DEsubs

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## 1. Package Setup

DEsubs is a network-based systems biology R package that extracts disease-perturbed subpathways within a pathway network as recorded by RNA-seq experiments. It contains an extensive and customizable framework with a broad range of operation modes at all stages of the subpathway analysis, enabling a case-specific approach. The operation modes refer to the pathway network construction and processing, the subpathway extraction, visualization and enrichment analysis with regard to various biological and pharmacological features. Its capabilities render it a valuable tool for both the modeler and experimentalist searching for the identification of more robust systems-level drug targets and biomarkers for complex diseases.

Before loading the package, please specify a user-accessible home directory using the following commands, which currently reflect the default directories for each architecture:

```r
if (.Platform[['OS.type']] == 'unix')
{
    options('DEsubsCACHE'=file.path(path.expand("-"), 'DEsubs'))
}
if (.Platform[['OS.type']] == 'windows')
{
    options('DEsubsCACHE'=file.path(
        gsub("\\", "/", Sys.getenv("USERPROFILE")), "AppData/DEsubs"))
}
```

Now the package, as well as the toy-data can be loaded as follows:

```r
library('DEsubs')
load(system.file('extdata', 'data.RData', package='DEsubs'))
```
2. User Input

DEsubs accepts RNA-seq expression paired case-control profile data. The following example in Table 1 shows the right structure for RNA-seq expression data input.

Table 1: Example of user input format

<table>
<thead>
<tr>
<th>Gene</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
<th>Case 4</th>
<th>Control 1</th>
<th>Control 2</th>
<th>Control 3</th>
<th>Control 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gene 1</td>
<td>1879</td>
<td>2734</td>
<td>2369</td>
<td>2636</td>
<td>2188</td>
<td>9743</td>
<td>9932</td>
<td>10099</td>
</tr>
<tr>
<td>Gene 2</td>
<td>97</td>
<td>124</td>
<td>146</td>
<td>114</td>
<td>126</td>
<td>33</td>
<td>19</td>
<td>31</td>
</tr>
<tr>
<td>Gene 3</td>
<td>485</td>
<td>485</td>
<td>469</td>
<td>428</td>
<td>475</td>
<td>128</td>
<td>135</td>
<td>103</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>Gene N-1</td>
<td>84</td>
<td>25</td>
<td>67</td>
<td>62</td>
<td>61</td>
<td>277</td>
<td>246</td>
<td>297</td>
</tr>
<tr>
<td>Gene N</td>
<td>120</td>
<td>312</td>
<td>78</td>
<td>514</td>
<td>210</td>
<td>324</td>
<td>95</td>
<td>102</td>
</tr>
</tbody>
</table>

3. Pathway network construction

KEGG signaling pathway maps have been downloaded and converted to pathway networks using CHRONOS package. Pathway networks for the seven supported organisms are included in the package itself (see Table 2).

Table 2: Supported KEGG organisms

<table>
<thead>
<tr>
<th>Supported Organisms</th>
<th>R command</th>
</tr>
</thead>
<tbody>
<tr>
<td>Homo sapiens</td>
<td>‘hsa’</td>
</tr>
<tr>
<td>Mus musculus</td>
<td>‘mmu’</td>
</tr>
<tr>
<td>Drosophila melanogaster</td>
<td>‘dme’</td>
</tr>
<tr>
<td>Saccharomyces cerevisiae</td>
<td>‘sce’</td>
</tr>
<tr>
<td>Arabidopsis thaliana</td>
<td>‘ath’</td>
</tr>
<tr>
<td>Rattus norvegicus</td>
<td>‘rno’</td>
</tr>
<tr>
<td>Danio rerio</td>
<td>‘dre’</td>
</tr>
</tbody>
</table>

DEsubs operates with Entrez ID labels, however twelve other label systems are supported after converting to Entrez IDs via a lexicon included in the package itself (see Table 3).

Table 3: Supported gene labels

<table>
<thead>
<tr>
<th>Supported Labels</th>
<th>R command</th>
</tr>
</thead>
<tbody>
<tr>
<td>Entrez</td>
<td>‘entrezgene’</td>
</tr>
<tr>
<td>Ensemble</td>
<td>‘ensembl_gene_id’, ‘ensembl_transcript_id’</td>
</tr>
<tr>
<td></td>
<td>‘ensembl_peptide_id’</td>
</tr>
<tr>
<td>HGNC</td>
<td>‘hgnc_id’, ‘hgnc_symbol’, ‘hgnc_transcript_name’</td>
</tr>
<tr>
<td>Refseq</td>
<td>‘refseq_mrna’, ‘refseq_peptide’</td>
</tr>
</tbody>
</table>
4. Pathway network processing

Next, the RNA-seq data are mapped onto the nodes and edges of the pathway network and two the pruning rules are applied to isolate interactions of interest among statistically significant differentially expressed genes (DEGs). DEGs are identified using the differential expression analysis tools in Table 5 by considering the FDR-adjusted P-value of each gene (Q-value). Instead of selecting one tools in Table 5, the user can import a custom ranked list of genes accompanied by their Q-values (argument `rankedList`).

Based on this information, the NodeRule prunes the nodes of the original network $G=(V,E)$, where Q-threshold (argument `DEpar`) defaults to 0.05:

$$ Q_{value}(i) < Q_{threshold}, i \in V $$

Next, the interactions among the selected genes are pruned based on both prior biological knowledge and the expression profiles of neighbouring genes, , where C-threshold defaults to 0.6 (argument `CORpar`):

$$ cor(i, j) * reg(i, j) > C_{threshold}, , i, j \in V $$

If genes i,j are connected with an edge with an activation type, then reg is set to 1, while if it the activation type is inhibitory, it is set to -1. The correlation between the profiles of the two genes i, j is calculated using the measures in Table 4 (argument `CORtool`).

Table 4: Edge Rule options

<table>
<thead>
<tr>
<th>Type</th>
<th>R command</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pearson product-moment correlation coefficient</td>
<td>‘pearson’</td>
</tr>
<tr>
<td>Spearman rank correlation coefficient</td>
<td>‘spearman’</td>
</tr>
<tr>
<td>Kendall rank correlation coefficient</td>
<td>‘kedhall’</td>
</tr>
</tbody>
</table>

Table 5: Node Rule options

<table>
<thead>
<tr>
<th>Supported Labels</th>
<th>R command</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Robinson, McCarthy, and Smyth 2010)</td>
<td>‘edgeR’</td>
</tr>
<tr>
<td>(Anders and Huber 2010)</td>
<td>‘DESeq2’</td>
</tr>
<tr>
<td>(Leng et al. 2013)</td>
<td>‘EBSeq’</td>
</tr>
<tr>
<td>(Smyth 2004)</td>
<td>‘vst2+limma’</td>
</tr>
<tr>
<td>(Anders and Huber 2010); (Smyth 2004)</td>
<td>‘voom+limma’</td>
</tr>
<tr>
<td>(Di et al. 2011)</td>
<td>‘NBPSeg’</td>
</tr>
<tr>
<td>(Auer and Doerge 2011)</td>
<td>‘TSPM’</td>
</tr>
</tbody>
</table>
Subpathway extraction is based on five main categories, (i) components, (ii) communities, (iii) streams, (iv) neighborhoods, (v) cascades. Each one sketches different topological aspect within the network. Indicative examples and a short description of DEsubs five main subpathway categories can be found in Figure 1.

The component category extracts strongly connected group of genes indicating dense local areas within the network. The community category extracts linked genes sharing a common property within the network. Thus the user can observe local gene sub-areas with a specific role within the network. Cascade, stream and neighborhood categories are generated starting from a gene of interest (GOI) in order to view the local perturbations within the network from different points of interest. The generation is performed by traversing either the forward or the backward propagation that stems from the GOI and is illustrated via three different topological schemes, gene sequences (‘cascade’ category), gene streams (‘stream’ category) and gene direct neighbors (‘neighborhood’ category).

Gene of interest (GOI)

Genes having crucial topological or functional roles within the network are considered as GOIs (see Table 6). The topological roles are portrayed using various topological measures from igraph package (Csardi and Nepusz 2006) capturing the local as well as global aspects of the network. Genes with crucial functional roles are considered as key genes for several biological and pharmacological features through \( f_{score} \), a measure which estimates how a gene acts as a bridge among specific
A high value of $f_{score}(i)$ for a condition $j$ indicates that gene $i$ participates to several functional terms of condition $j$ (for e.g. terms for diseases), hence it operates as a bridge for the terms of condition $j$ within the graph. As functional conditions we considered various biological and pharmacological features related with pathways, gene ontologies, diseases, drugs, microRNAs and transcription factors. External references are used to imprint gene associations by each feature separately. The references based on the approach of (Barneha, Jafari, and Mirzaie 2015); (Chen et al. 2013); (Li et al. 2011); (Vrahatis et al. 2016). Details are shown in Table 6.

Summarizing, a user-defined threshold (namely $top$) is used for the selection of GOIs. After the calculation of topological measures and functional measures by each condition (through $f_{score}$), the genes with the top best values are considered as GOIs. The parameter $top$ is user-defined with default value $top = 30$. In GOIs with crucial functional role we add and those having the lowest Q-value by each user-defined experimental dataset. Thus, the user is allowed to generate subpathways starting from the most statistical significant DEGs of his own experiment. A short description for all GOI types along with the corresponding parameters are shown in Table 6.

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
<th>R command</th>
</tr>
</thead>
<tbody>
<tr>
<td>Topological</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Degree</td>
<td>Number of adjacent interactions of the gene</td>
<td>‘degree’</td>
</tr>
<tr>
<td>Betweenness</td>
<td>Number of shortest paths from all vertices to all others that pass through that node</td>
<td>‘betweenness’</td>
</tr>
<tr>
<td>Closeness</td>
<td>Inverse of farness, which is the sum of distances to all other nodes</td>
<td>‘closeness’</td>
</tr>
<tr>
<td>Hub score</td>
<td>Kleinbergs hub centrality score</td>
<td>‘hub_score’</td>
</tr>
<tr>
<td>Eccentricity</td>
<td>Shortest path distance from the farthest node in the graph</td>
<td>‘eccentricity’</td>
</tr>
<tr>
<td>Page rank</td>
<td>Google Page Rank</td>
<td>‘page_rank’</td>
</tr>
<tr>
<td>Start Nodes</td>
<td>Nodes without any incoming links</td>
<td>‘start_nodes’</td>
</tr>
<tr>
<td>Functional</td>
<td></td>
<td></td>
</tr>
<tr>
<td>DEG</td>
<td>Genes highly differentially expressed according to the experimental data</td>
<td>‘deg’</td>
</tr>
<tr>
<td>Pathways</td>
<td>Genes acting as bridges among KEGG pathways</td>
<td>‘KEGG’</td>
</tr>
<tr>
<td>Biological Process</td>
<td>Genes acting as bridges among Gene Ontology Biological Process terms</td>
<td>‘GO_bp’</td>
</tr>
<tr>
<td>Cellular Component</td>
<td>Genes acting as bridges among Gene Ontology Cellular Component terms</td>
<td>‘GO_cc’</td>
</tr>
<tr>
<td>Molecular Function</td>
<td>Genes acting as bridges among Gene Ontology Molecular Function terms</td>
<td>‘GO_mf’</td>
</tr>
</tbody>
</table>
5.3. All subpathway options

Cascade, stream and neighborhood subpathway types can start from seventeen (17) different GOI types and their generation is performed either with forward or backward propagation. Thus, thirty-four (34) different types are created for each of the three types. Also, the component-based types are sixteen and the community-based types are six based on igraph package. DEmsubs therefore supports 124 subpathway types as described in Tables 7-11.

Table 7: Subpathway Options - STREAM

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
<th>R command</th>
</tr>
</thead>
<tbody>
<tr>
<td>Disease</td>
<td>Genes acting as bridges for OMIM targets</td>
<td>‘Disease_OMIM’</td>
</tr>
<tr>
<td>Disease</td>
<td>Genes acting as bridges for GAD targets</td>
<td>‘Disease_GAD’</td>
</tr>
<tr>
<td>Drug</td>
<td>Genes acting as bridges for DrugBank targets</td>
<td>‘Drug_DrugBank’</td>
</tr>
<tr>
<td>microRNA</td>
<td>Genes acting as bridges for microRNA targets</td>
<td>‘miRNA’</td>
</tr>
<tr>
<td>Transcription Factors</td>
<td>Genes acting as bridges for TF targets</td>
<td>‘TF’</td>
</tr>
</tbody>
</table>

Topological
Forward and backward streams starting from genes/nodes with crucial topological roles within the network
- ‘fwd.stream.topological.degree’
- ‘fwd.stream.topological.betweenness’
- ‘fwd.stream.topological.closeness’
- ‘fwd.stream.topological_hub_score’
- ‘fwd.stream.topological.eccentricity’
- ‘fwd.stream.topological.page_rank’
- ‘fwd.stream.topological.start_nodes’
- ‘bwd.stream.topological.degree’
- ‘bwd.stream.topological.betweenness’
- ‘bwd.stream.topological.closeness’
- ‘bwd.stream.topological_hub_score’
- ‘bwd.stream.topological.eccentricity’
- ‘bwd.stream.topological.page_rank’
- ‘bwd.stream.topological.start_nodes’

Functional
Forward and backward streams starting from genes/nodes with crucial functional roles within the network
- ‘fwd.stream.functional.GO_bp’
- ‘fwd.stream.functional.GO_cc’
- ‘fwd.stream.functional.GO_mf’
- ‘fwd.stream.functional.Disease_OMIM’
- ‘fwd.stream.functional.Disease_GAD’
- ‘fwd.stream.functional.Drug_DrugBank’
- ‘fwd.stream.functional.miRNA’
- ‘fwd.stream.functional.TF’
- ‘fwd.stream.functional.KEGG’
- ‘fwd.stream.functional.DEG’
- ‘bwd.stream.functional.GO_bp’
- ‘bwd.stream.functional.GO_cc’
<table>
<thead>
<tr>
<th>Description</th>
<th>R parameter</th>
</tr>
</thead>
</table>
| Topological | 'fwd.neighbourhood.topological.degree'  
|             | 'fwd.neighbourhood.topological.betweenness'  
|             | 'fwd.neighbourhood.topological.closeness'  
|             | 'fwd.neighbourhood.topological.hub_score'  
|             | 'fwd.neighbourhood.topological.eccentricity'  
|             | 'fwd.neighbourhood.topological.page_rank'  
|             | 'fwd.neighbourhood.topological.start_nodes'  
|             | 'bwd.neighbourhood.topological.degree'  
|             | 'bwd.neighbourhood.topological.betweenness'  
|             | 'bwd.neighbourhood.topological.closeness'  
|             | 'bwd.neighbourhood.topological.hub_score'  
|             | 'bwd.neighbourhood.topological.eccentricity'  
|             | 'bwd.neighbourhood.topological.page_rank'  
|             | 'bwd.neighbourhood.topological.start_nodes'  

| Functional | R parameter |
|           | 'fwd.neighbourhood.functional.GO_bp'  
|           | 'fwd.neighbourhood.functional.GO_cc'  
|           | 'fwd.neighbourhood.functional.GO_mf'  
|           | 'fwd.neighbourhood.functional.Disease_OMIM'  
|           | 'fwd.neighbourhood.functional.Disease_GAD'  
|           | 'fwd.neighbourhood.functional.Drug_DrugBank'  
|           | 'fwd.neighbourhood.functional.miRNA'  
|           | 'fwd.neighbourhood.functional.TF'  
|           | 'fwd.neighbourhood.functional.KEGG'  
|           | 'fwd.neighbourhood.functional.DEG'  
|           | 'bwd.neighbourhood.functional.GO_bp'  
|           | 'bwd.neighbourhood.functional.GO_cc'  
|           | 'bwd.neighbourhood.functional.GO_mf'  
|           | 'bwd.neighbourhood.functional.Disease_OMIM'  
|           | 'bwd.neighbourhood.functional.Disease_GAD'  
|           | 'bwd.neighbourhood.functional.Drug_DrugBank'  

Table 8: Subpathway Options - NEIGHBOURHOOD
<table>
<thead>
<tr>
<th>Description</th>
<th>R parameter</th>
</tr>
</thead>
</table>
| Topological | ‘fwd.cascade.topological.degree’  
|            | ‘fwd.cascade.topological.betweenness’  
|            | ‘fwd.cascade.topological.closeness’  
|            | ‘fwd.cascade.topological.hub_score’  
|            | ‘fwd.cascade.topological.eccentricity’  
|            | ‘fwd.cascade.topological.page_rank’  
|            | ‘fwd.cascade.topological.start_nodes’  
|            | ‘bwd.cascade.topological.degree’  
|            | ‘bwd.cascade.topological.betweenness’  
|            | ‘bwd.cascade.topological.closeness’  
|            | ‘bwd.cascade.topological.hub_score’  
|            | ‘bwd.cascade.topological.eccentricity’  
|            | ‘bwd.cascade.topological.page_rank’  
|            | ‘bwd.cascade.topological.start_nodes’  |
| Functional  | ‘fwd.cascade.functional.GO_bp’  
|            | ‘fwd.cascade.functional.GO_cc’  
|            | ‘fwd.cascade.functional.GO_mf’  
|            | ‘fwd.cascade.functional.Disease_OMIM’  
|            | ‘fwd.cascade.functional.Disease_GAD’  
|            | ‘fwd.cascade.functional.Drug_DrugBank’  
|            | ‘fwd.cascade.functional.mRNA’  
|            | ‘fwd.cascade.functional.TF’  
|            | ‘fwd.cascade.functional.KEGG’  
|            | ‘fwd.cascade.functional.DEG’  
|            | ‘bwd.cascade.functional.GO_bp’  
|            | ‘bwd.cascade.functional.GO_cc’  
|            | ‘bwd.cascade.functional.GO_mf’  
|            | ‘bwd.cascade.functional.Disease_OMIM’  
|            | ‘bwd.cascade.functional.Disease_GAD’  
|            | ‘bwd.cascade.functional.Drug_DrugBank’  
|            | ‘bwd.cascade.functional.mRNA’  
|            | ‘bwd.cascade.functional.TF’  
|            | ‘bwd.cascade.functional.KEGG’  
|            | ‘bwd.cascade.functional.DEG’  |

Table 9: Subpathway Options - CASCADE
Table 10: Subpathway Options - COMMUNITY

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
<th>R parameter</th>
</tr>
</thead>
<tbody>
<tr>
<td>Random Walk</td>
<td>Community structures that minimize the expected description length of a random walker trajectory</td>
<td>‘community.infomap’</td>
</tr>
<tr>
<td>Modular</td>
<td>Community structures via a modularity measure and a hierarchical approach</td>
<td>‘community.louvain’</td>
</tr>
<tr>
<td>Walktraps</td>
<td>Densely connected subgraphs via random walks</td>
<td>‘community.walktrap’</td>
</tr>
<tr>
<td>Leading eigen</td>
<td>Densely connected subgraphs based on the leading non-negative eigenvector of the modularity matrix</td>
<td>‘community.leading__eigen’</td>
</tr>
<tr>
<td>Betweenness</td>
<td>Community structures detection via edge betweenness</td>
<td>‘community.edge_betweenness’</td>
</tr>
<tr>
<td>Greedy</td>
<td>Community structures via greedy optimization of modularity</td>
<td>‘community.fast__greedy’</td>
</tr>
</tbody>
</table>

Table 11: Subpathway Options - COMPONENT

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
<th>R parameter</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cliques</td>
<td>A subgraph where every two distinct vertices in the clique are adjacent</td>
<td>‘component.3-cliques’</td>
</tr>
<tr>
<td>K-core</td>
<td>A maximal subgraph in which each vertex has at least degree k</td>
<td>‘component.3-coreness’</td>
</tr>
<tr>
<td>Max cliques</td>
<td>Largest of maximal cliques</td>
<td>‘component.9-coreness’</td>
</tr>
<tr>
<td>Components</td>
<td>All single components</td>
<td>‘component.max_cliques’</td>
</tr>
</tbody>
</table>

An example follows where *community.walktrap* is selected as the subpathway type.

```r
desub.run <- desubs(
  org='hsa',
  mRNAexpr=mRNAexpr,
  mRNAomnomenclature='entrezgene',
  pathways='All',
  DEtool=NULL, DEpar=0.05,
  CORtool='pearson', CORpar=0.7,
  subpathwayType='community.walktrap',
  rankedList=rankedList,
  verbose=FALSE)
```

10
6. Subpathway enrichment analysis

Eight different datasets with external resources are stored locally for further enrichment analysis of resulting subpathways. Each dataset is formed with a list of terms related to biological and pharmacological features and the respective associated genes for each term. A detailed description is shown in Table 12. Additionally, the user can supply a custom gene-set in the form of an .RData file storing a matrix, named targetsPerClass. The matrix should store the terms as rownames and the targets of each term at each row. Since some rows are bound to have more elements that others, empty cells should be filled with ‘0’ characters. Once the file is stored in a directory DESubs/Data, it will be permanently available along with the other eight resources, using the filename (without the .RData suffix) as the new functional feature type along with any of the default eight features.

The enrichment analysis is performed based on the cumulative hypergeometric distribution, where G is the number of genes in the user input list, l the number of those genes included in the subpathway, D the number of associated genes for a term and d the number of genes included in the subpathway (Li et al. 2013). Terms with P < 0.05 are regarded as terms with significant association with the respective subpathway.

\[
P = 1 - \sum_{x=0}^{d} \frac{\binom{D}{x} \binom{G-D}{l-x}}{\binom{G}{l}}
\]

Table 12: List of external databases

<table>
<thead>
<tr>
<th>Type</th>
<th>Description (# of terms)</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pathway Term</td>
<td>KEGG pathway maps (179)</td>
<td>(Chen et al. 2013)</td>
</tr>
<tr>
<td>GO Biological Process</td>
<td>Genes sharing a common biological process (5.192)</td>
<td>(Chen et al. 2013)</td>
</tr>
<tr>
<td>GO Cellular Component</td>
<td>Genes sharing a common cellular component (641)</td>
<td>(Chen et al. 2013)</td>
</tr>
<tr>
<td>GO Molecular Function</td>
<td>Genes sharing a common molecular level (1.136)</td>
<td>(Chen et al. 2013)</td>
</tr>
<tr>
<td>OMIM Disease</td>
<td>Disease related genes (90)</td>
<td>(Chen et al. 2013)</td>
</tr>
<tr>
<td>GAD Disease</td>
<td>Disease related genes (412)</td>
<td>(Li et al. 2011)</td>
</tr>
<tr>
<td>DrugBank Drug</td>
<td>Gene targets of drugs (1.488)</td>
<td>(Barneh, Jafari, and Mirzaie 2015)</td>
</tr>
<tr>
<td>Transcription Factor</td>
<td>Gene targetsof transcription factors (290)</td>
<td>(Chen et al. 2013)</td>
</tr>
</tbody>
</table>

Table 13: Example of custom gene set

<table>
<thead>
<tr>
<th>Term</th>
<th>Target 1</th>
<th>Target 2</th>
<th>Target 3</th>
<th>Target 4</th>
<th>Target 5</th>
<th>Target 6</th>
<th>Target 7</th>
</tr>
</thead>
<tbody>
<tr>
<td>Term 1</td>
<td>ACAD9</td>
<td>ACAD8</td>
<td>SH3GLB1</td>
<td>ESCO2</td>
<td>ESCO1</td>
<td>ADH1C</td>
<td>'0'</td>
</tr>
<tr>
<td>Term 2</td>
<td>ADH1C</td>
<td>ADH1B</td>
<td>ADHFE1</td>
<td>ADH1A</td>
<td>ADH6</td>
<td>ADH7</td>
<td>ADH4</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>Term N</td>
<td>PTPN1</td>
<td>RHOA</td>
<td>ACTN4</td>
<td>ACTN3</td>
<td>ACTN2</td>
<td>'0'</td>
<td>'0'</td>
</tr>
</tbody>
</table>
Figure 2: Subpathway extraction options consist of five main categories. The three of them (cascade, neighborhood, stream) are sub-categorized according to features (topological or functional) and the direction of propagation (forward or backward) of the gene of interest where each subpathway is starting. The other two (component, community) are sub-categorized according to various topological properties.
7. Visualization

DEsubs visualizes its results at a gene, subpathway and organism level through various schemes such as bar plots, heat maps, directed weighted graphs, circular diagrams (Gu et al. 2014) and dot plots. Indicative examples are illustrated in figures 2-8 based on DEsubs executions using the human pathway network and a synthetic dataset. Bar plots show the genes with the best Q-value from the user-selected DE analysis tool (the user defines the desired gene number). The figures are exported in the directory Output within the user specified location. Heat maps show the genes with the highest values either in our topological or functional measures (see Table 6).

Figure 3: Bar plots show the genes with the best Q-value from the user-selected DE analysis tool (the user defines the desired gene number). Heat maps show the genes with the highest values either in our topological or functional measures (see Table 6). Each extracted subpathway is illustrated though a directed graph by imprinting the degree of DE and correlation among the respective gene members. Subpathway enrichment in association with biological and pharmacological features (such as pathway terms, gene ontologies, regulators, diseases and drug targets) is depicted through circular diagrams. The total picture of the enriched subpathways is performed with dot plots.
7.1 Gene Level Visualization

```r
res <- geneVisualization(
  DEsubs.out="DEsubs.out", top=10,
  measures.topological=c('degree', 'betweenness', 'closeness',
                          'eccentricity', 'page_rank'),
  measures.functional=c('KEGG', 'GO_bp', 'GO_cc', 'GO_mf',
                        'Disease_OMIM', 'Disease_GAD',
                        'Drug_DrugBank', 'miRNA', 'TF'),
  size.topological=c(5,4),
  size.functional=c(7,4),
  size.barplot=c(5,6),
  export='plot', verbose=FALSE)
```

![Gene Level Visualization](image)

Figure 4: Bars illustrate the genes with the highest Q-values.
Figure 5: Heat map represents the twelve genes with the highest values of functional measures. The values are scaled and the red graduation indicates the value degree.

Figure 6: Heat map represents the twelve genes with the highest values of topological measures. The values are scaled and the red graduation indicates the value degree.
7.2. Subpathway Level Visualization

Each extracted subpathway is illustrated though a directed graph by imprinting the degree of differential expression and correlation among the respective gene members. Additionally, it can be extracted in a variety of formats so that it can be used by external software, such as .txt, .json, .gml, .ncol, .lgl, .graphml and .dot formats.

```r
res <- subpathwayToGraph(
  DEsubs.out=DEsubs.out,
  submethod='community.walktrap',
  subname='sub6', verbose=FALSE,
  export='plot'
)
```

Figure 7: Graph illustrates the links of a subpathway. Red graduation in nodes indicate the Q-value degree, the edge width indicates the correlation degree between the respective genes. Green or red color in edges indicates the positive or negative correlation respectively.

Subpathway enrichment in association with biological and pharmacological features (such as pathway terms, gene ontologies, regulators, diseases and drug targets) is depicted through circular diagrams.

```r
res <- subpathwayVisualization(
  DEsubs.out=DEsubs.out,
  references=c('GO', 'TF'),
  submethod='community.walktrap',
  subname='sub1',
  scale=c(1, 1),
  export='plot',
  verbose=FALSE
)
```
Figure 8: Circular Diagram shows the associations among genes including in a subpathway and Gene Ontology terms where are enriched

Figure 9: Circular Diagram shows the associations among genes included in a subpathway and enriched Transcription Factors.
7.3. Organism Level Visualization

The total picture of the enriched subpathways is performed with dot plots. The number of features represented are selected using `topTerms` argument.

```r
res <- organismVisualization(
  DEsubs.out=DEsubs.out,
  references='KEGG',
  topSubs=10,
  topTerms=20,
  export='plot',
  verbose=FALSE)
```

Figure 10: Dot plot shows the enriched associations among experiment-specific extracted subpathways and pathways from KEGG database. Twenty pathways were selected as the desired number of terms.
References


