Expanded_Design_Factor_SpeciesHomologs_Outputs_SpeciesN

Load required libraries

```
library(plyr)
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:plyr':
##
##
       arrange, count, desc, failwith, id, mutate, rename, summarise,
##
       summarize
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
library(tibble)
library(stringr)
library(biomaRt)
library(genefilter)
library(DESeq2)
## Loading required package: S4Vectors
## Loading required package: stats4
## Loading required package: BiocGenerics
## Loading required package: parallel
##
## Attaching package: 'BiocGenerics'
## The following objects are masked from 'package:parallel':
##
##
       clusterApply, clusterApplyLB, clusterCall, clusterEvalQ,
##
       clusterExport, clusterMap, parApply, parCapply, parLapply,
       parLapplyLB, parRapply, parSapply, parSapplyLB
## The following objects are masked from 'package:dplyr':
##
       combine, intersect, setdiff, union
##
## The following objects are masked from 'package:stats':
##
##
       IQR, mad, sd, var, xtabs
## The following objects are masked from 'package:base':
##
##
       anyDuplicated, append, as.data.frame, basename, cbind,
##
       colMeans, colnames, colSums, dirname, do.call, duplicated,
##
       eval, evalq, Filter, Find, get, grep, grepl, intersect,
```

```
##
       is.unsorted, lapply, lengths, Map, mapply, match, mget, order,
##
       paste, pmax, pmax.int, pmin, pmin.int, Position, rank, rbind,
##
       Reduce, rowMeans, rownames, rowSums, sapply, setdiff, sort,
##
       table, tapply, union, unique, unsplit, which, which.max,
##
       which.min
##
## Attaching package: 'S4Vectors'
## The following objects are masked from 'package:dplyr':
##
##
       first, rename
## The following object is masked from 'package:plyr':
##
##
       rename
## The following object is masked from 'package:base':
##
##
       expand.grid
## Loading required package: IRanges
##
## Attaching package: 'IRanges'
## The following objects are masked from 'package:dplyr':
##
##
       collapse, desc, slice
## The following object is masked from 'package:plyr':
##
##
       desc
## Loading required package: GenomicRanges
## Loading required package: GenomeInfoDb
## Loading required package: SummarizedExperiment
## Loading required package: Biobase
## Welcome to Bioconductor
##
##
       Vignettes contain introductory material; view with
##
       'browseVignettes()'. To cite Bioconductor, see
       'citation("Biobase")', and for packages 'citation("pkgname")'.
##
## Loading required package: DelayedArray
## Loading required package: matrixStats
## Attaching package: 'matrixStats'
## The following objects are masked from 'package:Biobase':
##
##
       anyMissing, rowMedians
## The following objects are masked from 'package:genefilter':
##
##
       rowSds, rowVars
```

```
## The following object is masked from 'package:dplyr':
##
##
       count
## The following object is masked from 'package:plyr':
##
##
## Loading required package: BiocParallel
##
## Attaching package: 'DelayedArray'
## The following objects are masked from 'package:matrixStats':
##
       colMaxs, colMins, colRanges, rowMaxs, rowMins, rowRanges
##
## The following objects are masked from 'package:base':
##
       aperm, apply
library(gplots)
##
## Attaching package: 'gplots'
## The following object is masked from 'package: IRanges':
##
##
       space
## The following object is masked from 'package:S4Vectors':
##
       space
## The following object is masked from 'package:stats':
##
##
       lowess
library(ggplot2)
library(RColorBrewer)
library(stringr)
library(viridis)
## Loading required package: viridisLite
library(devtools)
library(ggrepel)
library(reshape2)
library(data.table)
## Attaching package: 'data.table'
## The following objects are masked from 'package:reshape2':
##
##
       dcast, melt
## The following object is masked from 'package:SummarizedExperiment':
##
       shift
```

```
## The following object is masked from 'package:GenomicRanges':
##
       shift
##
## The following object is masked from 'package: IRanges':
##
##
## The following objects are masked from 'package:S4Vectors':
##
##
       first, second
## The following objects are masked from 'package:dplyr':
##
##
       between, first, last
library(purrr)
##
## Attaching package: 'purrr'
## The following object is masked from 'package:data.table':
##
##
       transpose
## The following object is masked from 'package:DelayedArray':
##
##
       simplify
## The following object is masked from 'package:GenomicRanges':
##
##
       reduce
## The following object is masked from 'package: IRanges':
##
       reduce
##
## The following object is masked from 'package:plyr':
##
##
       compact
```

Purpose

To compare the DGE profiles created in "Expanded_Design_Factor_SpeciesHomologs_Outputs_SpeciesMapped".

```
dplyr::filter(abs(log2FoldChange) >= 3)
 У
}
DGEs_filtered <- llply(DGE_readin, filtering)</pre>
llply(DGEs_filtered, function(x) nrow(x))
## $bonobo
## [1] 191
## $chimpanzee
## [1] 166
##
## $gorilla
## [1] 146
##
## $olive_baboon
## [1] 341
##
## $orangutan
## [1] 227
## $pigtailed_macaque
## [1] 285
##
## $rhesus macaque
## [1] 321
##
## $squirrel_monkey
## [1] 461
##Are there genes in common across all of these?
common_DGEs_all <- map(DGEs_filtered, ~.$X) %>% purrr::reduce(intersect)
common_DGEs_all
## [1] "ENSG00000084628" "ENSG00000118113" "ENSG00000141668" "ENSG00000155629"
## [5] "ENSG00000167105" "ENSG00000168334" "ENSG00000243232"
##What are the expression patterns of these common_DGEs_all across species?
common_DGE_values <- llply(DGEs_filtered, function(x) x[x$X %in% common_DGEs_all,])</pre>
sapply(common_DGE_values, '[[', 'SYMBOL')
##
        bonobo
                  chimpanzee gorilla
                                       olive_baboon orangutan
## [1,] "NKAIN1" "NKAIN1"
                             "NKAIN1"
                                       "NKAIN1"
                                                     "NKAIN1"
## [2,] "MMP8"
                  "MMP8"
                             "MMP8"
                                        "MMP8"
                                                     "MMP8"
## [3,] "CBLN2"
                  "CBLN2"
                             "CBLN2"
                                        "CBLN2"
                                                     "CBLN2"
## [4,] "PIK3AP1" "PIK3AP1"
                             "PIK3AP1" "PIK3AP1"
                                                     "PIK3AP1"
## [5,] "TMEM92" "TMEM92"
                             "TMEM92"
                                       "TMEM92"
                                                     "TMEM92"
## [6,] "XIRP1"
                  "XIRP1"
                             "XIRP1"
                                        "XIRP1"
                                                     "XIRP1"
## [7,] "PCDHAC2" "PCDHAC2" "PCDHAC2"
                                                     "PCDHAC2"
       pigtailed_macaque rhesus_macaque squirrel_monkey
##
## [1,] "NKAIN1"
                          "NKAIN1"
                                         "NKAIN1"
## [2,] "MMP8"
                          "MMP8"
                                          "MMP8"
## [3,] "CBLN2"
                          "CBLN2"
                                          "CBLN2"
```

```
## [4,] "PIK3AP1"
                          "PIK3AP1"
                                         "PIK3AP1"
## [5,] "TMEM92"
                          "TMEM92"
                                         "TMEM92"
## [6,] "XIRP1"
                          "XIRP1"
                                         "XIRP1"
## [7,] "PCDHAC2"
                          "PCDHAC2"
                                         "PCDHAC2"
sapply(common_DGE_values, '[[', 'log2FoldChange')
##
           bonobo chimpanzee gorilla olive_baboon orangutan
## [1,] -4.418091 -5.140629 -3.542261 -7.543956 -7.768750
## [2,] -3.043676 -3.149779 -4.434808
                                         -4.584951 -6.665350
## [3,] -4.541821 -5.454281 -4.647792
                                          -5.865005 -5.006622
## [4,] -4.741324 -7.637673 -3.041881 -9.114815 -8.970609
## [5,] -3.373311 -3.682827 -3.027787 -3.452867 -5.665877
## [6,] -6.307787 -5.383328 -4.841533
                                         -6.584460 -5.005451
## [7,] -5.332360 -5.352416 -3.135160
                                          -5.704674 -5.208426
##
       pigtailed_macaque rhesus_macaque squirrel_monkey
               -6.876403
## [1,]
                              -6.923015
                                               -6.388892
## [2,]
               -5.837492
                              -4.434709
                                               -3.997017
## [3,]
               -4.418456
                               -3.356138
                                               -4.681466
## [4,]
               -8.829510
                             -10.470970
                                               -3.619223
## [5,]
               -4.414538
                              -3.970958
                                               -4.906825
## [6,]
               -4.560612
                               -6.426313
                                               -4.979683
## [7,]
               -4.387420
                               -4.314616
                                               -4.815433
##All the values are negative, indicating these genes' expression is less than what is observed in huma
zoo <- llply(DGEs_filtered, function(x) dplyr::select(x, X))</pre>
names(zoo) <- c("bonobo", "chimp", "gorilla", "baboon", "orangutan", "pigtailed", "rhesus", "squirrel_m</pre>
zoo_log2FC <- llply(DGEs_filtered, function(x) dplyr::select(x, X, log2FoldChange))</pre>
names(zoo_log2FC) <- c("bonobo", "chimp", "gorilla", "baboon", "orangutan", "pigtailed", "rhesus", "squ</pre>
zoo_log2FC_df <- zoo_log2FC %>%
 purrr::reduce(full_join, by = "X")
## Warning: Column `X` joining factors with different levels, coercing to
## character vector
## Warning: Column `X` joining character vector and factor, coercing into
## character vector
## Warning: Column `X` joining character vector and factor, coercing into
## character vector
## Warning: Column `X` joining character vector and factor, coercing into
## Warning: Column `X` joining character vector and factor, coercing into
## character vector
## Warning: Column `X` joining character vector and factor, coercing into
## character vector
## Warning: Column `X` joining character vector and factor, coercing into
## character vector
```

```
colnames(zoo_log2FC_df) <- c("X", names(zoo))</pre>
zoo_log2FC_df$`Average Expression` <- rowMeans(zoo_log2FC_df[,2:9], na.rm=TRUE)</pre>
zoo_log2FC_df[is.na(zoo_log2FC_df)] <- 0</pre>
counter <- function(x) {</pre>
 y <- x %>% add_count(X)
 У
}
zoo_counter <- llply(zoo, counter)</pre>
for (i in seq_along(zoo_counter)){
  colnames(zoo_counter[[i]]) <- c("X", names(zoo_counter[i]))</pre>
zoo_df <- zoo_counter %>%
 purrr::reduce(full_join, by = "X")
## Warning: Column `X` joining factors with different levels, coercing to
## character vector
## Warning: Column `X` joining character vector and factor, coercing into
## character vector
## Warning: Column `X` joining character vector and factor, coercing into
## character vector
## Warning: Column `X` joining character vector and factor, coercing into
## character vector
## Warning: Column `X` joining character vector and factor, coercing into
## character vector
## Warning: Column `X` joining character vector and factor, coercing into
## character vector
## Warning: Column `X` joining character vector and factor, coercing into
## character vector
zoo_df[is.na(zoo_df)] <- 0</pre>
zoo_df <- zoo_df[c("X", "chimp", "bonobo", "gorilla", "orangutan", "baboon", "rhesus", "pigtailed", "sq
zoo_avg_df \leftarrow full_join(zoo_log2FC_df[c(1, 10)], zoo_df, by = "X")
write.csv(zoo_avg_df, paste(Sys.Date(), "SpeciesMapped_SummaryCrossSpeciesExpression.csv"))
##Now need to load in the DGEs for each species treated vs mock (not relative to human)
##and then the corresponding human DGEs limited to the one-to-one orthologs for that species.
SpeciesMapped_DGE <- "Expanded_Design_Factor_SpeciesHomologs_Outputs_SpeciesMapped"
output_dir <- "Expanded_Design_Factor_SpeciesHomologs_Outputs_SpeciesMapped"</pre>
sampleFiles_SpeciesMapped <- list.files(basename(Sys.glob(file.path(SpeciesMapped_DGE))),</pre>
                  pattern = "*treated dds1 DGE results.txt|*related.*.txt")
sampleNames_SpeciesMapped <- sub('_treated_v_mock|.treatmenttreated.*', '', sampleFiles_SpeciesMapped)</pre>
```

```
sub('\\d+-\\d+-\\d+\\s', '', .) %>%
  sub("SpeciesMapped.*", "", .) %>%
  sub('species', '', .)
length(sampleFiles_SpeciesMapped)
## [1] 16
reader <- function(files) {</pre>
 d <- read.delim(files)</pre>
}
feature human innate <- read.csv("2019-10-20 InnateDBGeneFeatures.csv")
Species_DGEs <- 1lply(file.path(SpeciesMapped_DGE, sampleFiles_SpeciesMapped), reader)</pre>
names(Species_DGEs) <- sampleNames_SpeciesMapped</pre>
##Sorting the DGEs by species so that a given NHP species DGEs are with the corresponding
##human DGEs for that species.
bab_cts <- Species_DGEs[grep("baboon", names(Species_DGEs))]</pre>
bonobo_cts <- Species_DGEs[grep("bonobo", names(Species_DGEs))]</pre>
orangutan_cts <- Species_DGEs[grep("orangutan", names(Species_DGEs))]
chimp_cts <- Species_DGEs[grep("chimpanzee", names(Species_DGEs))]</pre>
sqm_cts <- Species_DGEs[grep("squirrel", names(Species_DGEs))]</pre>
rhmac_cts <- Species_DGEs[grep("rhesus", names(Species_DGEs))]</pre>
ptmac_cts <- Species_DGEs[grep("pigtailed", names(Species_DGEs))]</pre>
gorilla cts <- Species DGEs[grep("gorilla", names(Species DGEs))]</pre>
##Pulling the genes only differentially expressed compared to human for each NHP species
names(zoo_df)
## [1] "X"
                                                                "gorilla"
                          "chimp"
                                             "bonobo"
## [5] "orangutan"
                                             "rhesus"
                          "baboon"
                                                                "pigtailed"
## [9] "squirrel_monkey"
chimp <- zoo_df %>% filter_at(., 3:9, all_vars(. == 0))
bonobo \leftarrow zoo_df %>% filter_at(., c(2, 4:9), all_vars(. == 0))
gorilla <- zoo_df %>% filter_at(., c(2:3, 5:9), all_vars(. == 0))
orang <- zoo_df %>% filter_at(., c(2:4, 6:9), all_vars(. == 0))
bab <- zoo_df %>% filter_at(., c(2:5, 7:9), all_vars(. == 0))
rhesus <- zoo_df %>% filter_at(., c(2:6, 8:9), all_vars(. == 0))
ptmac <- zoo_df %>% filter_at(., c(2:7, 9), all_vars(. == 0))
sqm <- zoo_df %>% filter_at(., c(2:8), all_vars(. == 0))
zoo_list <- list(bonobo, orang, chimp, gorilla, sqm, rhesus, ptmac, bab)
##Our different groups we want to examine
OWMs <- zoo_df %>% filter_at(., c(6:8), all_vars(. == 1 )) %>%
```

```
filter_at(., c(2:5, 9), all_vars(. == 0))
monkeys <- zoo_df %>% filter_at(., c(6:9), all_vars(. == 1 )) %>%
  filter_at(., c(2:5), all_vars(. == 0))
all_species <- zoo_df %>% filter_at(., c(2:9), all_vars(. == 1))
#################################
##Genes that are differentially expressed compared to human in ALL species
all_cts <- list(bonobo_cts, orangutan_cts, chimp_cts, gorilla_cts, sqm_cts, rhmac_cts, ptmac_cts, bab_c
names(all_cts) <- c("bonobo", "orangutan", "chimpanzee", "gorilla", "squirrel_monkey", "rhesus_macaque"</pre>
                     "olive_baboon")
all_DGE <- llply(all_cts, llply, function(x) (x[x$X %in% all_species$X, ])) %>%
  llply(., llply, function(x) dplyr::select(x, X, SYMBOL, padj, log2FoldChange))
all_DGE_names <- all_DGE
for (i in 1:length(all_DGE)) {
 all_DGE[[i]] <- all_DGE[[i]] %>% purrr::reduce(full_join, by = "X")
all_DGE_limited <- all_DGE %>% llply(., function(x) x[c(1:4, 6:7)]) ##then select columns we care about
                                                  ##and log2FoldChange for the NHP species and the relate
                                         ##each data frame within the list
##Now rename the columns within the data frames in our list so that they make sense once we melt it dow.
##for plotting.
colnames <- lapply(seq_along(all_DGE_limited), function(i)</pre>
  colnames(all_DGE_limited[[i]]) <-</pre>
    c("X", "SYMBOL", paste("padj_species"), names(all_DGE_names[[i]][1]),
      paste("padj_human"), names(all_DGE_names[[i]][2])))
for (i in 1:length(all_DGE_limited)) {
  names(all_DGE_limited[[i]]) <- colnames[[i]]</pre>
}
##Order the symbols as desired, with the human DGEs related to a given species in
## order from highest to lowest log2FoldChange.
significance <- function(x) {</pre>
  y <- dplyr::arrange(x, desc(x[,4]))
  y$padj_species <- ifelse((y$padj_species > 0.05), "ns", "sig")
  y$padj_human <- ifelse((y$padj_human > 0.05), "ns", "sig")
}
all_DGE_ordered <- llply(all_DGE_limited, significance)</pre>
names(all_DGE_ordered) <- names(all_cts)</pre>
##Melt it all down.
all_species.m <- melt(all_DGE_ordered)</pre>
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
```

```
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
##Break apart all_species.m into the constituent melted species so we can then re-set the symbol order
##we want based on what we did for all_DGE_ordered above.
leveler_species <- function(DGE_set, X) {</pre>
species.m <- filter(all_species.m, L1 == !!X)</pre>
species.m$SYMBOL <- factor(species.m$SYMBOL, levels = DGE_set$SYMBOL[order(DGE_set[,4])])</pre>
species.m
}
common_species <- mapply(leveler_species, all_DGE_ordered, names(all_cts), SIMPLIFY = FALSE) %>%
  do.call("rbind", .)
rownames(common_species) <- c()</pre>
common_species$padj_species <- ifelse(grepl("related", common_species$variable), "",</pre>
                                       common_species$padj_species)
common_species$padj_human <- ifelse(!grepl("related", common_species$variable), "",</pre>
                                     common_species$padj_human)
write.csv(common_species, file.path(output_dir,
        paste(Sys.Date(), "DifferFromHuman_SpeciesMapped_CommonAllSpecies.csv")))
##Need a "copy" of 'all.m_species' to create a geom_text plot along with our heatmap later. This idea c
##https://github.com/tidyverse/ggplot2/issues/2656 and the rest of it was modified and implemented belo
##making the faceted heat maps.
common_species_text <- common_species</pre>
##For every gene that is in our feature_human_innate data frame derived from InnateDB, we want to highl
##in our heatmap by making them in bold, italic face.
common_species_text$faces <- ifelse((common_species_text$X %in% feature_human_innate$hsapiens_homolog_e.
                                    'bold.italic', 'plain')
##Since we are making faceted plots, we only need one set of SYMBOLs shown per facet, so in our geom_te.
##we will discard any of the rows for variables that have "related" in its name, i.e. getting rid of th
##of symbols for a given species.
common_species_text <- dplyr::filter(common_species_text, !grepl(".*related",variable))</pre>
############################
##All monkeys
##Using the list above to limit our DGE profiles for each species
monkeys_cts <- list(sqm_cts, rhmac_cts, ptmac_cts, bab_cts)</pre>
names(monkeys_cts) <- c("squirrel_monkey", "rhesus_macaque", "pigtailed_macaque", "olive_baboon")</pre>
monkeys_DGE <- llply(monkeys_cts, llply, function(x) (x[x$X %in% monkeys$X, ])) %>%
  llply(., llply, function(x) dplyr::select(x, X, SYMBOL, padj, log2FoldChange))
##Could not for the life of me find an easier way to do this, just joining each list of data frames for
##each species so that you now have one data frame (instead of the original two) per species, containing
## the DGE for that species AND the DGE for the related human file.
monkeys_joined <- list((monkeys_DGE[[1]] %>% purrr::reduce(full_join, by = "X")), (monkeys_DGE[[2]] %>%
```

```
purrr::reduce(full_join, by = "X")), monkeys_DGE[[3]] %>%
                          purrr::reduce(full_join, by = "X"), monkeys_DGE[[4]] %>%
                          purrr::reduce(full_join, by = "X")) %>%
  liply(., function(x) x[c(1:4, 6:7)]) ##then select columns we care about (corresponds to X, SYMBOL, p
                                                  ##and log2FoldChange for the NHP species and the relate
                                         ##each data frame within the list
##Now rename the columns within the data frames in our list so that they make sense once we melt it dow.
##for plotting.
colnames <- lapply(seq_along(monkeys_joined), function(i)</pre>
    c("X", "SYMBOL", paste("padj_species"), names(monkeys_DGE[[i]][1]),
      paste("padj_human"), names(monkeys_DGE[[i]][2])))
for (i in 1:length(monkeys_joined)) {
  names(monkeys_joined[[i]]) <- colnames[[i]]</pre>
}
##Order the symbols as desired, with the human DGEs related to a given species in
## order from highest to lowest log2FoldChange.
  significance <- function(x) {</pre>
  y <- dplyr::arrange(x, desc(x[,4]))
 y$padj_species <- ifelse((y$padj_species > 0.05), "ns", "sig")
 y$padj_human <- ifelse((y$padj_human > 0.05), "ns", "sig")
}
monkeys_ordered <- llply(monkeys_joined, significance)</pre>
names(monkeys ordered) <- names(monkeys DGE)</pre>
##Melt it all down.
monkeys.m <- melt(monkeys_ordered)</pre>
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
leveler_monkeys <- function(DGE_set, X) {</pre>
species.m <- filter(monkeys.m, L1 == !!X)</pre>
species.m$SYMBOL <- factor(species.m$SYMBOL, levels = DGE_set$SYMBOL[order(DGE_set[,4])])</pre>
species.m
}
monkeys <- mapply(leveler_monkeys, monkeys_ordered, names(monkeys_ordered), SIMPLIFY = FALSE) %>%
  do.call("rbind", .)
rownames(monkeys) <- c()</pre>
monkeys$padj_species <- ifelse(grepl("related", monkeys$variable), "", monkeys$padj_species)
monkeys$padj_human <- ifelse(!grepl("related", monkeys$variable), "", monkeys$padj_human)</pre>
write.csv(monkeys, file.path(output_dir, paste(Sys.Date(), "DifferFromHuman_SpeciesMapped_AllMonkeys.cs
monkeys_text <- monkeys
monkeys_text$faces <- ifelse((monkeys_text$X %in% feature_human_innate$hsapiens_homolog_ensembl_gene),
                                    'bold.italic', 'plain')
monkeys_text <- dplyr::filter(monkeys_text, !grepl(".*related",variable))</pre>
```

```
##Only Old World monkeys
OWMs_cts <- list(rhmac_cts, ptmac_cts, bab_cts)</pre>
names(OWMs_cts) <- c("rhesus_macaque", "pigtailed_macaque", "olive_baboon")</pre>
OWMs_DGEs <- llply(OWMs_cts, llply, function(x) (x[x$X %in% OWMs$X, ])) %>%
  llply(., llply, function(x) dplyr::select(x, X, SYMBOL, padj, log2FoldChange))
##Could not for the life of me find an easier way to do this, just joining each list of data frames for
##each species so that you now have one data frame (instead of the original two) per species, containing
## the DGE for that species AND the DGE for the related human file.
OWMs_joined <- list((OWMs_DGEs[[1]] %>% purrr::reduce(full_join, by = "X")), (OWMs_DGEs[[2]] %>% purrr:
          OWMs_DGEs[[3]] %>% purrr::reduce(full_join, by = "X")) %>%
 liply(., function(x) x[c(1:4, 6:7)]) ##then select columns we care about (corresponds to X, SYMBOL and
                                       ##log2FoldChange for the NHP species and the related human file)
                                         ##each data frame within the list
##Now rename the columns within the data frames in our list so that they make sense once we melt it dow.
##for plotting.
colnames <- lapply(seq_along(OWMs_joined), function(i)</pre>
    c("X", "SYMBOL", paste("padj_species"), names(OWMs_DGEs[[i]][1]),
      paste("padj_human"), names(OWMs_DGEs[[i]][2])))
for (i in 1:length(OWMs_joined)) {
  names(OWMs_joined[[i]]) <- colnames[[i]]</pre>
##Order the symbols as desired, with the human DGEs related to a given species in
## order from highest to lowest log2FoldChange.
  significance <- function(x) {</pre>
  y <- dplyr::arrange(x, desc(x[,4]))
 y$padj_species <- ifelse((y$padj_species > 0.05), "ns", "sig")
 y$padj_human <- ifelse((y$padj_human > 0.05), "ns", "sig")
OWMs_ordered <- llply(OWMs_joined, significance)</pre>
names(OWMs ordered) <- names(OWMs DGEs)</pre>
##Melt it all down.
OWMs.m <- melt(OWMs_ordered)</pre>
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
leveler OWMs <- function(DGE set, X) {</pre>
species.m <- filter(OWMs.m, L1 == !!X)</pre>
species.m$SYMBOL <- factor(species.m$SYMBOL, levels = DGE_set$SYMBOL[order(DGE_set[,4])])</pre>
species.m
}
OWMs_all <- mapply(leveler_OWMs, OWMs_ordered, names(OWMs_ordered), SIMPLIFY = FALSE) %>%
 do.call("rbind", .)
```

```
rownames(OWMs_all) <- c()</pre>
OWMs_all$padj_species <- ifelse(grepl("related", OWMs_all$variable), "", OWMs_all$padj_species)
OWMs_all$padj_human <- ifelse(!grepl("related", OWMs_all$variable), "", OWMs_all$padj_human)
##Break apart OWMs.m into the constituent melted species so we can then re-set the symbol order to what
##we want based on what we did for OWMs ordered above.
write.csv(OWMs_all, file.path(output_dir, paste(Sys.Date(), "DifferFromHuman_SpeciesMapped_OWMsOnly.csv
OWMs all text <- OWMs all
OWMs_all_text$faces <- ifelse((OWMs_all_text$X %in% feature_human_innate$hsapiens_homolog_ensembl_gene)
OWMs_all_text <- dplyr::filter(OWMs_all_text, !grepl(".*related",variable))</pre>
##Pulling out the DGEs relative to human that are found in only one species at a time as determined
##above in the lines getting "chimp <-, "bonobo <-", etc
ind_differ_DGEs <- function(ctsX, speciesX) {</pre>
species_DGE <- llply(ctsX, function(x) x[x$X %in% speciesX$X,]) %>%
 llply(., function(x) dplyr::select(x, X, SYMBOL, padj, log2FoldChange)) %>%
 purrr::reduce(full_join, by = "X") %>%
    dplyr::select(X, SYMBOL.x, padj.x, padj.y, log2FoldChange.x, log2FoldChange.y)
colnames(species_DGE) <- c("X", "SYMBOL", "padj_species", "padj_human", names(ctsX))</pre>
species_DGE$SYMBOL <- str_replace(species_DGE$SYMBOL, ",.*", "")</pre>
species_DGE_ordered <- dplyr::arrange(species_DGE, desc(species_DGE[,5])) ##order by column containing
                                          ##the log2FC values for the human samples
species_DGE_ordered$padj_species <- ifelse((species_DGE_ordered$padj_species > 0.05), "ns", "sig")
species_DGE_ordered$padj_human <- ifelse((species_DGE_ordered$padj_human > 0.05), "ns", "sig")
species_DGE_ordered
all_DGEs <- mapply(ind_differ_DGEs, all_cts, zoo_list, SIMPLIFY = FALSE)
all_DGEs.m <- melt(all_DGEs)</pre>
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj species, padj human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
Na_Values <- all_DGEs.m[is.na(all_DGEs.m$SYMBOL),] ##get positions of NA values in the SYMBOL column
Na_Values_chimp <- all_DGEs$chimpanzee[is.na(all_DGEs$chimpanzee$SYMBOL),] ##get positions of NA
                        ##values in the SYMBOL column
levels(all_DGEs$chimpanzee$SYMBOL) <- c(levels(all_DGEs$chimpanzee$SYMBOL), "RN7SKP175")</pre>
all_DGEs$chimpanzee[36,2] = c("RN7SKP175")
```

```
Na_Values_bonobo <- all_DGEs$bonobo[is.na(all_DGEs$bonobo$SYMBOL),] ##get positions of NA values in the
levels(all_DGEs$bonobo$SYMBOL) <- c(levels(all_DGEs$bonobo$SYMBOL), "RNU1-125P")</pre>
all DGEs\$bonobo[34,2] = c("RNU1-125P")
Na Values baboon <- all DGEs$olive baboon[is.na(all DGEs$olive baboon$SYMBOL),] ##get positions of NA
                         ##values in the SYMBOL column
levels(all DGEs$olive baboon$SYMBOL) <- c(levels(all DGEs$olive baboon$SYMBOL), "AL592183.1")
all DGEs\$olive baboon[64,2] = c("AL592183.1")
Na_Values_gorilla <- all_DGEs$gorilla[is.na(all_DGEs$gorilla$SYMBOL),] ##get positions of NA values in
                                                          ##the SYMBOL column
levels(all_DGEs$gorilla$SYMBOL) <- c(levels(all_DGEs$gorilla$SYMBOL), "AC244197.3", "RF01233")</pre>
all_DGEs$gorilla[6,2] = c("AC244197.3")
all_DGEs$gorilla[55,2] = c("RF01233")
Na_Values_sqm <- all_DGEs$squirrel_monkey[is.na(all_DGEs$squirrel_monkey$SYMBOL),] ##get positions of
                                               ##NA values in the SYMBOL column
levels(all_DGEs$squirrel_monkey$SYMBOL) <- c(levels(all_DGEs$squirrel_monkey$SYMBOL), "RF01986")</pre>
all DGEs$squirrel monkey[155,2] = c("RF01986")
Na Values orangutan <- all DGEs$orangutan[is.na(all DGEs$orangutan$SYMBOL),] ##get positions of NA
                               ##values in the SYMBOL column
levels(all DGEs$orangutan$SYMBOL) <- c(levels(all DGEs$orangutan$SYMBOL), "ACO25283.2")</pre>
all_DGEs$orangutan[38,2] = c("ACO25283.2")
all_DGEs.m <- melt(all_DGEs)</pre>
## Using X, SYMBOL, padj species, padj human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
## Using X, SYMBOL, padj_species, padj_human as id variables
write.csv(all_DGEs.m, file.path(output_dir,
                   paste(Sys.Date(), "DifferFromHuman_SpeciesMapped_EachSpeciesUnique.csv")))
##Break apart all_DGEs.m into the constituent melted species so we can then re-set the symbol order to
##what we want based on what we did for all_DGEs above.
leveler <- function(DGE_set, X) {</pre>
species.m <- filter(all_DGEs.m, L1 == !!X)</pre>
species.m$SYMBOL <- factor(species.m$SYMBOL, levels = DGE set$SYMBOL[order(DGE set[,5])])</pre>
species.m
}
fewer <- mapply(leveler, all_DGEs, names(all_DGEs), SIMPLIFY = FALSE) %>%
 do.call("rbind", .)
```

```
rownames(fewer) <- c()</pre>
fewer$padj_species <- ifelse(grepl("related", fewer$variable), "", fewer$padj_species)
fewer$padj_human <- ifelse(!grepl("related", fewer$variable), "", fewer$padj_human)</pre>
##Making a fewer_text as done above
fewer_text <- fewer</pre>
fewer text$faces <- ifelse((fewer text$X %in% feature human innate$hsapiens homolog ensembl gene),
                           'bold.italic', 'plain')
fewer_text <- dplyr::filter(fewer_text, !grepl(".*related",variable))</pre>
output dir <- "Expanded Design Factor SpeciesHomologs Outputs SpeciesMapped"
##Genes for 6/8 species that were uniquely (i.e. only observed in each of these species
##and no other) differentially expressed upon poly(I:C) treatment compared to human.
fewer_filtered <- dplyr::filter(fewer, !grepl("baboon|squirrel", variable)) %>% droplevels()
fewer_text_filtered <- dplyr::filter(fewer_text, !grepl("baboon|squirrel", variable)) %>% droplevels()
fewer_filtered$variable <- with(fewer_filtered,</pre>
    factor(variable,levels = c("pigtailed_macaque","human_related_pigtailed_macaque ",
                                "rhesus_macaque", "human_related_rhesus_macaque ",
                               "gorilla", "human_related_gorilla ",
                              "orangutan", "human_related_orangutan ",
                              "bonobo", "human_related_bonobo ",
                              "chimpanzee", "human related chimpanzee ")))
fewer_filtered$L1 <- with(fewer_filtered, factor(L1,</pre>
   levels = c("chimpanzee", "bonobo", "gorilla", "orangutan", "rhesus_macaque", "pigtailed_macaque")))
pdf(file = file.path(output_dir, paste(Sys.Date(), "differFromHuman_shortList_SpeciesMapped.pdf")),
   height =4.7, width =9.3)
ggplot(fewer filtered, aes(SYMBOL, variable, fill = value)) +
  geom tile(color = 'black') +
  theme_bw(base_size = 6) +
  theme(panel.spacing = unit(2.5, "lines")) +
  geom_text(data = fewer_text_filtered,
            aes(fontface = faces, label = SYMBOL, x = SYMBOL), y = 0.05, size = rel(2), hjust = 1,
            angle = 90) +
  ylab("") +
  xlab("\n") +
  facet_wrap(~L1, scales = "free", nrow = 6) +
  scale_fill_gradient2(low = "#e66101", high = "#542788", mid = "white", midpoint = 0,
                       limits = c(-6, 11)) +
  theme(axis.text.x = element blank(),
        axis.title.x = element_text(margin = margin(b = 25))) +
  coord cartesian(clip = "off") +
  theme(strip.background = element_blank(), strip.text = element_blank())
dev.off()
## pdf
##
##Genes from all the monkeys (Old and New World) that were uniquely (i.e. only observed in this
##group of species and no other) differentially expressed upon poly(I:C) treatment compared to human.
##Unwrapped plot
monkeys filtered <- filter(monkeys,
grepl("^human_related_olive_baboon|^squirrel_monkey|^rhesus_macaque|^olive_baboon|^pigtailed_macaque",
```

```
variable)) %>%
  droplevels(.)
monkeys_filtered$variable <- with(monkeys_filtered,</pre>
                        factor(variable,levels = c("squirrel_monkey", "rhesus_macaque",
                                                    "pigtailed_macaque", "olive_baboon",
                                                    "human related olive baboon ")))
pdf(file = file.path(output_dir,
                     paste(Sys.Date(), "differfromHuman Monkeys SpeciesMapped unwrapped.pdf")),
   height = 1.5, width = 5)
ggplot(monkeys_filtered, aes(SYMBOL, variable, fill = value)) +
  geom_tile(color = 'black', aes(height =1)) +
 theme_bw(base_size = 6) +
  geom_text(data = monkeys_text,
            aes(fontface = faces, label = SYMBOL, x = SYMBOL), y = 0.05,
            size = rel(2),
           hjust = 1, angle = 90) +
  ylab("") +
  xlab("\n") +
  scale_fill_gradient2(low = "#e66101", high = "#542788", mid = "white",
                       midpoint = 0, limits = c(-6, 11)) +
  theme(axis.text.x = element_blank(),
        axis.title.x = element_text(margin = margin(b = 25))) +
    coord cartesian(clip = "off")
dev.off()
## pdf
##
##Genes from all the monkeys (Old and New World) that were uniquely (i.e. only observed in this
##group of species and no other) differentially expressed upon poly(I:C) treatment compared to human.
##Wrapped plot
pdf(file = file.path(output_dir, paste(Sys.Date(), "differfromHuman_Monkeys_SpeciesMapped.pdf")),
   height = 4, width = 5)
ggplot(monkeys, aes(SYMBOL, variable, fill = value)) +
  geom_tile(color = 'black', aes(height =1)) +
  theme bw(base size = 6) +
  theme(panel.spacing = unit(2.2, "lines")) +
  geom_text(data = monkeys_text,
            aes(fontface = faces, label = SYMBOL, x = SYMBOL), y = 0.05,
            size = 0.8*6/.pt, # match font size to theme
           hjust = 1, angle = 90) +
  vlab("") +
  xlab("\n") +
  facet_wrap(~L1, scales = "free", nrow = 4) +
  scale_fill_gradient2(low = "#e66101", high = "#542788") +
  theme(axis.text.x = element_blank(),
        axis.title.x = element_text(margin = margin(b = 25))) +
  coord cartesian(clip = "off") +
  theme(strip.background = element rect(colour="black", fill="white", linetype="solid"))
dev.off()
## pdf
##
```

```
##Genes from baboon that were uniquely (i.e. only observed in baboon
##and no other) differentially expressed upon poly(I:C) treatment compared to human.
fewer_baboon <- dplyr::filter(fewer, grepl("baboon", variable)) %>% droplevels()
fewer_text_baboon <- dplyr::filter(fewer_text, grepl("baboon", variable)) %>% droplevels()
fewer_baboon$variable <- with(fewer_baboon,</pre>
                        factor(variable,levels = c("olive baboon",
                                                   "human_related_olive_baboon ")))
pdf(file = file.path(output_dir, paste(Sys.Date(), "differfromHuman_olivebaboon_SpeciesMapped.pdf")),
   height =0.8, width = 15)
ggplot(fewer_baboon, aes(SYMBOL, variable, fill = value)) +
  geom_tile(color = 'black') +
 theme_bw(base_size = 6) +
  geom_text(data = fewer_text_baboon,
            aes(fontface = faces, label = SYMBOL, x = SYMBOL), y = 0.02,
            size = rel(2), hjust = 1, angle = 90) +
  vlab("") +
  xlab("\n") +
  theme(legend.position = "none") +
  scale_fill_gradient2(low = "#e66101", high = "#542788", mid = "white",
                       midpoint = 0, limits = c(-6, 11)) +
  theme(axis.text.x = element_blank(),
        axis.title.x = element_text(margin = margin(b = 25))) +
  coord_cartesian(clip = "off") +
  theme(strip.background = element_blank(), strip.text = element_blank())
dev.off()
## pdf
##
##Genes from squirrel monkey that were uniquely (i.e. only observed in baboon
##and no other) differentially expressed upon poly(I:C) treatment compared to human.
fewer_sqm <- dplyr::filter(fewer, grep1("squirre1", variable)) %>% droplevels()
fewer_text_sqm <- dplyr::filter(fewer_text, grepl("squirrel", variable)) %>% droplevels()
fewer_sqm$variable <- with(fewer_sqm,</pre>
                        factor(variable,levels = c("squirrel_monkey",
                                                    "human related squirrel monkey ")))
pdf(file = file.path(output_dir, paste(Sys.Date(), "differfromHuman_squirrelmonkey_SpeciesMapped.pdf"))
   height =0.8, width = 20)
ggplot(fewer_sqm, aes(SYMBOL, variable, fill = value)) +
  geom_tile(color = 'black') +
  theme_bw(base_size = 6) +
  geom_text(data = fewer_text_sqm,
            aes(fontface = faces, label = SYMBOL, x = SYMBOL), y = 0.02,
            size = rel(2), hjust = 1, angle = 90) +
  ylab("") +
  xlab("\n") +
  theme(legend.position = "none") +
  scale_fill_gradient2(low = "#e66101", high = "#542788", mid = "white",
                       midpoint = 0, limits = c(-6.5, 11)) +
  theme(axis.text.x = element_blank(),
        axis.title.x = element_text(margin = margin(b = 25))) +
  coord_cartesian(clip = "off") +
  theme(strip.background = element_blank(), strip.text = element_blank())
dev.off()
```

```
## pdf
##
##Genes from all the Old World monkey that were uniquely (i.e. only observed in this
##group of species and no other) differentially expressed upon poly(I:C) treatment compared to human.
##Wrapped plot
pdf(file = file.path(output_dir, paste(Sys.Date(), "differfromHuman_OWMs_SpeciesMapped.pdf")),
   height = 3, width = 5)
ggplot(OWMs all, aes(SYMBOL, variable, fill = value)) +
  geom_tile(color = 'black', aes(height =1)) +
  theme bw(base size = 6) +
  theme(panel.spacing = unit(2.2, "lines")) +
  geom_text(data = OWMs_all_text,
            aes(fontface = faces, label = SYMBOL, x = SYMBOL), y = 0.05,
            size = 0.8*6/.pt, # match font size to theme
           hjust = 1, angle = 90) +
  ylab("") +
  xlab("\n") +
  facet_wrap(~L1, scales = "free", nrow = 3) +
  scale_fill_gradient2(low = "#e66101", high = "#542788") +
  theme(axis.text.x = element_blank(),
        axis.title.x = element_text(margin = margin(b = 25))) +
  coord_cartesian(clip = "off") +
  theme(strip.background = element_rect(colour="black", fill="white", linetype="solid"))
dev.off()
## pdf
##
##Genes from all the Old World monkey that were uniquely (i.e. only observed in this
##group of species and no other) differentially expressed upon poly(I:C) treatment compared to human.
##Unwrapped plot
OWMs all filtered <- filter(OWMs all,
grepl("^human_related_olive_baboon|^rhesus_macaque|^olive_baboon|^pigtailed_macaque", variable))
OWMs_all_filtered$variable <- with(OWMs_all_filtered,</pre>
                        factor(variable,levels = c("rhesus_macaque",
                                                    "pigtailed macaque", "olive baboon",
                                                   "human related olive baboon ")))
pdf(file = file.path(output_dir, paste(Sys.Date(), "differfromHuman_OWMs_SpeciesMapped_unwrapped.pdf"))
   height = 1.2, width = 5)
ggplot(OWMs_all_filtered, aes(SYMBOL, variable, fill = value)) +
  geom_tile(color = 'black', aes(height =1)) +
  theme_bw(base_size = 6) +
  geom_text(data = OWMs_all_text,
            aes(fontface = faces, label = SYMBOL, x = SYMBOL), y = 0.05,
            size = rel(2),
           hjust = 1, angle = 90) +
  ylab("") +
  xlab("\n") +
  scale fill gradient2(low = "#e66101", high = "#542788", mid = "white",
                       midpoint = 0, limits = c(-6, 11)) +
  theme(axis.text.x = element_blank(),
        axis.title.x = element_text(margin = margin(b = 25))) +
    coord cartesian(clip = "off")
dev.off()
```

```
## pdf
##
##Genes common to all the NHP species that were differentially expressed upon poly(I:C) treatment
##compared to human.
##Wrapped plot
common_species_filtered <- filter(common_species,</pre>
grepl("^human_related_olive|^rhesus|^olive|^pigtailed|^bonobo|^orangutan|^squirrel|^chimp|^gorilla",
      variable)) %>%
      droplevels()
common_species_filtered$variable <- with(common_species_filtered,</pre>
              factor(variable,levels = c("squirrel_monkey", "pigtailed_macaque", "rhesus_macaque",
                                         "olive_baboon", "orangutan", "gorilla", "bonobo",
                                          "chimpanzee", "human_related_olive_baboon ")))
pdf(file = file.path(output_dir, paste(Sys.Date(),
      "differfromHuman_all_species_SpeciesMapped_unwrapped.pdf")), height = 1.5, width = 2.7)
ggplot(common_species_filtered, aes(SYMBOL, variable, fill = value)) +
  geom_tile(color = 'black', aes(height = 1)) +
  theme_bw(base_size = 6) +
  geom_text(data = common_species_text,
            aes(fontface = faces, label = SYMBOL, x = SYMBOL), y = 0.05,
            size = rel(2),
           hjust = 1, angle = 90) +
  ylab("") +
  xlab("\n") +
  scale fill gradient2(low = "#e66101", high = "#542788", mid = "white", midpoint = 0,
                       limits = c(-6, 11) +
  theme(axis.text.x = element blank(),
        axis.title.x = element_text(margin = margin(b = 25))) +
    coord_cartesian(clip = "off")
dev.off()
## pdf
##Genes common to all the NHP species that were differentially expressed upon poly(I:C) treatment
##compared to human.
##Wrapped plot
pdf(file = file.path(output_dir, paste(Sys.Date(),
"differfromHuman_all_species_SpeciesMapped_wrapped.pdf")), height = 8, width = 4)
ggplot(common_species, aes(SYMBOL, variable, fill = value)) +
  geom_tile(color = 'black', aes(height =1)) +
  theme_bw(base_size = 6) +
  theme(panel.spacing = unit(2.2, "lines")) +
  geom_text(data = common_species_text,
            aes(fontface = faces, label = SYMBOL, x = SYMBOL), y = 0.05,
            size = 0.8*6/.pt, # match font size to theme
           hjust = 1, angle = 90) +
 ylab("") +
  xlab("\n") +
  facet_wrap(~L1, scales = "free", nrow = 8) +
  scale_fill_gradient2(low = "#e66101", high = "#542788") +
  theme(axis.text.x = element_blank(),
        axis.title.x = element text(margin = margin(b = 25))) +
  coord_cartesian(clip = "off") +
```

```
theme(strip.background = element_rect(colour="black", fill="white", linetype="solid"))
dev.off()
## pdf
##
    2
## Default upset() arguments
sets_of_interest <- list(list("squirrel_monkey"), list("baboon"), list("orangutan"), list("rhesus"),</pre>
                    list("bonobo"), list("chimp"), list("gorilla"), list("pigtailed"),
                    list("pigtailed", "rhesus", "baboon"),
                    list("pigtailed", "rhesus", "baboon", "squirrel_monkey"),
list("pigtailed", "rhesus", "baboon", "squirrel_monkey", "gorilla", "orangutan", "chimp", "bonobo"),
list("pigtailed", "rhesus"), list("rhesus", "squirrel_monkey"),
list("pigtailed", "rhesus", "squirrel_monkey"), list("orangutan", "squirrel_monkey"),
list("chimp", "bonobo"), list("orangutan", "pigtailed", "rhesus", "baboon", "squirrel monkey"))
dot colors <- c("purple", "darkgreen", "#bee1f4", "forestgreen", "#0088ce",
"#7dc3e8", "darkolivegreen3", "#00659c", "grey55", "orange", "orange", "grey55", "grey55",
 "grey55", "grey55", "grey55", "orange")
nsets = 8; nintersects = 40; sets = NULL; keep.order = F; set.metadata = NULL;
intersections = sets_of_interest; matrix.color = "grey24"; main.bar.color = "gray55";
mainbar.y.label = ""; mainbar.y.max = NULL; sets.bar.color = "grey24"; sets.x.label = "Set Size";
point.size = 2.5; line.size = 0.4; mb.ratio = c(0.70,0.30); expression = NULL; att.pos = NULL;
att.color = dot_colors; order.by = 'freq'; decreasing = T; show.numbers = "yes";
number.angles = 0; group.by = "degree"; cutoff = NULL; queries = NULL; query.legend = "none";
shade.color = "gray88"; shade.alpha = 0.25; matrix.dot.alpha = 0.5; sempty.intersections = NULL;
color.pal = 1; boxplot.summary = c("Average Expression"); attribute.plots = NULL;
scale.intersections = "identity"; scale.sets = "identity"; text.scale = 1.4; set_size.angles = 0 ;
set_size.show = FALSE; set_size.numbers_size = NULL; set_size.scale_max = NULL
data = as.data.frame(zoo_avg_df[2:10]); matrix.color="blue";
sets.bar.color = c("purple", "darkgreen", "forestgreen", "darkolivegreen3", "#bee1f4", "#0088ce",
                   "#00659c", "#7dc3e8")
## Modified internal upset() code
startend <- UpSetR:::FindStartEnd(data)</pre>
 first.col <- startend[1]</pre>
  last.col <- startend[2]</pre>
  if(color.pal == 1){
   palette <- c("#1F77B4", "#FF7F0E", "#2CA02C", "#D62728", "#9467BD", "#8C564B", "#E377C2",
                 "#7F7F7F", "#BCBD22", "#17BECF")
   palette <- c("#E69F00", "#56B4E9", "#009E73", "#F0E442", "#0072B2", "#D55E00",
                 "#CC79A7")
  if(is.null(intersections) == F){
   Set_names <- unique((unlist(intersections)))</pre>
   Sets to remove <- UpSetR:::Remove(data, first.col, last.col, Set names)
   New_data <- UpSetR:::Wanted(data, Sets_to_remove)</pre>
   Num of set <- UpSetR:::Number of sets(Set names)</pre>
    if(keep.order == F){
```

```
Set_names <- UpSetR:::order_sets(New_data, Set_names)</pre>
    }
    All_Freqs <- UpSetR:::specific_intersections(data, first.col, last.col, intersections, order.by, gr
                                           cutoff, main.bar.color, Set_names)
  } else if(is.null(intersections) == T){
    Set names <- sets
    if(is.null(Set_names) == T || length(Set_names) == 0 ){
      Set_names <- UpSetR:::FindMostFreq(data, first.col, last.col, nsets)</pre>
    Sets_to_remove <- UpSetR:::Remove(data, first.col, last.col, Set_names)</pre>
    New_data <- UpSetR:::Wanted(data, Sets_to_remove)</pre>
    Num_of_set <- UpSetR:::Number_of_sets(Set_names)</pre>
    if(keep.order == F){
    Set_names <- UpSetR:::order_sets(New_data, Set_names)</pre>
    All_Freqs <- UpSetR:::Counter(New_data, Num_of_set, first.col, Set_names, nintersects, main.bar.col
                          order.by, group.by, cutoff, empty.intersections, decreasing)
  }
  Matrix_setup <- UpSetR:::Create_matrix(All_Freqs)</pre>
  labels <- UpSetR:::Make_labels(Matrix_setup)</pre>
  #Chose NA to represent NULL case as result of NA being inserted when at least one contained both x an
  #i.e. if one custom plot had both x and y, and others had only x, the y's for the other plots were NA
  #if I decided to make the NULL case (all x and no y, or vice versa), there would have been alot more
  #NA can be indexed so that we still get the non NA y aesthetics on correct plot. NULL cant be indexed
  att.x \leftarrow c(); att.y \leftarrow c();
  if(is.null(attribute.plots) == F){
    for(i in seq_along(attribute.plots$plots)){
      if(length(attribute.plots$plots[[i]]$x) != 0){
        att.x[i] <- attribute.plots$plots[[i]]$x</pre>
      }
      else if(length(attribute.plots$plots[[i]]$x) == 0){
        att.x[i] <- NA
      if(length(attribute.plots$plots[[i]]$y) != 0){
        att.y[i] <- attribute.plots$plots[[i]]$y</pre>
      else if(length(attribute.plots$plots[[i]]$y) == 0){
        att.y[i] <- NA
    }
  }
## Generate boxplot summary plots
BoxPlotsPlot <- function(bdat, att, att_color){</pre>
  ylab <- element_blank()</pre>
  col <- match(att, colnames(bdat))</pre>
  colnames(bdat)[col] <- "attribute"</pre>
  upper_xlim <- as.numeric((max(bdat$x) + 1))</pre>
  #upper_ylim <- max((data$`Average Expression`) + 1)</pre>
  #lower_ylim <- min((data$`Average Expression`) - 1)</pre>
  plot_lims <- as.numeric(0:upper_xlim)</pre>
  bdat$x <- as.factor(bdat$x)</pre>
  boxplotter <- ggplot(data = bdat, aes(x=x, y=attribute, fill=x, color = x)) +
```

```
geom_point()
boxplots <- boxplotter +</pre>
  geom_dotplot(binaxis = "y", stackdir="center", method = "histodot", binwidth = 0.3) +
  scale_x_discrete(limits = plot_lims, expand = c(0,0)) +
                          ## scale_y_continuous(breaks = seq(lower_ylim, upper_ylim, 2), limits = c(lowe
                          theme(plot.margin = unit(c(-0.7,0,0,3.15), "cm"),
                                   axis.title.y = element_blank(),
                                   axis.ticks.x = element_blank(),
                                   axis.text.x = element_blank(),
                                  panel.border = element_blank(),
                                  panel.background = element_rect(fill = NA),
                                   panel.grid.major = element_line(colour = "gray"),
                                   legend.position = "none",
                                   axis.title.x = element_blank()) +
                 scale_fill_manual(values = att_color) + scale_color_manual(values = att_color)
 return(boxplots)
}
  BoxPlots <- NULL
  if(is.null(boxplot.summary) == F){
    BoxData <- UpSetR:::IntersectionBoxPlot(All_Freqs, New_data, first.col, Set_names)</pre>
    BoxPlots <- list()</pre>
    for(i in seq_along(boxplot.summary)){
      BoxPlots[[i]] <- BoxPlotsPlot(BoxData, boxplot.summary[i], att.color)</pre>
    }
  }
  customAttDat <- NULL</pre>
  customQBar <- NULL</pre>
  Intersection <- NULL
  Element <- NULL</pre>
  legend <- NULL</pre>
  EBar_data <- NULL
  if(is.null(queries) == F){
    custom.queries <- UpSetR:::SeperateQueries(queries, 2, palette)
    customDat <- UpSetR:::customQueries(New_data, custom.queries, Set_names)</pre>
    legend <- UpSetR:::GuideGenerator(queries, palette)</pre>
    legend <- UpSetR:::Make_legend(legend)</pre>
    if(is.null(att.x) == F && is.null(customDat) == F){
      customAttDat <- UpSetR:::CustomAttData(customDat, Set_names)</pre>
    }
    customQBar <- UpSetR:::customQueriesBar(customDat, Set_names, All_Freqs, custom.queries)</pre>
  if(is.null(queries) == F){
    Intersection <- UpSetR:::SeperateQueries(queries, 1, palette)</pre>
    Matrix_col <- UpSetR:::intersects(QuerieInterData, Intersection, New_data, first.col, Num_of_set,
                              All_Freqs, expression, Set_names, palette)
    Element <- UpSetR:::SeperateQueries(queries, 1, palette)</pre>
    EBar_data <-UpSetR:::ElemBarDat(Element, New_data, first.col, expression, Set_names,palette, All_Fr
  } else{
    Matrix_col <- NULL</pre>
```

```
Matrix_layout <- UpSetR:::Create_layout(Matrix_setup, matrix.color, Matrix_col, matrix.dot.alpha)</pre>
  for(i in 1:8) {
      j <- which(Matrix_layout$y == i & Matrix_layout$value == 1)</pre>
      if(length(j) > 0) Matrix_layout$color[j] <- c("purple", "darkgreen", "forestgreen", "darkolivegre
                    "#00659c", "#7dc3e8")[i]
  }
## continuing with upset()
  Set_sizes <- UpSetR:::FindSetFreqs(New_data, first.col, Num_of_set, Set_names, keep.order)</pre>
  Bar_Q <- NULL</pre>
  if(is.null(queries) == F){
    Bar_Q <- UpSetR:::intersects(QuerieInterBar, Intersection, New_data, first.col, Num_of_set, All_Fre</pre>
  QInter_att_data <- NULL
  QElem_att_data <- NULL
  if((is.null(queries) == F) & (is.null(att.x) == F)){
    QInter_att_data <- UpSetR:::intersects(QuerieInterAtt, Intersection, New_data, first.col, Num_of_se
                                   expression, Set_names, palette)
    QElem_att_data <- UpSetR:::elements(QuerieElemAtt, Element, New_data, first.col, expression, Set_na
                                palette)
  AllQueryData <- UpSetR:::combineQueriesData(QInter_att_data, QElem_att_data, customAttDat, att.x, att
  ShadingData <- NULL
  if(is.null(set.metadata) == F){
    ShadingData <- get_shade_groups(set.metadata, Set_names, Matrix_layout, shade.alpha)
    output <- Make_set_metadata_plot(set.metadata, Set_names)</pre>
    set.metadata.plots <- output[[1]]</pre>
    set.metadata <- output[[2]]</pre>
    if(is.null(ShadingData) == FALSE){
    shade.alpha <- unique(ShadingData$alpha)</pre>
  } else {
    set.metadata.plots <- NULL</pre>
  if(is.null(ShadingData) == TRUE){
  ShadingData <- UpSetR:::MakeShading(Matrix_layout, shade.color)</pre>
  Main_bar <- suppressMessages(UpSetR:::Make_main_bar(All_Freqs, Bar_Q, show.numbers, mb.ratio, customQ
                                                         mainbar.y.label, mainbar.y.max, scale.intersection
Make_matrix_plot <- function(Mat_data,Set_size_data, Main_bar_data, point_size, line_size, text_scale,
                              shading_data, shade_alpha){
  if(length(text_scale) == 1){
    name_size_scale <- text_scale</pre>
  if(length(text_scale) > 1 && length(text_scale) <= 6){</pre>
    name_size_scale <- text_scale[5]</pre>
  }
```

```
Mat_data$line_col <- 'black'</pre>
  Matrix_plot <- (ggplot()</pre>
                  + theme(panel.background = element rect(fill = "white"),
                          plot.margin=unit(c(-0.2,0.5,0.5,0.5), "lines"),
                          axis.text.x = element_blank(),
                          axis.ticks.x = element_blank(),
                          axis.ticks.y = element blank(),
                          axis.text.y = element_text(colour = "gray0",
                                                      size = 8*name_size_scale, hjust = 1),
                          panel.grid.major = element_blank(),
                          panel.grid.minor = element_blank())
                  + xlab(NULL) + ylab("
                                          ")
                  + scale_y_continuous(breaks = c(1:nrow(Set_size_data)),
                                       limits = c(0.5, (nrow(Set_size_data) +0.5)),
                                       labels = labels, expand = c(0,0))
                  + scale_x_continuous(limits = c(0,(nrow(Main_bar_data)+1)), expand = c(0,0))
                  + geom_rect(data = shading_data, aes_string(xmin = "min", xmax = "max",
                                                               ymin = "y_min", ymax = "y_max"),
                              fill = shading_data$shade_color, alpha = shade_alpha)
                  + geom_line(data= Mat_data, aes_string(group = "Intersection", x="x", y="y",
                                                          colour = "line_col"), size = line_size)
                 + geom_point(data= Mat_data, aes_string(x= "x", y= "y"), colour = Mat_data$color,
                     size= point_size, alpha = Mat_data$alpha, shape=16)
                  + scale_color_identity())
  Matrix_plot <- ggplot_gtable(ggplot_build(Matrix_plot))</pre>
  return(Matrix_plot)
}
Matrix <- Make_matrix_plot(Matrix_layout, Set_sizes, All_Freqs, point.size, line.size,
                             text.scale, labels, ShadingData, shade.alpha)
  Sizes <- UpSetR:::Make_size_plot(Set_sizes, sets.bar.color, mb.ratio, sets.x.label, scale.sets, text.
                          set_size.scale_max, set_size.numbers_size)
  ## UpSetR:::Make base plot(Main bar, Matrix, Sizes, labels, mb.ratio, att.x, att.y, New data,
                    expression, att.pos, first.col, att.color, AllQueryData, attribute.plots,
      ##
                    legend, query.legend, BoxPlots, Set_names, set.metadata, set.metadata.plots)
pdf(file.path(DGE_source, paste(Sys.Date(), "UpSetPlot_DGE_SpeciesMapped_vs_Humans.pdf")), width = 15,
structure(class = "upset",
    .Data=list(
      Main_bar = Main_bar,
      Matrix = Matrix,
      Sizes = Sizes,
      labels = labels,
      mb.ratio = mb.ratio,
      att.x = att.x,
      att.y = att.y,
      New_data = New_data,
      expression = expression,
      att.pos = att.pos,
```

```
first.col = first.col,
      att.color = att.color,
      AllQueryData = AllQueryData,
      attribute.plots = attribute.plots,
      legend = legend,
      query.legend = query.legend,
     BoxPlots = BoxPlots,
     Set_names = Set_names,
     set.metadata = set.metadata,
      set.metadata.plots = set.metadata.plots)
  )
dev.off()
## pdf
##
Session Info
sessionInfo()
## R version 3.5.2 (2018-12-20)
## Platform: x86_64-apple-darwin15.6.0 (64-bit)
## Running under: macOS Sierra 10.12.6
##
## Matrix products: default
## BLAS: /Library/Frameworks/R.framework/Versions/3.5/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/3.5/Resources/lib/libRlapack.dylib
##
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
## attached base packages:
## [1] parallel stats4
                                     graphics grDevices utils
                                                                    datasets
                           stats
## [8] methods
                 base
##
## other attached packages:
## [1] bindrcpp_0.2.2
                                    purrr_0.2.5
## [3] data.table_1.12.0
                                    reshape2_1.4.3
## [5] ggrepel_0.8.0
                                    usethis_1.4.0
## [7] devtools_2.0.1
                                    viridis_0.5.1
## [9] viridisLite_0.3.0
                                    RColorBrewer_1.1-2
## [11] ggplot2_3.1.0
                                    gplots_3.0.1
## [13] DESeq2_1.22.2
                                    SummarizedExperiment_1.12.0
## [15] DelayedArray_0.8.0
                                    BiocParallel_1.16.5
## [17] matrixStats_0.54.0
                                    Biobase_2.42.0
## [19] GenomicRanges 1.34.0
                                    GenomeInfoDb 1.18.1
## [21] IRanges_2.16.0
                                    S4Vectors_0.20.1
## [23] BiocGenerics 0.28.0
                                    genefilter 1.64.0
## [25] biomaRt_2.38.0
                                    stringr_1.3.1
                                    dplyr_0.7.8
## [27] tibble_2.0.1
## [29] plyr_1.8.4
## loaded via a namespace (and not attached):
## [1] fs_1.2.6
                               bitops_1.0-6
                                                      bit64_0.9-7
## [4] progress_1.2.0
                               httr_1.4.0
                                                      UpSetR_1.4.0
```

```
[7] rprojroot_1.3-2
                               tools_3.5.2
                                                       backports 1.1.3
## [10] R6_2.3.0
                               rpart_4.1-13
                                                       KernSmooth_2.23-15
                               DBI 1.0.0
## [13] Hmisc 4.1-1
                                                       lazyeval 0.2.1
## [16] colorspace_1.4-0
                               nnet_7.3-12
                                                       withr_2.1.2
## [19] processx_3.2.1
                               tidyselect_0.2.5
                                                       gridExtra 2.3
## [22] prettyunits 1.0.2
                               bit 1.1-14
                                                       compiler 3.5.2
## [25] cli 1.0.1
                               htmlTable 1.13.1
                                                       desc 1.2.0
                               caTools_1.17.1.1
## [28] labeling 0.3
                                                       scales_1.0.0
## [31] checkmate_1.9.1
                               callr_3.1.1
                                                       digest_0.6.18
## [34] foreign_0.8-71
                               rmarkdown_1.11
                                                       XVector_0.22.0
## [37] base64enc_0.1-3
                               pkgconfig_2.0.2
                                                       htmltools_0.3.6
## [40] sessioninfo_1.1.1
                               htmlwidgets_1.3
                                                       rlang_0.3.1
## [43] rstudioapi_0.9.0
                               RSQLite_2.1.1
                                                       bindr_0.1.1
## [46] gtools_3.8.1
                               acepack_1.4.1
                                                       RCurl_1.95-4.11
## [49] magrittr_1.5
                               GenomeInfoDbData_1.2.0 Formula_1.2-3
## [52] Matrix_1.2-15
                               Rcpp_1.0.0
                                                       munsell_0.5.0
## [55] stringi_1.2.4
                               yam1_2.2.0
                                                       zlibbioc_1.28.0
## [58]
       pkgbuild_1.0.2
                               grid_3.5.2
                                                       blob 1.1.1
## [61] gdata_2.18.0
                               crayon_1.3.4
                                                       lattice_0.20-38
## [64] splines 3.5.2
                                annotate_1.60.0
                                                       hms 0.4.2
## [67] locfit_1.5-9.1
                               ps_1.3.0
                                                       knitr_1.21
## [70] pillar 1.3.1
                               pkgload_1.0.2
                                                       geneplotter_1.60.0
## [73] XML_3.98-1.16
                               glue_1.3.0
                                                       evaluate_0.12
## [76] latticeExtra 0.6-28
                               remotes 2.0.2
                                                       gtable 0.2.0
## [79] assertthat_0.2.0
                               xfun 0.4
                                                       xtable_1.8-3
## [82] survival 2.43-3
                               AnnotationDbi_1.44.0
                                                       memoise_1.1.0
## [85] cluster_2.0.7-1
```