# dGSEAview

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dGSEAview

Function to view enrichment results in a sample-specific manner

#### **Description**

dGSEAview is supposed to view results of gene set enrichment analysis but for a specific sample.

## Usage

```
dGSEAview(eTerm, which_sample = 1, top_num = 10, sortBy = c("adjp",
   "gadjp", "ES", "nES", "pvalue", "FWER", "FDR", "qvalue"), decreasing =
NULL,
details = F)
```

# **Arguments**

eTerm an object of class "eTerm" which\_sample which sample will be viewed

top\_num the maximum number of gene sets will be viewed

sortBy which statistics will be used for sorting and viewing gene sets. It can be "adjp"

for adjusted p value, "gadjp" for globally adjusted p value, "ES" for enrichment score, "nES" for normalised enrichment score, "pvalue" for p value, "FWER" for family-wise error rate, "FDR" for false discovery rate, "qvalue" for q value

decreasing logical to indicate whether to sort in a decreasing order. If it is null, it would be

true for "ES" or "nES"; otherwise it would be false

details logical to indicate whether the detail information of gene sets is also viewed. By

default, it sets to false for no inclusion

#### Value

a data frame with following components:

• setID: term ID

• ES: enrichment score

· nES: normalised enrichment score

• pvalue: nominal p value

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• adjp: adjusted p value

• gadjp: globally adjusted p value

• FDR: false discovery rate

• qvalue: q value

• setSize: the number of genes in the set; optional, it is only appended when "details" is true

• name: term name; optional, it is only appended when "details" is true

• namespace: term namespace; optional, it is only appended when "details" is true

• distance: term distance; optional, it is only appended when "details" is true

#### Note

none

### See Also

dGSEA

## **Examples**

#dGSEAview(eTerm, which\_sample=1, top\_num=10, sortBy="adjp", decreasing=FALSE, details=TRUE)