# Random Walk with Restart on Multiplex and Heterogeneous Network

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#### October 30, 2018

#### Abstract

This vignette describes how to use the *RandomWalkRestartMH* package to run Random Walk with Restart algorithms on monoplex, multiplex, heterogeneous and multiplex-heterogeneous networks. It is based on the work we presented on the following article:

https://www.biorxiv.org/content/early/2017/08/30/134734

#### Package

RandomWalkRestartMH 1.2.0

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## 1 Introduction

*RandomWalkRestartMH* (Random Walk with Restart on Multiplex and Heterogeneous Networks) is an R package built to provide easy access to the Random Walk with Restart (RWR) algorithm on different types of networks: i) Monoplex networks, ii) Multiplex networks, iii) Heterogeneous networks and iv) Multiplex-Heterogeneous networks. It is based on the work we presented in the article: https://www.biorxiv.org/content/early/2017/08/30/134734.

RWR simulates an imaginary particle that starts on a seed(s) node(s) and follows randomly the edges of a network. At each step, there is a restart probability, r, meaning that the particle can come back to the seed(s) [1]. This imaginary particle can explore the following types of networks:

- A monoplex or single network, which contains solely nodes of the same nature. In addition, all the edges belong to the same category.
- A multiplex network, defined as a collection of monoplex networks considered as layers of the multiplex network. In a multiplex network, the different layers share the same set of nodes, but the edges represent relationships of different nature [2]. In this case, the RWR particle can jump from one node to its counterparts on different layers.
- A heterogeneous network, which is composed of two monoplex networks containing nodes of different nature. These different kind of nodes can be connected thanks to bipartite edges, allowing the RWR particle to jump between the two networks.
- A multiplex and heterogeneous network, which is built by linking the nodes in every layer of a multiplex network to nodes of different nature thanks to bipartite edges. The RWR particle can now explore the full multiplex-heterogeneous network.

The user can introduce up to six single networks (monoplex networks) to create a multiplex network. The multiplex network can also be integrated, thanks to bipartite relationships, with a network containing nodes of different nature. Proceeding this way, a network both multiplex and heterogeneous will be generated.

Please note that this first version of the package deals only with undirected and unweighted networks. New functionalities will be included in future updated versions of *RandomWalkRestartMH*.

# 2 Installation of the *RandomWalkRestartMH* package

First of all, you need a current version of R (www.r-project.org). RandomWalkRestartMH is a freely available package deposited on http://bioconductor.org/. You can install it by running the following commands on an R console:

```
> if (!requireNamespace("BiocManager", quietly=TRUE))
```

```
+ install.packages("BiocManager")
```

> BiocManager::install("RandomWalkRestartMH")

## 3 A Detailed Workflow

In the following paragraphs, we describe how to use the *RandomWalkRestartMH* package to perform RWR on different types of biological networks. Concretely, we use a protein-protein interaction (PPI) network, a pathway network, a disease-disease similarity network and combinations thereof. These networks are obtained as detailed in [3]. The PPI and the Pathway network were reduced by only considering genes/proteins expressed in the adipose tissue, in order to reduce the computation time of this vignette.

The goal in the example presented here is, as described in [3], to find candidate genes potentially associated with diseases by a guilt-by-association approach. This is based on the fact that genes/proteins with similar functions or similar phenotypes tend to lie closer in biological networks. Therefore, the larger the RWR score of a gene, the more likely it is to be functionally related with the seeds.

We focus on a real biological example: the SHORT syndrome (MIM code: 269880) and its causative gene PIK3R1 as described in [3]. We will see throughout the following paragraphs how the RWR results evolve due to the the integration and exploration of additional networks.

### 3.1 Random Walk with Restart on a Monoplex Network

RWR has usually been applied within the framework of single PPI networks in bioinformatics[4]. A gene or a set of genes, so-called seed(s), known to be implicated in a concrete function or in a specific disease, are chosen as the starting point(s) of the algorithm. The RWR particle explores the neighbourhood of the seeds and the algorithm computes a score for all the nodes of the network. The larger it is the score of a node, the closer it is to the seed(s).

Let us generate an object of the class *Multiplex*, even if it is a monoplex network, with our PPI network.

```
> library(RandomWalkRestartMH)
> library(igraph)
> data(PPI_Network) # We load the PPI_Network
> ## We create a Multiplex object composed of 1 layer (It's a Monoplex Network)
> ## and we display how it looks like
> PPI_MultiplexObject <- create.multiplex(PPI_Network,Layers_Name=c("PPI"))</pre>
> PPI_MultiplexObject
Number of Layers:
[1] 1
Number of Nodes:
[1] 4317
IGRAPH 1573a63 UN-- 4317 18062 --
+ attr: name (v/c), type (e/c)
+ edges from 1573a63 (vertex names):
 [1] AAMP
            --VPS52
                         AAMP
                                --BHLHE40
                                            AAMP
                                                    --GABARAPL2 AAMP
                                                                       --MAP1LC3B
                                                                       --PRKAA1
 [5] VPS52 --TXN2
                         VPS52
                                            VPS52
                                                   --MFAP1
                                                                VPS52
                               - - DDX6
 [9] VPS52
           --LM04
                         VPS52 --STX11
                                            VPS52
                                                  --KANK2
                                                                VPS52
                                                                       --PPP1R18
[13] VPS52
            --TXLNA
                         VPS52
                               --KIAA1217
                                            VPS52
                                                   --VPS28
                                                                VPS52
                                                                       --ATP6V1D
[17] VPS52 --TPM3
                         VPS52 --KIF5B
                                            VPS52
                                                   --N0P2
                                                                VPS52
                                                                       --RNF41
```

[21] VPS52WTAP	VPS52MAPK3	VPS52ZMAT2	VPS52VPS51
[25] BHLHE40AES	BHLHE40 PRKAA1	BHLHE40 CCNK	BHLHE40 RBPMS
[29] BHLHE40COX5B	BHLHE40UBE2I	BHLHE40MAGED1	BHLHE40 PLEKHB2
+ omitted several	edges		

To apply the RWR on a monoplex network, we need to compute the adjacency matrix of the network and normalize it by column [4], as follows:

```
> AdjMatrix_PPI <- compute.adjacency.matrix(PPI_MultiplexObject)
> AdjMatrixNorm_PPI <- normalize.multiplex.adjacency(AdjMatrix_PPI)</pre>
```

Then, we need to define the seed(s) before running the RWR algorithm on this PPI network. As commented above, we are focusing on the example of the SHORT syndrome. Therefore, we take the *PIK3R1* gene as seed, and we execute RWR.

```
> SeedGene <- c("PIK3R1")</pre>
> ## We launch the algorithm with the default parameters (See details on manual)
> RWR_PPI_Results <- Random.Walk.Restart.Multiplex(AdjMatrixNorm_PPI,
                           PPI_MultiplexObject,SeedGene)
+
> # We display the results
> RWR_PPI_Results
Top 10 ranked Nodes:
   NodeNames
                    Score
1
        GRB2 0.006845881
2
        EGFR 0.006169129
3
         CRK 0.005674261
4
        ABL1 0.005617041
5
         FYN 0.005611086
6
       CDC42 0.005594680
7
        SHC1 0.005577900
8
        CRKL 0.005509182
9
     KHDRBS1 0.005443541
       TYR03 0.005441887
10
Seed Nodes used:
[1] "PIK3R1"
```

Finally, we can create a network (an *igraph* object) with the top scored genes. Visualize the top results within their interaction network is always a good idea in order to prioritize genes, since we can have a global view of all the potential candidates. The results are presented in Figure 1.

```
> ## In this case we selected to induce a network with the Top 15 genes.
```

```
> TopResults_PPI <-</pre>
```

+

```
+ create.multiplexNetwork.topResults(RWR_PPI_Results,PPI_MultiplexObject,
```

k=15)

```
> par(mar=c(0.1,0.1,0.1,0.1))
```

```
> plot(TopResults_PPI, vertex.label.color="black",vertex.frame.color="#ffffff",
```

```
+ vertex.size= 20, edge.curved=.2,
```

+ vertex.color = ifelse(igraph::V(TopResults\_PPI)\$name == "PIK3R1","yellow",

#### Random Walk with Restart on Multiplex and Heterogeneous Network



Figure 1: RWR on a monoplex PPI Network

Network representation of the top 15 ranked genes when the RWR algorithm is executed using the *PIK3R1* gene as seed (yellow node). Blue edges represent PPI interactions.

### 3.2 Random Walk with Restart on a Heterogeneous Network

A RWR on a heterogeneous (RWR-H) biological network was described by [5]. They connected a PPI network with a disease-disease similarity network using known gene-disease associations. In this case, genes and/or diseases can be used as seed nodes for the algorithm. In the following example, we also use a heterogeneous network integrating a PPI and a disease-disease similarity network. However, the procedure to obtain these networks is different to the one proposed in [5], and the details are described in our article [3].

To generate a PPI-disease heterogeneous network object, we load the disease-disease network, and combine it with our previously defined *Multiplex* object containing the PPI network, thanks to the gene-diseases associations obtained from OMIM [6]. A *MultiplexHet* object will be created, even if we are dealing with a monoplex-heterogeneous network.

- > data(Disease\_Network) # We load our disease Network
- > ## We load a data frame containing the gene-disease associations.
- > ## See ?create.multiplexHet for details about its format
- > data(GeneDiseaseRelations)
- > ## We keep gene-diseases associations where genes are present in the PPI
- > ## network
- > GeneDiseaseRelations\_PPI <-</pre>
- + GeneDiseaseRelations[which(GeneDiseaseRelations\$hgnc\_symbol %in%
- + PPI\_MultiplexObject\$Pool\_of\_Nodes),]
- > ## We create the MultiplexHet object.
- > PPI\_Disease\_Net <- create.multiplexHet(PPI\_MultiplexObject,

```
Disease_Network, GeneDiseaseRelations_PPI, c("Disease"))
> ## The results look like that
> PPI_Disease_Net
Number of Layers:
[1] 1
Number of Nodes Multiplex:
[1] 4317
IGRAPH 1573a63 UN-- 4317 18062 --
+ attr: name (v/c), type (e/c)
+ edges from 1573a63 (vertex names):
 [1] AAMP
           - - VPS52
                        AAMP
                               --BHLHE40
                                           AAMP
                                                  --GABARAPL2 AAMP
                                                                     --MAP1LC3B
 [5] VPS52 --TXN2
                        VPS52 --DDX6
                                           VPS52 --MFAP1
                                                              VPS52 -- PRKAA1
 [9] VPS52 --LMO4
                        VPS52 --STX11
                                           VPS52 --KANK2
                                                              VPS52 -- PPP1R18
[13] VPS52 --TXLNA
                        VPS52 --KIAA1217 VPS52 --VPS28
                                                              VPS52 --ATP6V1D
[17] VPS52 --TPM3
                        VPS52 --KIF5B
                                           VPS52 --N0P2
                                                              VPS52 -- RNF41
[21] VPS52 --WTAP
                        VPS52 --MAPK3
                                           VPS52 -- ZMAT2
                                                              VPS52 --VPS51
[25] BHLHE40--AES
                        BHLHE40 - - PRKAA1
                                           BHLHE40 - - CCNK
                                                              BHLHE40 - - RBPMS
[29] BHLHE40--COX5B
                        BHLHE40--UBE2I
                                           BHLHE40--MAGED1
                                                              BHLHE40 - - PLEKHB2
+ ... omitted several edges
Number of Nodes of the second network:
[1] 6947
Second Network
IGRAPH 3eed1b5 UN-- 6947 28246 --
+ attr: name (v/c), type (e/c)
+ edges from 3eed1b5 (vertex names):
 [1] 100050--122470 100050--227330 100050--259200 100050--305400 100050--601803
 [6] 100070--105800 100070--105805 100070--107550 100070--120000 100070--130090
[11] 100070--132900 100070--154750 100070--180300 100070--192310 100070--208060
[16] 100070--210050 100070--219100 100070--252350 100070--277175 100070--300537
[21] 100070--309520 100070--600459 100070--604308 100070--606519 100070--608967
[26] 100070--609192 100070--610168 100070--610380 100070--611788 100070--613780
[31] 100070--613834 100070--614042 100070--614437 100070--614980 100070--610655
[36] 100070--615436 100070--616166 100100--192350 100100--236700 100100--236730
+ ... omitted several edges
```

To apply the RWR-H on a heterogeneous network, we need to compute a matrix that accounts for all the possible transitions of the RWR particle within that network [5].

> PPIHetTranMatrix <- compute.transition.matrix(PPI\_Disease\_Net)</pre>

Before running RWR-H on this PPI-disease heterogeneous network, we need to define the seed(s). As in the previous paragraph, we take PIK3R1 as a seed gene. In addition, we can now set the SHORT syndrome itself as a seed disease.

```
> SeedDisease <- c("269880")
> ## We launch the algorithm with the default parameters (See details on manual)
> RWRH_PPI_Disease_Results <-</pre>
```

Random.Walk.Restart.MultiplexHet(PPIHetTranMatrix, + PPI\_Disease\_Net,SeedGene,SeedDisease) + > # We display the results > RWRH\_PPI\_Disease\_Results Top 10 ranked Multiplex nodes: NodeNames Score 1479 GRB2 0.001965500 1081 EGFR 0.001754048 797 CRK 0.001636329 CDC42 0.001630575 603 19 ABL1 0.001623304 798 CRKL 0.001605543 1360 FYN 0.001598567 3405 SHC1 0.001597720 2583 PDGFRB 0.001596830 4027 TYR03 0.001589911 Multiplex Seed Nodes used: [1] "PIK3R1" Top 10 ranked Second Network Nodes: SecondNet\_node Score 6352 615214 0.020817435 6705 616005 0.020785895 1699 194050 0.005868407 3625 309000 0.005687206 2901 262500 0.005681162 138920 0.005655559 686 2150 223370 0.005655117 4770 608612 0.005649291 4464 606176 0.005641777 1411 180500 0.005639919 Second Network Seed Nodes used:

[1] "269880"

Finally, we can create a heterogeneous network (an *igraph* object) with the top scored genes and the top scored diseases. The results are presented in Figure 2.

> ## In this case we select to induce a network with the Top 10 genes

```
> ## and the Top 10 diseases.
```

- > TopResults\_PPI\_Disease <-</pre>
- + create.multiplexHetNetwork.topResults(RWRH\_PPI\_Disease\_Results,
- + PPI\_Disease\_Net, GeneDiseaseRelations\_PPI, k=10)

> par(mar=c(0.1,0.1,0.1,0.1))

- > plot(TopResults\_PPI\_Disease, vertex.label.color="black",
- + vertex.frame.color="#ffffff",
- + vertex.size= 20, edge.curved=.2,
- + vertex.color = ifelse(V(TopResults\_PPI\_Disease)\$name == "PIK3R1"
- + | V(TopResults\_PPI\_Disease)\$name == "269880","yellow",
- + ifelse(V(TopResults\_PPI\_Disease)\$name %in% PPI\_Disease\_Net\$Pool\_of\_Nodes,

```
608612

608612

606176

223370

180500

180500

138920

138920

138920

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```

**Figure 2:** RWR-H on a heterogeneous PPI-Disease Network Network representation of the top 10 ranked genes and the top 10 ranked diseases when the RWR-H algorithm is executed using the *PIK3R1* gene and the SHORT syndrome disease (MIM code: 269880) as seeds (yellow nodes). Circular nodes represent genes and rectangular nodes show diseases. Blue edges are PPI interactions and black edges are similarity links between diseases. Dashed edges are the bipartite gene-disease associations.

### 3.3 Random Walk with Restart on a Multiplex Network

Some limitations can arise when single networks are used to represent and describe systems whose entities can interact through more than one type of connections [2]. This is the case of social interactions, transportation networks or biological systems, among others. The Multiplex framework provides an appealing approach to describe these systems, since they are able to integrate this diversity of data while keeping track of the original features and topologies of the different sources.

Consequently, algorithms able to exploit the information stored on multiplex networks should improve the results provided by methods operating on single networks. In this context, we extended the random walk with restart algorithm to multiplex networks (RWR-M) [3].

In the following example, we create a multiplex network integrated by our PPI network and a network derived from pathway databases [3].

```
> data(Pathway_Network) # We load the Pathway Network
> ## We create a 2-layers Multiplex object
> PPI_PATH_Multiplex <- create.multiplex(PPI_Network,Pathway_Network,
                         Layers_Name=c("PPI","PATH"))
+
> PPI_PATH_Multiplex
Number of Layers:
[1] 2
Number of Nodes:
[1] 4899
IGRAPH ed22f9b UN-- 4899 18062 --
+ attr: name (v/c), type (e/c)
+ edges from ed22f9b (vertex names):
           --VPS52
 [1] AAMP
                        AAMP
                               --BHLHE40
                                           AAMP
                                                  --GABARAPL2 AAMP
                                                                     --MAP1LC3B
 [5] VPS52 --TXN2
                        VPS52 --DDX6
                                           VPS52 --MFAP1
                                                              VPS52 -- PRKAA1
 [9] VPS52 --LMO4
                        VPS52 --STX11
                                           VPS52 --KANK2
                                                              VPS52 -- PPP1R18
[13] VPS52 --TXLNA
                        VPS52 --KIAA1217 VPS52 --VPS28
                                                              VPS52 --ATP6V1D
[17] VPS52 --TPM3
                        VPS52 --KIF5B
                                           VPS52 --N0P2
                                                              VPS52 -- RNF41
[21] VPS52 --WTAP
                        VPS52 --MAPK3
                                           VPS52 --ZMAT2
                                                              VPS52 --VPS51
[25] BHLHE40--AES
                        BHLHE40 - - PRKAA1
                                           BHLHE40 - - CCNK
                                                              BHLHE40 - - RBPMS
[29] BHLHE40--C0X5B
                        BHLHE40 - - UBE2I
                                           BHLHE40 - - MAGED1
                                                              BHLHE40 - - PLEKHB2
+ ... omitted several edges
IGRAPH 71919ca UN-- 4899 62602 --
+ attr: name (v/c), type (e/c)
+ edges from 71919ca (vertex names):
 [1] BANF1--PSIP1
                   BANF1--HMGA1
                                  BANF1--PPP2R1A BANF1--PPP2CA
                                                                 BANF1--KPNA1
 [6] BANF1--TPR
                   BANF1--NUP62
                                   BANF1--NUP153 BANF1--RANBP2
                                                                 BANF1 - - NUP54
[11] BANF1--POM121 BANF1--NUP85
                                   BANF1--PPP2R2A BANF1--EMD
                                                                 BANF1--LEMD2
[16] BANF1--ANKLE2 PSIP1--HMGA1
                                   PSIP1--KPNA1 PSIP1--TPR
                                                                 PSIP1--NUP62
[21] PSIP1--NUP153 PSIP1--RANBP2 PSIP1--NUP54
                                                  PSIP1--POM121
                                                                 PSIP1--NUP85
[26] HMGA1--TP53
                   HMGA1--KPNA1
                                   HMGA1--TPR
                                                  HMGA1--NUP62
                                                                 HMGA1 - - NUP153
                                                                 HMGA1--MYC
[31] HMGA1--RANBP2 HMGA1--NUP54
                                   HMGA1--POM121 HMGA1--NUP85
                   HMGA1--MAX
                                   HMGA1--RPS6KB1 XRCC6--XRCC5
                                                                 XRCC6--IRF3
[36] HMGA1--RB1
+ ... omitted several edges
```

Afterwards, as in the monoplex case, we have to compute and normalize the adjacency matrix of the multiplex network.

> AdjMatrix\_PPI\_PATH <- compute.adjacency.matrix(PPI\_PATH\_Multiplex)</pre>

> AdjMatrixNorm\_PPI\_PATH <- normalize.multiplex.adjacency(AdjMatrix\_PPI\_PATH)

Then, we set again as seed the *PIK3R1* gene and we perform RWR-M on this new multiplex network.

> ## We launch the algorithm with the default parameters (See details on manual)

- > RWR\_PPI\_PATH\_Results <- Random.Walk.Restart.Multiplex(AdjMatrixNorm\_PPI\_PATH,
  - PPI\_PATH\_Multiplex,SeedGene)
- > # We display the results

```
> RWR_PPI_PATH_Results
```

Тор	10	ranke	d Nodes:			
NodeNames Scor						
1		GRB2	0.001662893			
2		FYN	0.001517786			
3		HCST	0.001506594			
4		EGFR	0.001494459			
5		SHC1	0.001492456			
6		PTK2	0.001430282			
7		JAK2	0.001401867			
8		HRAS	0.001396854			
9		CRKL	0.001389391			
10	F	PDGFRB	0.001369533			
Seed	d No	odes u	sed:			

[1] "PIK3R1"

Finally, we can create a multiplex network (an *igraph* object) with the top scored genes. The results are presented in Figure 3.

```
> ## In this case we select to induce a multiplex network with the Top 15 genes.
> TopResults_PPI_PATH <-</pre>
+
      create.multiplexNetwork.topResults(RWR_PPI_PATH_Results,
                                           PPI_PATH_Multiplex, k=15)
+
> par(mar=c(0.1,0.1,0.1,0.1))
> plot(TopResults_PPI_PATH, vertex.label.color="black",
      vertex.frame.color="#ffffff", vertex.size= 20,
+
      edge.curved= ifelse(E(TopResults_PPI_PATH)$type == "PPI",
+
                       0.4,0),
+
+
      vertex.color = ifelse(igraph::V(TopResults_PPI_PATH)$name == "PIK3R1",
                       "yellow", "#00CCFF"), edge.width=0.8,
+
      edge.color=ifelse(E(TopResults_PPI_PATH)$type == "PPI",
+
                         "blue","red"))
```

## 3.4 Random Walk with Restart on a Multiplex-Heterogeneous Network

RWR-H and RWR-M remarkably improve the results obtained by classical RWR on monoplex networks, as we demonstrated in the particular case of retrieving known gene-disease associations [3]. Therefore, an algorithm able to execute a random walk with restart on both, multiplex and heterogeneous networks, is expected to achieve an even better performance. We extended our RWR-M approach to heterogeneous networks, defining a random walk with restart on multiplex-heterogeneous networks (RWR-MH) [3].

Let us integrate all the networks described previously (PPI, Pathways and disease-disease similarity) into a multiplex and heterogeneous network. To do so, we connect genes in both multiplex layers (PPI and Pathways) to the disease network, if a bipartite gene-disease relation exists.



Figure 3: RWR-M on a multiplex PPI-Pathway Network Network representation of the top 15 ranked genes when the RWR-M algorithm is executed using the

> ## We keep gene-diseases associations where genes are present in the PPI

*PIK3R1* gene (yellow node). Blue curved edges are PPI interactions and red straight edges are Pathways links. All the interactions are aggregated into a monoplex network only for visualization purposes.

```
> ## or in the pathway network
> GeneDiseaseRelations_PPI_PATH <-</pre>
      GeneDiseaseRelations[which(GeneDiseaseRelations$hqnc_symbol %in%
+
      PPI_PATH_Multiplex$Pool_of_Nodes),]
+
> ## We create the MultiplexHet object.
> PPI_PATH_Disease_Net <- create.multiplexHet(PPI_PATH_Multiplex,
      Disease_Network, GeneDiseaseRelations_PPI_PATH, c("Disease"))
+
> ## The results look like that
> PPI_PATH_Disease_Net
Number of Layers:
[1] 2
Number of Nodes Multiplex:
[1] 4899
IGRAPH ed22f9b UN-- 4899 18062 --
+ attr: name (v/c), type (e/c)
+ edges from ed22f9b (vertex names):
 [1] AAMP
            --VPS52
                        AAMP
                                --BHLHE40
                                            AAMP
                                                   --GABARAPL2 AAMP
                                                                       --MAP1LC3B
 [5] VPS52 --TXN2
                        VPS52 --DDX6
                                            VPS52 --MFAP1
                                                               VPS52
                                                                      --PRKAA1
 [9] VPS52 --LMO4
                        VPS52 --STX11
                                            VPS52 --KANK2
                                                               VPS52 -- PPP1R18
[13] VPS52
           --TXLNA
                        VPS52 --KIAA1217
                                            VPS52 --VPS28
                                                               VPS52
                                                                      --ATP6V1D
[17] VPS52 --TPM3
                        VPS52 --KIF5B
                                            VPS52 --N0P2
                                                               VPS52 -- RNF41
[21] VPS52 --WTAP
                        VPS52 --MAPK3
                                            VPS52 --ZMAT2
                                                               VPS52 --VPS51
[25] BHLHE40--AES
                        BHLHE40 - - PRKAA1
                                            BHLHE40 - - CCNK
                                                               BHLHE40 - - RBPMS
[29] BHLHE40--C0X5B
                        BHLHE40--UBE2I
                                            BHLHE40--MAGED1
                                                               BHLHE40 - - PLEKHB2
```

```
+ ... omitted several edges
IGRAPH 71919ca UN-- 4899 62602 --
+ attr: name (v/c), type (e/c)
+ edges from 71919ca (vertex names):
 [1] BANF1--PSIP1 BANF1--HMGA1
                                   BANF1--PPP2R1A BANF1--PPP2CA
                                                                BANF1--KPNA1
 [6] BANF1--TPR
                   BANF1--NUP62
                                  BANF1--NUP153 BANF1--RANBP2
                                                                BANF1--NUP54
[11] BANF1--POM121 BANF1--NUP85
                                  BANF1--PPP2R2A BANF1--EMD
                                                                BANF1--LEMD2
                                  PSIP1--KPNA1 PSIP1--TPR
[16] BANF1--ANKLE2 PSIP1--HMGA1
                                                                PSIP1--NUP62
[21] PSIP1--NUP153 PSIP1--RANBP2 PSIP1--NUP54
                                                 PSIP1--POM121
                                                                PSIP1--NUP85
[26] HMGA1--TP53
                   HMGA1--KPNA1
                                  HMGA1--TPR
                                                  HMGA1--NUP62
                                                                HMGA1 - - NUP153
[31] HMGA1--RANBP2 HMGA1--NUP54
                                  HMGA1--POM121 HMGA1--NUP85
                                                                HMGA1--MYC
                                  HMGA1--RPS6KB1 XRCC6--XRCC5
                                                                XRCC6--IRF3
[36] HMGA1--RB1
                   HMGA1--MAX
+ ... omitted several edges
Number of Nodes of the second network:
[1] 6947
Second Network
IGRAPH 04739c8 UN-- 6947 28246 --
+ attr: name (v/c), type (e/c)
+ edges from 04739c8 (vertex names):
 [1] 100050--122470 100050--227330 100050--259200 100050--305400 100050--601803
 [6] 100070--105800 100070--105805 100070--107550 100070--120000 100070--130090
[11] 100070--132900 100070--154750 100070--180300 100070--192310 100070--208060
[16] 100070--210050 100070--219100 100070--252350 100070--277175 100070--300537
[21] 100070--309520 100070--600459 100070--604308 100070--606519 100070--608967
[26] 100070--609192 100070--610168 100070--610380 100070--611788 100070--613780
[31] 100070--613834 100070--614042 100070--614437 100070--614980 100070--610655
[36] 100070--615436 100070--616166 100100--192350 100100--236700 100100--236730
+ ... omitted several edges
```

To apply the RWR-MH on a multiplex and heterogeneous network, we need to compute a matrix that accounts for all the possible transitions of the RWR particle within this network [3].

> PPI\_PATH\_HetTranMatrix <- compute.transition.matrix(PPI\_PATH\_Disease\_Net)

As in the RWR-H situation, we can take as seeds both, the *PIK3R1* gene and the the SHORT syndrome disease.

> ## We launch the algorithm with the default parameters (See details on manual)
> RWRH\_PPI\_PATH\_Disease\_Results <-</pre>

+ Random.Walk.Restart.MultiplexHet(PPI\_PATH\_HetTranMatrix,

+ PPI\_PATH\_Disease\_Net,SeedGene,SeedDisease)

> # We display the results

> RWRH\_PPI\_PATH\_Disease\_Results

Top 10 ranked Multiplex nodes:

NodeNames Score

1616 GHR 0.0005153889

```
4867 ZMPSTE24 0.0004781314
```

```
1706
          GRB2 0.0004642946
1625
          GJA1 0.0004457777
1781
          HCST 0.0004408101
1249
          EGFR 0.0004239882
          FYN 0.0004193109
1564
3883
          SHC1 0.0004161564
2990
       PDGFRB 0.0004079884
          PTK2 0.0004069179
3361
Multiplex Seed Nodes used:
[1] "PIK3R1"
Top 10 ranked Second Network Nodes:
     SecondNet_node
                          Score
             615214 0.020797695
6352
6705
             616005 0.020775842
            194050 0.005868202
1699
4770
             608612 0.005691321
2901
             262500 0.005691027
3625
            309000 0.005683228
686
            138920 0.005653962
2150
             223370 0.005653358
4464
             606176 0.005639004
1411
             180500 0.005631733
Second Network Seed Nodes used:
[1] "269880"
```

Finally, we can create a multiplex and heterogeneous network (an *igraph* object) with the top scored genes and the top scored diseases. The results are presented in Figure 4.

```
> ## In this case we select to induce a network with the Top 10 genes.
> ## and the Top 10 diseases.
> TopResults_PPI_PATH_Disease <-</pre>
      create.multiplexHetNetwork.topResults(RWRH_PPI_PATH_Disease_Results,
+
      PPI_PATH_Disease_Net, GeneDiseaseRelations_PPI_PATH, k=10)
+
> par(mar=c(0.1,0.1,0.1,0.1))
> plot(TopResults_PPI_PATH_Disease, vertex.label.color="black",
      vertex.frame.color="#ffffff",
+
      vertex.size= 20,
+
      edge.curved=ifelse(E(TopResults_PPI_PATH_Disease)$type == "PATH",
+
                      0, 0.3),
+
      vertex.color = ifelse(V(TopResults_PPI_PATH_Disease)$name == "PIK3R1"
+
      V(TopResults_PPI_Disease)$name == "269880","yellow",
+
      ifelse(V(TopResults_PPI_PATH_Disease)$name %in%
+
                 PPI_PATH_Disease_Net$Pool_of_Nodes,
+
+
      "#00CCFF","Grey75")),
      edge.color=ifelse(E(TopResults_PPI_PATH_Disease)$type == "PPI","blue",
+
      ifelse(E(TopResults_PPI_PATH_Disease)$type == "PATH", "red",
+
      ifelse(E(TopResults_PPI_PATH_Disease)$type == "Disease", "black", "grey50"))),
+
```

+ edge.width=0.8,

#### Random Walk with Restart on Multiplex and Heterogeneous Network

```
+ edge.lty=ifelse(E(TopResults_PPI_PATH_Disease)$type ==
```

- + "bipartiteRelations", 2,1),
- + vertex.shape= ifelse(V(TopResults\_PPI\_PATH\_Disease)\$name %in%
- + PPI\_PATH\_Disease\_Net\$Pool\_of\_Nodes,"circle","rectangle"))



Figure 4: RWR-MH on a multiplex and heterogeneous network (PPI-Pathway-Disease)

Network representation of the top 10 ranked genes and the top 10 ranked diseases when the RWR-H algorithm is executed using the *PIK3R1* gene and the SHORT syndrome disease (MIM code: 269880) as seeds (yellow nodes). Circular nodes represent genes and rectangular nodes show diseases. Blue curved edges are PPI interactions and red straight edges are Pathways links. Black edges are similarity links between diseases. Dashed edges are the bipartite gene-disease associations. Multiplex interactions are aggregated into a monoplex network only for visualization purposes.

# References

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- [5] Yongjin Li and Jagdish C. Patra. Genome-wide inferring gene-phenotype relationship by walking on the heterogeneous network. *Bioinformatics*, 26(9):1219–1224, 2010. doi:10.1093/bioinformatics/btq108.
- [6] Ada Hamosh, Alan F. Scott, Joanna S. Amberger, Carol A. Bocchini, and Victor A. McKusick. Online Mendelian Inheritance in Man (OMIM), a knowledgebase of human genes and genetic disorders. *Nucleic Acids Research*, 33(DATABASE ISS.):514–517, 2005. doi:10.1093/nar/gki033.

# A Session info

- R version 3.5.1 Patched (2018-07-12 r74967), x86\_64-pc-linux-gnu
- Locale: LC\_CTYPE=en\_US.UTF-8, LC\_NUMERIC=C, LC\_TIME=en\_US.UTF-8, LC\_COLLATE=C, LC\_MONETARY=en\_US.UTF-8, LC\_MESSAGES=en\_US.UTF-8, LC\_PAPER=en\_US.UTF-8, LC\_NAME=C, LC\_ADDRESS=C, LC\_TELEPHONE=C, LC\_MEASUREMENT=en\_US.UTF-8, LC\_IDENTIFICATION=C
- Running under: Ubuntu 16.04.5 LTS
- Matrix products: default
- BLAS: /home/biocbuild/bbs-3.8-bioc/R/lib/libRblas.so
- LAPACK: /home/biocbuild/bbs-3.8-bioc/R/lib/libRlapack.so
- Base packages: base, datasets, grDevices, graphics, methods, stats, utils
- Other packages: RandomWalkRestartMH 1.2.0, igraph 1.2.2
- Loaded via a namespace (and not attached): BiocGenerics 0.28.0, BiocManager 1.30.3, BiocStyle 2.10.0, MASS 7.3-51, Matrix 1.2-14, Rcpp 0.12.19, Rgraphviz 2.26.0, ape 5.2, backports 1.1.2, compiler 3.5.1, digest 0.6.18, dnet 1.1.4, evaluate 0.12, graph 1.60.0, grid 3.5.1, hexbin 1.27.2, htmltools 0.3.6, knitr 1.20, lattice 0.20-35, magrittr 1.5, nlme 3.1-137, parallel 3.5.1, pkgconfig 2.0.2, rmarkdown 1.10, rprojroot 1.3-2, stats4 3.5.1, supraHex 1.20.0, tools 3.5.1, yaml 2.2.0