contrasts.fit exploration

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library("scp")

```
## Loading required package: QFeatures
## Loading required package: MultiAssayExperiment
## Loading required package: SummarizedExperiment
## Loading required package: MatrixGenerics
## Loading required package: matrixStats
##
## Attaching package: 'MatrixGenerics'
  The following objects are masked from 'package:matrixStats':
##
##
       colAlls, colAnyNAs, colAnys, colAvgsPerRowSet, colCollapse,
##
       colCounts, colCummaxs, colCummins, colCumprods, colCumsums,
##
       colDiffs, colIQRDiffs, colIQRs, colLogSumExps, colMadDiffs,
##
       colMads, colMaxs, colMeans2, colMedians, colMins, colOrderStats,
##
       colProds, colQuantiles, colRanges, colRanks, colSdDiffs, colSds,
##
       colSums2, colTabulates, colVarDiffs, colVars, colWeightedMads,
##
       colWeightedMeans, colWeightedMedians, colWeightedSds,
##
       colWeightedVars, rowAlls, rowAnyNAs, rowAnys, rowAvgsPerColSet,
##
       rowCollapse, rowCounts, rowCummaxs, rowCummins, rowCumprods,
       rowCumsums, rowDiffs, rowIQRDiffs, rowIQRs, rowLogSumExps,
##
       rowMadDiffs, rowMads, rowMaxs, rowMeans2, rowMedians, rowMins,
##
       rowOrderStats, rowProds, rowQuantiles, rowRanges, rowRanks,
##
##
       rowSdDiffs, rowSds, rowSums2, rowTabulates, rowVarDiffs, rowVars,
       rowWeightedMads, rowWeightedMeans, rowWeightedMedians,
##
       rowWeightedSds, rowWeightedVars
## Loading required package: GenomicRanges
## Loading required package: stats4
## Loading required package: BiocGenerics
##
## Attaching package: 'BiocGenerics'
## The following objects are masked from 'package:stats':
##
##
       IQR, mad, sd, var, xtabs
  The following objects are masked from 'package:base':
##
##
##
       anyDuplicated, aperm, append, as.data.frame, basename, cbind,
```

```
##
       colnames, dirname, do.call, duplicated, eval, evalq, Filter, Find,
##
       get, grep, grepl, intersect, is.unsorted, lapply, Map, mapply,
##
       match, mget, order, paste, pmax, pmax.int, pmin, pmin.int,
       Position, rank, rbind, Reduce, rownames, sapply, setdiff, table,
##
       tapply, union, unique, unsplit, which.max, which.min
## Loading required package: S4Vectors
##
## Attaching package: 'S4Vectors'
## The following object is masked from 'package:utils':
##
##
       findMatches
## The following objects are masked from 'package:base':
##
##
       expand.grid, I, unname
## Loading required package: IRanges
## Loading required package: GenomeInfoDb
## 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")'.
## Attaching package: 'Biobase'
## The following object is masked from 'package:MatrixGenerics':
##
##
       rowMedians
## The following objects are masked from 'package:matrixStats':
##
##
       anyMissing, rowMedians
##
## Attaching package: 'QFeatures'
## The following object is masked from 'package:MultiAssayExperiment':
##
##
       longFormat
## The following object is masked from 'package:base':
##
##
       sweep
library("msqrob2")
library("limma")
##
## Attaching package: 'limma'
## The following object is masked from 'package:BiocGenerics':
##
##
       plotMA
```

```
library("lme4")
## Loading required package: Matrix
##
## Attaching package: 'Matrix'
## The following object is masked from 'package:S4Vectors':
##
##
        expand
library("UpSetR")
msqrobLm_fix <- function (y, formula, data, robust = TRUE, maxitRob = 5)</pre>
    myDesign <- model.matrix(formula, data)</pre>
    models <- apply(y, 1, function(y, design) {</pre>
         obs <- is.finite(y)</pre>
         type <- "fitError"</pre>
         model <- list(coefficients = NA, vcovUnscaled = NA,</pre>
             sigma = NA, df.residual = NA, w = NA)
         if (sum(obs) > 0) {
             X <- design[obs, , drop = FALSE]</pre>
             qrX \leftarrow qr(X)
             X <- X[, qrX$pivot[seq_len(qrX$rank)], drop = FALSE] # change</pre>
             # => old version
             \# X \leftarrow X[, colMeans(X == 0) != 1, drop = FALSE]
             y <- y[obs]
             colnames_orig <- colnames(design)</pre>
             if (robust) {
                  mod <- try(MASS::rlm(X, y, method = "M", maxit = maxitRob),</pre>
                      silent = TRUE)
                  if (!is(mod, "try-error")) {
                      type <- "rlm"
                  }
             }
             else {
                  mod <- try(lm.fit(X, y))</pre>
                  if ((!is(mod, "try-error")) & mod$rank == ncol(X)) {
                      type <- "lm"
                  }
             if (type == "rlm") {
                  w <- mod$w
                  sigma <- sqrt(sum(mod$w * mod$resid^2)/(sum(mod$w) -</pre>
                      mod$rank))
                  df.residual <- sum(mod$w) - mod$rank
                  if (df.residual < 2L)</pre>
                      type <- "fitError"</pre>
             if (type == "lm") {
                  w <- NULL
                  sigma <- sqrt(sum(mod$residuals^2/mod$df.residual))</pre>
                  df.residual <- mod$df.residual</pre>
                  if (df.residual < 2L)</pre>
```

```
type <- "fitError"</pre>
            }
            if (type != "fitError") {
                coef <- rep(NA, length(colnames_orig))</pre>
                 names(coef) <- colnames_orig</pre>
                 coef[names(mod$coef)] <- mod$coef</pre>
                 vcovUnscaled <- matrix(NA, nrow = length(colnames_orig),</pre>
                     ncol = length(colnames_orig))
                 rownames(vcovUnscaled) <- colnames(vcovUnscaled) <- colnames_orig</pre>
                 vcovUnscaled[names(mod$coef), names(mod$coef)] <- msqrob2:::.vcovUnscaled(mod)</pre>
                 model <- list(coefficients = mod$coef, vcovUnscaled = msqrob2:::.vcovUnscaled(mod),</pre>
                     sigma = sigma, df.residual = df.residual,
                     w = w
            }
        }
        .StatModel(type = type, params = model, varPosterior = as.numeric(NA),
            dfPosterior = as.numeric(NA))
    }, design = myDesign)
    hlp <- limma::squeezeVar(var = vapply(models, getVar, numeric(1)),</pre>
        df = vapply(models, getDF, numeric(1)))
    for (i in seq_len(length(models))) {
        mydf <- hlp$df.prior + getDF(models[[i]])</pre>
        models[[i]]@varPosterior <- as.numeric(hlp$var.post[i])</pre>
        models[[i]]@dfPosterior <- as.numeric(mydf)</pre>
    }
    return(models)
}
### msqrob2::msqrob with the new msqrobLm
msqrob_fix <- .local <- function (object, formula, modelColumnName = "msqrobModels",</pre>
        overwrite = FALSE, robust = TRUE, ridge = FALSE, maxitRob = 1,
        tol = 1e-06, doQR = TRUE, lmerArgs = list(control = lmerControl(calc.derivs = FALSE)))
        if (ncol(colData(object)) == 0)
            stop("colData is empty")
        if ((modelColumnName %in% colnames(rowData(object))) &
            !overwrite) {
            stop("There is already a column named '", modelColumnName,
                 "' in the rowData of the SummarizedExperiment object, set the argument overwrite=TRUE t
        }
        if (length(formula) == 3) {
            formula <- formula[-2]</pre>
        if (any(all.vars(formula) %in% colnames(rowData(object)))) {
            stop("Use the msqrobAggregate function to use rowData variables")
        check_vars <- all.vars(formula) %in% colnames(colData(object))</pre>
        if (!all(check_vars)) {
            if (sum(!check_vars) > 1) {
                 vars_not_found <- paste0(all.vars(formula)[!check_vars],</pre>
                     collapse = ", ")
```

```
stop(paste("Variables", vars_not_found, "are not found in coldata"),
                     sep = "")
             }
             else {
                 vars_not_found <- all.vars(formula)[!check_vars]</pre>
                 stop(paste0("Variable ", vars_not_found, " is not found in coldata"),
                     sep = "")
             }
        }
        if (!ridge & is.null(findbars(formula))) {
             rowData(object)[[modelColumnName]] <- msqrobLm_fix(y = assay(object),</pre>
                 formula = formula, data = droplevels(colData(object)),
                 robust = robust, maxitRob = maxitRob)
        }
        else {
             rowData(object)[[modelColumnName]] <- msqrobLmer(y = assay(object),</pre>
                 formula = formula, data = droplevels(colData(object)),
                 rowdata = NULL, robust = robust, maxitRob = maxitRob,
                 tol = tol, doQR = doQR, ridge = ridge, lmerArgs = lmerArgs)
        }
        rowData(object)[[modelColumnName]] <- rowData(object)[[modelColumnName]][rownames(rowData(object))]</pre>
        return(object)
    }
.createMockBalanced <- function(k, runs) {</pre>
    out <- split(runs, runs)</pre>
    out <- lapply(out, function(x) {</pre>
        index <- 1:length(x)</pre>
        x[sample(index)] <- rep(1:k, length.out = length(index))</pre>
    })
    out <- S4Vectors::unname(do.call(c, out))</pre>
    as.factor(out)
}
sce <- readRDS("data/leduc_mock.rds")</pre>
set.seed(1234)
sce$Mock <- .createMockBalanced(2, sce$Set)</pre>
num_features <- 2000 # Number of features to sample</pre>
total_features <- nrow(sce)</pre>
subset <- sce[sample(1:total_features, num_features),]</pre>
colData(subset)$Set <- as.factor(colData(subset)$Set)</pre>
colData(subset) <- droplevels(colData(subset))</pre>
subset_msqrob <- msqrob(subset,</pre>
                             formula = ~ 1 + Channel + Set + Mock,
                             robust = FALSE,
                             ridge = FALSE)
subset_msqrob_fix <- msqrob_fix(subset,</pre>
                             formula = ~ 1 + Channel + Set + Mock,
                             robust = FALSE,
                             ridge = FALSE)
```

```
subset_limma <- limma::lmFit(assay(subset),</pre>
                              model.matrix(~ 1 + Channel + Set + Mock,
                                            colData(subset)))
## Warning: Partial NA coefficients for 1840 probe(s)
subset_scp <- scp::scpModelWorkflow(subset,</pre>
                                     formula = ~ 1 + Channel + Set + Mock)
##
design <- model.matrix(~ 1 + Channel + Set + Mock, colData(subset))</pre>
L_limma <- makeContrasts(SeteAL00220, levels = design)</pre>
## Warning in makeContrasts(SeteAL00220, levels = design): Renaming (Intercept) to
## Intercept
contrast_limma <- contrasts.fit(subset_limma, L_limma)</pre>
toptable <- eBayes(contrast_limma)</pre>
toptable <- topTable(toptable, sort.by = "none", number = 2000)</pre>
L_msqrob <- makeContrast(c("SeteAL00220 = 0"), parameterNames = colnames(design))</pre>
contrast_msqrob <- hypothesisTest(subset_msqrob, contrast = L_msqrob)</pre>
contrast_msqrob <- rowData(contrast_msqrob)$SeteAL00220</pre>
contrast msgrob fix <- hypothesisTest(subset msgrob fix, contrast = L msgrob)</pre>
contrast_msqrob_fix <- rowData(contrast_msqrob_fix)$SeteAL00220</pre>
try(contrast_scp <- scp::scpDifferentialAnalysis(subset_scp, contrasts = list(c("Set", "eAL00220", "eAL</pre>
## Error in t(1) %*% mm : non-conformable arguments
# contrast_scp <- contrast_scp[[1]] %>%
   as.data.frame()
# rownames(contrast_scp) <- contrast_scp$feature</pre>
retrieve_msqrob_coef <- function(msqrob_model, coef_name){</pre>
    msgrob object <- rowData(msgrob model)$msgrobModels</pre>
  coef <- unlist(lapply(msqrob_object, function(x) {</pre>
       tryCatch(x@params$coefficients[coef_name],
                  error = function(e) NA)
  }))
  names(coef) <- rownames(msqrob_model)</pre>
  coef
}
summary_df <- data.frame(</pre>
  limma = subset_limma$coefficients[,"SeteAL00220"],
  msqrob = retrieve_msqrob_coef(subset_msqrob, "SeteAL00220"),
  msqrob_fix = retrieve_msqrob_coef(subset_msqrob_fix, "SeteAL00220"),
  contrast_limma = contrast_limma$coefficients[,"SeteAL00220"],
  toptable_limma = toptable$logFC,
  contrast_msqrob = contrast_msqrob$logFC,
  contrast_msqrob_fix = contrast_msqrob_fix$logFC,
  row.names = rownames(subset)
```

```
sets <- list(</pre>
  limma = row.names(summary_df)[!is.na(summary_df$limma)],
  contrast_limma = row.names(summary_df)[!is.na(summary_df$contrast_limma)],
  toptable_limma = row.names(summary_df)[!is.na(summary_df$toptable_limma)],
  msqrob = row.names(summary_df)[!is.na(summary_df$msqrob)],
  contrast_msqrob = row.names(summary_df)[!is.na(summary_df$contrast_msqrob)],
  msqrob_fix = row.names(summary_df)[!is.na(summary_df$msqrob_fix)],
  contrast_msqrob_fix = row.names(summary_df)[!is.na(summary_df$contrast_msqrob_fix)]
)
upset(fromList(sets), nsets = 7, order.by = "freq")
                                 250
                                                  226
                                 200
                      Intersection Size
                                 150
                                                                143
                                 100
                                  50
                          msarob
                        contrast_msqrob
                         msqrob_fix
                        ontrast_msqrob_fix
                          limma
                        contrast_limma
                        toptable_limma
400
    300
          200
               100
        Set Size
to_test_msqrob <- rownames(summary_df)[!is.na(summary_df$msqrob)]</pre>
identical(summary_df[to_test_msqrob, "limma"], summary_df[to_test_msqrob, "msqrob"])
## [1] TRUE
identical(summary_df[to_test_msqrob, "limma"], summary_df[to_test_msqrob, "contrast_msqrob"])
## [1] TRUE
identical(summary_df[to_test_msqrob, "limma"], summary_df[to_test_msqrob, "msqrob_fix"])
## [1] TRUE
identical(summary_df[to_test_msqrob, "limma"], summary_df[to_test_msqrob, "contrast_msqrob_fix"])
```

[1] TRUE

```
identical(summary_df[to_test_msqrob, "limma"], summary_df[to_test_msqrob, "contrast_limma"])
## [1] TRUE
identical(summary_df[to_test_msqrob, "limma"], summary_df[to_test_msqrob, "toptable_limma"])
## [1] TRUE
to_test_fix <- rownames(summary_df)[!is.na(summary_df$msqrob_fix)]
identical(summary_df[to_test_fix, "limma"], summary_df[to_test_fix, "msqrob_fix"])
## [1] TRUE
identical(summary_df[to_test_fix, "limma"], summary_df[to_test_fix, "contrast_msqrob_fix"])
## [1] TRUE
identical(summary_df[to_test_fix, "limma"], summary_df[to_test_fix, "contrast_limma"])
## [1] TRUE
identical(summary_df[to_test_fix, "limma"], summary_df[to_test_fix, "toptable_limma"])
## [1] TRUE</pre>
```