Network Module Detection from Multi-Modal Node Features with a Greedy Decision Forest

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Input

For Network Module Detection we need a network and node feature matrices. Here, we use a PPI-Network and multi-omics node features from gene expression and methylation data. In addition to that, a binary vector needs to be specified reflecting the outcome class. Here, it reflects the patient group survived vs non-survived.

```
PPI <- read.table("~/LinkedOmics/KIRC/KIDNEY_PPI.txt")
mRNA <- read.table("~/LinkedOmics/KIRC/KIDNEY_mRNA_FEATURES.txt")
Methy <- read.table("~/LinkedOmics/KIRC/KIDNEY_Methy_FEATURES.txt")
TARGET <- read.table("~/LinkedOmics/KIRC/KIDNEY_SURVIVAL.txt")</pre>
```

Dimensions are:

```
dim(PPI)
```

```
## [1] 6926452 3
dim(mRNA)
```

```
## [1] 306 12029
dim(Methy)
```

```
## [1] 306 12029
dim(TARGET)
```

```
## [1] 1 306
```

The PPI Network has 6926452 edges and is organized as follows:

```
head(PPI,5)
```

```
##
     protein1 protein2 combined_score
## 1
         ARF5
                  CALM2
## 3
         ARF5
                   ERN1
                                     159
## 4
         ARF5
                                     606
                 CDKN2A
## 5
         ARF5
                   P4HB
                                     167
         ARF5
                  STX10
                                     267
```

The first two columns refer to the connected nodes. The last column indicate the confidence of the interaction between these nodes/protein.

The node feature matrices are organized as follows:

```
mRNA[1:5,1:5]
```

```
## TCGA.3Z.A93Z 10.1967 10.8407 11.0698 3.0921 8.4911 ## TCGA.6D.AA2E 10.4898 11.2592 11.4613 3.4214 5.9663 ## TCGA.A3.3357 10.8225 11.4032 11.5370 3.0013 4.6062 ## TCGA.A3.3358 11.6874 10.9420 12.8086 5.4678 5.1437 ## TCGA.A3.3367 11.3013 11.0082 11.8861 4.9567 6.2678
```

where the rows are reflecting the patients and the columns are the genes/nodes. In this case, we analyze 12029 node feature values of 306 patients.

The same for the second multi-omics:

```
Methy[1:5,1:5]
```

```
## TCGA.3Z.A93Z -0.4897 -0.4686 -0.0063 -0.4851 -0.4156

## TCGA.6D.AA2E -0.4885 -0.4574 0.1481 -0.4799 -0.3343

## TCGA.A3.3357 -0.4854 -0.4721 -0.0540 -0.4859 -0.3052

## TCGA.A3.3358 -0.4838 -0.4339 -0.1005 -0.4778 -0.3213

## TCGA.A3.3367 -0.4890 -0.4684 -0.0493 -0.4830 -0.2719
```

The rows of the multi-modal feature matrices should refer to the exact same patient.

Finally, we need the target vector specifying the survival (0) and non-survival (1) groups.

```
TARGET [1:5]
```

```
## TCGA.3Z.A93Z TCGA.6D.AA2E TCGA.A3.3357 TCGA.A3.3358 TCGA.A3.3367 ## 13 0 0 0 0 0 0
```

Creating a DFNET graph object

```
library(DFNET)
require(igraph)
require(ranger)
require(pROC)

DFNET_graph <- DFNET_generate_graph_Omics(PPI, list(mRNA, Methy), TARGET, 0.95)
summary(DFNET_graph)</pre>
```

```
## Length Class Mode
## graph 10 igraph list
## Feature_Matrix 2 -none- list
## gene.names 9033 -none- character
```

The DFNET_graph object is a list and consists of three slots. The first slot is the PPI network internally converted to a igraph object.

```
DFNET_graph$graph
```

```
## IGRAPH cb42a4f U--- 9032 322206 --
## + edges from cb42a4f:
## [1] 365--3411 365--1618 365--5723 365--8823 365--2469 365--4046 365--3793
## [8] 365--6680 365--8829 365--6551 365--7310 365--3929 365--2550 365--5588
## [15] 365--1359 365--7638 120-- 365 140-- 365 365--1158 365--2171 365-- 589
```

```
## [22] 365--7900 365--6420 365--1297 365--8699 365--5182 365--4736 365--5805 ## [29] 365--7215 365--1538 365--8643 365--6355 365--4688 365--6619 365--1073 ## [36] 365--2961 365--2038 239-- 365 365--4766 278-- 365 365--4778 365--1284 ## [43] 365--3950 365--1262 365--2285 102-- 365 365--6932 365--5808 365--449 ## [50] 365--6536 365--1022 365--3422 365--8567 365--9011 365--6928 365--2717 ## [57] 365--757 365--5728 365--2213 170-- 365 365--2813 365--2968 365--2889 ## + ... omitted several edges
```

The second slot is a list of feature matrices.

```
DFNET_graph$Feature_Matrix[[1]][1:5,1:5]
```

```
##
                   AN_1
                           AN 2
                                    AN 3
                                           AN 4
                                                  AN 5
## TCGA.3Z.A93Z 10.1967 10.8407 11.0698 3.0921 8.4911
## TCGA.6D.AA2E 10.4898 11.2592 11.4613 3.4214 5.9663
## TCGA.A3.3357 10.8225 11.4032 11.5370 3.0013 4.6062
## TCGA.A3.3358 11.6874 10.9420 12.8086 5.4678 5.1437
## TCGA.A3.3367 11.3013 11.0082 11.8861 4.9567 6.2678
DFNET_graph$Feature_Matrix[[2]][1:5,1:5]
##
                   BN_1
                           BN_2
                                   BN 3
                                            BN 4
                                                    BN 5
## TCGA.3Z.A93Z -0.4897 -0.4686 -0.0063 -0.4851 -0.4156
## TCGA.6D.AA2E -0.4885 -0.4574 0.1481 -0.4799 -0.3343
```

The third slot contains the node/gene names.

```
head(DFNET_graph$gene.names,5)
```

```
## [1] "RBL2" "VDAC3" "ACTN1" "ATP2A1" "SFRP1"
```

TCGA.A3.3357 -0.4854 -0.4721 -0.0540 -0.4859 -0.3052 ## TCGA.A3.3358 -0.4838 -0.4339 -0.1005 -0.4778 -0.3213 ## TCGA.A3.3367 -0.4890 -0.4684 -0.0493 -0.4830 -0.2719

A DFNET_graph object can thus be easily created. Note, the node names need to be as specified. A prefix letter followed with a "N" and than simply the node identifier. Node identifier should match the identifier used for the igraph network.

```
head(as_edgelist(DFNET_graph$graph, names = TRUE))
```

```
## [,1] [,2]
## [1,] 365 3411
## [2,] 365 1618
## [3,] 365 5723
## [4,] 365 8823
## [5,] 365 2469
## [6,] 365 4046
```

As seen from the above table "AN_365" and "AN_3411" are connected. The same is true for the second modality "BN_365" and "BN_3411".

DFNET for Subnetwork Detection

The main function for network module detection expects the number of trees (ntrees), the number of greedy iteration (niter), and the initial size of the module as an input. The ntrees parameter specifies the number of random works initialized. The niter parameter sets the total number of greedy iterations, and the init.mtry defines the depth of the random work, and thus the initial size of the modules.

```
DFNET_object <- DFNET(DFNET_graph, ntrees=100, niter=100, init.mtry=10)
summary(DFNET_object)</pre>
```

```
## Length Class Mode
## DFNET_graph 3 -none- list
## DFNET_trees 600 -none- list
## DFNET_MODULES 100 -none- list
## DFNET_MODULES_AUC 100 -none- numeric
```

The DFNET_object contains four slots. The first slot "DFNET_graph" is the igraph object storing the network topology. The second slot "DFNET_trees" contains the Decision Trees of the Decision Forest. The third slot "DFNET_MODULES" stores the detected Network Modules, and the "DFNET_MODULES_AUC" consists of the corresponding AUC values of the modules. The accuracy of the Decision Forest Classifier can be calculated as

```
DFNET_acc <- DFNET_accuracy(DFNET_graph, DFNET_object)</pre>
```

[1] 0.7667751

DFNET Edge Importance Scores

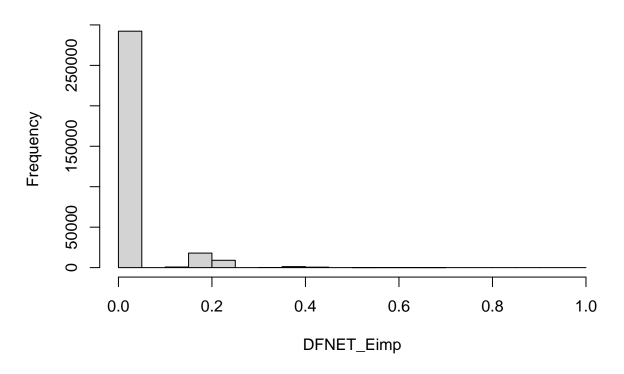
Edge Importance Scores can be calculated with the following function

```
DFNET_Eimp <- DFNET_Edge_Importance(DFNET_graph, DFNET_object)
```

```
length(DFNET_Eimp)
```

```
## [1] 322206
hist(DFNET_Eimp)
```

Histogram of DFNET_Eimp



DFNET Detected Modules

The detected modules can be retrieved via the "DFNET_modules" function

```
DFNET_mod <- DFNET_modules(DFNET_graph, DFNET_object, DFNET_Eimp)
```

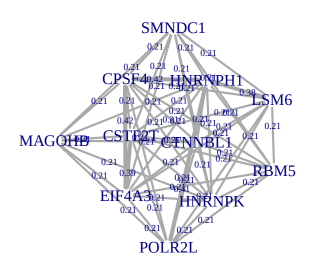
head(DFNET_mod)

```
##
                                                              Module EDGE IMP
## 2
             1532 4653 4917 5294 5495 6128 6297 6635 6784 8430 9013 2.182502
## 1 307 436 4276 4653 5301 5470 5583 6019 6379 6384 7723 8323 8883 2.200262
## 3
                             504 2813 4009 4880 5441 5972 7096 8890 1.964621
               11 3374 3771 4099 4292 4601 5952 7011 7371 7582 8592 1.684959
                            1869 4295 4379 5770 6038 6979 7453 8680 1.636016
## 5
## 6
             1068 1305 1749 4185 4420 4668 4973 5240 6133 7660 8807 1.591245
##
           AUC
                    IMP
## 2 0.7016790 2.884181
## 1 0.6684211 2.868683
## 3 0.6827068 2.647328
## 4 0.6961575 2.381117
## 5 0.7231853 2.359201
## 6 0.7532468 2.344492
```

The modules are ranked by their importance (last column). Note, node ids are shown, but the actual node names can be retrieved from "DFNET_graph\$gene.names".

Lets have a closer look at the top ranked module

```
<- as.numeric(strsplit(DFNET_mod[1,1]," ")[[1]])</pre>
DFNET_graph$gene.names[Nodes]
    [1] "SMNDC1"
                                                             "HNRNPH1" "EIF4A3"
                  "CTNNBL1" "CPSF4"
                                        "POLR2L"
                                                  "LSM6"
    [8] "MAGOHB"
                   "RBM5"
                             "HNRNPK"
                                        "CSTF2T"
##
The module is reflected by the following edges and nodes
Top_Module <- DFNET_get_module(Nodes, DFNET_graph, DFNET_Eimp)</pre>
head(Top_Module)
##
        GENE
                 GENE
                               EDGE_IMP
      SMNDC1 CTNNBL1 0.212290409880912
## 1
      SMNDC1
               CPSF4 0.212290409880912
## 3 CTNNBL1
               CPSF4 0.212290409880912
## 4 CTNNBL1 POLR2L 0.212290409880912
## 5
       CPSF4 POLR2L 0.212290409880912
                 LSM6 0.212290409880912
## 6 SMNDC1
We can visualize this subgraph using the function "DFNET_plot_module".
require(igraph)
DFNET_plot_module(Nodes, DFNET_graph, DFNET_Eimp)
```



DFNET Node Feature importance scores

The feature importances of the nodes of that module can be calculated as

```
DFNET_Fimp
             <- DFNET_calc_feature_importance(Nodes, DFNET_object, DFNET_graph)</pre>
DFNET_Fimp
              SMNDC1
                        CTNNBL1
                                      CPSF4
                                                 POLR2L
                                                               LSM6
                                                                        HNRNPH1
##
## omic1 0.01243056 0.01299474 0.11582528 0.02890176 0.01923387 0.08894694
## omic2 0.00000000 0.00000000 0.07120735 0.03436464 0.07839865 0.02436282
##
              EIF4A3
                         MAGOHB
                                       RBM5
                                                 HNRNPK
                                                             CSTF2T
## omic1 0.06299942 0.04710311 0.01071508 0.01244777 0.01919737
## omic2 0.06120851 0.04662184 0.00757414 0.00000000 0.03246349
## GGPLOT
library(ggplot2)
library(reshape)
RES1 <- cbind(colnames(DFNET_Fimp),DFNET_Fimp[1,])</pre>
RES2 <- cbind(colnames(DFNET_Fimp),DFNET_Fimp[2,])</pre>
RES1 <- cbind(RES1,"mRNA")</pre>
RES2 <- cbind(RES2, "Methylation")</pre>
RES <- rbind(RES1,RES2)</pre>
rownames(RES) <- NULL
colnames(RES) <- c("Gene","IMP","Type")</pre>
        <- as.data.frame(RES)</pre>
RES$IMP <- as.numeric(RES$IMP)</pre>
p <- ggplot(RES, aes(fill=Type, y=IMP, x=Gene)) +
    geom_bar(position="dodge", stat="identity") +
    ylab("Feature Importance") +
    theme(axis.text.x = element_text(angle = 90, vjust = 0.5, hjust=1))
plot(p)
```

