Introduction and Comparison On Random Graph Generation Algorithms

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**Abstract**

This article puts forward two novel user-grouping algorithms for grouped multi-carrier (MC)-code division multiple access (CDMA) systems. As is well known, the adaptive assignment for user-grouping plays an important role for link quality of multi-access transmissions. In the study, the capacity-maximizing problem of user-grouping is formulated. By using the Kuhn-Tucker condition, the optimal criterion is deduced and found to have a similar form with signal to noise plus interference (SINR). However SINR includes the signal power that can only be determined after user-grouping. Therefore the optimal criterion will lead to an impractical application. To deal with it, the user’s equivalent SINR for minimum mean square error (MMSE) detector is proposed and served as a suboptimal assignment criterion, based on which two kinds of user-grouping algorithms are proposed. In the algorithms, only partial channel information is needed at the base station, which saves a large part of the bandwidth occupied by feedback information. Computer simulations have evaluated an excellent performance of the proposed algorithms at both link quality and data rate. Meanwhile, the proposed algorithms have lower implementation complexity for practical reality.

**Keywords**  Random graph, Stochastic process

**1 Introduction[[1]](#footnote-1)**

In an era where quantity of information explodes and most of them are interconnected and correlative, data analysis on sophisticated graphs and complex networks is increasingly valued by researchers and business owners. A lot of data mining algorithms and models are developed to extract and exploit valuable and potentially profitable information from existing graphs. To verify them, tons of data has to be collected, well organized, and correctly labelled in order to construct a graph ready for analysis. However, such process involves with a large amount of manual work that is very painful and costly. In order to enable agile verification and iteration of new models, generation of heterogeneous random graphs has become very important. Fortunately, many effective algorithms are already proposed and broadly adopted.

This article would list a couple of typical algorithms of graph generation for random undirected graphs, discuss the most fundamental ideas under the hood, and then compare them on different and practical perspectives. Afterward, some real-world graphs are provided in comparison to samples of our random graphs.

**2 Random Graphs**

In mathematical points of view, a random graph is described by either probability distribution of vertices and edges or a stochastic process that generate it. Practically, two types of construction process are very common:

1. Create all nodes at once, and connect them upon a specific probability distribution
2. Add nodes by ones or batches, and wire(rewire) them on growth

Given some parameters determined, such as number of vertices and edges, a sample of the random graph could be created and put into use.

Before digging into concrete generation algorithms, some important indices for realistic graph evaluations are listed below:

1. **degree distribution**: probability distribution of degree of nodes in the graph
2. **average path length**: average smallest number of edges between any two nodes in the graph
3. **clustering coefficient**: an index that describe how nodes form communities

**3 Graph Generation Algorithms**

3.1 Erdős–Rényi model

Considering the first type of construction process, the Erdős–Rényi model further specify that the presence of edge between any two vertices is equally and independently probable, i.e. observes a uniform distribution. It may either be parameterized as or , where is the number of vertices, is the number of edges and is the probability of a connection to form.

In notation, the total number of edges is deterministic, i.e. . Thus the scale of the sample is known beforehand and spawn probability of edges may vary. However, it is obvious that the ML estimation of connection rate satisfy the formula:

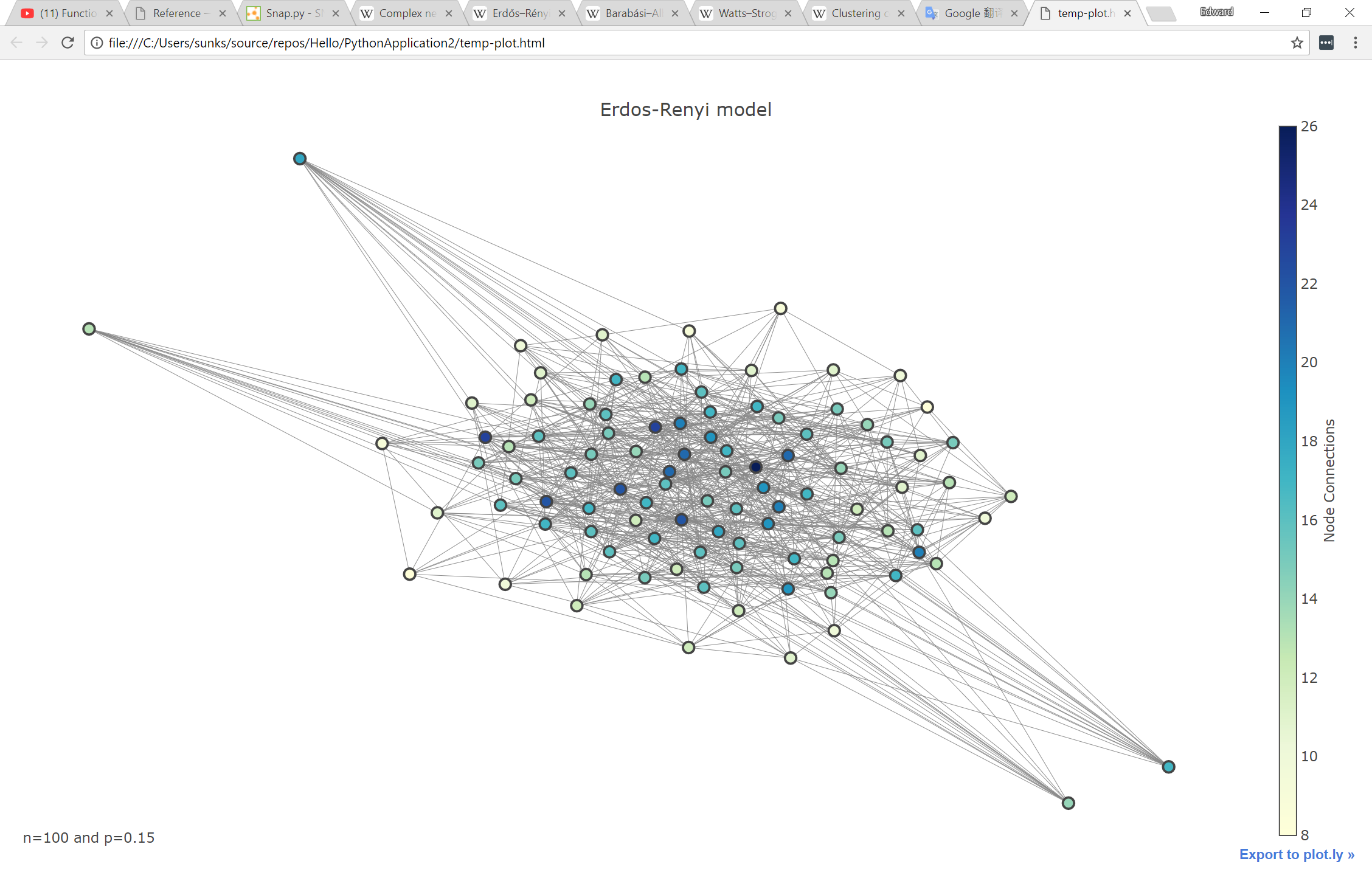
Meanwhile, in abstraction, the edge count is a random variable with a binomial probability distribution

and an expected value

In this article, the latter model is preferred to be referred as ER model. By definition, ER model has a binomial degree distribution

where k denotes the degree of a random vertex, and a fairly small clustering coefficient. The graph it samples is usually very dense and such topology is rare in reality. Also, because of no random nature of the graph, it is not guarantee one totally inter-connected graph is sampled instead of a forest. However, it is provable that there is usually a major component in branches of forest, if any.

A sample of ER model is given below, where and , and vertices are colored by its degree.



3.2 Barabási–Albert model

In real-world, many networks have the property of being scale-free; that is, their degree distribution follows a power law, instead of being binomial. Suppose P(k) denotes the proportion nodes that have k outgoing edges. The power law could be described as , where typically falls in the range .

The construction process of scale-free networks usually (but not necessarily) possess the following schemes:

1. Construct the whole graph by adding new nodes
2. Apply preferential attachment: vertices with more connections are more popular

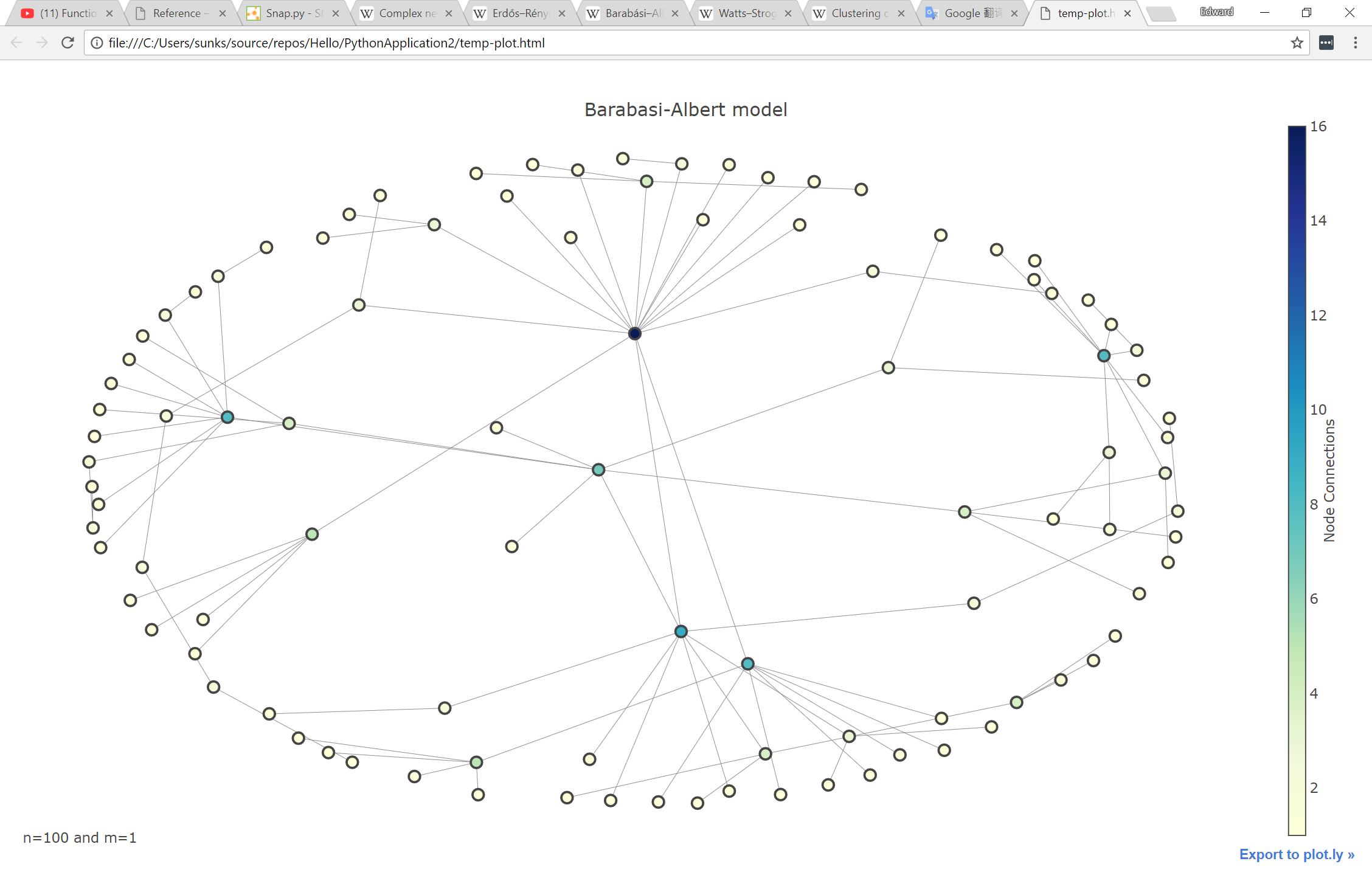
Similar to the ER model, Barabási–Albert model requires a linear preferential attachment probability. For an existing node , a new node would have a chance of to be attached or connected, noting is degree of the particular vertex. Usually, the number of edges to be attached for each new node is denoted as a constant m. Therefore, the model may be denoted as .

For a model starts with a single isolated vertex, the edge count is also a deterministic constant . Different from previously mentioned ER model, graph samples of BA model usually forms a certain number of central nodes and possess a very recognizable topology structure. And more importantly, they are scale-free, where degree distribution would be around

and average path length would be around

To be noted, preferential attachment is not a requirement. And different random distribution for new attachment might work better in some circumstances.

The following image shows a sample of BA model, where and . It is apparent that the network BA model generates is very centralized, and most end-points rely on a few inter-connected hubs to communicate with each other, which, in some sense, is very realistic.

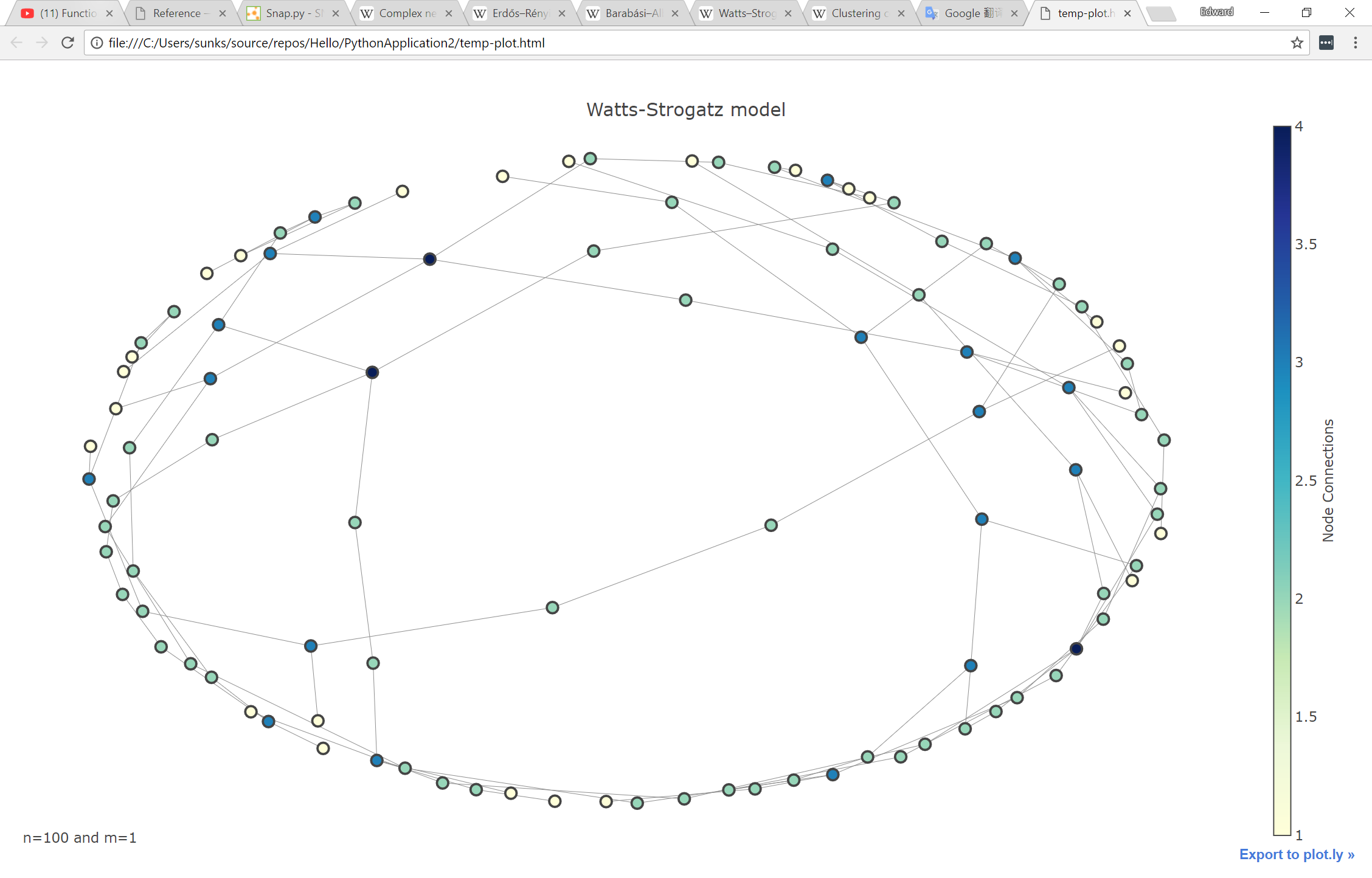


3.3 Watts–Strogatz model

On top of being scale-free, many real-world networks are also observed as following the analogy of small-world phenomenon, where length of path between any two nodes are relatively short in comparison to the scale of whole graph. Mathematically, a small-world network has a small average node distance and a significantly large clustering coefficient.

And the most trivial method for generating small-world graph is Watts-Strogatz model. This is also very simple an algorithm. It starts with a regular ring lattice graph with nodes connected to its neighbors. For each lattice edge from to where , there is a chance of that a rewire process occur and a random new node is selected in place of in uniform distribution. In addition, the algorithm should always avoid some that may lead to self loop or duplicate edges. After all edges are visited, a small-world graph is done.

Note properties such as degree distribution and average path length is rather complicated and rely heavily on the parameter \beta. Here, the formula would be skipped, and a simulation of the model is beneath:



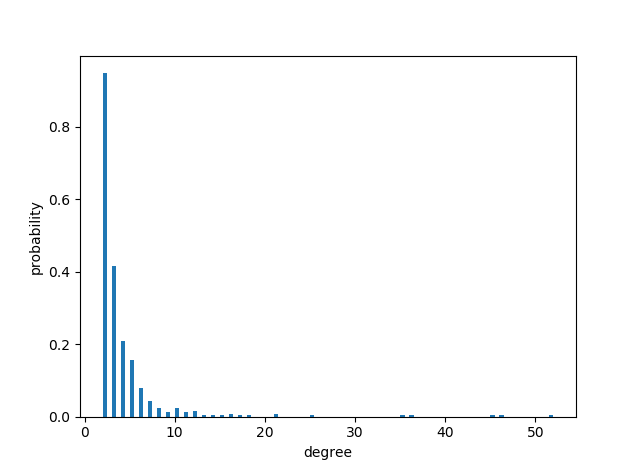
3.4 Ensemble model

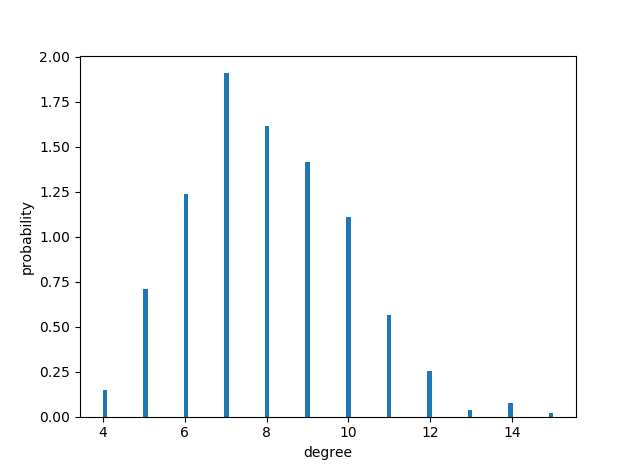
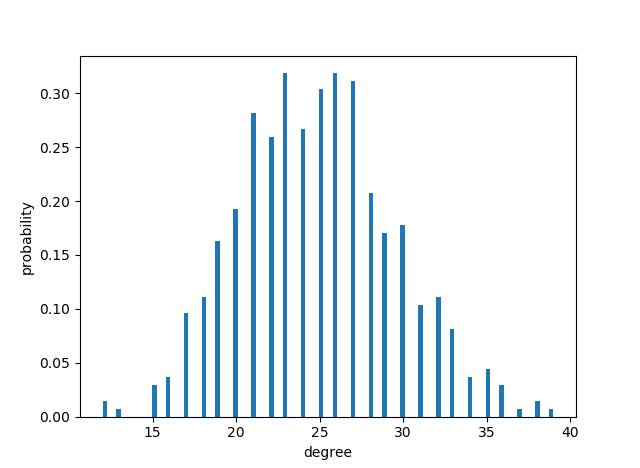
According to the discussion above, although being carefully designed, both BA model and WS model cannot perfectly abstract our real-world needs. However, as graph construction process is incremental, we may first construct several sub-graphs as an autonomous system and then merge them into a single large network.

**4 Monte Carlo Simulation On Statistical Properties**

To be noted, the simulation graph below is mostly smoothened by averaging results from multiple epochs. Also, the simulation results are always listed as, respectively, ER, BA and WS model.

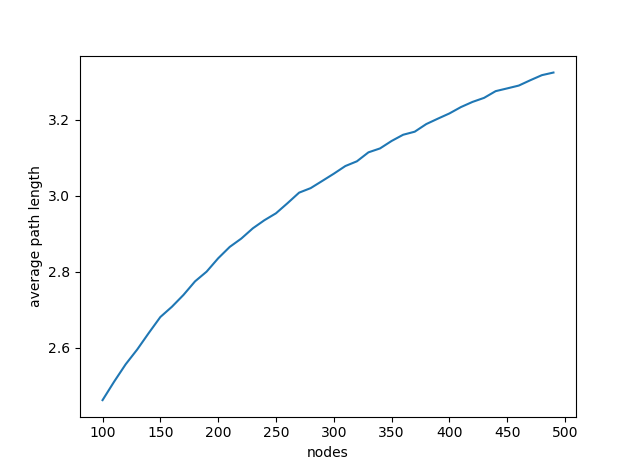
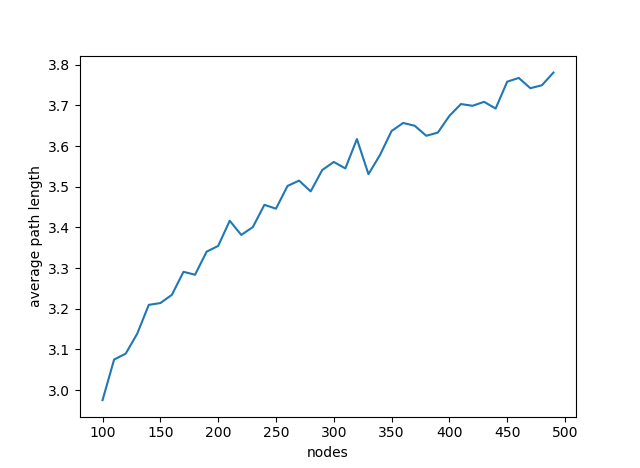
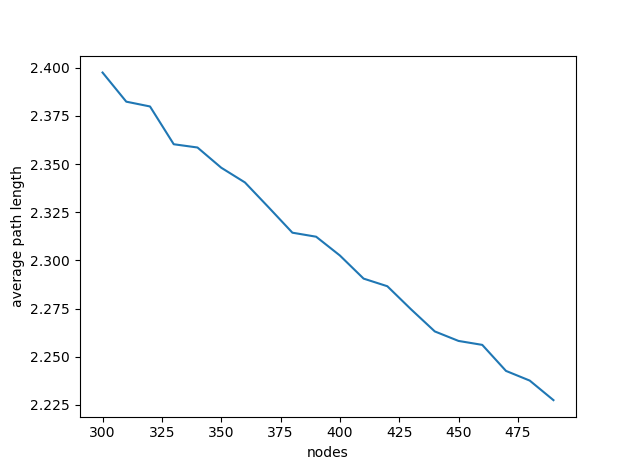
3.1 Degree Distribution





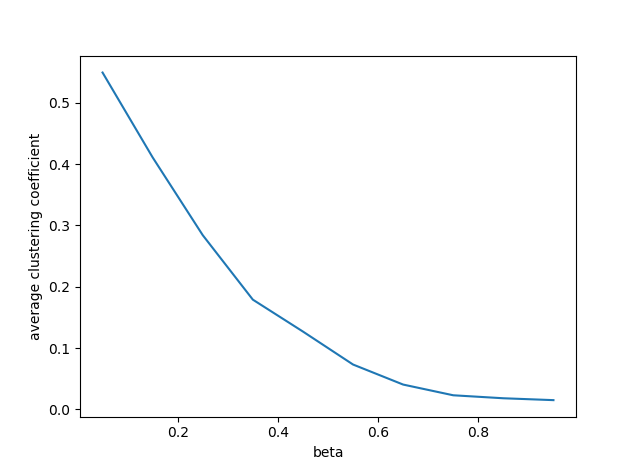
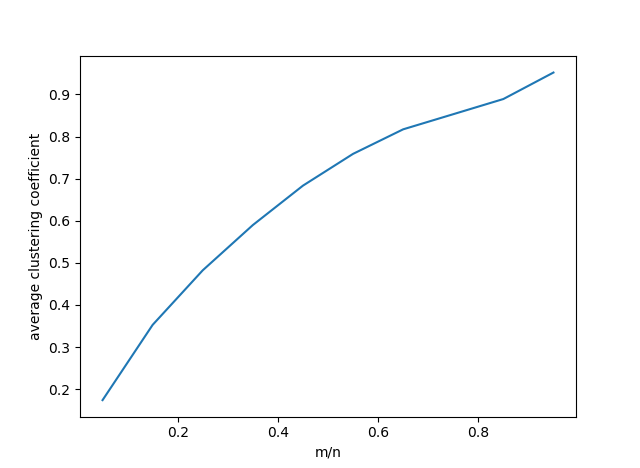
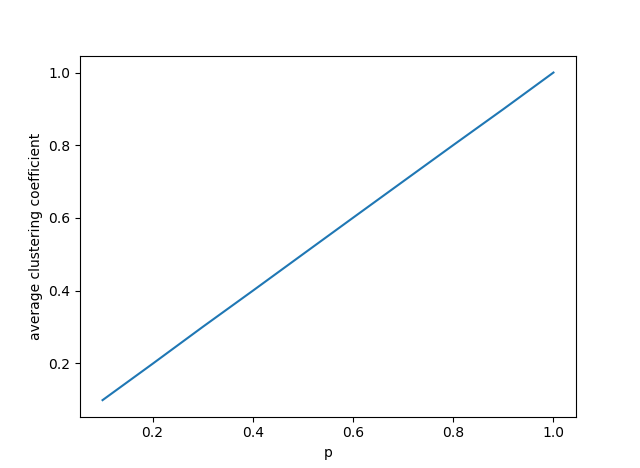
A Poisson distribution and power-law distribution can be easily observed.

3.2 Average Path Length



To be noted, for ER model, as p persists as a constant, the number of edges grows faster than the scale of the graph, so the average path length actually drops as the count of nodes increases. Apparently, these three models have an average path length that grows in logarithm as the graph scales.

3.3 Average Clustering Coefficient



Although ER model and BA model can have a significantly large average clustering coefficient, those parameter range is usually not typical and rarely happen. For example, in ER model is usually very small, especially when the graph generated is fairly large. In such situation, the average clustering coefficient remains tiny all the time.

**4 Some Real-world Networks Collected**

**5 Conclusions**

This study mainly considers the capacity optimization criterion of subcarrier allocation for the downlink grouped MC-CDMA. MAI was put

**References**

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1. [↑](#footnote-ref-1)