normalized.counts.Cytokines.Islet_94_Cytokines<- as.data.frame(normalized.counts.Cytoki
nes[,15])
normalized.counts.Cytokines.Islet 95 Cytokines<- as.data.frame(normalized.counts.Cytoki</pre>
nes[,16])
normalized.counts.Cytokines.Islet_96_Cytokines<- as.data.frame(normalized.counts.Cytoki
nes[,17])
normalized.counts.Cytokines.Islet 97 Cytokines<- as.data.frame(normalized.counts.Cytoki
nes[,18])
normalized.counts.Cytokines.Islet_99_Cytokines<- as.data.frame(normalized.counts.Cytoki
nes[,19])
colnames(normalized.counts.Cytokines.Islet_100_Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet 103 Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet_144_Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet 146 Cytokines) <- c("Expression")
colnames(normalized.counts.Cytokines.Islet_147_Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet_148_Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet_149_Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet_75_Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet 78 Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet_81_Cytokines) <- c("Expression")</pre>
```

colnames(normalized.counts.Cytokines.Islet 82 Cytokines) <- c("Expression")</pre>

```
colnames(normalized.counts.Cytokines.Islet 86 Cytokines) <- c("Expression")
colnames(normalized.counts.Cytokines.Islet_92_Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet_93_Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet 94 Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet_95_Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet_96_Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet_97_Cytokines) <- c("Expression")</pre>
colnames(normalized.counts.Cytokines.Islet_99_Cytokines) <- c("Expression")</pre>
normalized.counts.Cytokines.Islet_100_Cytokines$Islet<- c(colnames(normalized.counts.Cyt</pre>
okines)[1])
normalized.counts.Cytokines.Islet_103_Cytokines$Islet<- c(colnames(normalized.counts.Cyt
okines)[2])
normalized.counts.Cytokines.Islet_144_Cytokines$Islet<- c(colnames(normalized.counts.Cyt
okines)[3])
normalized.counts.Cytokines.Islet_146_Cytokines$Islet<- c(colnames(normalized.counts.Cyt
okines)[4])
normalized.counts.Cytokines.Islet_147_Cytokines$Islet<- c(colnames(normalized.counts.Cyt
okines)[5])
normalized.counts.Cytokines.Islet_148_Cytokines$Islet<- c(colnames(normalized.counts.Cyt
okines)[6])
normalized.counts.Cytokines.Islet_149_Cytokines$Islet<- c(colnames(normalized.counts.Cyt
okines)[7])
normalized.counts.Cytokines.Islet_75_Cytokines$Islet<- c(colnames(normalized.counts.Cyto
kines)[8])
normalized.counts.Cytokines.Islet_78_Cytokines$Islet<- c(colnames(normalized.counts.Cyto
kines)[9])
normalized.counts.Cytokines.Islet_81_Cytokines$Islet<- c(colnames(normalized.counts.Cyto
kines)[10])
normalized.counts.Cytokines.Islet_82_Cytokines$Islet<- c(colnames(normalized.counts.Cyto
kines)[11])
normalized.counts.Cytokines.Islet_86_Cytokines$Islet<- c(colnames(normalized.counts.Cyto
kines)[12])
normalized.counts.Cytokines.Islet_92_Cytokines$Islet<- c(colnames(normalized.counts.Cyto
kines)[13])
normalized.counts.Cytokines.Islet_93_Cytokines$Islet<- c(colnames(normalized.counts.Cyto
kines)[14])
normalized.counts.Cytokines.Islet_94_Cytokines$Islet<- c(colnames(normalized.counts.Cyto
kines)[15])
normalized.counts.Cytokines.Islet_95_Cytokines$Islet<- c(colnames(normalized.counts.Cyto</pre>
kines)[16])
normalized.counts.Cytokines.Islet_96_Cytokines$Islet<- c(colnames(normalized.counts.Cyto</pre>
kines)[17])
normalized.counts.Cytokines.Islet_97_Cytokines$Islet<- c(colnames(normalized.counts.Cyto
normalized.counts.Cytokines.Islet_99_Cytokines$Islet<- c(colnames(normalized.counts.Cyto
kines)[19])
normalized.counts.Cytokines.Islet_100_Cytokines$Genes<- rownames(normalized.counts.Cytok
normalized.counts.Cytokines.Islet_103_Cytokines$Genes<- rownames(normalized.counts.Cytok
ines)
```

```
normalized.counts.Cytokines.Islet_144_Cytokines$Genes<- rownames(normalized.counts.Cytok
ines)
normalized.counts.Cytokines.Islet_146_Cytokines$Genes<- rownames(normalized.counts.Cytok</pre>
normalized.counts.Cytokines.Islet_147_Cytokines$Genes<- rownames(normalized.counts.Cytok
ines)
normalized.counts.Cytokines.Islet_148_Cytokines$Genes<- rownames(normalized.counts.Cytok
ines)
normalized.counts.Cytokines.Islet_149_Cytokines$Genes<- rownames(normalized.counts.Cytok
ines)
normalized.counts.Cytokines.Islet_75_Cytokines$Genes<- rownames(normalized.counts.Cytoki
normalized.counts.Cytokines.Islet_78_Cytokines$Genes<- rownames(normalized.counts.Cytoki
nes)
normalized.counts.Cytokines.Islet_81_Cytokines$Genes<- rownames(normalized.counts.Cytoki
nes)
normalized.counts.Cytokines.Islet_82_Cytokines$Genes<- rownames(normalized.counts.Cytoki
nes)
normalized.counts.Cytokines.Islet_86_Cytokines$Genes<- rownames(normalized.counts.Cytoki
nes)
normalized.counts.Cytokines.Islet_92_Cytokines$Genes<- rownames(normalized.counts.Cytoki</pre>
nes)
normalized.counts.Cytokines.Islet_93_Cytokines$Genes<- rownames(normalized.counts.Cytoki
nes)
normalized.counts.Cytokines.Islet_94_Cytokines$Genes<- rownames(normalized.counts.Cytoki
nes)
normalized.counts.Cytokines.Islet_95_Cytokines$Genes<- rownames(normalized.counts.Cytoki
nes)
normalized.counts.Cytokines.Islet_96_Cytokines$Genes<- rownames(normalized.counts.Cytoki
nes)
normalized.counts.Cytokines.Islet_97_Cytokines$Genes<- rownames(normalized.counts.Cytoki
nes)
normalized.counts.Cytokines.Islet_99_Cytokines$Genes<- rownames(normalized.counts.Cytoki
nes)
final.df.Cytokines <- rbind(normalized.counts.Cytokines.Islet_100_Cytokines,</pre>
                            normalized.counts.Cytokines.Islet_103_Cytokines,
                            normalized.counts.Cytokines.Islet_144_Cytokines,
                            normalized.counts.Cytokines.Islet_146_Cytokines,
                            normalized.counts.Cytokines.Islet_147_Cytokines,
                            normalized.counts.Cytokines.Islet 148 Cytokines,
                            normalized.counts.Cytokines.Islet_149_Cytokines,
                            normalized.counts.Cytokines.Islet_75_Cytokines,
                            normalized.counts.Cytokines.Islet_78_Cytokines,
                            normalized.counts.Cytokines.Islet_81_Cytokines,
                            normalized.counts.Cytokines.Islet_82_Cytokines,
                            normalized.counts.Cytokines.Islet_86_Cytokines,
                            normalized.counts.Cytokines.Islet_92_Cytokines,
                            normalized.counts.Cytokines.Islet_93_Cytokines,
                            normalized.counts.Cytokines.Islet 94 Cytokines,
                            normalized.counts.Cytokines.Islet_95_Cytokines,
```

normalized.counts.Cytokines.Islet_96_Cytokines,

```
normalized.counts.Cytokines.Islet_97_Cytokines,
normalized.counts.Cytokines.Islet_99_Cytokines)
```

final.df.Cytokines\$Treatment <- c("D_Cytokines")</pre>


```
normalized.counts.Untreated.Islet_100_Untreated<- as.data.frame(normalized.counts.Untre
ated[,1])
normalized.counts.Untreated.Islet_103_Untreated<- as.data.frame(normalized.counts.Untre
ated[.2])
normalized.counts.Untreated.Islet_144_Untreated<- as.data.frame(normalized.counts.Untre
ated[,3])
normalized.counts.Untreated.Islet_146_Untreated<- as.data.frame(normalized.counts.Untre
ated[,4])
normalized.counts.Untreated.Islet 147 Untreated<- as.data.frame(normalized.counts.Untre
ated[,5])
normalized.counts.Untreated.Islet_148_Untreated<- as.data.frame(normalized.counts.Untre
ated[,6])
normalized.counts.Untreated.Islet 149 Untreated<- as.data.frame(normalized.counts.Untre
ated[,7])
normalized.counts.Untreated.Islet 75 Untreated<- as.data.frame(normalized.counts.Untrea
ted[,8])
normalized.counts.Untreated.Islet 78 Untreated<- as.data.frame(normalized.counts.Untrea
ted[,9])
normalized.counts.Untreated.Islet 81 Untreated<- as.data.frame(normalized.counts.Untrea
ted[,10])
normalized.counts.Untreated.Islet_82_Untreated<- as.data.frame(normalized.counts.Untrea
ted[,11])
normalized.counts.Untreated.Islet_86_Untreated<- as.data.frame(normalized.counts.Untrea
ted[,12])
normalized.counts.Untreated.Islet_92_Untreated<- as.data.frame(normalized.counts.Untrea
ted[,13])
normalized.counts.Untreated.Islet_93_Untreated<- as.data.frame(normalized.counts.Untrea
ted[,14])
normalized.counts.Untreated.Islet_94_Untreated<- as.data.frame(normalized.counts.Untrea
ted[,15])
normalized.counts.Untreated.Islet_95_Untreated<- as.data.frame(normalized.counts.Untrea
ted[.16])
normalized.counts.Untreated.Islet_96_Untreated<- as.data.frame(normalized.counts.Untrea
ted[,17])
normalized.counts.Untreated.Islet_97_Untreated<- as.data.frame(normalized.counts.Untrea
ted[.18])
normalized.counts.Untreated.Islet_99_Untreated<- as.data.frame(normalized.counts.Untrea
ted[,19])
colnames(normalized.counts.Untreated.Islet_100_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet 103 Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_144_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet 146 Untreated) <- c("Expression")</pre>
```

```
colnames(normalized.counts.Untreated.Islet 147 Untreated) <- c("Expression")
colnames(normalized.counts.Untreated.Islet_148_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_149_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_75_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_78_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_81_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_82_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_86_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_92_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_93_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet 94 Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_95_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_96_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_97_Untreated) <- c("Expression")</pre>
colnames(normalized.counts.Untreated.Islet_99_Untreated) <- c("Expression")</pre>
normalized.counts.Untreated.Islet_100_Untreated$Islet<- c(colnames(normalized.counts.Unt
normalized.counts.Untreated.Islet_103_Untreated$Islet<- c(colnames(normalized.counts.Unt
reated)[2])
normalized.counts.Untreated.Islet_144_Untreated$Islet<- c(colnames(normalized.counts.Unt
reated)[3])
normalized.counts.Untreated.Islet_146_Untreated$Islet<- c(colnames(normalized.counts.Unt
reated)[4])
normalized.counts.Untreated.Islet_147_Untreated$Islet<- c(colnames(normalized.counts.Unt
reated)[5])
normalized.counts.Untreated.Islet_148_Untreated$Islet<- c(colnames(normalized.counts.Unt
reated)[6])
normalized.counts.Untreated.Islet 149 Untreated$Islet<- c(colnames(normalized.counts.Unt
reated)[7])
normalized.counts.Untreated.Islet_75_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[8])
normalized.counts.Untreated.Islet_78_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[9])
normalized.counts.Untreated.Islet_81_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[10])
normalized.counts.Untreated.Islet_82_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[11])
normalized.counts.Untreated.Islet_86_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[12])
normalized.counts.Untreated.Islet_92_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[13])
normalized.counts.Untreated.Islet_93_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[14])
normalized.counts.Untreated.Islet_94_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[15])
normalized.counts.Untreated.Islet_95_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[16])
normalized.counts.Untreated.Islet_96_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[17])
normalized.counts.Untreated.Islet_97_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[18])
```

```
normalized.counts.Untreated.Islet_99_Untreated$Islet<- c(colnames(normalized.counts.Untr
eated)[19])
normalized.counts.Untreated.Islet_100_Untreated$Genes<- rownames(normalized.counts.Untre
ated)
normalized.counts.Untreated.Islet_103_Untreated$Genes<- rownames(normalized.counts.Untre
ated)
normalized.counts.Untreated.Islet_144_Untreated$Genes<- rownames(normalized.counts.Untre
ated)
normalized.counts.Untreated.Islet_146_Untreated$Genes<- rownames(normalized.counts.Untre
ated)
normalized.counts.Untreated.Islet_147_Untreated$Genes<- rownames(normalized.counts.Untre
ated)
normalized.counts.Untreated.Islet_148_Untreated$Genes<- rownames(normalized.counts.Untre
normalized.counts.Untreated.Islet_149_Untreated$Genes<- rownames(normalized.counts.Untre
ated)
normalized.counts.Untreated.Islet_75_Untreated$Genes<- rownames(normalized.counts.Untrea
normalized.counts.Untreated.Islet_78_Untreated$Genes<- rownames(normalized.counts.Untrea
ted)
normalized.counts.Untreated.Islet_81_Untreated$Genes<- rownames(normalized.counts.Untrea
ted)
normalized.counts.Untreated.Islet_82_Untreated$Genes<- rownames(normalized.counts.Untrea
ted)
normalized.counts.Untreated.Islet_86_Untreated$Genes<- rownames(normalized.counts.Untrea
ted)
normalized.counts.Untreated.Islet_92_Untreated$Genes<- rownames(normalized.counts.Untrea
ted)
normalized.counts.Untreated.Islet_93_Untreated$Genes<- rownames(normalized.counts.Untrea
ted)
normalized.counts.Untreated.Islet_94_Untreated$Genes<- rownames(normalized.counts.Untrea
ted)
normalized.counts.Untreated.Islet_95_Untreated$Genes<- rownames(normalized.counts.Untrea
ted)
normalized.counts.Untreated.Islet_96_Untreated$Genes<- rownames(normalized.counts.Untrea
ted)
normalized.counts.Untreated.Islet_97_Untreated$Genes<- rownames(normalized.counts.Untrea</pre>
normalized.counts.Untreated.Islet_99_Untreated$Genes<- rownames(normalized.counts.Untrea
ted)
final.df.Untreated <- rbind(normalized.counts.Untreated.Islet 100 Untreated,</pre>
                            normalized.counts.Untreated.Islet_103_Untreated,
                            normalized.counts.Untreated.Islet 144 Untreated,
                            normalized.counts.Untreated.Islet_146_Untreated,
                            normalized.counts.Untreated.Islet_147_Untreated,
                            normalized.counts.Untreated.Islet_148_Untreated,
                            normalized.counts.Untreated.Islet_149_Untreated,
                            normalized.counts.Untreated.Islet 75 Untreated,
                            normalized.counts.Untreated.Islet_78_Untreated,
                            normalized.counts.Untreated.Islet 81 Untreated,
```

```
normalized.counts.Untreated.Islet_82_Untreated,
                         normalized.counts.Untreated.Islet_86_Untreated,
                         normalized.counts.Untreated.Islet_92_Untreated,
                         normalized.counts.Untreated.Islet 93 Untreated,
                         normalized.counts.Untreated.Islet_94_Untreated,
                         normalized.counts.Untreated.Islet_95_Untreated,
                         normalized.counts.Untreated.Islet_96_Untreated,
                         normalized.counts.Untreated.Islet_97_Untreated,
                         normalized.counts.Untreated.Islet_99_Untreated)
final.df.Untreated$Treatment <- c("C Untreated")</pre>
final.df.INF <- rbind(final.df.Untreated,final.df.Cytokines)</pre>
# Generating dot-and-boxplot
final.df.ERS$Type <- c("ERS")</pre>
final.df.INF$Type <- c("INF")</pre>
final.df <- rbind(final.df.ERS,final.df.INF)</pre>
final.df$Expression <- log(final.df$Expression)</pre>
p <- ggboxplot(</pre>
 final.df, x = "Type", y = "Expression", color = "Treatment",
 palette = c("grey45", "forestgreen", "grey45","darkorange"), facet.by = "Genes", add =
"jitter", size=0.5,
) + ylab("Normalized Expression") + ylim(4.8,7.3)
ggarrange(p, nrow=1, ncol=2)
```

