# Package 'NetFrac'

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Type Package
Title Netfrac - Community distance calculation within sequence similarity networks
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Description The NetFrac package allows the users to calculate the eight following network distances, NetUniFrac, Motifs, Spp, Spep, Spelp, Spinp, and the Transfer distance (two variants), between species communities present in a given sequence similarity network. The shortest path in a phylogenetic tree is traditionally used to estimate evolutionary distances between species. Moreover, the shortest path concept can be extended to sequence similarity networks. The five following distances (NetUniFrac, Spp, Spep, Spelp, and Spinp) can be seen as network analogues of the traditional UniFrac distance used to assess distances between species communities in a phylogenetic tree, whereas the Transfer distance is designed for estimating the rate and direction of gene transfers, or species dispersal, between different phylogenetic, or ecological, species communities.
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dist\_paths

Shortest path distances

## Description

Takes as input the network as well as the species communities to be compared and gives as output the distance matrix between species communities. It is better to use the NetFrac function if the global distances are needed.

## Usage

```
dist_paths(x, col1, col2, distance = "paths", paths = "single", info = NULL, type = "graph", maxcores
```

#### **Arguments**

X	The igraph object to be analyzed
col1	Community one
col2	Community two
distance	Distances that are implemented in the NetFrac. See also the NetFrac function
paths	Take into account all possible shortest paths or only one
info	Information about the tree if the input is a tree
type	Tree or network
maxcores	Number of parallel cores to be used for calculation (one by default)

#### See Also

NetFrac

multicore multicore

#### Description

Uses parallel cores for bigger networks.

#### Usage

```
multicore(nc = 1)
```

## Arguments

nc

number of cores

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## Description

Title

#### Usage

```
NetFrac(x, distances = "UniFrac", paths = "single", mats = "",
  maxcores = 1, share_weight = 0)
```

#### **Arguments**

X	The network(or tree) that will be analyzed. The network must be in the igraph format, the edges being accessible with $E(x)$ and vertices with $V(x)$ . The communities are under $V(x)$ stax and the branch weights under $E(x)$ sweight. These functions are applicable with igraph objects only
distances	Distance(s) to calculate: Transfer, Transfer2, Spaths, UniFrac and Motifs. The transfer, paths and Unifrac distances can also be calculated for trees
paths	this parameter is used to decide whether all the shortest paths between network nodes should be calculated ("all"), or only one of them ("single"). The last option can significantly reduce the time of computation. For Motifs distance, decide the size of the motifs (i.e. 2 or 3)
mats	the similarity matrix used to reconnect the network. See also reconnect
maxcores	uses the multicore function to set up the number of cores to be used
share_weight	weight if there are mixed communities. For now, only 0 is implemented

## Value

```
$'Spp' A B A 0.0000000 0.5555556 B 0.5555556 0.0000000
$Spep A B A 0.0000000 0.6296296 B 0.6296296 0.0000000
$Spelp A B A 0.0000000 0.6296296 B 0.6296296 0.0000000
$Spinp A B A 0.0000000 0.9907407 B 0.9907407 0.0000000
$Transfer A B A 0.0000000 0.6666667 B 0.3333333 0.0000000
```

## **Examples**

```
NetFrac(net_a)
NetFrac(net_a, "Spaths", "all")
```

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NetUnifrac

NetUnifrac

#### Description

Calculates the network version of the UniFrac distance.

## Usage

```
NetUnifrac(igraph, tax1 = "", tax2 = "", weight = TRUE)
```

#### **Arguments**

igraph The igraph objecttax1 The first communitytax2 The second communityweight Use weights of the edges if existing

## **Examples**

```
NetUnifrac(net_a, "A", "B")
```

reconnect

Connect the network using a similarity matrix

## Description

Creates a fully connected network using a similarity matrix between nodes.

## Usage

```
reconnect(graph, matrice = "")
```

## Arguments

graph The disconnected igraph object

matrice The similarity matrix containing pairwise distances between the nodes

#### See Also

reconnect\_btw

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Connect the network a graph using betweenness centrality

#### **Description**

Creates a fully connected network using the betweenness centrality principle.

#### Usage

```
reconnect_btw(graph)
```

#### **Arguments**

graph

The disconnected igraph object

#### See Also

reconnect

set\_color

Set colors for the communities in the network

#### **Description**

Visualizes the network by adding different colors to different species communities, when plotting either with R own color list or a provided color list.

## Usage

```
set_color(graph, colors_list = "")
```

#### **Arguments**

graph The igraph object used to define colors. The nodes accessed with V() should

have a \$tax attribute which links them to a community, so a different color can

be assigned to each community.

colors\_list Optional; the colors can be defined in a list with the community names defined

with names(). Otherwise, the function will return the numbers already associ-

ated to R colors.

#### **Examples**

```
foo_CAT_env <- c("red","green","blue","grey")
names(foo_CAT_env) = c("host","ubiquitous","water","unknown")
CAT_env <- set_color(CAT_env, foo_CAT_env)
plot(net_a)</pre>
```

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subgroup\_graph

Create a subgraph based on chosen species communities

#### **Description**

Creates another igraph object containing only the nodes from some chosen species communities (two or more), and the edges connecting them.

#### Usage

```
subgroup_graph(graph, groups)
```

#### **Arguments**

graph

The igraph object

groups

The communities that should be isolated, indicated as a list of two items

#### **Examples**

```
CAT_env1 <- set_color(subgroup_graph(CAT_env, c("host", "ubiquitous")))
plot(CAT_env1)</pre>
```

Transfer

Algorithm for a fast calculation of the Transfer distance

#### **Description**

The transfer distance can be used to calculate the proportion of species/sequences of community X that have been affected by horizontal gene transfers from species/sequences of community Y. The Transfer distance is assymetric. If the threshold alpha (0<alpha<1) is satisfied, the distance is incremented. The operation is repeated for every node of the network. The result is a pairwise matrix reporting the Transfer distance between each pair of species communities.

#### Usage

```
Transfer(graph)
```

#### **Arguments**

graph

The igraph object

#### **Examples**

```
transfer2(net_a)
```

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