Comparison of liger with fgsea

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liger is just one of many methods for gene set enrichment analysis. fgsea is another similar method that uses a faster cumulative statistic calculation on preranked values. Here, we compare liger to fgsea.

Comparison

We will use the example data and example gene sets that come with the fgsea package. Note that exampleRanks is a numeric vector where each value is fold-change or differential expression z-score between two biological conditions (some type of metric used for ranking genes) and examplePathways is a list of lists where each entry is a gene name corresponding to exampleRanks.

```
# example pathways from fgsea
library(fgsea)
data(examplePathways)
data(exampleRanks)
head(examplePathways)
## $`1221633_Meiotic_Synapsis`
    [1] "12189"
                     "13006"
##
                                  "15077"
                                               "15078"
                                                            "15270"
    [6] "15512"
                     "16905"
                                  "16906"
                                               "19357"
                                                            "20842"
   [11] "20843"
                     "20957"
                                  "20962"
                                               "21749"
                                                            "21750"
##
##
   Г167
        "22196"
                     "23856"
                                  "24061"
                                               "28113"
                                                            "50878"
##
  [21] "56739"
                     "57321"
                                  "64009"
                                               "66654"
                                                            "69386"
##
   [26]
        "71846"
                     "74075"
                                  "77053"
                                               "94244"
                                                            "97114"
        "97122"
   [31]
                     "97908"
                                               "140557"
                                                            "223697"
##
                                  "101185"
##
   [36]
        "260423"
                     "319148"
                                  "319149"
                                               "319150"
                                                            "319151"
##
   [41]
        "319152"
                     "319153"
                                  "319154"
                                               "319155"
                                                            "319156"
##
   [46]
       "319157"
                     "319158"
                                  "319159"
                                               "319160"
                                                            "319161"
   [51] "319565"
                     "320332"
                                  "320558"
                                               "326619"
                                                            "326620"
   [56] "360198"
                                                            "667250"
##
                     "497652"
                                  "544973"
                                               "625328"
   [61] "100041230"
                     "102641229" "102641751"
                                               "102642045"
##
##
## $\`1368092_Rora_activates_gene_expression\`
  [1] "11865"
                 "12753" "12894" "18143" "19017" "19883"
                                                                 "20787"
                                                                           "217166"
## [9] "328572"
##
## $`1368110_Bmal1:Clock, Npas2_activates_circadian_gene_expression`
##
    [1] "11865"
                  "11998"
                           "12753"
                                     "12952"
                                               "12953"
                                                         "13170"
                                                                  "14068"
##
    [8] "18143"
                  "18626"
                            "18627" "19013" "19883"
                                                         "20893"
                                                                  "59027"
## [15] "79362"
                  "217166"
##
## $\int 1445146_Translocation_of_Glut4_to_the_Plasma_Membrane\int
##
    [1] "11461"
                     "11465"
                                  "11651"
                                               "11652"
                                                            "12313"
##
    [6] "12314"
                     "12315"
                                  "16568"
                                               "16569"
                                                            "16579"
  [11] "17274"
                     "17884"
                                  "17886"
                                               "17913"
                                                            "17918"
## [16] "19079"
                     "19082"
                                  "19325"
                                               "19341"
                                                            "20336"
```

```
## [21] "20528"
                                              "20912"
                     "20619"
                                  "20909"
                                                           "22318"
   [26] "22627"
                     "22628"
                                  "22629"
                                              "22630"
                                                           "22631"
   [31] "53413"
                                  "55948"
                     "54401"
                                              "56044"
                                                           "57915"
   [36] "66482"
                     "68328"
                                  "68365"
                                              "68938"
                                                           "69940"
##
   [41] "102058"
                     "105504"
                                  "107371"
                                              "108079"
                                                           "108097"
## [46] "108099"
                     "210789"
                                  "211446"
                                              "240028"
                                                           "241113"
## [51] "241694"
                     "100039786" "102634437" "102641200" "102641764"
##
## $`186574_Endocrine-committed_Ngn3+_progenitor_cells`
## [1] "18012" "18088" "18506" "53626"
## $`186589_Late_stage_branching_morphogenesis_pancreatic_bud_precursor_cells`
## [1] "11925" "15205" "21410" "246086"
head(exampleRanks)
##
      170942
                 109711
                            18124
                                       12775
                                                  72148
                                                            16010
## -63.33703 -49.74779 -43.63878 -41.51889 -33.26039 -32.77626
barplot(sort(exampleRanks, decreasing=TRUE), names.arg='')
4
0
We will test all gene sets of a particular size.
# filter pathways to certain size
size <- lapply(examplePathways, length)</pre>
vi <- size > 15 & size < 500
table(vi)
## vi
## FALSE TRUE
     702
           755
examplePathways <- examplePathways[vi]</pre>
Now, we run both methods.
# run fgsea
start_time <- Sys.time()</pre>
fgseaRes <- fgsea(pathways = examplePathways,</pre>
```

stats = exampleRanks,

Time difference of 3.118531 secs

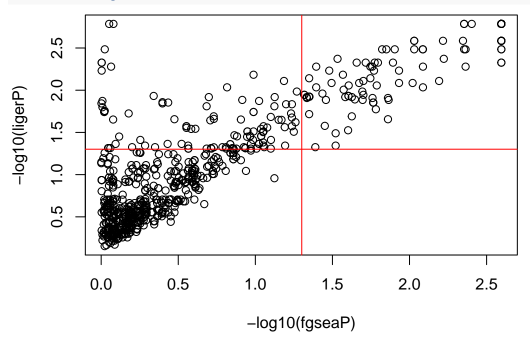
```
## initial: [1e+02 - 448] [1e+03 - 212] [1e+04 - 101] done
end_time <- Sys.time()
print(end_time - start_time)</pre>
```

Time difference of 3.253569 mins

We can plot the -log10(corrected p-values) for both approaches and assess their correspondence.

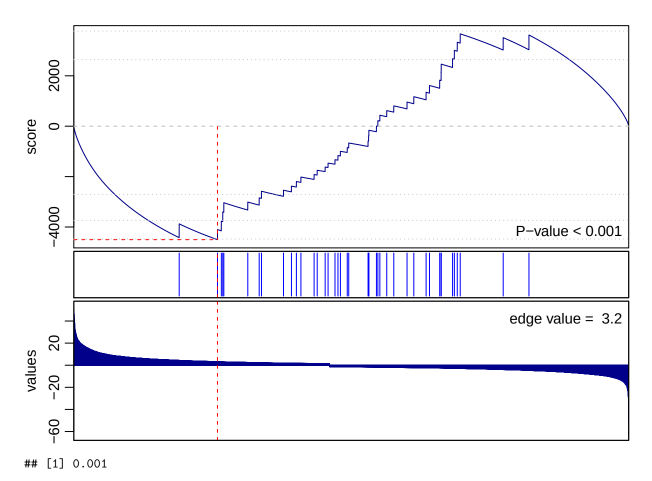
```
# compare
fgseaP <- fgseaRes$padj; names(fgseaP) <- fgseaRes$pathway
fgseaP <- fgseaP[names(examplePathways)]
ligerP <- ligerRes$q.val; names(ligerP) <- rownames(ligerRes)
ligerP <- ligerP[names(examplePathways)]

# plot
par(mfrow=c(1,1), mar=rep(5,4))
plot(-log10(fgseaP), -log10(ligerP))
abline(v = -log10(0.05), col='red')
abline(h = -log10(0.05), col='red')</pre>
```



Each dot here is a gene set. The x position is the -log10(p-value) of the gene set from fgsea while the y-axis is from liger. While there does appear to be a good general correspondence (strong diagonal), notice a set of gene sets that are very significant in liger but not in fgsea. Let's take a closer look at what are these gene sets.

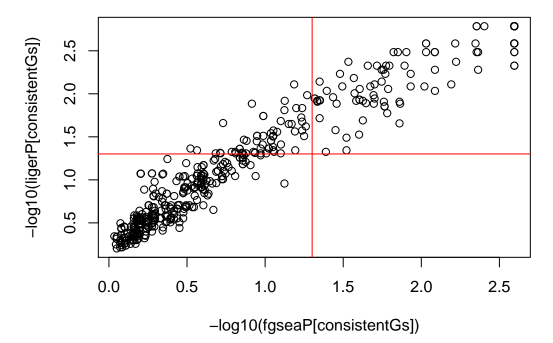
```
# maximal difference (most inconsistent) between methods
diff <- abs(-log10(fgseaP) - -log10(ligerP))</pre>
diff <- sort(diff, decreasing=TRUE)</pre>
# pick most inconsistent
gs <- names(diff)[1]
print(fgseaP[gs])
## 5991461_Peptide_chain_elongation
##
                             0.8874602
print(ligerP[gs])
## 5991461_Peptide_chain_elongation
                          0.001638967
##
# fgsea
plotEnrichment(examplePathways[[gs]], exampleRanks)
    0.2 -
    0.1 -
enrichment score
     0.0
   -0.1 -
    -0.2
                           2500
                                            5000
                                                             7500
                                                                              10000
                                                                                               12500
                                                   rank
# liger
gsea(exampleRanks, examplePathways[[gs]])
```



What we can see is that liger detected a significant lack of genes in this gene set among the most highly ranked genes as noted by the positive edge but negative sscore. This particular type of enrichment testing may be important if we want to make claims about certain gene sets never being highly differentially expressed (depleted in representation) but are not necessarily down-regulated.

As fgsea does not detect such patterns, to make our comparison between the two methods more appropriate, we will restrict to gene sets for which liger detects a consistent sscore and edge (ie. both positive suggesting upregulation or both negative suggesting downregulation).

```
# make comparable
vi <- ligerRes$sscore * ligerRes$edge > 0
consistentGs <- rownames(ligerRes)[vi]
par(mfrow=c(1,1), mar=rep(5,4))
plot(-log10(fgseaP[consistentGs]), -log10(ligerP[consistentGs]))
abline(v = -log10(0.05), col='red')
abline(h = -log10(0.05), col='red')</pre>
```



Now, results are highly consistent between the two approaches.

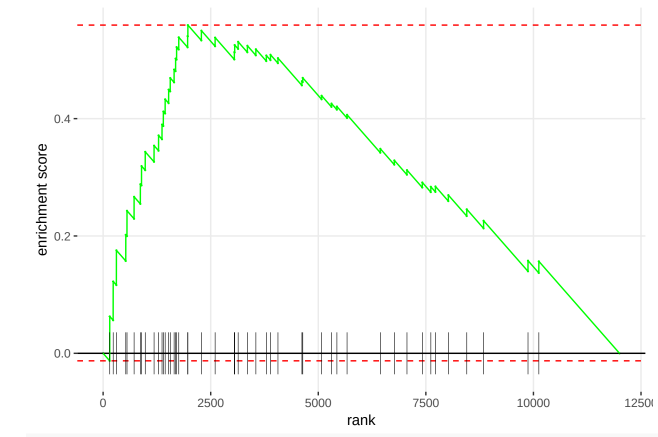
```
# pick a significant gene set
gs <- names(which(fgseaP[consistentGs]==min(fgseaP[consistentGs])))[1]
print(fgseaP[gs])

## 5990976_Assembly_of_the_pre-replicative_complex
## 0.002537457

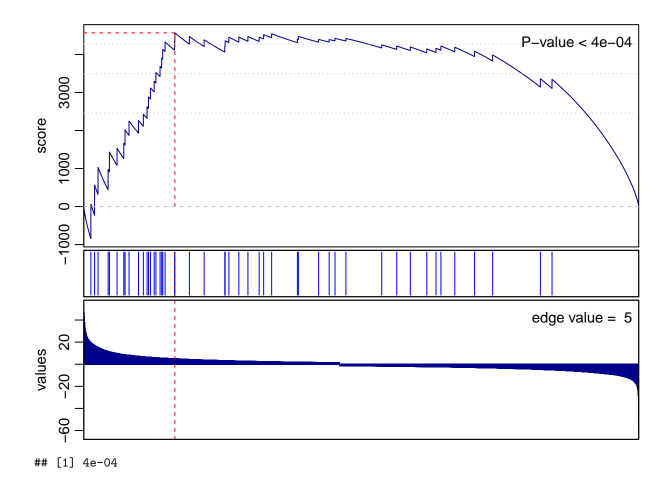
print(ligerP[gs])

## 5990976_Assembly_of_the_pre-replicative_complex
## 0.001638967

# fgsea
plotEnrichment(examplePathways[[gs]], exampleRanks)</pre>
```



liger
gsea(exampleRanks, examplePathways[[gs]])



Conclusion

In conclusion, fgsea provides a very fast test for gene sets where ranked gene values are appropriate. Both fgsea and liger offer very comparable results when looking for significantly upregulated or downregulated gene sets.

Ultimately, the appropriateness of gene set enrichment analysis approaches will depend on your question of interest. If you are only looking to test simply for over-representation of a set of genes, perhaps a hypergeometric test will be sufficient. If you care about the magnitude of the gene expression fold-change used in your gene ranking, a purely rank-based approach may be less optimal. If you are only interested in consistent upregulation and downregulation, significant results pointing to a depletion in representation among highly ranked genes may not be useful and should be ignored.

What ever gene set enrichment analysis you choose and whatever hypotheses they may help you generate, given the multitude of issues associated with gene sets, their accuracy, particularly as they pertain to your biological system of study, additional biological validation is always encouraged.