# **dEnricherView**

June 10, 2015

dEnricherView

Function to view enrichment results of dEnricher

#### **Description**

dEnricherView is supposed to view results of enrichment analysis by dEnricher.

### Usage

```
dEnricherView(eTerm, top_num = 10, sortBy = c("adjp", "pvalue",
"zscore",
"nAnno", "nOverlap", "none"), decreasing = NULL, details = F)
```

# Arguments

eTerm an object of class "eTerm"

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

sortBy which statistics will be used for sorting and viewing gene sets (terms). It can

be "adjp" for adjusted p value, "pvalue" for p value, "zscore" for enrichment z-score, "nAnno" for the number of sets (terms), "nOverlap" for the number in

overlaps, and "none" for ordering according to ID of gene sets (terms)

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

true for "zscore", "nAnno" or "nOverlap"; otherwise it would be false

details logical to indicate whether the detailed information of gene sets (terms) is also

viewed. By default, it sets to false for no inclusion

#### Value

a data frame with following components:

• setID: term ID

• nAnno: number in gene members annotated by a term

• nOverlap: number in overlaps

• zscore: enrichment z-score

• pvalue: nominal p value

• adjp: adjusted p value

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- 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
- members: members (represented as Gene Symbols) in overlaps; optional, it is only appended when "details" is true

#### Note

none

#### See Also

dEnricher

## **Examples**

#dEnricherView(eTerm, top\_num=10, sortBy="adjp", decreasing=FALSE, details=TRUE)