

# Lecture with Computer Exercises: Modelling and Simulating Social Systems with MATLAB

Project Report

# Network Based Modelling for the Spread of Scientific Ideas

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Network-Based Modelling for the Spread of Scientific Ideas

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# 1 Abstract

Using simulations, we investigated the changes in networks with nodes that held different 'scientific ideas' and influenced each other through a complex contagion mechanism. This was divided into two parts: by varying the network structure and observing features of the distribution of ideas in networks, and by varying the starting idea distributions of the nodes and observing features of the network structures. The update step included a rewiring probability, a complex contagion threshold, and a probability of innovation (producing a new idea). We found that both structures and idea distributions influenced each other's features. Values of the update parameter also played a role.

# 2 Individual contributions

This report represents a group effort by all members.

# 3 Introduction and Motivation

We live in a time in which aspects of our lives and of our world become more and more connected with each other. To understand one aspect, we must understand many other aspects, which all together form a large, complex system, or network. Globalization has changed the meaning of 'distance' and communication, allowing seemingly unrelated and unconnected individuals to share more than they ever could before. Fields of studies are overlapping with each other, creating new interdisciplinary domains and building a diverse playground for the sharing of ideas. But how do ideas spread? This question is especially interesting with the increase of technology that allows us to record and visualize the networks that connect individuals and ideas, and in particular the ability to 'see' how they change. This can lead to insight about why and when ideas spread and into the complexity of the matter. Research has shown that not only does the nature of information or innovation influence the diffusion of it, but also that the structure of a network influences the diffusion dynamics. Here, we try to simulate the spread of scientific ideas in different networks. The model presented is based on two studies: one that investigated critical parameter values for complex contagion (Centola and Macy, 2007) and another that investigated critical values of a rewiring parameter (Holme and Newman, 2006).

# 3.1 Fundamental Questions

The main goals of this simulation study are to investigate how network structure influences the distribution of ideas, and how the distribution of ideas influences network structure. For a list of the terminology that will be used throughout the paper, please refer to Table 1. By varying three parameters - probability of rewiring, rate of innovation, and complex contagion threshold ( $\phi$ ,  $\alpha$ , and  $\delta$  respectively) - and using different network structures and idea distributions, we observed how network structure characteristics changed. Similarly, we observed how the distribution of ideas and the connections between them changed. Below we describe our questions more specifically.

Table 1: List of terminology

Table 1. List of terminology					
	Term				
Neighborhood index	The fraction of the holders of the same idea				
	who are neighbor as well averaged over all ideas.				
Intra-idea distance	The average distance between				
	holders of the same idea in the network.				
Dominant frequency	The frequency of the dominant idea				
	in the network at each time steps of the simulation.				
Average dominance time The average number of time steps					
	in which the dominant idea keeps its dominance.				
Novelty index	The fraction of newly generated ideas.				
Average shortest path	The average number of steps				
	along the shortest paths for all possible pairs of network nodes.				
Clustering coefficient	A measure of degree to which				
	nodes in a graph tend to cluster together.				
Degree of connectivity	The number of edges incident to the vertex.				
Connected component	A subgraph in which any two vertices are				
	connected to each other by paths, and which is connected to no				
	additional vertices in the supergraph.				
Diameter of network	The longest of all the calculated shortest paths				
	in a network.				

#### 3.1.1 Effects of Network Structure on Idea Distribution

Given a starting network and a random idea distribution, how do different network structures (see Section 4.1) affect the distance between nodes that have the same idea (intra-idea distance)? How do they affect the neighbourhood index? How do they change the emergence of dominant ideas and their time of dominance? How do their effects depend on the values of  $\phi$ ,  $\alpha$  and  $\delta$ ?

More rigid network structures (those with less 'randomness', such as the caveman and the small world networks) may make it more difficult for 'like-minded' nodes (that is, nodes with the same idea) to connect and may thus have smaller neighbourhood indexes and larger intra-idea distances than the more random network structures (such as the random and scale-free networks). Their effects may be more sensitive to the values of  $\phi$  (because this affects how likely it is for their structure to change) and to values of  $\delta$  because being restricted to a more closed group of nodes makes it difficult to reach a threshold necessary to become similar to surrounding nodes. Values of  $\alpha$  may decrease the neighbourhood indexes by creating larger diversity

among neighbouring nodes.

If more rigid network structures do make it more difficult for like-minded nodes to connect, then it would be more difficult for a dominant idea to emerge in these networks. These effects may be smaller for larger values of  $\phi$  since these values would allow for the structure to change more. For larger values of  $\phi$  therefore one could expect that the effects of the network structures on the characteristics of the idea distribution are more similar since allowing to change the structure removes their initial influence.

# 3.1.2 Effects of Idea Distribution on Network Structure

Given a starting idea distribution and a caveman network structure, how do different idea distributions affect the average path length and diameter of the network? Do they change the number of connected components in the network? Do clusters form differently, and how does the clustering coefficient change? What does the distribution of node degree looks like? How do these effects depend on the values of  $\phi$ ,  $\alpha$ , and  $\delta$ ?

If the starting idea distribution is parallel to the caveman network structure (see Section 4.1.1 for a description of idea distributions), then like-minded nodes will already be connected and thus rewiring will probably not change much of the average path length, nor will it change the network diameter. Similarly, the clustering coefficient will remain high just like the starting value. The distribution of the node degree will also not change (nodes will have one of two values for their degree). In other words, if the idea distribution is parallel to the network structure, the structure will not change much. Changing  $\phi$  and  $\delta$  will not change these effects, and perhaps increasing  $\alpha$  will decrease the clustering coefficient and will increase the number of connected components because nodes will disconnect from nodes with novel ideas and will rewire to nodes with the same idea.

If the starting idea distribution is random, then the network's caves will disintegrate as nodes will rewire with other nodes outside of their caves. This will change the degree distribution by increasing its variance (because nodes will have a variety of different degree values). Depending on the value of  $\delta$  this disintegration may be reduced because nodes have a higher chance of forming dominant ideas within caves. Similarly, increasing  $\phi$  will increase the disintegration of caves. Thus, for this idea distribution the parameter values may play a larger role.

If the starting idea distribution is anti-parallel, nodes within each cave will initially be connected with nodes that do not hold the same idea as them. Therefore the threshold  $\delta$  will not be met in order for nodes to change their ideas, and they will rewire with other nodes outside of their cave. The clustering coefficient, as well as the number of connected components, the network diameter, and the average path length, will likely decrease since the structure will change significantly. The

degree distribution will increase in variance. Increasing  $\phi$  and  $\alpha$  and decreasing  $\delta$  will probably increase the magnitude of these effects. Thus, having an anti-parallel idea distribution will probably display the most changes in the characteristics of the network structure that are in question.

# 4 Description of the Model

The model used here is based on a study by Holme and Newman (2006). Each simulation begins with a specified network structure as well as a distribution of the 'idea' (or state) of the nodes. At each time step a node either changes its idea to that of one of its neighbours' ideas if its frequency surpasses a defined threshold, rewires to connect with a node that has the same idea, or generates a novel idea (this is the innovation parameter).

Given the network structure and node states, three parameters are introduced:  $\phi$  (probability of rewiring),  $\alpha$  (probability of innovation), and  $\delta$  (complex contagion threshold). As in Holme and Newman (2006),  $\phi$  is a value from zero to one, and is the probability that one of the edges of a randomly chosen node i will be changed to connect to another node j that i is unconnected with. We decided to add one more criterion to this definition: node j is a node that has the same idea as node i. This encourages the simulations to reflect a common tendency of individuals to seek out others who think like them. If there is no such node j, then the chosen node will do nothing.

At each time step a node may 'come up with a new idea' with a probability of  $\alpha$ . This value is small to reflect that novel ideas are not frequently observed.

We introduced a node threshold  $\delta$  to the general model in order to investigate the behaviour of complex contagion as opposed to simple contagion. This was motivated by a study by Centola and Macy (2007). Simple contagion is well suited for modeling the spread of diseases since they may often be passed on by a single contact with an infected individual. However, as our intuition may suggest, other kinds of innovations raise questions about the legitimacy and credibility of the innovations themselves, and may thus require exposure to multiple sources of the innovation. This is called complex contagion. Models usually represent it in two ways regarding the number of connected sources that must have adopted an innovation in order to influence the agent (or individual) in question: it is either a fixed number (greater than one) or a fraction (between zero and one, inclusive). Our simulation used the second formulation since the degree of individual nodes varied across network structures.

#### 4.1 Networks

For the purposes of our simulations, we used four network structures. \*\*\*\*\*NET-WORK STRUCTURE FIGURE\*\*\*\* illustrates them. These structures can be characterized by properties such as average shortest path lengths, clustering coefficients, and the degree of connectivity (see Table 1 for definitions). Below are short descriptions of each network structure. In order to compare between different structures, the network structure parameters were chosen such that the mean degree

of the networks were similar (approximately 25). For further details about parameter values, see Table 2.

Table 2: Table of parameters used in simulations.

Network	Parameter	Value		
	$\phi$	0.1	0.3	0.5
	$\alpha$	0.01	0.05	0.1
	$\delta$	0.001	0.01	0.05
	n	1000		
	$t_{end}$	1000		
Caveman	m	40		
	p	40		
Random (Erdos-Renyi)	prob	0.025		
Scale free	$m_0$	24		
	$m_1$	12		
Small world	$\kappa$	24		
	$\beta$	0.1		

## 4.1.1 Random Graph

Random graphs have a short average path length. The graph is defined by the total number of nodes, and by the probability of any two nodes to be connected. Thus, all connections are random. They typically have a small clustering coefficient. Here we used a variant of the Erdős-Rényi random graph model (Erdős and Rényi, 1960) as implemented by Brugger and Schwirzer (2011).

### 4.1.2 Caveman Graph

The caveman structure, as defined by Watts (2003), has k isolated and fully connected 'cliques' from which one link is changed to connect one clique to another, rendering all cliques to be connected. Thus, relative to random graphs, they have a high clustering coefficient and a large average shortest path length.

### 4.1.3 Small World Graph

Small world graphs have characteristics that lie in between random graphs and highly clustered graphs (such as caveman graphs): they have a high clustering coefficient similar to the latter, but also have a small average shortest path similar to the former. Many real-world networks have been observed to have a small world structure, and

thus we included it in our simulations. Here we used the graph as defined by Watts and Strogatz (1998), and implemented by Brugger and Schwirzer (2011).

## 4.1.4 Scale-Free Graph

Scale-free network structures are often found where new nodes are constantly being added, and they are connected to already well-connected nodes. Such a structure displays a scale-free power-law distribution of the degree (connectivity) of nodes. Thus, there are few nodes that are highly connected, and more nodes that are moderately or mildly connected. Compared to random graphs, they have a smaller average shortest path. This graph was implemented by Brugger and Schwirzer (2011) as defined by Barabási and Albert (1999).

#### 4.2 Ideas

After choosing a starting structure for our model, we then chose a distribution for the starting ideas (states) of nodes. Each node was randomly assigned one of these ideas, thus allowing for multiple nodes to have the same idea. For the caveman structure, however, there were two other options: to either distribute the starting ideas 'parallel' to the structure, i.e. such that all nodes in a cave shared the same idea, or 'anti-parallel' such that all nodes in a cave had a different idea. This was used for the analysis of the effect of the idea distribution on the network structure. \*\*\*\*FIGURE OF IDEA DISTRIBUTION\*\*\*\* illustrates these idea distributions. Why was the caveman structure investigated? There were two reasons: firstly, it was straightforward how to define idea distributions that are in accord or disaccord with the caves in the network. Secondly, in the scientific community, research teams may often be made up of closely-connected members that are only weakly connected to other research teams, and within these teams, members may or may not be interested in the same ideas for research.

As previously mentioned, each node had a small probability  $\alpha$  of adopting a novel idea from a virtually unlimited number of new ideas.

# 5 Implementation

Our simulation comprises mainly two parts: the phase 1 corresponding to the study of the effect of network structure on the idea distribution, and phase 2 where we study how the distribution of the ideas affect the topology of the network. These two phases are implemented in the same MATLAB file **mainscript.m**. Implementation of both phases is divided into three steps.

**Step 1:** Network structure is chosen and we generate the adjacency matrix corresponding to that structure. The initial idea distribution is also generated.

Step 2: The updating process is done onto the chosen network structure.

**Step 3:** At this step a series of functions are called to get the results.

# 5.1 Step 1: Generating network structures and initial idea distribution

For the first phase, one of the functions step1\_scalefree, step1\_caveman, step1\_smallworld and step1\_randomgraph is called, depending on our choice. Each function is found in the MATLAB files step1\_scalefree.m, step1\_caveman.m, step1\_randomgraph.m and step1\_smallworld.m correspondingly. Each of these functions generates the adjacency matrix corresponding to the chosen network structure. Also in this step, the initial idea distribution - a random distribution for the first phase - is applied onto the nodes.

For the second phase we generated just the adjacency matrix for the caveman structure network, and then we chose among three different initial idea distributions: random, parallel or anti-parallel. After choosing one of them, the initial idea distribution vector is generated.

## 5.2 Step 2: Update

This step is the same for both phases.

The updating rules in the simulation are implemented in the file **step2.m** and done by function **step2** which requires the following parameters:

• t\_end: number of iterations

• phi: network reorganization rate

• alpha: innovation rate

• mat: initial connectivity matrix

• vec: initial idea vector

• p: initial number of opinions

And has as outputs:

- mat: connectivity matrix after simulation
- vec: idea vector after simulation
- dominant\_freq: the vector holding the frequency of the dominant idea
- most\_freq: the vector holding the index of the dominating idea in each time step

The function **step2**, in which the updating process is executed, implements the following algorithm:

# Algorithm 1 Update process

```
for each of the iterations do
   choose a node x1 in the network at random
   generate a random number a1
   if a1 < phi then
                                    \triangleright With probability phi we reorganize the network
      eliminate the connection between x1 and one of its neighbours with a different
idea
      select another node at random among the nodes with the same idea as x1 and
that are not already neighbours of x1 and create a connection between them
   else change the idea of x1 to one of the ideas of its neighbours which meet the
threshold
   end if
   choose a node y in the random to come up with a novel idea
   generate a random number a2
   if a2 < \alpha then y comes up with a new idea \triangleright With probability alpha y comes up
with a new idea
   end if
end for
```

# 5.3 Step 3: Getting results

In the phase 1 we want to observe the influence of a certain network structure on the distribution of ideas after the updating process (step 2). For this, it is necessary to search for features that reflect the final distribution of ideas. In this study we observed the following features:

- n\_index: the average neighbor index of the network
- intra\_idea\_distance: the average of the average shortest distance between agents holding the same idea
- average\_dominance\_time: the average of the dominance time for different dominance periods

All these parameters are obtained in **step3a.m**, **step3b.m** and **step3d.m** which are called in **main\_script.m**.

In phase 2 we want to find out how the distribution of ideas changes the network structure. For this reason we look at a fixed network structure: the caveman structure. After step 2 we want to see how the initial structure was modified by the updating process, and in order to do this we observed the following features:

- clust\_coeff: the clustering coefficient of the network
- [dgr,frq]: degree vector and its corresponding frequency vector
- average\_path\_length: the average path length for the graph
- graph\_diameter: outputs the diameter of the graph

# 6 Simulation Results and Discussion

For the first section of the results, ideas were assigned randomly onto the nodes of four different network structures (caveman, small world, random, and scale-free). Chapter 4 describes the network structures and idea distributions that are discussed. Simulations were run for each network structure for 27 different parameter combinations (three values for each of  $\phi$ ,  $\delta$  and  $\alpha$ ). The effects of each network structure on the idea distribution was evaluated on five features: the average dominance time of dominant (i.e., most frequent) ideas, the novelty index, the intra-idea distance, the neighbourhood index, and the frequency of dominance of an idea (see Table 1). We also investigated how these effects varied with the three parameters  $(\phi, \delta, \alpha)$ .

The second section of the results investigated the opposite direction of effects: how changing the idea distribution affected the resulting network structure. To do this we applied three different idea distributions (random, parallel, and anti-parallel to the structure) to a caveman network (see Table 1) with 40 caves. There were again 27 different parameter combinations for each idea distribution, as in the previous analysis. The five features of the network structure that we evaluated were: the clustering coefficient, the degree distribution of the nodes, the number of connected components, the average path length, and the network diameter.

We found that there were different behaviours for some features of the idea distribution depending on the network structure, and that there were different behaviours for the structure features depending on the idea distribution. Both of these results were at least partly influenced by the values of  $\phi$ ,  $\delta$ , and  $\alpha$ .

#### 6.1 Effects of Network Structure on Idea Distribution

The results of the average dominance time and the novelty index were too dependent on the parameter values and were not included; the network structure does not seem to play a strong enough role over all parameter combinations in influencing these two features. It is not surprising, however, that the novelty index was highly dependent on the value of  $\alpha$ , but it is not so intuitive why the average dominance time was not correlated with  $\alpha$  at all: increasing the number of novel ideas decreases the possible number of 'followers' for already-established ideas, which should affect the frequency of dominance and therefore the dominance time. Perhaps the value of  $\alpha$  would need to be increased to observe this behaviour.

Furthermore, we observed two different results for the intra-idea distance and the neighbourhood index: those from the caveman and small world structures, and those from the random and scale-free structures. All three parameter values affected these results.

## 6.1.1 Intra-Idea Distance and Neighbourhood Index

For any parameter combination, the caveman and small world structures resulted in larger intra-idea distances (respectively) than those of the random and scale-free structures, which were very similar to each other (Figure 1). Interestingly, a similar influence was found on the neighbourhood index: the caveman structure held the largest index regardless of parameter combination, followed by the small world, random, and scale-free structures (Figure 1). It seems that the caveman structure encourages nodes to be within the direct neighbourhood of like-minded nodes (nodes with the same idea) and at a farther distance from like-minded nodes that are not in their direct neighbourhood, whereas the random and scale-free structures have a tendency to keep like-minded nodes in each other's direct neighbourhood but to also keep those like-minded nodes not in their direct neighbourhood at a shorter distance. The difference in intra-idea distance between network structures could be a result of the general smaller average path distance that random and scale-free networks have as compared to the caveman and small world graphs.

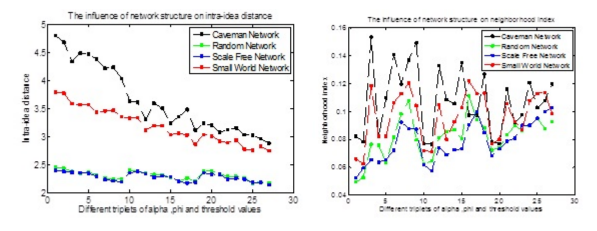


Figure 1: The influence of network structure on intra idea distance (left) and neighborhood index (right) for 27 distinct sets of parameters (triplets of alpha-phi-threshold).

**Dependence on**  $\alpha$  For all network structures, increasing the level of innovation  $\alpha$  decreased the intra-idea distance. This effect was more pronounced in the scale-free and random networks (Figure 2). Does innovation bring people together in scientific communities? Perhaps, and perhaps not. It is possible that these results are due to the construction of our model: because the mechanism of influence involves complex

contagion, new ideas would not be able to spread (since they are held only by the originator) and therefore these ideas would technically have an intra-idea distance of zero.

Higher values of  $\alpha$  also increased the neighbourhood index for all network structures. This correlation was again most visible for the scale-free network, followed by the random network (Figure 3). One possible explanation for this correlation is that the more novel ideas nodes create, the less likely it is that the contagion threshold is met for other ideas, and thus nodes will only be rewiring instead of also changing their ideas. Thus more like-minded nodes will be connected, and the index increases.

**Dependence on**  $\phi$  By increasing  $\phi$ , the intra-idea distance decreased for all network structures. This correlation is quite an intuitive result since  $\phi$  is the probability of deleting a connection between two nodes with different ideas and the formation of a new connection between two nodes with the same idea, and thus, by increasing  $\phi$  the distance between like-minded nodes decreases. Unlike the  $\alpha$  parameter, the effects of  $\phi$  are more pronounced for the caveman and small world structures rather than for the random and scale-free networks (Figure 4).

On the other hand, the effects of  $\phi$  on the neighborhood index varied between networks. Increasing  $\phi$  increased the neighbourhood indexes of the random and scale-free networks, while it decreased the neighbourhood index of the caveman network and had no correlation with changes in the small world network (Figure 5).

**Dependence on**  $\delta$  Increasing values of the complex contagion threshold  $\delta$  slightly decreased the intra-idea distance for the caveman and small world network structures (Figure 6).

No correlation was found between the neighbourhood index of the small world, scale-free, and random network structures and values of  $\delta$ . However, the neighbourhood index for the caveman network increased as  $\delta$  increased (Figure 7). This is an interesting result. On one hand, it is intuitive since requiring more like-minded nodes to be in the direct neighbourhood for a node to adopt their idea automatically increases the number of like-minded nodes in the neighbourhood (if the node adopts their idea). However, increasing  $\delta$  could have the opposite effect: if the threshold is too high, nodes will not adopt their neighbours' ideas and thus the neighbourhood index would remain small. The influence of  $\delta$  only on the caveman network could be due to this structure's higher degree per node: almost all nodes have the same degree, whereas the other structures have the same average amount but with greater variance.

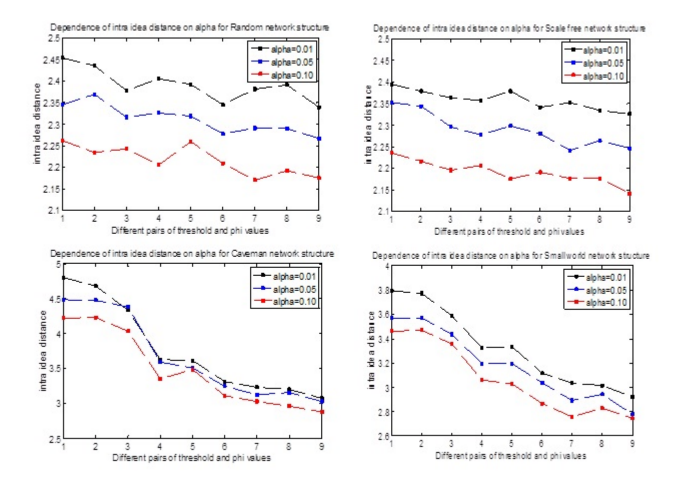


Figure 2: Dependence of intra idea distance on alpha for each network structure. Horizontal axis corresponds to distinct pairs of phi-threshold and vertical axis is the intra idea distance. Each curve corresponds to a different value of alpha. Black curve corresponds to the smallest value of alpha and it always has the highest intra idea distance while the red curve corresponds to the largest value of alpha which always has the lowest value of intra idea distance.

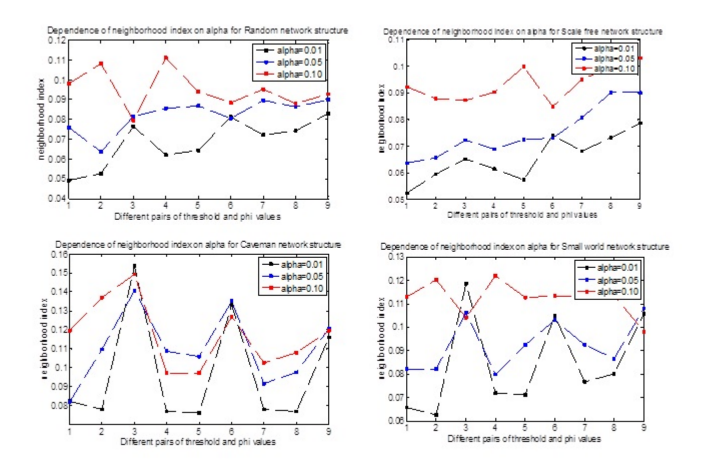


Figure 3: Dependence of neighborhood index on alpha for each network structure. Horizontal axis corresponds to distinct pairs of phi-threshold and vertical axis is the neighborhood index. Each curve corresponds to a different value of alpha. Red curve corresponds to the largest value of alpha and it has the highest neighborhood index while the black curve corresponds to the smallest value of alpha which has the lowest value of intra idea distance.

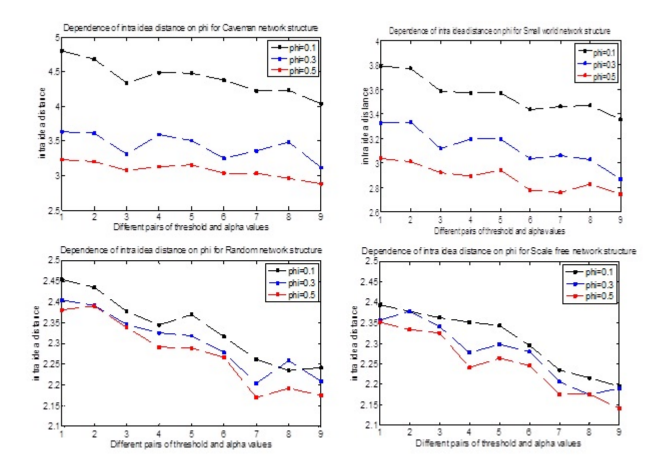


Figure 4: Dependence of intra idea distance on phi for each network structure. Horizontal axis corresponds to distinct pairs of alpha-threshold and vertical axis is the intra idea distance. Each curve corresponds to a different value of phi. Black curve corresponds to the smallest value of phi and it always has the highest intra idea distance while the red curve corresponds to the largest value of phi which always has the lowest value of intra idea distance.

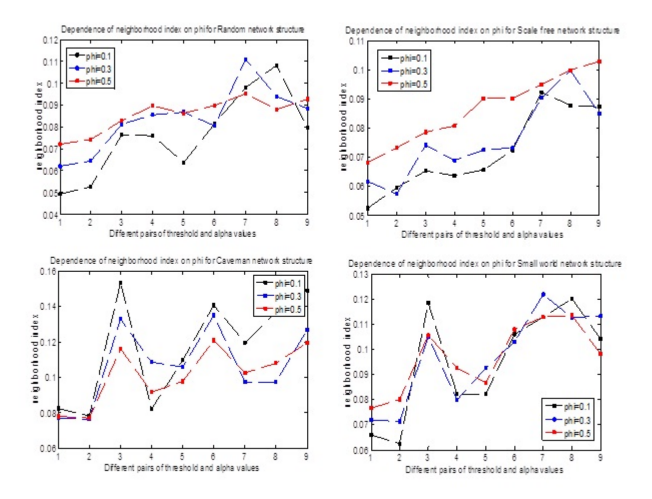


Figure 5: Dependence of neighborhood index on phi for each network structure. Horizontal axis corresponds to distinct pairs of alpha-threshold and vertical axis is the neighborhood index. Each curve corresponds to a different value of phi. Increasing phi leads to increase neighborhood distance for Random and Scale free networks while increasing phi causes the neighborhood index to be decreased for Cavemen Network. For Small world network there is no significant dependence of neighborhood index on phi.

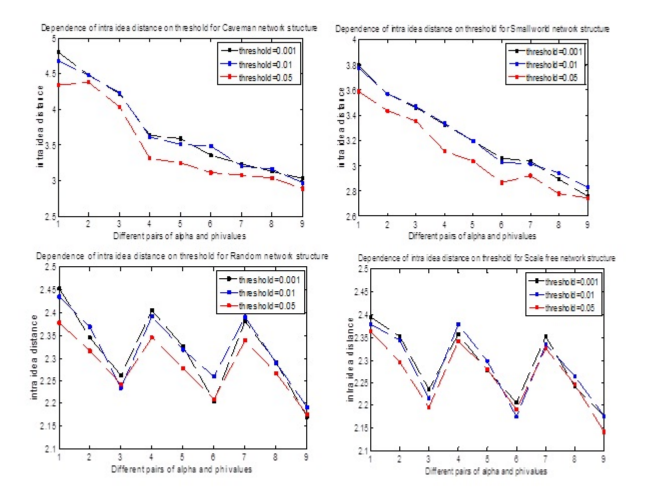


Figure 6: Dependence of intra idea distance on complex contagion threshold for each network structure. Horizontal axis corresponds to distinct pairs of alpha-phi and vertical axis is the intra idea distance. Each curve corresponds to a different value of complex contagion threshold. Black curve corresponds to the smallest value of complex contagion threshold and it has the highest intra idea distance while the red curve corresponds to the largest value of complex contagion threshold which has the lowest value of intra idea distance.

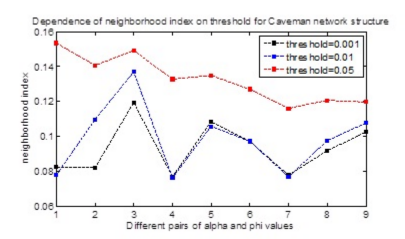


Figure 7: Dependence of neighborhood index on complex contagion threshold. Horizontal axis corresponds to distinct pairs of alpha-phi and vertical axis is neighborhood index. Each curve corresponds to a different value of complex contagion threshold. Red curve corresponds to the largest value of complex contagion threshold which has the largest value of neighborhood index while black curve corresponds to the smallest value of complex contagion threshold and it has the smallest neighborhood index.

#### 6.1.2 Frequency of Dominance

This feature is interesting if one considers scientific society. How dominant are dominant ideas in the scientific community? Does the structure of the community influence this dominance? For all parameter combinations and for all network structures in our simulations, the frequency of dominance of ideas increased with time. This may suggest that none of these structures impede the adoption of new ideas. More surprisingly, however, the increase in dominance frequency progressed more quickly for the caveman network structure (Figure 8). Could it be that this structure encourages nodes to adopt dominant ideas more easily? This may not be generalizable because the results are quite sensitive to parameter values due to the stochasticity of the simulations. For example, Figure 8 shows a different combination of parameters, and here the faster increase in the dominance is not observed for the caveman structure.

## 6.2 Effects of Idea Distribution on Network Structure

We observed that the results of applying a parallel idea distribution on the features of the network structure were quite different to the results of the random and anti-parallel idea distributions. It is interesting that for all parameter combinations and for each idea distribution, the network always remained fully connected. This could be because of the nature of our model: in order to disconnect with one node, there must be another node with the same idea to connect with. The nodes in a caveman network structure are, in a sense, 'saturated' since they are fully connected to their caves, and thus they remain connected to at least one of their original cave members.

While the networks always remained fully connected (Figure 9), the remaining features changed. These effects did not change with the parameter  $\alpha$  for any of the idea distributions (see Figure 10, Figure 11 and Figure 12). Given these simulation results, perhaps a caveman-structured scientific community would also manage certain levels of innovativity without changing its fundamental structure.

## 6.2.1 Clustering Coefficient

The clustering coefficient of resulting networks varied with the type of idea distribution. When nodes in the same cave shared the same idea (parallel distribution), the clustering coefficient was larger (Figure 13). Additionally, the clustering coefficients for both the random and the anti-parallel idea distributions decreased in a similar manner regardless of the parameter combinations. Both of these results are intuitive since less rewiring would have taken place for the parallel distribution case, and the high clustering coefficient of the caveman structure would have been conserved,

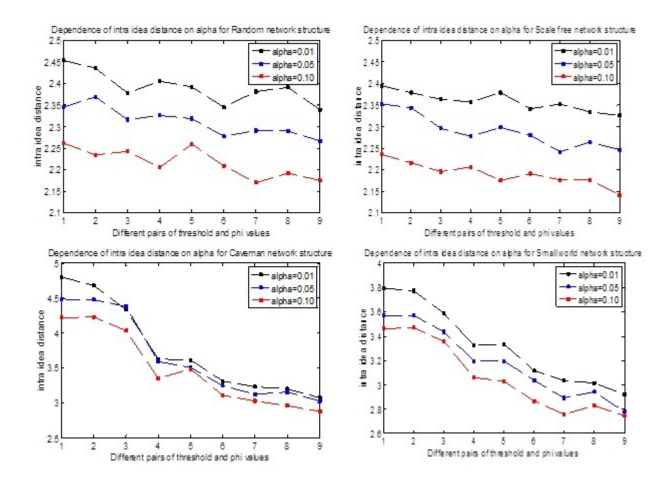


Figure 8: Frequency of the dominant idea at each time for four distinct network structures and two different sets of parameters: phi=0.5, alpha=0.01, threshold=0.05 (top) and phi=0.3, alpha=0.1, threshold=0.05 (bottom).

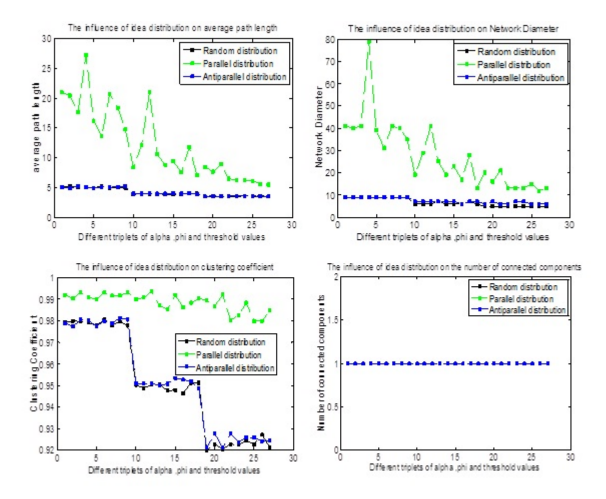


Figure 9: The influence of idea distribution on average path length (upper left), network diameter (upper right), clustering coefficient (bottom left) and number of connected components (bottom right) for 27 distinct sets of parameters (triplets of alpha-phi-threshold).

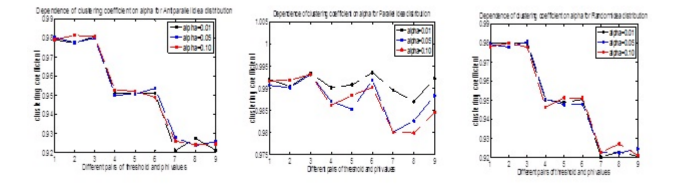


Figure 10: Dependence of clustering coefficient on alpha for each idea distribution. Horizontal axis corresponds to distinct pairs of phi-threshold and vertical axis is the clustering coefficient. Each curve corresponds to a different value of alpha. There is no dependence on alpha for clustering coefficient of the networks with any idea distribution.

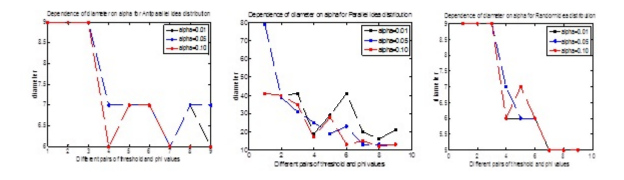


Figure 11: Dependence of network diameter on alpha for each idea distribution. Horizontal axis corresponds to distinct pairs of phi-threshold and vertical axis is the network diameter. Each curve corresponds to a different value of alpha. There is no dependence on alpha for the diameters of the networks with any idea distribution

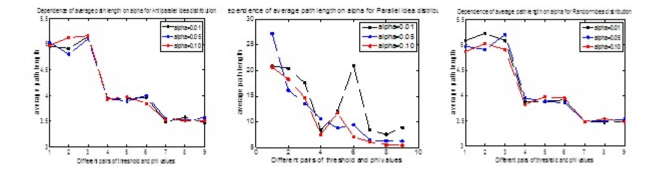


Figure 12: Dependence of average path length on alpha for each idea distribution. Horizontal axis corresponds to distinct pairs of phi-threshold and vertical axis is the average path length. Each curve corresponds to a different value of alpha. There is no dependence on alpha for average path length of the networks with any idea distribution.

whereas more rewiring would have ocurred for the two other distributions, thus decreasing the clustering coefficients.

**Dependence on parameters** Increasing values of the rewiring parameter  $\phi$  decreased the clustering coefficient for all starting distributions, especially for the random and anti-parallel distributions (Figure 14). This again is intuitive since  $\phi$  increases the chances of changing connections in a highly clustered network. Increasing values of  $\delta$ , however, only slightly increased the clustering coefficient when using a parallel idea distribution (Figure 15).

### 6.2.2 Average Path Length

The average path length of resulting networks was larger when a parallel idea distribution was used (Figure 9). This is not surprising, since the structure of the caveman network was more preserved (because most nodes were already connected to like-minded nodes and did not need to rewire), and its average path length is larger than that of more randomized networks, such as the ones resulting from a larger amount of rewiring.

**Dependence on parameters** Increasing  $\phi$ , regardless of the idea distribution, decreased the average path length. This effect was more prononced for the random and anti-parallel idea distribution cases (Figure 16). This is another intuitive result since more rewiring naturally disturbs the rigid structure of a caveman network,

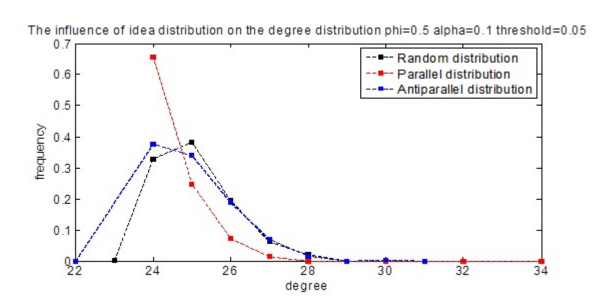


Figure 13: The influence of idea distribution on the degree distribution of the network. Horizontal axis corresponds to the degree of the nodes and vertical axis corresponds to the relative frequency of the nodes with that certain degree.

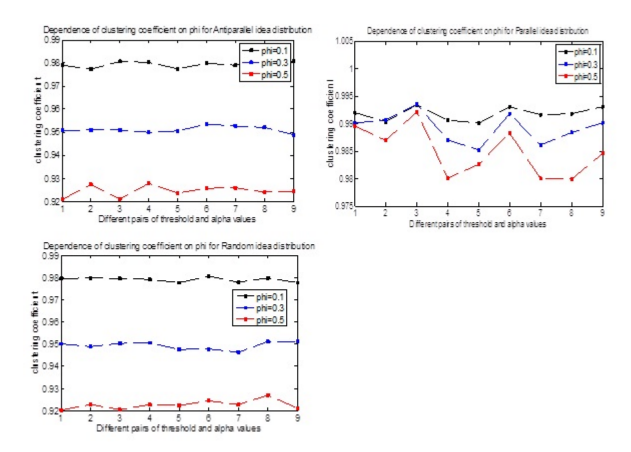


Figure 14: Dependence of average path length on phi for each idea distribution. Horizontal axis corresponds to distinct pairs of alpha-threshold and vertical axis is the average path length. Each curve corresponds to a different value of phi.

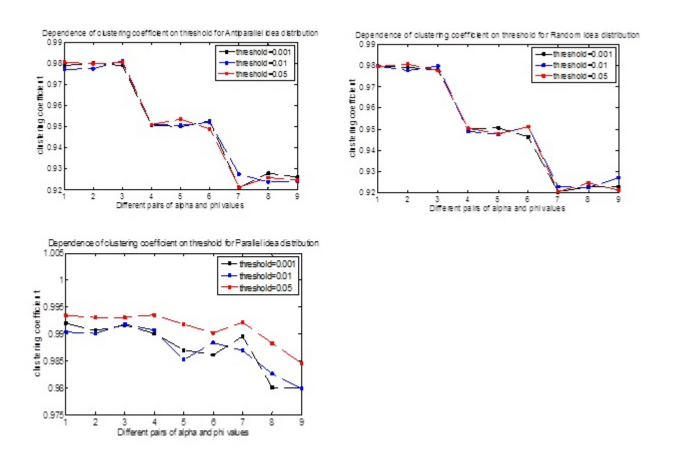


Figure 15: Dependence of clustering coefficient on complex contagion threshold for each idea distribution. Horizontal axis corresponds to distinct pairs of phi-alpha and vertical axis is the clustering coefficient. Each curve corresponds to a different value of complex contagion threshold.

thereby decreasing its large average path length; rewiring also occurs less frequently in the parallel idea distribution case because most nodes are already connected to like-minded nodes. Changing values of  $\delta$  (which requires more than one neighbouring node to have the same idea in order to influence the chosen node) did not change the effects of idea distributions on the average path length (Figure 17). This is natural, since nodes were either already connected to a significant number of nodes with the same idea (in the case of the parallel distribution), not connected to any other node with the same idea (the anti-parallel case), or connected to nodes with random assignment of ideas. Thus the threshold  $\delta$  would either already be fulfilled from the start, would definitely not be fulfilled, or would have a very small chance of being fulfilled, respectively.

#### 6.2.3 Network Diameter

Similar to the clustering coefficient and the average path length, the network diameter was larger when the parallel idea distribution was applied (Figure 9). This distribution encouraged the structure of the caveman network to remain mostly unchanged, and thus the farthest distance between two nodes was larger than for the case of random or anti-parallel distributions, where more rewiring occurred.

**Dependence on parameters** As  $\phi$  increased, the network diameter decreased for all idea distributions, and slightly less for the parallel distribution (Figure 19). Rewiring a caveman network intuitively may decrease the network diameter by connecting more of its caves. Similar to the average path length, values of  $\delta$  did not change the behaviour of the network diameter given the idea distributions (Figure 18).

#### 6.2.4 Degree Distribution

Random graphs have a somewhat normally distributed degree distribution. Scale-free graphs, on the other hand, have a degree distribution that follows a scale-free power-law. We observed that the degree distribution of the networks depended on the idea distribution (Figure 13). A parallel idea distribution resulted in a degree distribution similar to that of a scale-free graph, which is not surprising. Caveman graphs have two or three different degrees for their nodes, and allowing for some rewiring would 'smooth' out this discrete distribution. Similar to previous results, the random and anti-parallel idea distributions behaved similarly: their degree distribution was similar to that of random graphs. It is interesting to see such a visible difference in these distributions over relatively few time steps (1000 steps), regardless of the parameter combinations.

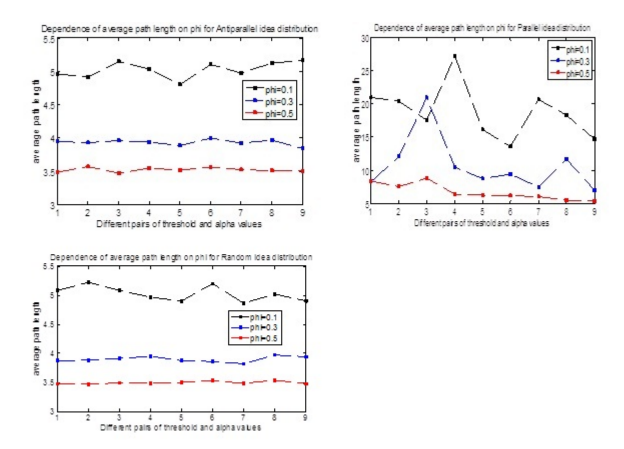


Figure 16: Dependence of clustering coefficient on phi for each idea distribution. Horizontal axis corresponds to distinct pairs of alpha- threshold and vertical axis is the clustering coefficient. Each curve corresponds to a different value of phi.

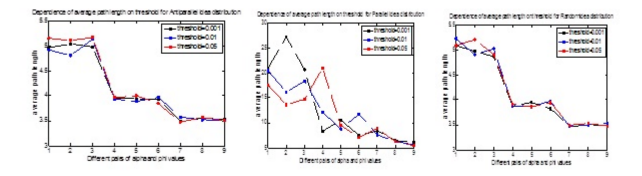


Figure 17: Dependence of average path length on complex contagion threshold for each idea distribution. Horizontal axis corresponds to distinct pairs of alpha-phi and vertical axis is the average path length. Each curve corresponds to a different value of complex contagion threshold. There is no dependence on complex contagion for average path length with any idea distribution.

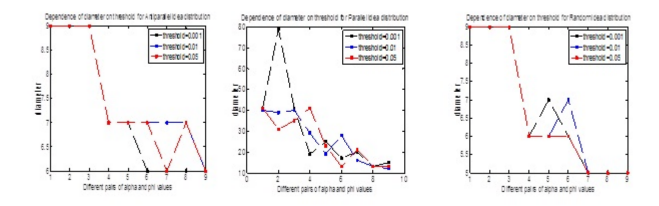


Figure 18: Dependence of network diameter on complex contagion threshold for each idea distribution. Horizontal axis corresponds to distinct pairs of phi-alpha and vertical axis is the network diameter. Each curve corresponds to a different value of complex contagion threshold. There is no dependence on complex contagion for network diameter with any idea distribution.

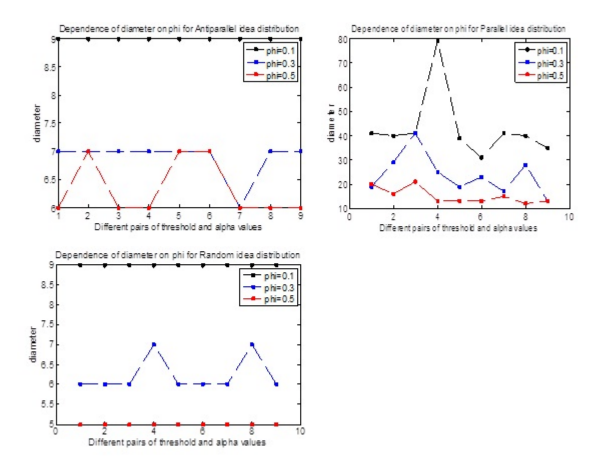


Figure 19: Dependence of network diameter on phi for each idea distribution. Horizontal axis corresponds to distinct pairs of alpha-threshold and vertical axis is the network diameter. Each curve corresponds to a different value of phi.

#### 6.3 Discussion

As previously mentioned, we observed that some features of the idea distribution varied with the type of network structure. We also observed that some features of the network structure varied with the kind of idea distribution implemented. Some of these results were intuitive and expected, while others were not.

Network structure mainly influenced two features of the idea distribution: the intra-idea distance and the neighbourhood index. The intra-idea distance was larger for the caveman and small world networks, as anticipated. However, contrary to expectations, the neighbourhood index was larger for these two structures, and the caveman structure did not decrease the frequency of dominance (and neither did any other structure). Additionally, the values of  $\phi$  and  $\delta$  did correlate with changes in the values of the intra-idea distance (by decreasing it) and the neighbourhood index (with different effects depending on the structure), but not always:  $\delta$  did not vary the results of the intra-idea distance or the neighbourhood index of the random and scale-free networks, and  $\phi$  did not vary the neighbourhood index in the small world network. Lastly, and also contrary to expectations, increasing the  $\alpha$  values increased the neighbourhood index of networks.

Four of the five features of the network structure changed with different idea distributions. The connected component interestingly remained one, regardless of the idea distribution and parameter values. Feature values of the anti-parallel and random idea distributions differed from those of the parallel distribution. Most features changed as expected: the clustering coefficient, average path length, and network diameter were larger when the parallel idea distribution was used as compared to the other two, and their values were similar to those of a caveman network structure. The clustering coefficient of the anti-parallel idea distribution networks did decrease, as expected. The degree distribution did increase in variance (becoming more similar to a normal distribution) for the networks that were initiated with a random or antiparallel idea distribution, but the degree distribution of the parallel idea distribution networks resembled that of a scale-free power law. Values of  $\phi$  influenced the features of the random and antiparallel distribution networks the most. However, the structure features of the networks with the random idea distribution were not more sensitive to parameter values as would have been expected; they behaved similar to those of the networks with the anti-parallel idea distribution. Values of  $\delta$  only correlated with changes in the clustering coefficient of the networks with a parallel idea distribution, and only weakly. Values of  $\alpha$  did not correlated with any changes of feature values.

# 7 Summary and Outlook

To conclude our simulation study, our results support the general idea that network structure and qualities of the ideas held in them may mutually influence each other. The more rigid the structure of a network was, the more likely that the intra-idea distance and neighbourhood index of the network was larger. These two features varied more with the innovation rate  $\alpha$  for less structured networks, and varied more with the complex contagion threhold  $\delta$  for the most rigid structure - the caveman network. The average dominance time seemed to vary more with parameter values  $(\phi, \delta, \text{ and } \alpha)$  than with the network structures, whereas the frequency of dominance of ideas did not decrease on average in the long run with any network structure.

A network in which the pattern of ideas held by the nodes are 'in accord' within the clusters of the caveman network maintained more of the structure features of a caveman network. These values varied with values of rewiring probabilities  $\phi$  and complex contagion threholds  $\delta$ . Networks with a random pattern of ideas or with a pattern with more 'disaccord' within the clusters resulted in a structure that began to resemble a random graph more than a caveman network. These networks' values varied more with the parameter  $\phi$  of rewiring. Nodes given the chance to rewire with nodes that share the same idea had more opportunity to do so in the random and 'disaccord' idea patterns than in the pattern which already had much accord.

Thus, in addition to the structure of networks and the pattern of ideas in them, complex contagion thresholds, innovation rates, and the probability of creating new connections (while reflecting 'preferences' of being connected with like-minded others) tend to influence features of the network. Several extensions to the proposed model could be investigated to better understand the relationships between network structure and idea distribution.

Rewiring criteria Future simulations may compare the emerging features of idea distributions not just between network structures, but also between different rewiring criteria. It is not clear how much of the features of the idea distribution in this simulation study was a result of the network structure or of the 'preference' that nodes had in rewiring to like-minded nodes. Therefore, these results could be compared to (1) random rewiring or, to allow structure to play a larger role, (2) to allow random rewiring to nodes that are at most a distance of three nodes away. This is somewhat more realistic since most people connect with individuals who are somewhat in their vicinity through mutual connections.

Complex contagion and innovation Considering complex contagion, novel ideas in our model were at a disadvantage for spreading in the network. Allowing novel

ideas to have a larger influence weight may be one way to grant the ability of them to spread to other nodes. Alternatively, the complex contagion threshold for novel ideas may be lowered.

Random idea adoption Future simulations may investigate the effects of network structure on idea distributions by comparing to a 'benchmark' model. This model would update the ideas of nodes at random, and thus the effects of connections may be better observed within network structures and then compared between network structures.

# 8 References

#### A MATLAB code

## A.1 Main script

```
\scriptstyle 1 %% The first phase: Simulation to study the influence of network ...
     2 %%% The following parameters will remain constant during this study,
3 %%% so we won't play with them %%%
4 n=1000; %% the number of agents %%%
5 m=40; %% initial number of clusters for caveman matrix%%%
6 p=40; %% initial number of opinions%%
7 t_end=1000; %%% number of iterations%%
8 %%% for the following parameters we'll run different simulations based on
9 %%% conmbinatorial complexity of the parameters%%%
phi_choices=[0.1,0.3,0.5]; %%% network reorganization rate%%
11 alpha_choices=[0.01,0.05,0.10]; %%% innovation rate %%%
12 threshold_choices=[0.001,0.01,0.05];%%% threshold for complex contagion %%%
13
15 %%% a totally random idea distribution , independent of the connectivity
16 %%% matrix, so applicable for every network structure is defined for this
17 %%% phase %%%
18 vec1=zeros(1,n);
19 for i=1:n
20
      vec1(i) = ceil(rand()*p);
21 end
23 for choice1=1:4
24 %% Step1: Definition of Different initial matrices ...
     switch choice1
26
         case 1
27
          %%% option 1: Caveman Connectivity Matrix
28
          mat1=step1_caveman(n,m);
29
          s1='Caveman';
31
         case 2
         %%% option 2: Random Connectivity Matrix
         prob=0.025; %%% probability of edge formation between any pairs ...
            of edges
         mat1=step1_randomgraph(n,prob);
34
         s1='Random';
35
         case 3
36
         %%% option 3: Scale Free Connectivity Matrix
37
```

```
38
          m0=24; % number of initially placed nodes
          m1=12; % number of nodes a new added node is connected to, 1 \leq ...
39
             m1 < m0
          mat1=step1_scalefree(n, m0, m1);
40
          s1='Scale_free';
          case 4
42
43
          %%% option 4: Small world Connectivity Matrix
          ka=24; %% mean degree (assumed to be an even integer)
44
          beta=0.01; %% rewiring probability
45
          mat1= step1_smallworld(n, ka, beta);
46
47
          s1='Small_world';
      end
      %% Step2: Simulation ...
50
          for choice2=1:3
51
          phi=phi_choices(choice2);
52
53
          for choice3=1:3
54
              alpha=alpha_choices(choice3);
             for choice4=1:3
55
56
               threshold=threshold_choices(choice4);
57
               [mat2, vec2, dominant_freq, most_freq] = step2 (t_end, phi, alpha, mat1, vec1, p, threshold);
58
                  %%% obtaining the final matrix and vector after running ...
                  simulation.
               %%%% We need step4c here, since it's outputs will be the input
               %%%% for step 3b
61
               62
63
               sp_mat2=sparse(mat2);
               [s,c]=graphconncomp(sp_mat2); %% matlab built in function ...
64
                  'Bioinformatics Toolbox'
               %%% s: number of connected components
65
               %%% c: vector which assigns each node to a connected component
66
67
               %% Step3: Results for structure to idea ...
68
                  70
               %%%%%% step3a: defining the average intra_idea ...
                  neighbourhood index %%%%%%
               neighbor_index=step3a(mat2, vec2);
71
72
               %%%%%% step3b: defining the average intra_idea distance ...
73
                  응응응응응응
               intra_idea_distance=step3b (mat2, vec2, s, c);
75
               %%%%%% step3c: frequency of dominant idea with respect to ...
76
                  time %%%%%%%
               %%% is the third output of the step2 function ...
77
                  (dominant_freq)%%%
```

```
78
                dominant_freq;
79
                %%%%%% step3d: Fraction of novel ideas (novelity index) %%%%%%
80
                nov_index=(length(find(vec2>p)))/(length(vec2)); %%% ...
81
                   indicates the fraction of agents holding the newly ...
                   generated ideas
82
                %%%%%% step3e: defining the average dominance time (the ...
83
                   average amount of time in which the dominating idea ...
                   keeps it's dominance over differnt dominance periods)
                average_dominance_time=step3e(most_freq);
                %%% naming the file which saves the results
               s2=int2str(choice2);
87
               s3=int2str(choice3);
               s4=int2str(choice4);
89
               name=['phase1_',s1,'_',s2,'_',s3,'_',s4];
90
91
                save(name);
              end
93
           end
94
       end
95 end
96
97 clear;
  100 %% The second phase: Simulation to study the influence of opinion on the ...
       101
102 %%% The following three parameters will remain constant during this study,
103 %%% so won't play with them %%%
104 n=1200; %% the number of agents
105 m=40; %% initial number of clusters for caveman matrix
106 p=40; %% initial number of opinions
107 t_end=1000; %%% number of iterations%%
108 %% for the following parameters we'll run different simulations based on
109 %%% conmbinatorial complexity of the parameters%%%
|110 phi_choices=[0.1,0.3,0.5]; %%% network reorganization rate%%
|111 alpha_choices=[0.01,0.05,0.10]; %%% innovation rate %%%
112 threshold_choices=[0.001,0.01,0.05]; %%% threshold for complex contagion %%%
113
114
115
116 %%%%%in this phase we'll keep connectivity matrix constant, so we only use
117 %%%%Caveman connectivity matrix %%%
118 mat1=step1_caveman(n,m);
119
120
121 for choice1=1:3
```

```
122 %% Step1: Definition of Different initial idea vectors ...
       %%%%%%%%%%%%%%%%%%%%%%
       switch choice1
123
              case 1
124
               %%%%%% option1: Random idea vector %%%%%%%
125
               %%% a totally random idea distribution , independent of the \dots
126
                  connectivity
               %%%% matrix, so applicable for every network structure
127
              vec1=zeros(1,n);
128
              for i=1:n
129
130
                    vec1(i) = ceil(rand()*p);
131
              end
132
              s1='Random';
133
              case 2
134
              %%%%%% option2: Parallel idea vector %%%%%%%
              %%% This idea vector is applicable only for caveman ...
135
                  connectivity matrix in
136
               %%% which every agents inside a cluster have the same idea
137
               vec1=zeros(1,n);
138
               for i=1:(m-1) %%% for each cluster
139
                  for j=1:ceil(n/m)
                      vec1((i*ceil(n/m))+j)=i; %%% all agents will hold the ...
140
                          i-th idea
141
                  end
142
              end
143
              s1='Parallel';
144
              case 3
145
               %%%%%% option3: Antiparallel idea vector %%%%%%%
               %%% This idea vector is applicable only for caveman ...
146
                  connectivity matrix in
               %%% which every agents inside a cluster have different idea
147
148
               vec1=zeros(1,n);
               for i=1:(m-1) %%% for each cluster
149
150
                   for j=1:ceil(n/m)
                      vec1((i*ceil(n/m))+j)=j; %%% all agents will hold ...
151
                         different idea
152
                   end
153
               end
154
               s1='Antiparallel';
155
       end
       %% Step2: Simulation ...
156
           for choice2=1:3
157
           phi=phi_choices(choice2);
158
159
            for choice3=1:3
160
                alpha=alpha_choices(choice3);
161
162
             for choice4=1:3
               threshold=threshold_choices(choice4);
163
164
```

```
165
              [mat2, vec2, dominant_freq, most_freq] = step2 (t_end, phi, alpha, mat1, vec1, p, threshold);
                  %%% obtaining the final matrix and vector after running ...
                  simulation.
166
167
              %% Step3: Results for idea to structure ...
168
                  169
              %%%%%% step4a: clustering coefficient of the final network ...
170
                  응응응응응응응
171
              clust_coefficient=step4a(mat2);
172
173
              %%%%%% step4b: degree distribution of the final network ...
                  응응응응응응응응응응
174
              [dgr,frq]=step4b(mat2);
175
              average_degree=sum(dgr.*frq)/sum(frq);
176
177
              178
              sp_mat2=sparse(mat2);
179
              [s,c]=graphconncomp(sp_mat2); %% matlab built in function ...
                  'Bioinformatics Toolbox'
              %%% s: number of connected components
180
              %%% c: vector which assigns each node to a connected component
181
182
183
              %%%%%% step4d: average path length for the final network ...
                  응응응응응응응응응
              average_path_length = step4d( mat2,s,c );
184
185
              186
              diam=step4e(mat2,s,c);
187
188
189
              %%%naming and saving
              s2=int2str(choice2);
190
              s3=int2str(choice3);
191
              s4=int2str(choice4);
192
              name=['phase2_',s1,'_',s2,'_',s3,'_',s4];
193
              save(name);
194
195
            end
196
           end
       end
197
198 end
199
200 clear;
```

## A.2 Step 1: generation of structure networks

### randomgraph.m

```
2 %%% code adopted from %%%
3 %%% Modeling and Simulating Social Systems with MATLAB %%%
4 %%% http://www.soms.ethz.ch/teaching/MatlabFall2012 %%%
5 %%% Authors: Stefan Brugger and Cristoph Schwirzer, 2011 %%%
8 function final= step1_randomgraph(n, p)
9 % Generates an undirected random graph (without self-loops) of size n (as
10 % described in the Erdoes-Renyi model)
11 %
12 % INPUT
13 % n: [1]: number of nodes
14 % p: [1]: probability that node i and node j, i != j, are connected by ...
      an edge
15 %
16 % OUTPUT
17 % final: [n n] full symmetric adjacency matrix representing the ...
      generated graph
18
19 % Note: A generation based on sprandsym(n, p) failed (for some values of p
20 % sprandsym was far off from the expected number of n*n*p non-zeros), ...
      therefore
21 % this longish implementation instead of just doing the following:
23 % B = sprandsym(n, p);
24 % A = (B-diag(diag(B)) \neq 0);
26
27 % Idea: first generate the number of non-zero values in every row for a ...
      general
28 % 0-1-adjacency matrix. For every row this number is distributed ...
      binomially with
29 % parameters n and p.
31 % The following lines calculate "rowsize = binoinv(rand(1, n), n, p)", ...
      just in a
32 % faster way for large values of n.
34 % generate a vector of n values chosen u.a.r. from (0,1)
35 v = rand(1, n);
36 % Sort them and calculate the binomial cumulative distribution function with
37 % parameters n and p at values 0 to n. Afterwards match the sorted random
38 % 0-1-values to those cdf-values, i.e. associate a binomial distributed ...
      value
```

```
39 % with each value in r. Each value in v also corresponds to a value in r:
40 % permute the values in rowSize s.t. they correspond to the order given \dots
       in v.
41 [r index] = sort(v); % i.e. v(index) == r holds
42 rowSize = zeros(1, n);
43 \quad \dot{j} = 0;
44 binoCDF = cumsum(binopdf(0:n, n, p));
45 for i = 1:n
46
     while j \le n \& \& binoCDF(j+1) \le r(i)
47
       j = j + 1;
       end
       rowSize(i) = j;
50 end
51 rowSize(index) = rowSize;
53 % for every row choose the non-zero entries in it
54 nNZ = sum(rowSize);
I = zeros(1, nNZ);
J = zeros(1, nNZ);
57 \quad j = 1;
58 \text{ for i} = 1:n
       I(j:j+rowSize(i)-1) = i;
59
       J(j:j+rowSize(i)-1) = randsample(n, rowSize(i));
61
       j = j + rowSize(i);
62 end
64 % restrict I and J to indices that correspond to entries above the main ...
      diagonal
65\, % and finally construct a symmetric sparse matrix using I and J
66 upperTriu = find(I<J);
67 I = I(upperTriu);
68 J = J(upperTriu);
69 A = sparse([I;J], [J;I], ones(1, 2*size(I, 2)), n, n);
70 final=full(A);
71 end % random_graph(...)
```

#### scalefree.m

```
2 %%% code adopted from
3 %%% Modeling and Simulating Social Systems with MATLAB
                                                         응응응
4 %%% http://www.soms.ethz.ch/teaching/MatlabFall2012
5 %%% Authors: Stefan Brugger and Cristoph Schwirzer, 2011 %%%
8 function A = scalefree(n, m0, m)
9 % Use the Barabasi-Albert model to generate a scale free graph of size n (as
10 % described in Albert-Laszlo Barabasi & Reka Albert: "Emergence of scaling
11 % in random networks")
12 %
13 % INPUT
14 % n: [1]: number of nodes
15 % m0: [1]: number of initially placed nodes
  % m: [1]: number of nodes a new added node is connected to, 1 \le m < m0
18 % OUPUT
19 % A: [n n] sparse symmetric adjacency matrix representing the generated ...
      graph
21\, % Start with a graph of size m0 and add edges to this graph. Each of ...
22 % nodes is connected to at least m nodes.
B = zeros(m0, m0);
24 \text{ for } i = 1:m0
      neighbors = randsample(m0-1, m);
      neighbors = neighbors + (neighbors≥i);
      B(i, neighbors) = 1;
      B(neighbors, i) = 1;
28
29 end
30
31 % Create a vector of edges added so far, i.e. nodes edge(2*i) and ...
      edge(2*i-1),
32 % 1 \le i \le nEdges, are connected by an edge.
33 [rows, columns] = find(triu(B));
34 nEdges = size(rows, 1);
35 edges = reshape([rows';columns'], 2*nEdges, 1);
36 edges = [edges; zeros(2*(n-m0)*m,1)];
38 % Add nodes m0+1:n, one at a time. Each node is connected to m existing ...
39 % where each of the existing nodes is chosen with a probability that is
40 % proportional to the number of nodes it is already connected to.
41 used = zeros(n, 1); % is a node already used in a timestep?
42 for i = m0+1:n
      neighbors = zeros(1, m);
43
```

```
for j=1:m
44
          k = edges(randi(2*nEdges));
45
          while used(k)
46
              k = edges(randi(2*nEdges));
47
          used(k) = 1;
49
50
          neighbors(j) = k;
       end
51
       used(neighbors) = 0;
52
       edges(2*nEdges+1:2*nEdges+2*m) = reshape([repmat(i, 1, m); ...
          neighbors], ...
        1, 2*m);
55
       nEdges = nEdges+m;
56 end
57
58 % finally construct a symmetric adjacency matrix using the vector of edges
59 edges = edges(1:2*nEdges);
60 first = edges(1:2:end);
61 second = edges(2:2:end);
62 A = sparse([first; second], [second; first], ones(2*nEdges, 1), n, n);
63
64 end % scale_free(...)
```

#### smallworld.m

```
2 %%% code adopted from
                                                    응응응
3 %%% Modeling and Simulating Social Systems with MATLAB
4 %% http://www.soms.ethz.ch/teaching/MatlabFall2012
5 %%% Authors: Stefan Brugger and Cristoph Schwirzer, 2011 %%%
8 function A = smallworld(n, k, beta)
9 % Generate a small world graph using the "Watts and Strogatz model" as
10 % described in Watts, D.J.; Strogatz, S.H.: "Collective dynamics of
11 % 'small-world' networks."
12 % A graph with n*k/2 edges is constructed, i.e. the nodal degree is n*k for
13 % every node.
14 %
15 % INPUT
16 % n: [1]: number of nodes of the graph to be generated
17 % k: [1]: mean degree (assumed to be an even integer)
18 % beta: [1]: rewiring probability
19 %
20 % OUPUT
21 % A: [n n] sparse symmetric adjacency matrix representing the generated \dots
     graph
```

```
23 % Construct a regular lattice: a graph with n nodes, each connected to k
24 % neighbors, k/2 on each side.
25 kHalf = k/2;
26 rows = reshape(repmat([1:n]', 1, k), n*k, 1);
27 columns = rows+reshape(repmat([[1:kHalf] [n-kHalf:n-1]], n, 1), n*k, 1);
28 columns = mod(columns-1, n) + 1;
29 B = sparse(rows, columns, ones(n*k, 1));
30 A = sparse([], [], [], n, n);
32 % With probability beta rewire an edge avoiding loops and link duplication.
33 % Until step i, only the columns 1:i are generated making implicit use ...
      of A's
34 % symmetry.
35 \text{ for i} = [1:n]
       \mbox{\%} The i-th column is stored full for fast access inside the \dots
           following loop.
       col=[full(A(i, 1:i-1))'; full(B(i:end, i))];
37
38
       for j = i+find(col(i+1:end))'
39
           if (rand()<beta)</pre>
40
               col(j)=0;
               k = randi(n);
41
               while k==i \mid \mid col(k)==1
42
                    k = randi(n);
               end
               col(k) = 1;
           end
       end
47
       A(:,i) = col;
48
49 end
50
51 % A is not yet symmetric: to speed things up, an edge connecting i and ...
       j, i < j
52 % implies A(i,j) ==1, A(i,j) might be zero.
T = triu(A);
54 A = T+T';
56 end % small_world(...)
```

#### step1\_caveman.m

```
1 function cave_mat = step1_caveman(n,m)
3 %%% n: the number of agents
4 %% m: initial number of clusters
5 %% p: initial maximum index of opinions
6 cave_mat=zeros(n,n); %Caveman Matrix
8 for i=1:n
      x=ceil(i/(n/m));
      for j=1:n
10
          y=ceil(j/(n/m));
11
          if x==y
12
              if i≠j
13
              cave_mat(i,j)=1; %definition of intracluster edges
14
15
              end
16
          end
17
      end
18 end
19 %%% x1 and x2 for each cluster represent the two agents who interact with
20 %%% nearby clusters%%
x1 = zeros(1, m);
x2 = zeros(1, m);
23 for i=1:m
      x1(i) = ceil(rand() * (n/m) + (n/m) * (i-1));
24
      ind=0;
25
      while(ind==0) %% This loop is used to prevent x1 and x2 to make the ...
26
          same numbers
          x2(i) = ceil(rand() * (n/m) + (n/m) * (i-1));
28
          if x2(i) \neq x1(i)
              ind=1;
29
30
          end
      end
31
32 end
33 %%% definition of intercluster edges
34 for i=1: (m-1)
      cave_mat (x2(i), x1(i+1))=1;
      cave_mat (x1(i+1), x2(i))=1;
36
37 end
38 cave_mat(x1(1), x2(m))=1;
39 cave_mat(x2(m), x1(1))=1;
40 end
```

## step1\_randomgraph.m

```
3 \quad \text{????} \\
4 %%% code adopted from %%%
5 %%% Modeling and Simulating Social Systems with MATLAB %%%
6 %%% http://www.soms.ethz.ch/teaching/MatlabFall2012 %%%
7 %%% Authors: Stefan Brugger and Cristoph Schwirzer, 2011 %%%
10 function final= step1_randomgraph(n, p)
11 % Generates an undirected random graph (without self-loops) of size n (as
12 % described in the Erdoes-Renyi model)
13 %
14 % INPUT
15 % n: [1]: number of nodes
16 % p: [1]: probability that node i and node j, i != j, are connected by ...
      an edge
17 %
18 % OUTPUT
19 % final: [n n] full symmetric adjacency matrix representing the ...
      generated graph
20
21 % Note: A generation based on sprandsym(n, p) failed (for some values of p
22 % sprandsym was far off from the expected number of n*n*p non-zeros), ...
      therefore
23 % this longish implementation instead of just doing the following:
25 \% B = sprandsym(n, p);
26 % A = (B-diag(diag(B)) \neq 0);
28
29 % Idea: first generate the number of non-zero values in every row for a ...
      general
30 % 0-1-adjacency matrix. For every row this number is distributed ...
     binomially with
31 % parameters n and p.
33 % The following lines calculate "rowsize = binoinv(rand(1, n), n, p)", ...
      just in a
34 % faster way for large values of n.
36 % generate a vector of n values chosen u.a.r. from (0,1)
37 v = rand(1, n);
38 % Sort them and calculate the binomial cumulative distribution function with
39 % parameters n and p at values 0 to n. Afterwards match the sorted random
40 % 0-1-values to those cdf-values, i.e. associate a binomial distributed ...
      value
```

```
41 % with each value in r. Each value in v also corresponds to a value in r:
42 % permute the values in rowSize s.t. they correspond to the order given \dots
      in v.
43 [r index] = sort(v); % i.e. v(index) == r holds
44 rowSize = zeros(1, n);
45 \quad \dot{j} = 0;
46 binoCDF = cumsum(binopdf(0:n, n, p));
47 for i = 1:n
48
      while j<n && binoCDF(j+1)<r(i)</pre>
49
       j = j + 1;
      end
       rowSize(i) = j;
52 end
53 rowSize(index) = rowSize;
55 % for every row choose the non-zero entries in it
56 nNZ = sum(rowSize);
I = zeros(1, nNZ);
J = zeros(1, nNZ);
59 \quad j = 1;
60 for i = 1:n
61
       I(j:j+rowSize(i)-1) = i;
       J(j:j+rowSize(i)-1) = randsample(n, rowSize(i));
       j = j + rowSize(i);
64 end
66~\% restrict I and J to indices that correspond to entries above the main ...
      diagonal
67 % and finally construct a symmetric sparse matrix using I and J
68 upperTriu = find(I<J);
69 I = I(upperTriu);
70 J = J(upperTriu);
71 A = sparse([I;J], [J;I], ones(1, 2*size(I, 2)), n, n);
72 final=full(A);
73 end % random_graph(...)
```

## $step1\_scalefree.m$

```
2 %%% code adopted from %%%
 3 %%% Modeling and Simulating Social Systems with MATLAB %%%
 4 %%% http://www.soms.ethz.ch/teaching/MatlabFall2012 %%%
 5 %%% Authors: Stefan Brugger and Cristoph Schwirzer, 2011 %%%
 6 \quad \text{ as a second of the content 
 8 function final = step1_scalefree(n, m0, m)
 9 % Use the Barabasi-Albert model to generate a scale free graph of size n (as
10 % described in Albert-Laszlo Barabasi & Reka Albert: "Emergence of scaling
11 % in random networks")
12 %
13 % INPUT
14 % n: [1]: number of nodes
15 % m0: [1]: number of initially placed nodes
     % m: [1]: number of nodes a new added node is connected to, 1 \le m < m0
18 % OUPUT
19 % final: [n n] full symmetric adjacency matrix representing the ...
              generated graph
21\, % Start with a graph of size m0 and add edges to this graph. Each of ...
22 % nodes is connected to at least m nodes.
B = zeros(m0, m0);
24 \text{ for } i = 1:m0
              neighbors = randsample(m0-1, m);
              neighbors = neighbors + (neighbors≥i);
              B(i, neighbors) = 1;
              B(neighbors, i) = 1;
28
29 end
30
31 % Create a vector of edges added so far, i.e. nodes edge(2*i) and ...
              edge(2*i-1),
32 % 1 \le i \le nEdges, are connected by an edge.
33 [rows, columns] = find(triu(B));
34 nEdges = size(rows, 1);
35 edges = reshape([rows';columns'], 2*nEdges, 1);
36 edges = [edges; zeros(2*(n-m0)*m,1)];
38 % Add nodes m0+1:n, one at a time. Each node is connected to m existing ...
39 % where each of the existing nodes is chosen with a probability that is
40 % proportional to the number of nodes it is already connected to.
41 used = zeros(n, 1); % is a node already used in a timestep?
42 for i = m0+1:n
              neighbors = zeros(1, m);
```

```
for j=1:m
44
          k = edges(randi(2*nEdges));
45
          while used(k)
46
              k = edges(randi(2*nEdges));
47
          used(k) = 1;
49
50
          neighbors(j) = k;
       end
51
       used(neighbors) = 0;
52
       edges(2*nEdges+1:2*nEdges+2*m) = reshape([repmat(i, 1, m); ...
53
           neighbors], ...
        1, 2*m);
       nEdges = nEdges + m;
55
56 end
57
58 % finally construct a symmetric adjacency matrix using the vector of edges
59 edges = edges(1:2*nEdges);
60 first = edges(1:2:end);
61 second = edges(2:2:end);
62 A = sparse([first; second], [second; first], ones(2*nEdges, 1), n, n);
63 final=full(A);
64 end % scale_free(...)
```

#### $step1\_smallworld.m$

```
2 %%% code adopted from %%%
3 %%% Modeling and Simulating Social Systems with MATLAB %%%
4 %%% http://www.soms.ethz.ch/teaching/MatlabFall2012 %%%
5 %%% Authors: Stefan Brugger and Cristoph Schwirzer, 2011 %%%
8 function final = step1_smallworld(n, k, beta)
9 % Generate a small world graph using the "Watts and Strogatz model" as
10 % described in Watts, D.J.; Strogatz, S.H.: "Collective dynamics of
11 % 'small-world' networks."
12 % A graph with n*k/2 edges is constructed, i.e. the nodal degree is n*k for
13 % every node.
14 %
15 % INPUT
16 % n: [1]: number of nodes of the graph to be generated
17 % k: [1]: mean degree (assumed to be an even integer)
18 % beta: [1]: rewiring probability
19 %
20 % OUPUT
21 % A: [n n] sparse symmetric adjacency matrix representing the generated \dots
     graph
```

```
23 % Construct a regular lattice: a graph with n nodes, each connected to k
24 % neighbors, k/2 on each side.
25 kHalf = k/2;
26 rows = reshape(repmat((1:n)', 1, k), n*k, 1);
27 columns = rows+reshape(repmat([(1:kHalf) (n-kHalf:n-1)], n, 1), n*k, 1);
28 columns = mod(columns-1, n) + 1;
29 B = sparse(rows, columns, ones(n*k, 1));
30 A = sparse([], [], [], n, n);
32 % With probability beta rewire an edge avoiding loops and link duplication.
33 % Until step i, only the columns 1:i are generated making implicit use ...
      of A's
34 % symmetry.
35 \text{ for i} = 1:n
       \mbox{\%} The i-th column is stored full for fast access inside the \dots
           following loop.
       col=[full(A(i, 1:i-1))'; full(B(i:end, i))];
37
38
       for j = i+find(col(i+1:end))'
39
           if (rand()<beta)</pre>
40
               col(j)=0;
               k = randi(n);
41
               while k==i \mid \mid col(k)==1
42
                   k = randi(n);
               end
               col(k) = 1;
           end
       end
47
       A(:,i) = col;
48
49 end
50
51 % A is not yet symmetric: to speed things up, an edge connecting i and ...
       j, i < j
52 % implies A(i,j) ==1, A(i,j) might be zero.
T = triu(A);
54 A = T+T';
55 final=sparse(A);
57 end % small_world(...)
```

## A.3 Step 2: Rewiring process

### step2.m

```
1 function [mat, vec, dominant_freq, most_freq] = ...
      step2 (t_end, phi, alpha, mat, vec, p, threshold)
3 %%% t_end: number of iterations
4 %%% phi: network reorganization rate
5 %%% alpha: innovation rate
6 %%% mat: initial connectivity matrix
7 %%% vec: initial idea vector
8 %% p: initial number of opinions
9 %%% then it outputs:
10 %%% mat: connectivity matrix after simulation
11 %%% vec: idea vector after simulation
12 %%% dominant_ferq: the vector holding the frequency of dominant idea
13 %%% most_freq: the vector holding the index of dominating idea in each time
15 most_freq=zeros(1,t_end); %%% Vector for storing the index of the ...
      dominating idea in each time step.
16 dominant_freq=zeros(1,t_end); %%% Vector for storing the frequency of ...
      dominant idea in each time step.
17 a=size(mat);
n=a(1); %%% number of agents
20 for t=1:t_end
      x1=ceil(rand()*n); %%% choosing one person randomly for network ...
          reorganization or changing idea
      a1=rand();
      b1=phi;
      if al <b1 %%% i.e with probability phi to reorganize the network
         v00=find(mat(x1,:)==1); %%% defining neighbours of x1
25
         v=find(vec(v00)\neq vec(x1)); %%% define neighbours of x1 that do not ...
26
             have the same idea as x1
         if ¬isempty(v)
27
            x2=v (ceil(rand()*length(v))); %%% choosing one neighbour with ...
                different idea randomly to remove connection with
            mat(x1,x2)=0; %%% deletion of the edge between x1 and x2
            mat(x2,x1)=0; %% deletion of the edge between x2 and x1
30
            similar_idea=find(vec==vec(x1)); %%% define the agents with ...
31
                the same idea as x1
            non_neighbor_sim_idea=setdiff(similar_idea,v00); %%% the ...
                agents with similar idea as x1 who are not neighbor of x1
            if ¬isempty(non_neighbor_sim_idea)
                x3=non_neighbor_sim_idea(ceil(rand()*length(non_neighbor_sim_idea))); ...
34
                    %%% choose x3 randomly among the agents with the same ...
                    idea and non-neighbor with x1 as the newly connected ...
                    agent to x
```

```
mat(x1,x3)=1; %%% formation of new edge between x1 and x3
35
                 mat(x3,x1)=1; %%% formation of new edge between x3 and x1
36
             else %%% we'll forget about network reorganization in this ...
37
                 time by reforming the deleted edges.
                 mat(x1, x2) = 1;
                 mat(x2, x1) = 1;
39
40
             end
          end
41
42
       else %%% otherwise change the idea of x1 to that of one of it's ...
43
           randomly chosen neighbours
          v2=find(mat(x1,:)==1); %%% defining the neighbours of x1
45
          if ¬isempty(v2)
             vv=vec(v2); %%% the corresponding ideas of the neighbours
46
             vvv=unique(vv); %%% vector of all the distinct ideas
47
             freq=zeros(1,length(vvv)); %%% vector for frequencies of the ideas
48
             for i=1:length(vvv) %%% to test for all distinct ideas
49
                if length(find(vv==vvv(i)))>(threshold*length(v2)) ...
                    %%%whether the frequency is larger than the threshold ...
                    [To include complex contagion definition]
                    freq(i)=1;
51
                end
52
             end
53
             candidates=vvv(find(freq==1)); %%% The ideas meeting the threshold
             candidates_size=length(candidates); %%% The number of ideas ...
                 meeting the threshold
             if candidates_size>0
56
                chosen=ceil(rand()*candidates_size);%%% Randomly choose one ...
57
                    of the candidates
                vec(x1)=candidates(chosen); %%% change the idea of x1 to ...
58
                    the chosen idea
59
             end
60
          end
61
       end
62
       y=ceil(rand()*n); %%%choosing one person randomly for coming up with ...
63
           a new idea
64
       a2=rand();
       b2=alpha;
       if a2<b2 %%% i.e. with probability alpha to generate a novel idea
           bound=10^6; %%% to limit the index of new ideas to 10^6 which ...
67
               simulates nearly boundryless pool of ideas
           new_idea=ceil(rand()*bound)+p; %%% the index of new idea
68
           vec(y)=new_idea; %%% changing the idea of agent y to the novel one
69
       end
70
       응응응응
71
       most_freq(t)=mode(vec); %%% the dominant idea at time t
72
       dominant_freq(t) = length(find(vec==most_freq(t))); %%% the frequency ...
73
           of the dominant idea at time t
74 end
```

#### A.4 Step 3: Results

#### step3a.m

```
1 function n_index = step3a(mat, vec)
_2 %%%%%% step3a: defining the neighbourhood index between similar ideas ...
       응응응응응응
3 %%% mat: adjacency matrix
4 %%% vec: idea vector
5 %%% it outputs: n_index which is the average neighbour index of the network
6 u=unique(vec); %%% collection of distinct ideas
7 w=zeros(1, length(u)); %%% initializing the vector for storing the ...
       neighbourhood index for each distinct idea
s \, s=zeros(1,length(u)); %%% initializing the vector for storing the number \dots
      of agents holding each distinct ideas
9 for i=1:length(u)
       x=find(vec=u(i)); %%% defining the set of agents holding the idea u(i)
11
       s(i) = length(x);
12
       if s(i) > 1
       sum1=0; %%% number of agents with idea i which are in direct ...
13
           neighbourhood
       for k=1:(s(i)-1) %%% These two for loops are used to test the ...
14
           neighborhoods of all distinct pairs
           for j = (k+1) : s(i)
15
               if mat(x(k), x(j)) == 1 \%\% to check if they are neighbours
16
                    sum1=sum1+1;
17
               end
18
           end
19
20
       end
21
       \max_{i=1}^{n} (s(i) * (s(i)-1)/2); %%% normalizing factor (i.e., the ...
           maximum number of pairs of distinct agents)
       w(i)=sum1/max_neighbors; %%% neighborhood index for idea 'i'
22
       else
23
           w(i) = 0;
24
       end
25
26 end
27 norml=sum(s); %%% Normalizing factor
28 count=sum(w.*s); %%% The weighted sum of neighborhood index
29 n_index=count/norml; %%% average neighbour index of the whole network
30 end
```

#### steb3b.m

```
1 function intra_idea_distance = step3b(mat, vec, s, c)
2 %%%%%%%%%% step3b: Defining the average of the average shortest ...
      3 %%% mat: adjacency matrix
4 %%% vec: idea vector
5 %% s: number of connected components
6 %%% c: vector which assigns each node to a connected component
7 %%% and it outputs the average of average intra_distance between agents
9 ung=unique(vec); %%% collection of distinct ideas
10 mean_sum=zeros(1,length(ung)); %%% initializing the vector for storing ...
      the average distance between agents holding a particular idea
11 n=zeros(1,length(unq)); %%% initializing the corresponding vector for ...
      storing the number of agents holding a certain idea
12 mats=sparse(mat);
13
  for i=1:length(ung)
      x=find(vec==unq(i)); %%% defining the set of agents holding the ...
          idea unq(i)
      n(i) = length(x); %%% number of agents holding the idea unq(i)
16
17
      nn=zeros(1,s); %%% initializing the vector which will store the ...
          number of agents holding the idea unq(i) who belong to the m-th ...
          connected component
18
      mean_sum0=zeros(1,s); %%% initializing the vector which stores the ...
          mean of the distances between agents holding the idea unq(i) who ...
          belong to each distinct connected component
19
      for m=1:s %%% for each connected component
          y=x(find(c(x)==m)); %% the set of agents holding the idea ...
              ung(i) who belong to the m-th connected component
          nn(m) = length(y); %%% number of agents holding the idea ung(i) ...
21
              who belong to the m-th connected component
          sum0=0;
22
          if nn(m) > 1
23
             for j=1: (nn(m)-1)
24
25
                 for k=(j+1):nn(m)
                     dis=graphshortestpath(mats,y(j),y(k)); %%% distance ...
                        of each agent with similar idea from agent j ...
                        inside the group [needs the bioinformatics ...
                        toolbox to be installed]
                     sum0=sum0+dis; %%% sum of the distances between ...
27
                        agents holding the idea unq(i) who belong to m-th ...
                        connected component
28
                 end
29
             norm10=nn(m)*(nn(m)-1)/2; %%% normalizing factor
30
             mean_sum0(m)=sum0/norm10; %%% The avarage intra_idea distance ...
31
                 between agents holding the idea unq(i) who belong to the ...
```

```
m-th connected component
           else
32
              mean_sum0(m)=0;
33
34
           end
       end
36
37
       if n(i) > 1
          mean\_sum(i) = (sum(mean\_sum0.*nn))/(sum(nn)); %%% The avarage ...
38
              intra_idea distance between agents holding the idea unq(i)
       else
39
          mean_sum(i)=0;
       end
42 end
43
44 intra_idea_distance=(sum(mean_sum.*n))/(sum(n)); %%% The avarage ...
       intra_idea distance for the whole network
45 end
```

#### step3e.m

```
1 function [ average_dominance_time ] = step3e( most_freq )
2 %%%% this function calculates the average of dominance time for the ...
      dominating idea during the simulation
_3 %%%% most_freq: is the vector obtained from simulation which holds the ...
      index of dominating ideas for each time steps of the simulation
4 %%%% it outputs the average_time which is the average of the dominance ...
      time for different dominance periods
5 temp=zeros(1,length(most_freq)); %%% a temporary array which will hold ...
      the dominating period for dominating idea
6 count=1; %%% is the number of consecutive time steps in which an idea is \dots
      considered as dominating
7 ind=0; %%% is the index of domination period
  for i=2:length(most_freq)
       if most_freq(i) == most_freq(i-1)
           count=count+1; %%% the number of consecutive steps is conted
10
11
           ind=ind+1; %%% as soon as another domination period gets tarted ...
12
               the index of domination period adds by one
           temp(ind)=count; %%% the number of consecutive steps will be ...
               stored in the ind-th index of tmp
           count=0; %%% and count will be reset to zero to count the ...
               duration of the new domination period.
       end
15
16 end
17
18 if ind==0 %%% in case during the simulation, just one special idea ...
      remains dominating forever
```

```
ind=ind+1;
temp(ind)=count;

end

end

make ind=ind+1;
temp(ind)=count;

end

make ind=ind+1;

temp(ind)=count;

end

make ind=ind+1;

ma
```

#### step4a.m

```
2 function clust_coeff = step4a( mat )
3 % This function calculates the clustering coefficient of a network with
4 % Corresponding connectivity matrix: mat
5 % The approach used for calculation of clustering coefficient was the one ...
      which was proposed by Watts and Strogatz
_{6} % in which the clustering coefficent of the whole network equals to the
7 % average of the local clustering coeefient of all nodes:
8 % D. J. Watts and Steven Strogatz (June 1998). "Collective dynamics of ...
       'small-world' networks". Nature 393 (6684): 440 442.
9 a=size(mat);
10 local_clust_coeff=zeros(1,a(1)); %%% the vector to store the local ...
       clustering coefficent of each node of the network
11 for i=1:a(1) %%% for each agent in the network
       x=find(mat(i,:)==1); %%% find all the neighbours of agent i
       count=0;
13
       l=length(x);
       if 1>1
       for j=1:(1-1) %%% loop to investigate the neighbour relationship for ...
          all possible pairs among the neighbors of node i
           for k = (j+1):1
17
               if mat(x(j), x(k)) == 1
                   count=count+1; % to count the number of neighbour ...
                       relationships
21
           end
       end
22
       norml=1*(l-1)/2; %% normalizing factor (i.e. the number of distinct ...
23
       local_clust_coeff(i) = count/norml; %% local clustering coefficient of ...
24
          node i
```

```
25    else
26    local_clust_coeff(i)=0;
27    end
28   end
29    clust_coeff=sum(local_clust_coeff)/a(1); %% the average clustering ...
        coefficient of entire network
30   end
```

#### step4b.m

```
1 function [dgr,frq] = step4b(mat)
2 %% this function outputs the degree vector (dgr) and its corresponding
_3 %%% frequency vector (frq) from the connectivity matrix (mat). In other ...
      words,
4 %%% first the degree of all nodes in the network will be calculated and
5 %%% then the set of unique degrees will be stored in the dgr vector and the
6 %%% corresponding frequency will be stored in vector frq. For example,
7 %%% degree x will be stored in the i-th element of the dgr vector, then \dots
      the number of nodes whose degree equals x
8 %% will be calculated and will be stored in the i-th element of vector frq.
9 a=size(mat);
10 degr=zeros(1,a(1)); %%% the vector which stores the degree of each node ...
      of the network
11 for i=1:a(1)
       degr(i) = sum (mat(i,:)); %%% degree for each node
12
13 end
14 dgr=unique(degr); %%% set of unique degrees of the nodes of the network
15 frq=zeros(1,length(dgr)); %%% the corresponding frequency vector
16 for j=1:length(dgr)
       frq(j) = length(find(degr = -dgr(j))); %%% frq(j) is the number of nodes ...
           in the network whose degree equals to dgr(j)
18 end
20 end
```

# step4d.m

```
1 function average_path_length = step4d( mat,s,c )
2 %This function outputs the average path length for the graph
3 %mat: connectivity matrix
4 %s: number of connected components
5 %c: the connected component vector
6 av_pl=zeros(1,s); %%% the vector which will store the average path ...
       length for each connected component of the 'mat' network
7 n=zeros(1,s); %% the vector which will store the number of nodes ...
      belonging to each connected component.
8 sp=sparse(mat);
10 for i=1:s %%% for each connected component
       x=find(c==i); %%% charachterizing the nodes which belong to the i-th ...
11
           connected component
       a=length(x); %%% the number of nodes in the i-th connected component
13
       n(i)=a;
14
       count=0;
       if a>1
15
       for k=1:(a-1)
16
           for j=(k+1):a
17
           count = count + graph shortest path (sp, x(k), x(j)); %% sum of the path ...
               length between all pairs of node
19
           end
20
       end
       norml=a*(a-1)/2; %%% normalization factor
21
       av_p_l(i)=count/norml; %%% average of path length
22
23
       else
24
       av_p_1(i) = 0;
25
       end
26 end
27 average_path_length=sum(av_p_l.*n)/sum(n); %%% weighted average path length
```

## step4e.m

```
1 function graph_diameter = step4e( mat,s,c )
2 %This function outputs the diameter of the graph 'mat'
3 %mat: connectivity matrix
4 %s: number of connected components
5 %c: the connected component vector
6 sp=sparse(mat);
7 max_path=zeros(1,s); %%% vector storing diameter for each connected ...
      components.
8 for i=1:s %%% for each connected component
       x=find(c==i); %%% charachterizing the nodes which belong to the i-th ...
          connected component
       a=length(x); %%% the number of nodes in the i-th connected component
10
       path_length=zeros(1,(a*(a-1)/2)); %%% The vector storing the ...
11
           shortest path between each pairs of nodes belonging to the ...
          connected component i
12
       count=0;
13
       if a>1
       for k=1:(a-1)
14
           for j=(k+1):a
15
               count=count+1;
16
17
               path_length(count) = graphshortestpath(sp, x(k), x(j)); %%% the ...
                   vector storing the shortest path between pairs k and j
18
           end
19
       end
       max_path(i)=max(path_length); %% the maximum shortest path for ...
20
          connected component i
21
       else
22
       max_path(i)=0;
23
       end
24 end
25 graph_diameter=max(max_path); %% the maximum shortest path for the hole ...
      network (Diameter of the network)
26 end
```

#### A.5 Plote codes

#### plots\_phase1.m

```
3 %%% Extracting information from the saved Mat files %%%%%%%
4 neighbor_ind=zeros(1,108);
5 int_id_dis=zeros(1,108);
6 nov_ind=zeros(1,108);
7 av_dom_tim=zeros(1,108);
8 dom_freq=zeros(108,1000);
9 alpha=[0.01,0.05,0.1];
10 phi=[0.1,0.3,0.5];
11 threshold=[0.001,0.01,0.05];
12 count=0;
13
   f=figure();
14 for i=1:4
15
      switch i
16
          case 1
          s1='Caveman';
17
          case 2
18
          s1='Random';
19
          case 3
          s1='Scale_free';
          case 4
          s1='Small_world';
23
24
      end
      for j=1:3
25
          s2=int2str(j);
26
          for k=1:3
27
              s3=int2str(k);
28
              for 1=1:3
29
                  count=count+1;
30
                  s4=int2str(1);
31
                  name=['phase1_',s1,'_',s2,'_',s3,'_',s4];
32
                  a=load(name);
                  neighbor_ind(count) = a.neighbor_index;
                  int_id_dis(count) = a.intra_idea_distance;
36
                  nov_ind(count) = a.nov_index;
                  av_dom_tim(count) = a.average_dominance_time;
37
                  dom_freq(count,:) = a.dominant_freq;
38
              end
39
40
          end
41
      end
42 end
43
44 %%% phase diagrams for each parameter pairs for the results of the effects
45 %% of network structure on idea distribution.
```

```
47 for i=1:4
       switch i
48
           case 1
49
           s1='Caveman';
50
           val=0;
51
52
           case 2
           s1='Random';
53
           val=27;
54
           case 3
55
           s1='Scale free';
           val=54;
           case 4
59
           s1='Small world';
60
           val=81;
       end
61
       for j=1:4
62
63
           switch j
64
           case 1
65
           s2='neighbor index';
           main_vec=neighbor_ind;
66
           case 2
67
           s2='intra idea distance';
68
           main_vec=int_id_dis;
           case 3
           s2='nov index';
           main_vec=nov_ind;
           case 4
73
           s2='average dominance time';
74
           main_vec=av_dom_tim;
75
           end
76
           for k=1:3
77
                 switch k
78
                 case 1
79
                 s3='(alpha:phi)';
80
                 x=0:0.5:1;
81
                 xl='alpha';
82
                 y=0:0.5:1;
                 yl='phi';
                 for 1=1:3
                     M=[main_vec(val+1), main_vec(val+1+9), main_vec(val+1+18); main_vec(val+1+3), main_vec(val+1+3)
86
87
                     [xlab,ylab]=meshgrid(x,y);
88
                     hold on;
89
90
                     view(0,90);
91
                     surf(xlab, ylab, M, 'EdgeColor', 'none');
92
                     colorbar;
93
                     set(gca, 'FontSize', 14)
94
                     xlabel(xl);
95
                     ylabel(yl);
```

```
switch 1
96
                           case 1
97
                                s5='first';
98
                            case 2
99
100
                                s5='second';
101
                           case 3
102
                                s5='third';
103
                       end
104
                       name2=['phase diagram of ',s2,' for alpha versus phi',' ...
                           and', 'the', s5,' threshold', 'obtained ...
                           for',s1,'network structure'];
105
                       title(name2);
                       name=['plotting1_','Pdiag_',s1,'_',s2,'_',s3,'_',s4,'AND','the', ...
106
                           s5, 'threshold'];
107
                       saveas(f, name);
                       hold off
108
109
110
                  end
111
                  case 2
112
                  s3='(alpha:threshold)';
                  x=0:0.5:1;
113
                  xl='alpha';
1114
                  y=0:0.5:1;
1115
116
                  yl='threshold';
117
                  for 1=1:3
                       M = [main\_vec(val+9*(l-1)+1), main\_vec(val+9*(l-1)+2), main\_vec(val+9*(l-1)+3); main\_vec(val+9*(l-1)+3)]
118
119
120
                       [xlab,ylab]=meshgrid(x,y);
                       hold on;
121
                       view(0,90);
122
                       surf(xlab,ylab,M,'EdgeColor','none');
123
124
                       colorbar;
125
                       set(gca, 'FontSize', 14)
                       xlabel(xl);
126
127
                       ylabel(yl);
                        switch 1
128
129
                           case 1
130
                                s5='first';
131
                           case 2
132
                                s5='second';
133
                           case 3
134
                                s5='third';
135
                       end
                       name2=['phase diagram of ',s2,' for alpha versus ...
136
                           threshold', 'and', 'the', s5, 'phi', 'obtained ...
                           for', s1, 'network structure'];
                       title(name2);
137
138
                       name=['plotting1_','Pdiag_',s1,'_',s2,'_',s3,'_',s4,' ...
                           and', 'the', s5, 'phi'];
                       saveas(f, name);
139
```

```
hold off
140
141
142
                 end
143
144
                 case 3
145
                 s3='(phi:threshold)';
146
                 x=0:0.5:1;
147
                 xl='phi';
148
                 y=0:0.5:1;
                 yl='threshold';
149
150
                 for 1=1:3
151
                     M = [main\_vec(val+3*(1-1)+1), main\_vec(val+3*(1-1)+2), main\_vec(val+3*(1-1)+3); main\_vec(val+3*(1-1)+3)]
152
153
                     [xlab,ylab]=meshgrid(x,y);
154
                     hold on;
                     view(0,90);
155
                     surf(xlab,ylab,M,'EdgeColor','none');
156
157
                     colorbar;
158
                     set(gca, 'FontSize', 14)
159
                     xlabel(xl);
160
                     ylabel(yl);
161
                      switch 1
162
                         case 1
163
                             s5='first';
164
                         case 2
165
                             s5='second';
166
                         case 3
                             s5='third';
167
168
                     end
                     name2=['phase diagram of ',s2,' for phi versus ...
169
                         threshold',' and','the', s5,' threshold','obtained ...
                         for',s1,'network structure'];
                     title(name2);
170
                     name=['plotting1_','Pdiag_',s1,'_',s2,'_',s3,'_',s4,' ...
171
                         and', 'the', s5,' threshold'];
                     saveas(f, name);
172
                     hold off
173
174
175
                 end
176
                 end
177
            end
178
        end
179 end
180
182 %% Dependence of the property of each network to the parameters
183
184 for i=1:4
        switch i
185
            case 1
186
```

```
187
            s1='Caveman';
188
            val=0;
189
            case 2
            s1='Random';
190
191
            val=27;
192
             case 3
193
            s1='Scale free';
194
            val=54;
195
            case 4
            s1='Small world';
196
197
            val=81;
198
        end
199
        for j=1:4
200
            switch j
201
            case 1
            s2='neighborhood index';
202
            main_vec=neighbor_ind;
203
204
            case 2
205
            s2='intra idea distance';
206
            main_vec=int_id_dis;
207
            case 3
            s2='novelity index';
208
            main_vec=nov_ind;
209
210
            case 4
211
            s2='average dominance time';
212
            main_vec=av_dom_tim;
213
            end
214
            for k=1:3
215
                 switch k
216
                     case 1
                          s3='threshold';
217
218
                          f=figure();
219
                          for 1=1:3
                              line=main_vec(3*(0:8)+val+1);
220
221
                              set(gca, 'FontSize', 14)
222
                              xlabel('Different pairs of alpha and phi values');
                              ylabel(s2);
223
                              namek=['Dependence of',' ',s2,' ','on',' ',s3,' ...
224
                                  ','for',' ',s1,' ','network structure'];
225
                              title(namek);
226
                              switch 1
227
228
                                   case 1
                                        plot(line, '-ks', 'LineWidth', 2, 'MarkerEdgeColor', 'k', 'MarkerE
229
230
231
                                   case 2
                                        plot(line,'-bs','LineWidth',2,'MarkerEdgeColor','b','MarkerF
232
233
                                        hold on
234
                                   case 3
                                        plot(line, '--rs', 'LineWidth', 2, 'MarkerEdgeColor', 'r', 'MarkerE
235
```

```
236
237
                              legend('threshold=0.001', 'threshold=0.01', 'threshold=0.05')
                              name=['plotting1_',s1,'_',s2,'_','threshold_sensitivity'];
238
                              saveas(f, name);
239
240
                          end
241
242
                     case 2
243
                         s3='alpha';
244
                         f=figure();
                         for 1=1:3
245
246
                              line=main_vec([1,2,3,10,11,12,19,20,21]+val+3*([1-1));
247
                              set (gca, 'FontSize', 14)
248
                              xlabel('Different pairs of threshold and phi ...
                                  values');
                              ylabel(s2);
249
                              namek=['Dependence of',' ',s2,' ','on',' ',s3,' ...
250
                                  ','for',' ',s1,' ','network structure'];
251
                              title(namek);
252
253
                              switch 1
254
                                  case 1
                                        plot(line,'-ks','LineWidth',2,'MarkerEdgeColor','k','MarkerF
255
                                        hold on
256
257
258
                                        plot(line,'--bs','LineWidth',2,'MarkerEdgeColor','b','MarkerF
259
                                  case 3
260
                                        plot(line, '-rs', 'LineWidth', 2, 'MarkerEdgeColor', 'r', 'MarkerE
261
262
                              end
                              legend('alpha=0.01', 'alpha=0.05', 'alpha=0.10')
263
                              name=['plotting1_',s1,'_',s2,'_','alpha_sensitivity'];
264
265
                              saveas(f, name);
                         end
266
267
                     case 3
268
                         s3='phi';
269
                         f=figure();
270
                         for 1=1:3
271
272
                              line=main_vec((1:9)+val+9*(1-1));
                              set (gca, 'FontSize', 14)
273
                              xlabel('Different pairs of threshold and alpha ...
274
                                  values');
                              ylabel(s2);
275
                               namek=['Dependence of',' ',s2,' ','on',' ',s3,' ...
276
                                   ','for',' ',s1,' ','network structure'];
                              title(namek);
277
278
                              switch 1
279
280
                                  case 1
                                        plot(line,'-ks','LineWidth',2,'MarkerEdgeColor','k','MarkerF
281
```

```
282
                                  hold on
283
                             case 2
                                  plot(line, '--bs', 'LineWidth', 2, 'MarkerEdgeColor', 'b', 'MarkerF
284
285
                                  hold on
286
                             case 3
                                  plot(line, '-rs', 'LineWidth', 2, 'MarkerEdgeColor', 'r', 'MarkerF
287
288
                         legend('phi=0.1','phi=0.3','phi=0.5')
289
                         name=['plotting1_',s1,'_',s2,'_','phi_sensitivity'];
290
                         saveas(f, name);
291
292
                      end
293
              end
294
          end
       end
295
   end
296
297
   298
   299
300
   for i=1:4
301
       switch i
302
          case 1
              f=figure();
303
              line1=neighbor_ind(1:27);
304
305
              line2=neighbor_ind(28:54);
306
              line3=neighbor_ind(55:81);
307
              line4=neighbor_ind(82:108);
              308
309
              hold on
              plot(line2,'--gs','LineWidth',2,'MarkerEdgeColor','g','MarkerFaceColor','g','Marker
310
311
              hold on
              plot(line3,'-bs','LineWidth',2,'MarkerEdgeColor','b','MarkerFaceColor','b','Marke
312
313
              hold on
              plot(line4,'-rs','LineWidth',2,'MarkerEdgeColor','r','MarkerFaceColor','r','Marker
314
              set(gca, 'FontSize', 14)
315
              xlabel('Different triplets of alpha ,phi and threshold values')
316
              ylabel('Neighborhood index')
317
              title('The influence of network structure on neighborhood ...
318
                  index');
319
              legend('Caveman Network', 'Random Network', 'Scale Free ...
                  Network','Small World Network')
              name=('Plotting1_Network Comparison_Neighborhood_index');
320
              saveas(f, name);
321
          case 2
322
323
              f=figure();
              line1=int_id_dis(1:27);
324
              line2=int_id_dis(28:54);
325
              line3=int_id_dis(55:81);
326
              line4=int_id_dis(82:108);
327
              plot(line1,'-ks','LineWidth',2,'MarkerEdgeColor','k','MarkerFaceColor','k','Marker
328
329
              hold on
```

```
plot(line2,'--gs','LineWidth',2,'MarkerEdgeColor','g','MarkerFaceColor','g','Marker
330
331
               hold on
               plot(line3,'-bs','LineWidth',2,'MarkerEdgeColor','b','MarkerFaceColor','b','Marker
332
333
               hold on
               plot(line4,'--rs','LineWidth',2,'MarkerEdgeColor','r','MarkerFaceColor','r','Marker
334
335
               set(gca, 'FontSize', 14)
336
               xlabel('Different triplets of alpha ,phi and threshold values')
               ylabel('Intra-idea distance')
337
               title('The influence of network structure on intra-idea ...
338
                   distance')
               legend('Caveman Network', 'Random Network', 'Scale Free ...
339
                  Network', 'Small World Network')
               name=('Plotting1_Network Comparison_intra_idea_distance');
340
341
               saveas(f, name);
342
           case 3
               f=figure();
343
344
               line1=nov_ind(1:27);
345
               line2=nov_ind(28:54);
346
               line3=nov_ind(55:81);
347
               line4=nov_ind(82:108);
348
               plot(line1,'-ks','LineWidth',2,'MarkerEdgeColor','k','MarkerFaceColor','k','Marker
349
               hold on
               plot(line2,'--gs','LineWidth',2,'MarkerEdgeColor','g','MarkerFaceColor','g','Marker
350
351
               hold on
352
               plot(line3,'--bs','LineWidth',2,'MarkerEdgeColor','b','MarkerFaceColor','b','Marker
353
               plot(line4,'-rs','LineWidth',2,'MarkerEdgeColor','r','MarkerFaceColor','r','Marker
354
               set(gca, 'FontSize', 14)
355
               xlabel('Different triplets of alpha ,phi and threshold values')
356
357
               ylabel('Novelity Index')
               title('The influence of network structure on Novelity index')
358
359
               legend('Caveman Network', 'Random Network', 'Scale Free ...
                   Network','Small World Network')
               name=('Plotting1_Network Comparison_intra_novelity_index');
360
361
               saveas(f, name);
362
           case 4
               f=figure();
363
364
               line1=av_dom_tim(1:27);
365
               line2=av_dom_tim(28:54);
366
               line3=av_dom_tim(55:81);
367
               line4=av_dom_tim(82:108);
               plot(line1,'-ks','LineWidth',2,'MarkerEdgeColor','k','MarkerFaceColor','k','Marker
368
369
               hold on
               370
371
               hold on
               372
373
               hold on
               plot(line4,'--rs','LineWidth',2,'MarkerEdgeColor','r','MarkerFaceColor','r','Marker
374
               set(gca, 'FontSize', 14)
375
               xlabel('Different triplets of alpha ,phi and threshold values')
376
```

```
377
              ylabel('Average Dominance Time')
              title('The influence of network structure on Average \dots
378
                  Dominance Time')
              legend('Caveman Network', 'Random Network', 'Scale Free ...
379
                  Network','Small World Network')
              name=('Plotting1_Network Comparison_Average Dominance Time');
380
381
              saveas(f, name);
382
       end
383 end
384
385
  count=0;
389
   for choice1=1:3
390
       switch choice1
          case 1
391
              s1='phi=0.1';
392
393
              m1 = '1';
394
           case 2
              s1='phi=0.3';
395
396
              m1='2';
397
           case 3
398
              s1='phi=0.5';
399
              m1 = '3';
400
       end
401
       for choice2=1:3
402
           switch choice2
403
404
              case 1
                  s2='alpha=0.01';
405
406
                  m2 = '1';
407
              case 2
                  s2='alpha=0.05';
408
409
                  m2 = '2';
410
              case 3
                  s2='alpha=0.1';
411
412
                  m2 = '3';
413
           end
414
           for choice3=1:3
415
              switch choice3
416
                  case 1
417
                      s3='threshold=0.001';
418
                      m3 = '1';
419
420
                  case 2
421
                      s3='threshold=0.01';
422
                      m3='2';
423
                  case 3
                      s3='threshold=0.05';
424
```

```
425
                         m3 = '3';
                end
426
427
                count=count+1;
                line1=dom_freq(count,:);
428
429
                line2=dom_freq(27+count,:);
                line3=dom_freq(54+count,:);
430
431
                line4=dom_freq(81+count,:);
432
                f=figure();
                plot(line1,'-ks','LineWidth',2,'MarkerEdgeColor','k','MarkerFaceColor','k','Marker
433
                hold on
434
435
                plot(line2,'-gs','LineWidth',2,'MarkerEdgeColor','g','MarkerFaceColor','g','Marker
436
                hold on
437
                plot(line3,'-bs','LineWidth',2,'MarkerEdgeColor','b','MarkerFaceColor','b','Marker
                hold on
438
                plot(line4,'--rs','LineWidth',2,'MarkerEdgeColor','r','MarkerFaceColor','r','Marker
439
440
                set(gca, 'FontSize', 14)
                xlabel('time')
441
442
                ylabel('Frequency of the dominant idea')
443
                title(['The influence of network structure on the frequency ...
                    of the dominant idea',' ',s1,' ',s2,' ',s3]);
                legend('Caveman Network', 'Random Network', 'Scale Free ...
444
                    Network','Small World Network')
                name=(['Plotting1_Network ...
445
                    Comparison_frequency_dominant_idea-', m1, '_', m2, '_', m3]);
446
                saveas(f, name);
447
            end
        end
448
449 end
```

## plots\_phase2.m

```
2 응응응
3 %%% Extracting information from the saved Mat files %%%%%%
5 clust_coefficient=zeros(1,81);
6 \text{ s=zeros}(1,81);
7 average_path_length=zeros(1,81);
8 diam=zeros(1,81);
9 dgr=cell(1,81);
10 frq=cell(1,81);
11 alpha=[0.01,0.05,0.1];
12 phi=[0.1,0.3,0.5];
13 threshold=[0.001,0.01,0.05];
14
15
  count=0;
16
17
  for i=1:3
       switch i
18
          case 1
19
          s1='Random';
20
^{21}
          case 2
22
          s1='Parallel';
23
          case 3
          s1='Antiparallel';
24
      end
25
       for j=1:3
26
          s2=int2str(j);
27
          for k=1:3
29
              s3=int2str(k);
              for 1=1:3
30
                  count=count+1;
31
                  s4=int2str(1);
32
                  name=['phase2_',s1,'_',s2,'_',s3,'_',s4];
33
                  a=load(name);
34
                  clust_coefficient(count) = a.clust_coefficient;
                  s(count)=a.s;
                  average_path_length(count) = a.average_path_length;
37
                  diam(count) = a.diam;
38
                  dgr{1,count}=a.dgr;
39
                  frq{1,count}=a.frq;
40
41
              end
42
          end
43
       end
44 end
45
  f=figure();
46
47
```

```
48 %%% phase diagrams for each parameter pairs for the results of the effects
49 %%% of idea distribution on network structure.
51 for i=1:3
       switch i
52
           case 1
53
54
           s1='Random';
55
           val=0;
56
           case 2
            s1='Parallel';
57
           val=27;
           case 3
           s1='Antiparallel';
61
            val=54;
62
       end
       for j=1:4
63
64
            switch j
65
            case 1
            s2='Clusteing Coefficient';
66
67
            main_vec=clust_coefficient;
            case 2
68
            s2='Number of connected components';
69
            main_vec=s;
70
71
            case 3
72
            s2='Average path length';
           main_vec=average_path_length;
           s2='Network diameter';
75
           main_vec=diam;
76
            end
77
            for k=1:3
78
79
                 switch k
                 case 1
80
                 s3='(alpha:phi)';
81
82
                 x=0:0.5:1;
                 xl='alpha';
83
                 y=0:0.5:1;
84
                 yl='phi';
                 for 1=1:3
                     M = [main\_vec(val+1), main\_vec(val+1+9), main\_vec(val+1+18); main\_vec(val+1+3), main\_vec(val+1+3)]
88
                     [xlab, ylab] = meshgrid(x, y);
89
                     hold on;
90
                     view(0,90);
91
                      surf(xlab, ylab, M, 'EdgeColor', 'none');
92
93
                      colorbar;
                      set (gca, 'FontSize', 14)
94
95
                     xlabel(xl);
                     ylabel(yl);
96
97
                      switch 1
```

```
98
                           case 1
                                s5='first';
99
                           case 2
100
101
                                s5='second';
102
                           case 3
103
                                s5='third';
104
                       end
105
                       name2=['phase diagram of ',s2,' for alpha versus phi',' ...
                           and','the', s5,' threshold','obtained for',s1,'idea ...
                           distribution'];
106
                       title(name2);
107
                       name=['Plotting2_','Pdiag_',s1,'_',s2,'_',s3,'_',s4,'AND','the', ...
                           s5, 'threshold'];
108
                       saveas(f, name);
109
                       hold off
110
111
                  end
112
                  case 2
113
                  s3='(alpha:threshold)';
114
                  x=0:0.5:1;
                  xl='alpha';
115
                  y=0:0.5:1;
116
                  yl='threshold';
1117
118
                  for 1=1:3
119
                       M = [main\_vec(val+9*(1-1)+1), main\_vec(val+9*(1-1)+2), main\_vec(val+9*(1-1)+3); main\_vec(val+9*(1-1)+3)]
120
121
                       [xlab, ylab] = meshgrid(x, y);
122
                       hold on;
                       view(0,90);
123
                       surf(xlab,ylab,M,'EdgeColor','none');
124
                       colorbar;
125
126
                       set(gca, 'FontSize', 14)
127
                       xlabel(xl);
                       ylabel(yl);
128
129
                        switch 1
                           case 1
130
                                s5='first';
131
132
                           case 2
133
                                s5='second';
134
                           case 3
135
                                s5='third';
136
                       end
                       name2=['phase diagram of ',s2,' for alpha versus ...
137
                           threshold', 'and', 'the', s5, 'phi', 'obtained ...
                           for',s1,'idea distribution'];
                       title(name2);
138
                       name=['Plotting2_','Pdiag_',s1,'_',s2,'_',s3,'_',s4,' ...
139
                           and', 'the', s5, 'phi'];
                       saveas(f, name);
140
                       hold off
141
```

```
142
143
                   end
144
145
                  case 3
                  s3='(phi:threshold)';
146
147
                  x=0:0.5:1;
148
                  xl='phi';
149
                  y=0:0.5:1;
150
                  yl='threshold';
                   for 1=1:3
151
152
                       M = [main\_vec(val+3*(l-1)+1), main\_vec(val+3*(l-1)+2), main\_vec(val+3*(l-1)+3); main\_vec(val+3*(l-1)+3)]
153
154
                       [xlab,ylab]=meshgrid(x,y);
155
                       hold on;
156
                       view(0,90);
                       surf(xlab,ylab,M,'EdgeColor','none');
157
158
                       colorbar;
                       set(gca, 'FontSize', 14)
159
160
                       xlabel(xl);
161
                       ylabel(yl);
162
                        switch 1
163
                            case 1
164
                                s5='first';
165
                            case 2
166
                                s5='second';
167
                            case 3
168
                                s5='third';
169
                       end
                       name2=['phase diagram of ',s2,' for phi versus ...
170
                           threshold',' and','the', s5,' threshold','obtained ...
                            for',s1,'idea distribution'];
171
                       title(name2);
                       name=['Plotting2_','Pdiag_',s1,'_',s2,'_',s3,'_',s4,' ...
172
                           and', 'the', s5,' threshold'];
                       saveas(f, name);
173
                       hold off
174
175
176
                   end
177
                   end
178
             end
179
        end
180 end
181
   %%% Dependence of the property of each network to the parameters
182
183
184
   for i=1:3
185
        switch i
186
             case 1
187
             s1='Random';
188
```

```
189
            val=0;
190
            case 2
            s1='Parallel';
191
192
            val=27;
193
             case 3
194
             s1='Antiparallel';
195
             val=54;
196
        end
197
        for j=1:4
198
199
            switch j
200
            case 1
201
            s2='clust_coefficient';
202
            main_vec=clust_coefficient;
203
            case 2
            s2='s';
204
            main_vec=s;
205
206
            case 3
207
            s2='average_path_length';
208
            main_vec=average_path_length;
209
            case 4
            s2='diam';
210
            main_vec=diam;
211
212
            end
213
            for k=1:3
214
                 switch k
215
                     case 1
                          s3='threshold';
216
                          f=figure();
217
                          for 1=1:3
218
                              line=main_vec(3*(0:8)+val+1);
219
220
                              set(gca, 'FontSize', 14)
221
                              xlabel('Different pairs of alpha and phi values');
222
                              ylabel(s2);
223
                              namek=['Dependence of',' ',s2,' ','on',' ',s3,' ...
                                   ','for',' ',s1,' ','idea distribution'];
                              title(namek);
224
225
226
                              switch 1
227
                                        plot(line, '-ks', 'LineWidth', 2, 'MarkerEdgeColor', 'k', 'MarkerE
228
                                        hold on
229
230
                                   case 2
                                        plot(line, '--bs', 'LineWidth', 2, 'MarkerEdgeColor', 'b', 'MarkerF
231
232
                                        hold on
233
                                   case 3
                                        plot(line,'--rs','LineWidth',2,'MarkerEdgeColor','r','MarkerF
234
235
                              end
                              legend('threshold=0.001', 'threshold=0.01', 'threshold=0.05')
236
                              name=['plotting1_',s1,'_',s2,'_','threshold_sensitivity'];
237
```

```
238
                              saveas(f, name);
                          end
239
240
                     case 2
241
242
                          s3='alpha';
243
                          f=figure();
244
                          for 1=1:3
245
                              line=main_vec([1,2,3,10,11,12,19,20,21]+val+3*([1-1));
246
                              set(gca,'FontSize',14)
                              xlabel('Different pairs of threshold and phi ...
247
                                  values');
248
                              ylabel(s2);
                              namek=['Dependence of',' ',s2,' ','on',' ',s3,' ...
249
                                  ','for',' ',s1,' ','idea distribution'];
                              title(namek);
250
251
                              switch 1
252
253
                                  case 1
254
                                        plot(line, '-ks', 'LineWidth', 2, 'MarkerEdgeColor', 'k', 'MarkerE
255
                                        hold on
256
                                  case 2
                                        plot(line,'--bs','LineWidth',2,'MarkerEdgeColor','b','MarkerF
257
                                        hold on
258
259
                                  case 3
260
                                        plot(line,'--rs','LineWidth',2,'MarkerEdgeColor','r','MarkerF
261
                              legend('alpha=0.01', 'alpha=0.05', 'alpha=0.10')
262
                              name=['plotting1_',s1,'_',s2,'_','alpha_sensitivity'];
263
264
                              saveas(f, name);
265
                          end
266
267
                     case 3
                          s3='phi';
268
269
                          f=figure();
                          for 1=1:3
270
                              line=main_vec((1:9)+val+9*(1-1));
271
272
                              set (gca, 'FontSize', 14)
273
                              xlabel('Different pairs of threshold and alpha ...
                              vlabel(s2);
274
                               namek=['Dependence of',' ',s2,' ','on',' ',s3,' ...
275
                                   ','for',' ',s1,' ','idea distribution'];
                              title(namek);
276
277
278
                              switch 1
279
                                        plot(line,'-ks','LineWidth',2,'MarkerEdgeColor','k','MarkerF
280
281
                                        hold on
282
                                  case 2
                                        plot(line, '--bs', 'LineWidth', 2, 'MarkerEdgeColor', 'b', 'MarkerF
283
```

```
284
                                hold on
                           case 3
285
                                plot(line, '-rs', 'LineWidth', 2, 'MarkerEdgeColor', 'r', 'MarkerE
286
                        end
287
                        legend('phi=0.1','phi=0.3','phi=0.5')
288
                        name=['plotting1_',s1,'_',s2,'_','phi_sensitivity'];
289
290
                        saveas(f, name);
291
                    end
292
             end
          end
293
294
      end
295
   end
296
297
  %%%%%%% generating plots for each features of network structure for each
298
   299
300
301
   for i=1:4
302
      switch i
303
          case 1
304
             f=figure();
             line1=clust_coefficient(1:27);
305
             line2=clust_coefficient(28:54);
306
307
             line3=clust_coefficient(55:81);
308
             309
             hold on
             plot(line2,'-qs','LineWidth',2,'MarkerEdgeColor','q','MarkerFaceColor','q','Marker
310
311
             hold on
             plot(line3,'--bs','LineWidth',2,'MarkerEdgeColor','b','MarkerFaceColor','b','Marker
312
313
             set(gca, 'FontSize', 14)
             xlabel('Different triplets of alpha ,phi and threshold values')
314
315
             ylabel('Clustering Coefficient')
             title('The influence of idea distribution on clustering ...
316
                 coefficient');
             legend('Random distribution', 'Parallel ...
317
                distribution', 'Antiparallel distribution')
             name=('Plotting2_idea_Comparison_clustcoeff');
318
319
             saveas(f, name);
320
          case 2
321
             f=figure();
             line1=s(1:27);
322
             line2=s(28:54);
323
             line3=s(55:81);
324
             325
326
             hold on
             327
             hold on
328
             plot(line3,'-bs','LineWidth',2,'MarkerEdgeColor','b','MarkerFaceColor','b','Marker
329
             set(gca, 'FontSize', 14)
330
             xlabel('Different triplets of alpha ,phi and threshold values')
331
```

```
332
               ylabel('Number of connected components')
               title('The influence of idea distribution on the number of ...
333
                   connected components')
               legend('Random distribution','Parallel ...
334
                  distribution','Antiparallel distribution')
               name=('Plotting2_idea_Comparison_conncmp');
335
               saveas(f, name);
336
           case 3
337
               f=figure();
338
               line1=average_path_length(1:27);
339
340
               line2=average_path_length(28:54);
341
               line3=average_path_length(55:81);
342
               plot(line1, '-ks', 'LineWidth', 2, 'MarkerEdgeColor', 'k', 'MarkerFaceColor', 'k', 'Marker
343
               hold on
344
               hold on
345
               plot(line3,'-bs','LineWidth',2,'MarkerEdgeColor','b','MarkerFaceColor','b','Marker
346
347
               set(gca, 'FontSize', 14)
348
               xlabel('Different triplets of alpha ,phi and threshold values')
349
               ylabel('average path length')
350
               title('The influence of idea distribution on average path ...
                   length')
               legend('Random distribution', 'Parallel ...
351
                  distribution', 'Antiparallel distribution')
352
               name=('Plotting2_idea_Comparison_average_path_length');
353
               saveas(f, name);
354
           case 4
355
               f=figure();
               line1=diam(1:27);
356
357
               line2=diam(28:54);
358
               line3=diam(55:81);
               359
360
               hold on
               plot(line2,'--qs','LineWidth',2,'MarkerEdgeColor','g','MarkerFaceColor','g','Marker
361
362
               hold on
               plot(line3,'-bs','LineWidth',2,'MarkerEdgeColor','b','MarkerFaceColor','b','Marker
363
               set(gca, 'FontSize', 14)
364
365
               xlabel('Different triplets of alpha ,phi and threshold values')
366
               ylabel('Network Diameter')
               title('The influence of idea distribution on Network Diameter')
367
               legend('Random distribution', 'Parallel ...
368
                  distribution', 'Antiparallel distribution')
               name=('Plotting2_idea_Comparison_diam');
369
               saveas(f, name);
370
371
       end
372 end
373
374 %%%%%%% plots for degree distribution resulting from different idea
375 %%%%%%%%% distribution
376 count=0;
```

```
377 for choice1=1:3
        switch choice1
378
             case 1
379
                 s1='phi=0.1';
380
381
                 m1 = '1';
382
             case 2
383
                 s1='phi=0.3';
384
                 m1='2';
385
             case 3
                 s1='phi=0.5';
386
387
                 m1 = '3';
388
        end
389
390
        for choice2=1:3
391
             switch choice2
                 case 1
392
                      s2='alpha=0.01';
393
394
                      m2 = '1';
395
                 case 2
396
                      s2='alpha=0.05';
                      m2 = '2';
397
398
                 case 3
399
                      s2='alpha=0.1';
400
                      m2 = '3';
401
             end
402
403
             for choice3=1:3
404
                 switch choice3
                      case 1
405
                          s3='threshold=0.001';
406
                          m3 = '1';
407
408
                      case 2
409
                          s3='threshold=0.01';
                          m3 = '2';
410
411
                      case 3
412
                          s3='threshold=0.05';
                          m3 = '3';
413
414
                 end
415
                 count=count+1;
416
                 line1=dgr{count};
                 mine1=frq{count}/1000;
417
                 line2=dgr{27+count};
418
                 mine2=frq{27+count};
419
                 line3=dgr{54+count};
420
421
                 mine3=frq{54+count};
422
                 f=figure();
                 loglog(line1, mine1, '-ks', 'LineWidth', 2, 'MarkerEdgeColor', 'k', 'MarkerFaceColor', '}
423
424
                 hold on
                 plot(line2, mine2, '-rs', 'LineWidth', 2, 'MarkerEdgeColor', 'r', 'MarkerFaceColor', 'r',
425
                 hold on
426
```

```
plot(line3, mine3, '-bs', 'LineWidth', 2, 'MarkerEdgeColor', 'b', 'MarkerFaceColor', 'b',
427
428
                set(gca, 'FontSize', 14)
                xlabel('Log(degree)')
429
                ylabel('Log(frequency)')
430
                title(['The influence of idea distribution on the degree ...
431
                    distribution',' ',s1,' ',s2,' ',s3]);
                legend('Random distribution','Parallel ...
432
                    distribution','Antiparallel distribution')
                name=(['Plotting2_idea_comparison_degree_distribution-',m1,'_',m2,'_',m3]);
433
434
                saveas(f,name);
435
            end
        end
437 end
```

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