Understanding Diffusion with **netdiffuseR**Simulating data Supplet 2016 INSNA

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The plan

- 1. Review some concepts
- 2. Simulating diffusion networks
- 3. netdiffuseR's core functions
- 4. Bonus track (if we have time)

Introduction

Before start, a review of concepts that we will be using here

- 1. Exposure: What proportion/number of your neighbors has adopted an innovation.
- 2. Threshold: What was the proportion/number of your neighbors had adopted by the time you adopted.
- 3. Infectiousness: How much i's adoption affects her alters
- 4. Susceptibility: How much i's alters' adoption affects her.
- 5. Structural equivalence: How similar are *i* and *j* in terms of position on the network.

We will simulate a diffusion network with the following parameters:

- 1. Will have 1,000 vertices,
- 2. Will span 20 time periods,
- 3. The set of early adopters will be random,
- 4. Early adopters will be a 10% of the network,
- 5. The graph will be small-world,
- 6. Will use the WS algorithmwith p = .2 (probability of rewire).
- 7. Threshold levels will be uniformly distributed between [0.3, 0.7]

In netdiffuseR

To generate such diffusion network we can use the rdiffnet function included in the package:

```
# Loading the package
library(netdiffuseR)
# Setting the seed for the RNG
set.seed(1213)
# Generating a random diffusion network
net <- rdiffnet(
                 = 1e3.
                                                # 1.
 n
                 = 20.
                                                # 2.
 seed.nodes = "random",
                                                # 3.
 seed.p.adopt = .1,
 seed.graph = "small-world",
                                                # 5.
 rgraph.args = list(p=.2),
                                                # 6.
 threshold.dist = function(x) runif(1, .3, .7) # 7.
```

In **netdiffuseR** (cont. 1)

This output can be printed to see some information about this network

net

```
## Dynamic network of class -diffnet-
## # of nodes : 1000 (1, 2, 3, 4, 5, 6, 7, 8, ...)
## # of time periods : 20 (1 - 20)
## Type : directed
## Final prevalence : 0.90
## Static attributes : real_threshold (1)
## Dynamic attributes : -
```

In **netdiffuseR** (cont. 2)

Including some summary stats (we only will see times 1, 5, 10, 15 and 20).

```
summary(net, slices=c(1,5,10,15,20))
```

```
## Diffusion network summary statistics
   _____
##
  Period Adopters Cum Adopt. Cum Adopt. % Hazard Rate Density Moran's I
##
     1
          100
                 100
                         0.10
                                      0.00
                                            -0.00
           34
                  318
                         0.32 0.05
                                      0.00 0.01
##
                        0.56 0.11
##
   10
         56
                 558
                                      0.00 0.02
##
   15 39
                 784
                      0.78 0.15 0.00 0.01
##
     20
         17
                 899
                       0.90 0.14 0.00 0.02
```

Left censoring : 0.10 (100) ## Right centoring : 0.10 (101)

of nodes : 1000

In netdiffuseR changing the first adopters

Using different parameters for seed.nodes (who are the first adopters), we get different [theoretical] resuls:

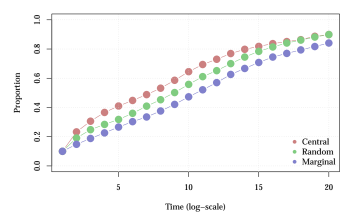


Figure 1: Cumulative Adopters for Different seed.nodes

Testing theoretical results

▶ Given dynamic graphs of size n=150, with density $\rho \sim 0.013$ and t=5 time periods. What should we expect from the following combinations:

```
(Bernoulli, Scale-free, Small-world) × (Marginal, Central, Random)
```

In which of these the diffusion process is fastest?

Lets run some monte carlo simulations to find out!

Testing theoretical results (cont.)

Recall that densities for these graphs can be computed as follow:

► Bernoulli

$$\rho = p$$

► Small-world

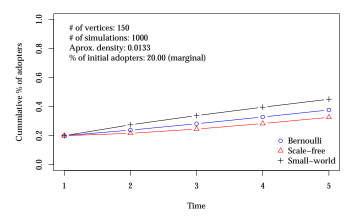
$$\rho = \frac{n \times k}{n \times (n-1)} = \frac{k}{n-1}$$

Scale-free

$$\rho = \frac{m \times t}{(m_0 + t)(m_0 + t - 1)}$$

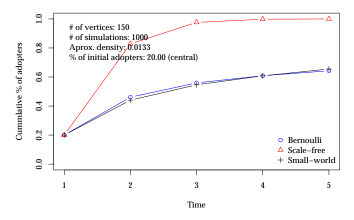
We use this to generate graphs with similar densities (see on the code).

Theoretically, what should we expect from... marginal



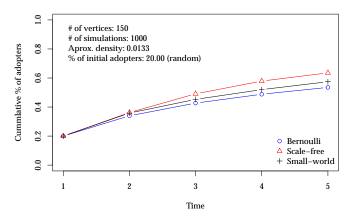
Small-worlds! In *better* connected networks, marginal nodes are closer to the entire graph.

Theoretically, what should we expect from... central



Scale-free! Central nodes have higher degree in these networks.

Theoretically, what should we expect from... random



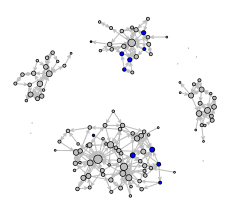
Any clues here?

- ▶ Now we will review some of the package's main functions.
- Among netdiffuseR features, we find three classical Diffusion Network Datasets:
 - brfarmersDiffNet Brazilian farmers and the innovation of Hybrid Corn Seed (1966).
 - medInnovationsDiffNet Doctors and the innovation of Tetracycline (1955).
 - kfamilyDiffNet Korean women and Family Planning methods (1973).
- ▶ For the review we will use the Medical Innovations dataset.

Medical Innovations

```
data("medInnovationsDiffNet")
plot(medInnovationsDiffNet)
```

Diffusion network in time 1



Structural equivalence

 \triangleright Structural equivalence between vertices (i, j) is defined as

$$SE_{ij} = \frac{(dmax_i - d_{ji})^v}{\sum_{k \neq i}^n (dmax_i - d_{ki})^v}$$

with the summation over $k \neq i$, and d_{ji} , eucledian distance in terms of geodesics, is defined as

$$d_{ji} = \left[(z_{ji} - z_{ij})^2 + \sum_{k}^{n} (z_{jk} - z_{ik})^2 + \sum_{k}^{n} (z_{ki} - z_{kj})^2 \right]^{\frac{1}{2}}$$

with z_{ij} as the geodesic (shortest path) from i to j, and $dmax_i$ equal to largest Euclidean distance between i and any other vertex in the network. All summations are made over $k \notin \{i, j\}$.

▶ This is a disimilarity measure! So as $SE_{ij} \rightarrow \infty$ it means that i and j are more *structurally* different.

Structural equivalence (cont.)

```
# Computing and printing
se <- struct equiv(medInnovationsDiffNet, groupvar="city")
se
## Structural equivalence for a dynamic graph
## # nodes : 125
## # of slices: 18
## Access elements via [[nslice]]$ (as a nested list). Available elements are:
           : Structural equivalence matrix (n x n)
          : Euclidean distances matrix (n x n)
   - gdist : Structural equivalence matrix (n x n)
# Taking a look at the first elements at time 1
se[[1]]$SE[1:5, 1:5]
## 5 x 5 sparse Matrix of class "dgCMatrix"
               1001
                          1002
                                                1004
                                                            1005
                                     1003
## 1001
                   0.00541279 0.02278506 0.00482712 0.003376139
## 1002 0,006802655 .
                              0.01433493 0.02272156 0.019716520
## 1003 0.013127450 0.01220834 .
                                          0.01234462 0.010268728
## 1004 0.006439334 0.02282702 0.01438956 .
                                                     0.018054677
## 1005 0.007146494 0.02248588 0.01567656 0.02059242 .
```

Exposure

The exposure for the vertex $i \in V$ can be computed as

$$e_{i,t} = \frac{\sum_{j \neq i} s_{i,j} \times x_j \times a_j}{\sum_{j \neq i} s_{i,j} \times x_j}$$

Where $s_{i,j} \in S_t$ is the ij-th element of the adjacency matrix, $x_j \in \mathbf{x}_t$ is j's attribute in time t, and $a_j \in A_t$ is a dichotomous scalar equal to 1 if j has adopted the innovation in time t. In matrix notation would be:

$$E_t = \left(S_t \times \left[\mathbf{x}_t \circ A_t\right]\right) / \left(S_t \times \mathbf{x}_t\right)$$

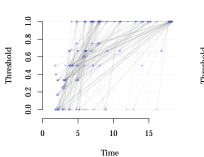
Exposure (cont. 1)

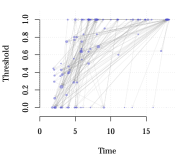
```
# Computing
e1 <- exposure(medInnovationsDiffNet)
e2 <- exposure(medInnovationsDiffNet, attrs="proage")

oldpar <- par(no.readonly = TRUE)
par(mfrowre(1,2))
plot_threshold(medInnovationsDiffNet, expo=e1, main="Plain exposure")
plot_threshold(medInnovationsDiffNet, expo=e2, main="Attribute weighted exposure")
```

Plain exposure

Attribute weighted exposure





Infectiousness and Susceptibility

Susceptibility of $i \in V$

$$S_{i} = \frac{\sum_{k=1}^{K} \sum_{j=1}^{n} x_{ij(t-k+1)} z_{j(t-k)} \times \frac{1}{w_{k}}}{\sum_{k=1}^{K} \sum_{j=1}^{n} x_{ij(t-k+1)} z_{j(1 \le t \le t-k)} \times \frac{1}{w_{k}}} \quad \text{for } i, j = 1, \dots, n \quad i \ne j$$

Infectiousness of $i \in V$

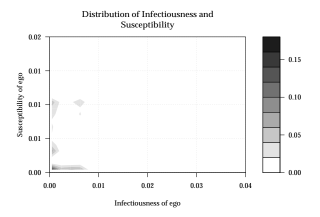
$$I_{i} = \frac{\sum_{k=1}^{K} \sum_{j=1}^{n} X_{ji(t+k-1)} Z_{j(t+k)} \times \frac{1}{w_{k}}}{\sum_{k=1}^{K} \sum_{j=1}^{n} X_{ji(t+k-1)} Z_{j(t+k \le t \le T)} \times \frac{1}{w_{k}}} \quad \text{for } i, j = 1, \dots, n \quad i \ne j$$

Normalized

$$S'_{i} = \frac{S_{i}}{\sum_{k=1}^{K} \sum_{j=1}^{n} z_{j(t-k)} \times \frac{1}{w_{k}}} \qquad I'_{i} = \frac{I_{i}}{\sum_{k=1}^{K} \sum_{j=1}^{n} z_{j(t-k)} \times \frac{1}{w_{k}}}$$

Infectiousness and Susceptibility (cont. 1)

```
# Computing and plotting both with K=5
is <- plot_infectsuscep(net, K=5L, logscale = FALSE)</pre>
```



Infectiousness and Susceptibility (cont. 2)

```
# Is there any thing?
t.test(is$infect, alternative = "greater", na.rm=TRUE)
   One Sample t-test
## data: is$infect
## t = 20.457, df = 898, p-value < 2.2e-16
## alternative hypothesis: true mean is greater than 0
## 95 percent confidence interval:
## 0.005340058
                        Tnf
## sample estimates:
## mean of x
## 0.00580749
t.test(is$suscept, alternative = "greater", na.rm=TRUE)
   One Sample t-test
## data: is$suscept
## t = 29.319, df = 898, p-value < 2.2e-16
## alternative hypothesis: true mean is greater than 0
## 95 percent confidence interval:
## 0.003700033
                        Tnf
## sample estimates:
## mean of x
## 0.00392019
```

- A novel statistical method (work-in-progress) that allows conducting inference.
- ► Included in the package, tests whether a particular network statistic actually depends on network structure
- ► Suitable to be applied to network thresholds (you can't use thresholds in regression-like models!)

Idea

Let $\mathcal{G} = (V, E)$ be a graph, γ a vertex attribute, and $\beta = f(\gamma, \mathcal{G})$, then

$$\gamma \perp \mathcal{G} \implies \mathbb{E} \left[\beta(\gamma, \mathcal{G}) | \mathcal{G} \right] = \mathbb{E} \left[\beta(\gamma, \mathcal{G}) \right]$$

- ▶ This is, if for example time of adoption is independent on the structure of the network, then the average threshold level will be independent from the network structure as well.
- Another way of looking at this is that the test will allow us to see how probable is to have this combination of network structure and network threshold (if it is uncommon then we say that the diffusion model is highly likely)

Structural dependence test: Not random TOA

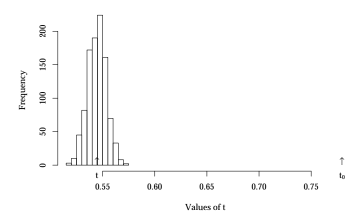
- ► To use this test, **netdiffuseR** has the struct_test function.
- Basically it simulates networks with the same density and computes a particular statistic every time, generating an EDF (Empirical Distribution Function) under the Null hyphothesis (p-values).

```
# Running the test
test <- struct_test(
  graph = net,
  statistic = function(x) mean(threshold(x), na.rm = TRUE),
  R = 1e3
)

# See the output
test</pre>
```

Structural dependence test: Not random TOA (cont. 2)

Empirical Distribution of Statistic



Structural dependence test: Random TOA (cont. 3)

```
# Resetting TOAs (now will be completely random)
diffnet.toa(net) <- sample(c(NA, 1:20), nnodes(net), TRUE)

# Running the test
test <- struct_test(
    graph = net,
    statistic = function(x) mean(threshold(x), na.rm = TRUE),
    R = 1e3
    )

# See the output
test</pre>
```

Structural dependence test: Random TOA (cont. 4)

