Cardiomyocyte interactome, additional KEGG & Gene Ontology analyses

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R script for KEGG pathway analysis of cardiomyocyte RNA interactome proteins

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Loading of the data tables.

```
load("data/kegg.brite.rda")
load("data/interactome.rda")
load("data/wcl.rda")
```

Fetching KEGG identifiers:

```
ids <- unlist(lapply(strsplit(kegg.brite$C, " "), function(x) x[1]))</pre>
rownames(kegg.brite) <- ids</pre>
total.keggIDs <- keggLink("mmu", "pathway")</pre>
save(total.keggIDs, file = "data/total.keggIDs.rda")
```

We have found a total of length(total.keggIDs) which are used for mapping the WCL and interactome

Now we proceed to mapping of WCL protein IDs to KEGG IDs and testing for enrichments against background of all KEGG proteins contained in KEGG pathways.

```
# retrieved Entrez IDs from Biomart
mouse <- useMart("ensembl", dataset = "mmusculus_gene_ensembl")</pre>
human <- useMart("ensembl", dataset = "hsapiens_gene_ensembl")</pre>
attribs <- listAttributes(mouse)</pre>
pages <- attributePages(mouse)</pre>
hsap.attribs <- listAttributes(human)
entrez_ids <- getBM(attributes = c("ensembl_gene_id", "entrezgene"), values = wcl[,</pre>
    "ensembl_gene_id"], filters = "ensembl_gene_id", mart = mouse)
wcl.human_homologs <- getBM(attributes = c("ensembl_gene_id", "hsapiens_homolog_ensembl_gene"),
    values = wcl[, "ensembl_gene_id"], filters = "ensembl_gene_id", mart = mouse)
# remove ensembl_gene_ids which have duplicated entrez_ids
entrez ids <- entrez ids[-which(duplicated(entrez ids$ensembl gene id)), ]</pre>
wcl <- merge(wcl, entrez_ids, by.x = "ensembl_gene_id", by.y = "ensembl_gene_id",
    all.x = T
```

```
wcl.entrezIDs <- unique(wcl[!is.na(wcl$entrezgene), ]$entrezgene)</pre>
# this retrieval is fairly slow, therefore the results were written to
# './data'
wcl.keggIDs <- keggConv.batch(wcl.entrezIDs)</pre>
save(wcl.keggIDs, file = "data/wcl.keggIDs.rda")
wcl.keggQ <- lapply(wcl.keggIDs, function(x) keggGet(x))</pre>
save(wcl.keggQ, file = "data/wcl.keggQ.rda")
wcl.pathways <- unique(unlist(lapply(strsplit(names(unlist(lapply(wcl.keggQ,
    function(x) x[[1]]$PATHWAY))), "\\."), function(x) x[3]))
save(wcl.pathways, file = "data/wcl.pathways.rda")
wcl.pathways.genes <- lapply(wcl.pathways, function(x) keggLink("genes", x))</pre>
names(wcl.pathways.genes) <- wcl.pathways</pre>
save(wcl.pathways.genes, file = "data/wcl.pathways.genes.rda")
wcl.pathways.genes.entrez_ids <- unique(gsub("mmu:", "", as.character(unlist(wcl.pathways.genes))))</pre>
wcl.df <- kegg.brite[gsub("mmu", "", wcl.pathways), ]</pre>
wcl.df$ID <- rownames(wcl.df)</pre>
wcl.df$total <- rep(0, nrow(wcl.df))</pre>
wcl.df$total <- sapply(rownames(wcl.df), function(x) length(wcl.pathways.genes[[paste("mmu",
    x, sep = "")]]))
wcl.df$count <- rep(0, nrow(wcl.df))</pre>
wcl.df$frac <- rep(0, nrow(wcl.df))</pre>
for (i in rownames(wcl.df)) {
    # print(i)
    kL1 <- keggLink("mmu", paste("mmu", i, sep = ""))</pre>
    wcl.df[i, ]$count <- length(which(wcl.keggIDs %in% kL1))</pre>
    wcl.df[i, ]$frac <- round(length(which(wcl.keggIDs %in% kL1))/length(kL1) *</pre>
        100, 2)
}
# extract list of IDs in pathway
wcl.in_path.IDs <- lapply(rownames(wcl.df), function(x) {</pre>
    kL1 <- keggLink("mmu", paste("mmu", x, sep = ""))</pre>
    in_path <- wcl.keggIDs[which(wcl.keggIDs %in% kL1)]</pre>
})
names(wcl.in_path.IDs) <- rownames(wcl.df)</pre>
# perform Fisher's Exact Test for each category
bkgd <- length(unique(total.keggIDs))</pre>
smpl <- length(wcl.keggIDs)</pre>
ftl <- apply(wcl.df[1, ], 1, function(x) {</pre>
    ct <- as.integer(x["count"])
    tt <- as.integer(x["total"])</pre>
    m1 <- matrix(c(ct, tt, smpl - ct, bkgd - tt), 2, 2)
    fisher.test(m1, alternative = alternative)
})
wcl.df$ft_pval <- unlist(lapply(ftl, function(x) {</pre>
```

```
x$p.value
}))
wcl.df$ft_OR <- unlist(lapply(ftl, function(x) {
    x$estimate
}))
wcl.df$ft_fdr <- p.adjust(wcl.df$ft_pval, method = "fdr")
save(wcl.df, file = "data/wcl.df.rda")</pre>
```

Mapping of Interactome protein IDs to KEGG IDs and testing for enrichments against background of WCL proteins contained in KEGG pathways.

```
interactome.entrez_ids <- getBM(attributes = c("ensembl_gene_id", "entrezgene"),</pre>
    values = interactome[, "ensembl_gene_id"], filters = "ensembl_gene_id",
    mart = mouse)
interactome.human_homologs <- getBM(attributes = c("ensembl_gene_id", "hsapiens_homolog_ensembl_gene"),
    values = interactome[, "ensembl_gene_id"], filters = "ensembl_gene_id",
    mart = mouse)
# remove ensembl_gene_ids which have duplicated entrez_ids
interactome.entrez_ids <- interactome.entrez_ids[-which(duplicated(interactome.entrez_ids$ensembl_gene_
interactome <- merge(interactome, interactome.entrez_ids, by.x = "ensembl_gene_id",
    by.y = "ensembl_gene_id", all.x = T)
interactome.entrezIDs <- unique(interactome[!is.na(interactome$entrezgene),</pre>
    ]$entrezgene)
save(interactome.entrezIDs, file = "data/interactome.entrezIDs")
interactome.keggIDs <- keggConv.batch(interactome.entrezIDs)</pre>
save(interactome.keggIDs, file = "data/interactome.keggIDs.rda")
interactome.keggQ <- lapply(interactome.keggIDs, function(x) keggGet(x))</pre>
save(interactome.keggQ, file = "data/interactome.keggQ.rda")
interactome.pathways <- unique(unlist(lapply(strsplit(names(unlist(lapply(interactome.keggQ,</pre>
    function(x) x[[1]]$PATHWAY))), "\\."), function(x) x[3]))
save(interactome.pathways, file = "data/interactome.pathways.rda")
interactome.pathways.genes <- lapply(interactome.pathways, function(x) keggLink("genes",</pre>
    x))
names(interactome.pathways.genes) <- interactome.pathways</pre>
save(interactome.pathways.genes, file = "data/interactome.pathways.genes.rda")
interactome.pathways.genes.entrez_ids <- unique(gsub("mmu:", "", as.character(unlist(interactome.pathwa
# create dataframe for counting hits in pathways
interactome.df <- kegg.brite[gsub("mmu", "", interactome.pathways), ]</pre>
interactome.df$source <- rep("Interactome", nrow(interactome.df))</pre>
interactome.df$ID <- rownames(interactome.df)</pre>
# we are now using WCL as background to test for enrichment
i1 <- intersect(rownames(interactome.df), rownames(wcl.df))</pre>
```

```
interactome.df$total <- rep(0, nrow(interactome.df))</pre>
interactome.df[i1, ]$total <- wcl.df[i1, ]$count</pre>
interactome.df$count <- rep(0, nrow(interactome.df))</pre>
interactome.df$frac <- rep(0, nrow(interactome.df))</pre>
for (i in rownames(interactome.df)) {
    kL1 <- keggLink("mmu", paste("mmu", i, sep = ""))</pre>
    interactome.df[i, ]$count <- length(which(interactome.keggIDs %in% kL1))</pre>
    interactome.df[i, ]$frac <- round(length(which(interactome.keggIDs %in%</pre>
        kL1))/length(kL1) * 100, 2)
}
# extract list of IDs in pathway
interactome.in_path.IDs <- lapply(rownames(interactome.df), function(x) {</pre>
    kL1 <- keggLink("mmu", paste("mmu", x, sep = ""))</pre>
    in_path <- interactome.keggIDs[which(interactome.keggIDs %in% kL1)]</pre>
})
# perform Fisher's Exact Test for each category
bkgd <- length(unique(wcl.keggIDs))</pre>
smpl <- length(interactome.keggIDs)</pre>
ftl <- apply(interactome.df, 1, function(x) {</pre>
    ct <- as.integer(x["count"])</pre>
    tt <- as.integer(x["total"])</pre>
    m1 <- matrix(c(ct, tt, smpl - ct, bkgd - tt), 2, 2)
    fisher.test(m1, alternative = alternative)
})
interactome.df$ft_pval <- unlist(lapply(ftl, function(x) {</pre>
    x$p.value
}))
interactome.df$ft_OR <- unlist(lapply(ftl, function(x) {</pre>
}))
interactome.df$ft_fdr <- p.adjust(interactome.df$ft_pval, method = p.adjust.method,
    n = nrow(wcl.df)
save(interactome.df, file = "data/interactome.df.rda")
```

KEGG contains three levels/hierarchies (A>B>C), here we summarize the enrichment at B level:

```
# -----interactome-summarizing data at 'B' level before doing Fisher's
# Exact test-----
interactome.B.df <- data.frame(matrix(ncol = 5, nrow = length(unique(interactome.df$B))))
colnames(interactome.B.df) <- c("B", "A", "total", "count", "source")
interactome.B.df$B <- unique(interactome.df$B)
interactome.B.df$A <- sapply(unique(interactome.df$B), function(x) {
    A <- unique(interactome.df[which(interactome.df$B %in% x), "A"])
})
interactome.B.df$source <- rep("Interactome", nrow(interactome.B.df))
interactome.B.df$total <- sapply(unique(interactome.df$B), function(x) {
    tot <- sum(interactome.df[which(interactome.df$B %in% x), "total"])
})</pre>
```

```
interactome.B.df$count <- sapply(unique(interactome.df$B), function(x) {</pre>
    count <- sum(interactome.df[which(interactome.df$B %in% x), "count"])</pre>
})
bkgd <- length(unique(wcl.keggIDs))</pre>
smpl <- length(interactome.keggIDs)</pre>
ftl <- apply(interactome.B.df, 1, function(x) {</pre>
    ct <- as.integer(x["count"])
    tt <- as.integer(x["total"])</pre>
    m1 <- matrix(c(ct, tt, smpl - ct, bkgd - tt), 2, 2)
    fisher.test(m1, alternative = alternative)
})
interactome.B.df$ft_pval <- unlist(lapply(ftl, function(x) {</pre>
    x$p.value
}))
interactome.B.df$ft_OR <- unlist(lapply(ftl, function(x) {</pre>
    x$estimate
}))
interactome.B.df$ft_fdr <- p.adjust(interactome.B.df$ft_pval, method = p.adjust.method,
    n = nrow(wcl.df)
save(interactome.B.df, file = "data/interactome.B.df.rda")
```

We are splitting the Interactome proteins into RNA-related and un-related proteins, based on annotation analysis (upstream of these steps): First, RNA-unrelated

```
#-----GO RNA unrelated-----
# subset the interactome table
interactome.go_rna_unrelated <- interactome[which(interactome$GO == "unrelated"),]</pre>
interactome.go_rna_unrelated.entrezIDs <- unique(interactome.go_rna_unrelated[!is.na(interactome.go_rna
interactome.go_rna_unrelated.keggIDs <- keggConv.batch(interactome.go_rna_unrelated.entrezIDs)
# dataframe for count data
interactome.go_rna_unrelated.df <- interactome.df</pre>
interactome.go_rna_unrelated.df$source <- rep("GO_RNA_unrelated", nrow(interactome.go_rna_unrelated.df)</pre>
interactome.go_rna_unrelated.df$ID <- rownames(interactome.go_rna_unrelated.df)</pre>
interactome.go_rna_unrelated.df$total <- rep(0, nrow(interactome.go_rna_unrelated.df))</pre>
# we are now using WCL as background to test for enrichment
i1 <- intersect(rownames(interactome.go_rna_unrelated.df), rownames(wcl.df))</pre>
interactome.go_rna_unrelated.df[i1,]$total <- wcl.df[i1,]$count</pre>
interactome.go_rna_unrelated.df$count <- rep(0, nrow(interactome.go_rna_unrelated.df))</pre>
for (i in rownames(interactome.go_rna_unrelated.df)) {
 kL1 <- keggLink("mmu", paste("mmu", i, sep = ""))</pre>
  interactome.go_rna_unrelated.df[i, ]$count <- length(which(interactome.go_rna_unrelated.keggIDs %in%)
# extract list of IDs in pathway
interactome.go_rna_unrelated.in_path.IDs <- lapply(rownames(interactome.go_rna_unrelated.df), function(
  kL1 <- keggLink("mmu", paste("mmu", x, sep = ""))</pre>
```

in_path <- interactome.go_rna_unrelated.keggIDs[which(interactome.go_rna_unrelated.keggIDs %in% kL1)]

```
names(interactome.go_rna_unrelated.in_path.IDs) <- rownames(interactome.go_rna_unrelated.df)</pre>
# perform Fisher's Exact Test for each category
# Using WCL as background
bkgd <- length(unique(wcl.keggIDs))</pre>
smpl <- length(interactome.go_rna_unrelated.keggIDs)</pre>
ftl <- apply(interactome.go_rna_unrelated.df, 1, function (x) {
  ct <- as.integer(x["count"])</pre>
 tt <- as.integer(x["total"])</pre>
 m1 <- matrix(c(ct, tt, smpl - ct, bkgd - tt), 2, 2)
  fisher.test(m1, alternative = alternative)
})
interactome.go_rna_unrelated.df$ft_pval <- unlist(lapply(ftl, function(x) {x$p.value}))</pre>
interactome.go_rna_unrelated.df$ft_OR <- unlist(lapply(ftl, function(x) {x$estimate}))</pre>
interactome.go_rna_unrelated.df$ft_fdr <- p.adjust(interactome.go_rna_unrelated.df$ft_pval, method = p.
save(interactome.go_rna_unrelated.df, file = "data/interactome.go_rna_unrelated.df.rda")
# summarizing data at "B" level before doing Fisher's Exact test
interactome.go_rna_unrelated.B.df <- data.frame(matrix(ncol = 5, nrow = length(unique(interactome.go_rn
colnames(interactome.go_rna_unrelated.B.df) <- c("B", "A", "total", "count", "source")</pre>
interactome.go_rna_unrelated.B.df$B <- unique(interactome.go_rna_unrelated.df$B)
interactome.go_rna_unrelated.B.df$A <- sapply(unique(interactome.go_rna_unrelated.df$B), function(x) {A
interactome.go_rna_unrelated.B.df$source <- rep("GO_RNA_unrelated", nrow(interactome.go_rna_unrelated.B
interactome.go_rna_unrelated.B.df$total <- sapply(unique(interactome.go_rna_unrelated.df$B), function(x
interactome.go_rna_unrelated.B.df$count <- sapply(unique(interactome.go_rna_unrelated.df$B), function(x
# using WCL as background
ftl <- apply(interactome.go_rna_unrelated.B.df, 1, function (x) {
  ct <- as.integer(x["count"])</pre>
 tt <- as.integer(x["total"])</pre>
 m1 <- matrix(c(ct, tt, smpl - ct, bkgd - tt), 2, 2)
  fisher.test(m1, alternative = alternative)
})
interactome.go_rna_unrelated.B.df$ft_pval <- unlist(lapply(ftl, function(x) {x$p.value}))
interactome.go_rna_unrelated.B.df$ft_OR <- unlist(lapply(ftl, function(x) {x$estimate}))</pre>
interactome.go_rna_unrelated.B.df$ft_fdr <- p.adjust(interactome.go_rna_unrelated.B.df$ft_pval, method
save(interactome.go_rna_unrelated.B.df, file = "data/interactome.go_rna_unrelated.B.df")
Now, RNA-related:
#-----GO RNA related-----
interactome.go_rna_related <- interactome[-which(interactome$GO == "unrelated"),]</pre>
interactome.go_rna_related.entrezIDs <- unique(interactome.go_rna_related[!is.na(interactome.go_rna_rel
interactome.go_rna_related.keggIDs <- keggConv.batch(interactome.go_rna_related.entrezIDs)</pre>
# we are testing this subset of "interactome", therefore we include all the pathways from "interactome"
interactome.go_rna_related.df <- interactome.df</pre>
i1 <- intersect(rownames(interactome.go_rna_related.df), rownames(wcl.df))</pre>
```

```
interactome.go_rna_related.df$total <- rep(0, nrow(interactome.go_rna_related.df))</pre>
interactome.go_rna_related.df[i1,]$total <- wcl.df[i1,]$count</pre>
interactome.go_rna_related.df$source <- rep("GO_RNA_related", nrow(interactome.go_rna_related.df))</pre>
interactome.go_rna_related.df$ID <- rownames(interactome.go_rna_related.df)</pre>
interactome.go_rna_related.df$count <- rep(0, nrow(interactome.go_rna_related.df))</pre>
interactome.go_rna_related.df$frac <- rep(0, nrow(interactome.go_rna_related.df))</pre>
for (i in rownames(interactome.go_rna_related.df)) {
  kL1 <- keggLink("mmu", paste("mmu", i, sep = ""))</pre>
  interactome.go_rna_related.df[i, ]$count <- length(which(interactome.go_rna_related.keggIDs %in% kL1)
}
# extract list of IDs in pathway
interactome.go_rna_related.in_path.IDs <- lapply(rownames(interactome.go_rna_related.df), function(x) {
  kL1 <- keggLink("mmu", paste("mmu", x, sep = ""))</pre>
  in_path <- interactome.go_rna_related.keggIDs[which(interactome.go_rna_related.keggIDs %in% kL1)]</pre>
})
names(interactome.go_rna_related.in_path.IDs) <- rownames(interactome.go_rna_related.df)</pre>
# perform Fisher's Exact Test for each category
# Using WCL as background
bkgd <- length(unique(wcl.keggIDs))</pre>
smpl <- length(interactome.go_rna_related.keggIDs)</pre>
ftl <- apply(interactome.go_rna_related.df, 1, function (x) {
  ct <- as.integer(x["count"])</pre>
 tt <- as.integer(x["total"])</pre>
 m1 <- matrix(c(ct, tt, smpl - ct, bkgd - tt), 2, 2)
  fisher.test(m1, alternative = alternative)
})
interactome.go_rna_related.df$ft_pval <- unlist(lapply(ftl, function(x) {x$p.value}))</pre>
interactome.go_rna_related.df$ft_OR <- unlist(lapply(ftl, function(x) {x$estimate}))</pre>
interactome.go_rna_related.df$ft_fdr <- p.adjust(interactome.go_rna_related.df$ft_pval, method = p.adju
save(interactome.go_rna_related.df, file = "data/interactome.go_rna_related.df")
# summarizing data at "B" level before doing Fisher's Exact test
interactome.go_rna_related.B.df <- data.frame(matrix(ncol = 5, nrow = length(unique(interactome.go_rna_
colnames(interactome.go_rna_related.B.df) <- c("B", "A", "total", "count", "source")</pre>
interactome.go\_rna\_related.B.df\$B <- \ unique(interactome.go\_rna\_related.df\$B)
interactome.go_rna_related.B.df$A <- sapply(unique(interactome.go_rna_related.df$B), function(x) {A <-
interactome.go_rna_related.B.df$source <- rep("GO_RNA_related", nrow(interactome.go_rna_related.B.df))</pre>
interactome.go_rna_related.B.df$total <- sapply(unique(interactome.go_rna_related.df$B), function(x) {t
interactome.go_rna_related.B.df$count <- sapply(unique(interactome.go_rna_related.df$B), function(x) {c
ftl <- apply(interactome.go_rna_related.B.df, 1, function (x) {</pre>
  ct <- as.integer(x["count"])</pre>
  tt <- as.integer(x["total"])</pre>
 m1 \leftarrow matrix(c(ct, tt, smpl - ct, bkgd - tt), 2, 2)
 fisher.test(m1, alternative = alternative)
})
interactome.go_rna_related.B.df$ft_pval <- unlist(lapply(ftl, function(x) {x$p.value}))
```

```
interactome.go_rna_related.B.df$ft_OR <- unlist(lapply(ftl, function(x) {x$estimate}))
interactome.go_rna_related.B.df$ft_fdr <- p.adjust(interactome.go_rna_related.B.df$ft_pval, method = p.
save(interactome.go_rna_related.B.df, file = "data/interactome.go_rna_related.B.df.rda")</pre>
```

Plotting the results of the enrichment analysis:

```
library(ggplot2)
library(grid)
library(scales)
load("data/interactome.df.rda")
load("data/interactome.go_rna_unrelated.df.rda")
load("data/interactome.go_rna_related.df")
dfC <- rbind(interactome.df[, c("A", "B", "C", "ft_OR", "ft_fdr", "source", "count")],</pre>
              interactome.go_rna_related.df[, c("A", "B", "C", "ft_OR", "ft_fdr", "source", "count")],
              interactome.go_rna_unrelated.df[, c("A", "B", "C", "ft_OR", "ft_fdr", "source", "count")]
)
dfC$source <- as.factor(dfC$source)</pre>
dfC$source <- factor(dfC$source, levels = levels(dfC$source)[c(3,1,2)])</pre>
select1 <- unique(as.character(dfC[which(dfC$ft_fdr <= 0.1 & dfC$ft_OR > 1),]$C))
select1.pathIDs <- paste("mmu", unlist(lapply(strsplit(select1, "\\ "), function(x) x[1])), sep = "")</pre>
dfC <- dfC[which(dfC$C %in% select1),]</pre>
dfC$C <- as.factor(as.character(dfC$C))</pre>
dfCf_0R.cut \leftarrow cut(log2(dfCf_0R), breaks = c(-Inf,-4:4), right = F)
dfC$C <- factor(dfC$C, levels = levels(dfC$C)[dfC[dfC$source == "Interactome", "C"][order(dfC[which(dfC
# formatting labels etc for plotting
11 <- levels(dfC$ft_OR.cut)</pre>
11 <- gsub("\\[", "", 11)</pre>
11 <- gsub("\\)", "", 11)</pre>
levels(dfC$ft OR.cut) <- 11</pre>
11 <- as.character(levels(dfC$C))</pre>
11 <- unlist(lapply(strsplit(l1, " "), function(x) {</pre>
  for (i in 2:length(x)){
    if (i == 2){
      v \leftarrow x[i]
    } else {
      v <- paste(v, x[i])</pre>
    }
  }
  return(v)
}))
levels(dfC$C) <- 11</pre>
levels(dfC$source)[2:3] <- c("RNA-related", "RNA-unrelated")</pre>
levels(dfC$C)[3] <- "Ribosome biogenesis"</pre>
```

```
levels(dfC$C)[6] <- "TCA cycle"</pre>
levels(dfC$C)[7] <- "mRNA surveillance"</pre>
levels(dfC$C)[11] <- "AA biosynthesis"</pre>
levels(dfC$C)[8] <- "H. simplex infection"</pre>
levels(dfC$C)[9] <- "Antibiotic biosynthesis"</pre>
levels(dfC$C)[12] <- "Glycolysis/Gluconeogenesis"</pre>
flevels <- levels(dfC$source)</pre>
11 \leftarrow factor(dfC\$C, levels = levels(dfC\$C)[c(1,2,3,7,4,5,6,12,8,9,10,11)])
dfC$C <- 11
p1 <- ggplot(data = dfC, aes(y = source, x = C)) +
      geom_tile(aes(fill = ft_OR.cut), colour = "white") +
      scale_fill_manual(values = brewer_pal(pal = "PuOr")(8), labels = levels(dfC\ft_OR.cut)) + #
      theme(axis.text.y = element_text(angle = 0, size = 8), axis.title = element_blank()) +
      guides(fill = guide_legend(label.position = "bottom", direction = "horizontal")) +
      theme(axis.text.x = element_text(angle = 45, vjust = 0.9, hjust = 0.8, size = 10)) +
      labs(fill = "Log2 OR") +
      scale_y_discrete(limits = rev(flevels)) +
      theme(legend.position = c(0.4, -1.92),
            legend.text = element_text(size = 4),
            legend.text.align = 0.5,
            legend.title = element_text(size = 4, vjust = 5),
            legend.key.size = unit(3.5, "mm"),
            legend.key.width = unit(3.5, "mm"),
            legend.margin = unit(0, "mm"),
            panel.margin = unit(1, "mm")) +
      ggtitle("KEGG pathway enrichment")
plot(p1)
```

