Quick Detection of Nodes with Large Degrees

Konstantin Avrachenkov\(^\ast\), Nelly Litvak\(^\dagger\), Marina Sokol\(^\ddagger\), Don Towsley\(^\S\)

Abstract

Our goal is to quickly find top \(k\) lists of nodes with the largest degrees in large complex networks. If the adjacency list of the network is known (not often the case in complex networks), a deterministic algorithm to find the top \(k\) list of nodes with the largest degrees requires an average complexity of \(O(n)\), where \(n\) is the number of nodes in the network. Even this modest complexity can be very high for large complex networks. We propose to use the random walk based method. We show theoretically and by numerical experiments that for large networks the random walk method finds good quality top lists of nodes with high probability and with computational savings of orders of magnitude. We also propose stopping criteria for the random walk method which requires very little knowledge about the structure of the network.

1 Introduction

We are interested in quickly detecting nodes with large degrees in very large networks. Firstly, node degree is one of centrality measures used for the analysis of complex networks. Secondly, large degree nodes can serve as proxies for central nodes corresponding to the other centrality measures as betweenness centrality or closeness centrality [10, 11]. In the present work we restrict ourself to undirected networks or symmetrized versions of directed networks. In particular, this assumption is well justified in social networks. Typically, friendship or acquaintance is a symmetric relation. If the adjacency list of the network is known (not often the case in complex networks), a deterministic algorithm to find the top \(k\) list of nodes with the largest degrees requires an average complexity of \(O(n)\), where \(n\) is the number of nodes in the network. For instance, if HeapSort is used to find the top \(k\) list of nodes with the largest degrees, the complexity estimation can be specified as \(O(n + k \log(n))\). We assume that the degree is available

\(^\ast\)Inria Sophia Antipolis, France, k.avrachenkov@sophia.inria.fr
\(^\dagger\)University of Twente, The Netherlands, n.litvak@utwente.nl
\(^\ddagger\)Inria Sophia Antipolis, France, marina.sokol@sophia.inria.fr
\(^\S\)University of Massachusetts Amherst, USA, towsley@cs.umass.edu
when accessing a node (if this is not the case, the complexity should be counted in terms of links). However, even linear complexity can be very high for very large, possibly varying, complex networks. Furthermore, when crawling some online social networks like Facebook or Twitter, a crawler is constrained by a certain limit on the speed of crawling. For example, Twitter has the limit of one access per minute for the rate of crawling for one standard account. Thus, to crawl the entire network with more than 500 million users we need more than 950 years. Certainly we would like to discover nodes with largest degrees well before the entire network is crawled.

In the present work we suggest using random walk based methods for detecting a small number of nodes with the largest degree. The main idea is that the random walk very quickly comes across large degree nodes. Thus, the analysis of our approach is equivalent to the analysis of hitting times of a random walk. In our numerical experiments random walks outperform the standard deterministic algorithms by orders of magnitude in terms of computational complexity. For instance, in our experiments with the web graph of the UK domain (about 18500000 nodes) the random walk method spends on average only about 5400 steps to detect the largest degree node. Potential memory savings are also significant since the method does not require knowledge of the entire network. In many practical applications we do not need a complete ordering of the nodes and even can tolerate some errors in the top list of nodes. We observe that the random walk method obtains many nodes in the top list correctly and even those nodes that are erroneously placed in the top list have large degrees. Therefore, as typically happens in randomized algorithms [14, 15], we trade off exact results for very good approximate results or for exact results with high probability and gain significantly in computational efficiency.

The paper is organized as follows: in the next section we introduce our basic random walk with uniform jumps and demonstrate that it is able to quickly find large degree nodes. Then, in Section 3 using configuration model we provide an estimate for the necessary number of steps for the random walk. In Section 4 we propose stopping criteria that use very little information about the network. In Section 5 we show the benefits of allowing few erroneous elements in the top $k$ list. Finally, we conclude the paper in Section 6.

2 Random walk with uniform jumps

Let us consider a random walk with uniform jumps which serves as a basic algorithm for quick detection of large degree nodes. The random walk with uniform jumps is described by the following transition probabilities [1]

$$p_{ij} = \begin{cases} \frac{\alpha/n+1}{d_i+\alpha}, & \text{if } i \text{ has a link to } j, \\ \frac{\alpha/n}{d_i+\alpha}, & \text{if } i \text{ does not have a link to } j. \end{cases}$$ (1)
where \( d_i \) is the degree of node \( i \). The random walk with uniform jumps can be regarded as a random walk on a modified graph where all the nodes in the graph are connected by artificial edges with a weight \( \alpha/n \). The parameter \( \alpha \) controls the rate of jumps. Introduction of jumps helps in a number of ways. As was shown in [1], it reduces the mixing time to stationarity. It also solves a problem encountered by a random walk on a graph consisting of two or more components, namely the inability to visit all nodes. The random walk with jumps also reduces the variance of the network function estimator [1]. This random walk resembles the PageRank random walk. However, unlike the PageRank random walk, the introduced random walk is reversible. One important consequence of the reversibility of the random walk is that its stationary distribution is given by a simple formula

\[
\pi_i(\alpha) = \frac{d_i + \alpha}{2|E| + n\alpha} \quad \forall i \in V,
\]

from which the stationary distribution of the unperturbed random walk can easily be retrieved. We observe that the modification preserves the monotonicity of the stationary distribution with respect to the node degree, which is particularly important for our application.

We illustrate on several network examples how the random walk helps us quickly detect large degree nodes. We consider as examples one synthetic network generated by the preferential attachment rule and two natural large networks. The Preferential Attachment (PA) network combines 100,000 nodes. It has been generated according to the generalized preferential attachment mechanism [8]. The average degree of the PA network is two and the power law exponent is 2.5. The first natural example is the symmetrized web graph of the whole UK domain crawled in 2002 [5]. The UK network has 18,520,486 nodes and its average degree is 28.6. The second natural example is the network of co-authorships of DBLP [6]. Each node represents an author and each link represents a co-authorship of at least one article. The DBLP network has 986,324 nodes and its average degree is 6.8.

We carry out the following experiment: we initialize the random walk (1) at a node chosen according to the uniform distribution and continue the random walk until we hit the largest degree node. The largest degrees for the PA, UK and DBLP networks are 138, 194,955, and 979, respectively. For the PA network we have made 10,000 experiments and for the UK and DBLP networks we performed 1,000 experiments (these networks were too large to perform more experiments).

In Figure 1 we plot the histograms of hitting times for the PA network. The first remarkable observation is that when \( \alpha = 0 \) (no restart) the average hitting time, which is equal to 123,000 steps, is nearly three orders of magnitude larger than 3,720, the hitting time when \( \alpha = 2 \). The second remarkable observation is that 3,720 is of the same order of magnitude as the value \( 1/\pi_{\text{max}}(\alpha) = (2|E| + n\alpha)/(d_{\text{max}} + \alpha) = 2,857 \), which corresponds
to the average return time to the largest degree node in the random walk with jumps.

![Histograms of hitting times in the PA network.](image)

Figure 1: Histograms of hitting times in the PA network.

We were not able to collect a representative number of experiments for the UK and DBLP networks when $\alpha = 0$. The reason for this is that the random walk gets stuck either in disconnected or weakly connected components of the networks. For the UK network we were able to make 1000 experiments with $\alpha = 0.001$ and obtain the average hitting time 30750. Whereas if we take $\alpha = 28.6$ for the UK network, we obtain the average hitting time 5800. Note that the expected return time to the largest degree node in the UK network is given by $1/\pi_{\text{max}}(\alpha) = (2|E| + n\alpha)/(d_{\text{max}} + \alpha) = 5432$. For the DBLP graph we conducted 1000 experiments with $\alpha = 0.00001$ and obtained an average hitting time of 41131. Whereas if we take $\alpha = 6.8$, we obtain an average hitting time of 14200. The expected return time to the largest degree node in the DBLP network is given by $1/\pi_{\text{max}}(\alpha) = (2|E| + n\alpha)/(d_{\text{max}} + \alpha) = 13607$. The two natural network examples confirm our guess that the average hitting time for the largest degree node is fairly close to the average return time to the largest degree node, which is reciprocal to the value of the stationary distribution at the largest degree node. Next, using asymptotic analysis, we show that if $\alpha$ is sufficiently large, the principal term in the asymptotic expansion for the expected hitting time is close to the expected return time.

**Theorem 1** Without loss of generality, index the nodes such that node 1 is a node under consideration, $(1, i) \in E, i = 2, ..., s, s = d_1 + 1$, and let $\nu$ denote the initial distribution of the random walk with jumps. Then, for sufficiently large $\alpha$ and small $\alpha/n$, the expected hitting time to node 1 starting from an arbitrary initial distribution $\nu$ is given by

$$E_\nu[H_1] = \frac{\sum_{i=2}^{n} d_i + (n - 1)\alpha}{d_1 + 2\alpha(1 - 1/n)} + O(1).$$  \hspace{1cm} (3)
**Proof:** The expected hitting time from distribution \( \nu \) to node 1 is given by the formula

\[
E_\nu[H_1] = \nu[I - P_{-1}]^{-1}1,
\]

(4)

where \( P_{-1} \) is a taboo probability matrix (i.e., matrix \( P \) with the 1-st row and 1-st column removed). The matrix \( P_{-1} \) is substochastic but is very close to stochastic. Let us represent it as a stochastic matrix minus some perturbation term:

\[
P_{-1} = \tilde{P} - \varepsilon Q = \tilde{P} - \begin{bmatrix}
\frac{1+2\alpha/n}{d_2+\alpha} & 0 & 0 \\
0 & \ddots & \ddots \\
0 & \frac{2\alpha/n}{d_{s+1}+\alpha} & 0 \\
0 & 0 & \frac{2\alpha/n}{d_n+\alpha}
\end{bmatrix}
\]

We add missing probability mass to the diagonal of \( \tilde{P} \), which corresponds to an increase in the weights for self-loops. The matrix \( \tilde{P} \) represents a reversible Markov chain with the stationary distribution

\[
\tilde{\pi}_j = \frac{d_j + \alpha}{\sum_{i=2}^{n} d_i + (n-1)\alpha}.
\]

Now we can use the following result from the perturbation theory (see Lemma 1 in [2]):

\[
[I - \tilde{P} + \varepsilon Q]^{-1} = \frac{1}{\tilde{\pi}(\varepsilon Q)1}X_0 + \varepsilon X_1 + ...,
\]

(5)

where \( \tilde{\pi} \) is the stationary distribution of the stochastic matrix \( \tilde{P} \). In our case, the quantity \( \max_{i=2,\ldots,s}\{1/(d_i + \alpha), 1/n\} \) will play the role of \( \varepsilon \). We apply the series (5) to approximate the expected hitting time. Towards this goal, we calculate

\[
\tilde{\pi}(\varepsilon Q)1 = \sum_{j=2}^{n}\tilde{\pi}_j\varepsilon q_{jj}
\]

\[
= \sum_{j=2}^{s} \frac{d_j + \alpha}{\sum_{i=2}^{n} d_i + (n-1)\alpha} \frac{1+2\alpha/n}{d_j + \alpha} + \sum_{j=s+1}^{n} \frac{d_j + \alpha}{\sum_{i=2}^{n} d_i + (n-1)\alpha} \frac{2\alpha/n}{d_j + \alpha}
\]

\[
= \frac{d_1(1+2\alpha/n) + (n-d_1-1)(2\alpha/n)}{\sum_{i=2}^{n} d_i + (n-1)\alpha} = \frac{d_1 + 2\alpha(1 - 1/n)}{\sum_{i=2}^{n} d_i + (n-1)\alpha}.
\]

Observing that \( \nu 1 \tilde{\pi} 1 = 1 \), we obtain (3).

\[\Box\]
Indeed, the asymptotic expression (3) is very close to 
\( (2|E|+n\alpha)/(d_1+\alpha) \), which is the expected return time to node 1.

Based on the notion of the hitting time we propose an efficient method for quick detection of the top \( k \) list of largest degree nodes. The algorithm maintains a top \( k \) candidate list. Note that once one of the \( k \) nodes with the largest degrees appears in this candidate list, it remains there subsequently. Thus, we are interested in hitting events. We propose the following algorithm for detecting the top \( k \) list of largest degree nodes.

**Algorithm 1 Random walk with jumps and candidate list**

1. Set \( k \), \( \alpha \) and \( m \).

2. Execute a random walk step according to (1). If it is the first step, pick the initial node arbitrarily (in particular, the initial node can be chosen by the uniform distribution).

3. Check if the current node has a larger degree than one of the nodes in the current top \( k \) candidate list. If it is the case, insert the new node in the top-\( k \) candidate list and remove the worst node out of the list.

4. If the number of random walk steps is less than \( m \), return to Step 2 of the algorithm. Stop, otherwise.

The value of parameter \( \alpha \) is not crucial. In our experiments, we have observed that as long as the value of \( \alpha \) is neither too small nor too big, the algorithm performs well. According to our observations, a good option for the choice of \( \alpha \) is a value around the average node degree. Let us explain this choice.

Consider a random walk \( \{W_t\}_{t=0}^{\infty} \) with transition probabilities (1). We denote by \( P_\nu(\cdot) \) the probability distribution of this Markov chain with initial distribution \( \nu \). Now assume that the Markov chain is in a stationary regime (the stationary regime is achieved quickly when the parameter \( \alpha \) is not too small [1]). Then by the Bayes formula we derive two remarkable equations:

\[
P_\pi[W_t = i | \text{jump}] = \frac{P_\pi[W_t = i, \text{jump}]}{P_\pi[\text{jump}]} = \frac{P_\pi[W_t = i]P_\pi[\text{jump}|W_t = i]}{\sum_{j=1}^n P_\pi[W_t = j]P_\pi[\text{jump}|W_t = j]} = \frac{d_i + \alpha}{2|E| + n\alpha} \frac{\alpha}{\sum_{j=1}^n d_j + \alpha} = \frac{1}{n},
\]

\[
P_\pi[W_t = i | \text{no jump}] = \frac{P_\pi[W_t = i, \text{no jump}]}{P_\pi[\text{no jump}]} = \frac{P_\pi[W_t = i]P_\pi[\text{no jump}|W_t = i]}{\sum_{j=1}^n P_\pi[W_t = j]P_\pi[\text{no jump}|W_t = j]} = \frac{d_i + \alpha}{2|E| + n\alpha} \frac{d_i}{\sum_{j=1}^n d_j + \alpha}
\]

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Thus, in a stationary distribution, given that no jump occurred, the probability that \( W_t = i \) is exactly \( \pi_i(0) \).

Next observe that \( W_t \) is a regenerative process, where regeneration points are the jumps to the uniform distribution, and the regenerating cycles are independent. Concerning the choice of \( \alpha \), there is a clear trade-off: if \( \alpha \) is too small, then regenerating cycles are long and a random walk can get entangled in some part of the network; but if \( \alpha \) is too large, then the cycle will often consist only of one step corresponding to a jump. Thus, we would like to maximize the long-run fraction of independent observations from \( \pi(0) \). To this end, we note that given \( m' \) cycles, the mean total number of steps is

\[
m'\mathbb{E}[\text{cycle length}] = m'(P_\pi[\text{jump}])^{-1}.
\]

Out of the random walk run with \( m' \) cycles, \( m' \) independent observations from \( \pi \) are generated, from which on average \( m'P_\pi[\text{jump}] \) observations coincide with a jump. As will be discussed in Section 4, we need to maximize the long-run fraction of independent observations, that are not a jump, in a sample compared to the number of steps of a random walk:

\[
\frac{m' - m'P_\pi[\text{jump}]}{m'(P_\pi[\text{jump}])^{-1}} = P_\pi[\text{jump}](1 - P_\pi[\text{jump}]) \to \max.
\]

Obviously, the maximum is achieved when

\[
P_\pi[\text{jump}] = \frac{1}{2}.
\]

It remains to rewrite \( P_\pi[\text{jump}] \) in terms of the algorithm parameters:

\[
P_\pi[\text{jump}] = \sum_{j=1}^{n} P_\pi[W_t = j]P_\pi[\text{jump}|W_t = j]
\]

\[
= \sum_{j=1}^{n} \frac{d_j + \alpha}{2|E| + n\alpha} \frac{\alpha}{d_j + \alpha} = \frac{n\alpha}{2|E| + n\alpha} = \frac{\alpha}{d + \alpha},
\]

where \( d := 2|E|/n \) is the average degree. For the maximal efficiency, the last fraction above must be equal to \( 1/2 \), which gives the optimal value for the parameter \( \alpha \)

\[
\alpha_* = \bar{d}.
\]

With this choice of \( \alpha \), the random walk contains the maximal possible fraction of independent observations from the distribution \( \pi_i(0) \).

The average degree is not necessarily known in advance. However, we may chose \( \alpha \) based on our knowledge of samples of similar nature, and
then estimate the average degree using (8) and the observed cycle length. Specifically, we can use the equation

\[ E_u[T] = \frac{1}{P_{\pi}[\text{jump}]} = \frac{2|E|/n + \alpha}{\alpha}. \] (9)

Then we can adjust \( \alpha \) to its optimal value.

Theorem 1 demonstrates that the expected hitting time to a large degree node is approximately equal to the reciprocal of the stationary probability. Next, under technical but natural assumptions we show that in fact the reciprocal of the stationary probability is an upper bound on the expected hitting time. Without loss of generality, let us consider node \( k \) from the top-\( k \) list (\( d_1 \geq ... \geq d_k \geq d_{k+1} \geq ... \)). Assume also that the initial node is chosen according to the uniform distribution. Let \( H_k \) be the hitting time to node \( k \) and let \( T \) be the time of the first jump (to the uniform distribution). Then, using Wald’s identity, we can write

\[ E_u[H_k] = E_u[\#\text{jumps on } [0, H_k]] E_u[\min\{T, H_k\}], \] (10)

where \( E_u[\cdot] \) is the expectation given the random walk starts from the uniform distribution. We note that

\[ E_u[\min\{T, H_k\}] \leq E_u[T]. \] (11)

Next, we also note that

\[ E_u[\#\text{jumps on } [0, H_k]] = \frac{1}{P_u[H_k \leq T]}. \] (12)

To estimate the probability \( P_u[H_k \leq T] \), we assume that node \( k \) can be visited at most once during any cycle. This is a natural technical assumption, if \( \alpha \) is not too small, and consequently, the cycles are not too large. In particular, this is the case if we choose the value of \( \alpha \) as the average degree. Then, we have

\[ P_u[H_k \leq T] = P_u[\text{node } k \text{ is on a cycle}] = E_u[\#\text{visits to } k \text{ before and incl. } T] = \pi_k(\alpha) E_u[T]. \]

Combining the above equation with (10), (11) and (12), we obtain

\[ E_u[H_k] \leq \frac{1}{\pi_k(\alpha)} = \frac{2|E| + n\alpha}{d_k + \alpha}. \] (13)

In particular, choosing \( \alpha = \bar{d} \) in (13) yields

\[ E_u[H_k] \leq \frac{2\bar{d}n}{d_k + \bar{d}}. \] (14)
The number of random walk steps, \( m \), is a crucial parameter. Our experiments indicate that we obtain a top \( k \) list with many correct elements with high probability if we take the number of random walk steps to be twice or thrice as large as the expected hitting time of the nodes in the top \( k \) list. This observation can be made rigorous thanks to the result from [7, Ch.9,p.333] that we can adapt for our situation as follows.

**Proposition 1** Let \( H_1, \ldots, H_k \) denote the hitting times to the top-\( k \) nodes with the largest degrees \((d_1 \geq \ldots \geq d_k \geq d_{k+1} \geq \ldots)\). Then, the expected time, \( E_u[\tilde{H}] \), for the random walk with transition probabilities (1) and starting from the uniform distribution to detect a fraction \( \beta \) of top-\( k \) nodes is bounded by

\[
E_u[\tilde{H}] \leq \frac{1}{1 - \beta} E_u[H_k]. \tag{15}
\]

From Theorem 1 or bound (14), we know that the expected hitting time of a large degree node is related to the value of the node’s degree. Thus, the problem of choosing \( m \) reduces to the problem of estimating the values of the largest degrees. We address this problem in the following section.

3 Estimating the largest degrees in the configuration network model

The estimations for the values of the largest degrees can be derived in the configuration network model [9] with a power law degree distribution. In some applications the knowledge of the power law parameters might be available to us. For instance, it is known that web graphs have power law degree distribution and we know typical ranges for the power law parameters (see e.g., [4]).

We assume that the node degrees \( D_1, \ldots, D_n \) are i.i.d. random variables with a power law distribution \( F \) and finite expectation \( E[D] \). Let us determine the number of links contained in the top \( k \) nodes. Denote

\[
F(x) = P[D \leq x], \quad \bar{F}(x) = 1 - F(x), \quad x \geq 0.
\]

Further let \( D_{(1)} \geq \ldots \geq D_{(n)} \) be the order statistics of \( D_1, \ldots, D_n \). Under the assumption that \( D_j \)'s obey a power law, we use the results from the extreme value theory as presented in [13], to state that there exist sequences of constants \( (a_n) \) and \( (b_n) \) and a constant \( \delta \) such that

\[
\lim_{n \to \infty} n\bar{F}(a_n x + b_n) = (1 + \delta x)^{-1/\delta}. \tag{16}
\]

This implies the following approximation for high quantiles of \( F \), with exceedance probability close to zero [13]:

\[
x_p \approx a_n \frac{(pm)^{-\delta} - 1}{\delta} + b_n.
\]
For the $j$th largest degree, where $j = 2, \ldots, k$, the estimated exceedance probability equals $(j - 1)/n$, and thus we can use the quantile $x_{(j-1)/n}$ to approximate the degree $D_{(j)}$ of this node:

$$D_{(j)} \approx a_n \frac{(j - 1)^{-\delta} - 1}{\delta} + b_n. \quad (17)$$

The sequences $(a_n)$ and $(b_n)$ are easy to find for a given shape of the tail of $F$. Below we derive the corresponding results for the commonly accepted Pareto tail distribution of $D$, that is,

$$\bar{F}(x) = Cx^{-\gamma} \text{ for } x > x', \quad (18)$$

where $\gamma > 1$ and $x'$ is a fixed sufficiently large number so that the power law degree distribution is observed for nodes with degree larger than $x'$. In that case we have

$$\lim_{n \to \infty} n\bar{F}(a_n x + b_n) = \lim_{n \to \infty} nC(a_n x + b_n)^{-\gamma}$$

$$= \lim_{n \to \infty} (C^{-1/\gamma}n^{-1/\gamma}a_n x + C^{-1/\gamma}n^{-1/\gamma}b_n)^{-\gamma},$$

which directly gives (16) with

$$\delta = 1/\gamma, \quad a_n = \delta C^\delta n^\delta, \quad b_n = C^\delta n^\delta. \quad (19)$$

Substituting (19) into (17) we obtain the following prediction for $D_{(j)}$, $j = 2, \ldots, k$, in the case of the Pareto tail of the degree distribution:

$$D_{(j)} \approx C^{1/\gamma}(j - 1)^{-1/\gamma}n^{1/\gamma}. \quad (20)$$

It remains to find an approximation for $D_{(1)}$, the maximal degree in the graph. From the extreme value theory it is well known that if $D_1, \ldots, D_n$ obey a power law then

$$\lim_{n \to \infty} P\left( \frac{D_{(1)} - b_n}{a_n} \leq x \right) = H_\delta(x) = \exp(-(1 + \delta x)^{-1/\delta}),$$

where, for Pareto tail, $a_n, b_n$ and $\delta$ are defined in (19). Thus, as an approximation for the maximal node degree we can choose $a_n x + b_n$ where $x$ can be chosen as either a mean, a median or a mode of $H_\delta(x)$. If we choose the mode, $((1 + \delta)^{-\delta} - 1)/\delta$, then we obtain an approximation, which is smaller than the one for the 2nd largest degree. Further, the mean $(\Gamma(1 - \delta) - 1)/\delta$ is very sensitive to the value of $\delta = 1/\gamma$, especially when $\gamma$ is close to one, which is often the case in complex networks. Besides, the parameter $\gamma$ is hard to estimate with high precision. Thus, we suggest to choose the median $\log(2)^{-\delta} - 1)/\delta$, which is less sensitive to the value of $\delta$. This yields

$$D_{(1)} \approx a_n \frac{\log(2)^{-\delta} - 1}{\delta} + b_n = C^{1/\gamma}\log(2)^{-1/\gamma}n^{1/\gamma}. \quad (21)$$
For instance, in the PA network $\gamma = 2.5$ and $C = 3.7$, which gives according to (21) $D(1) \approx 195$. (This is a reasonably good prediction even though the PA network is not generated according to the configuration model. We also note that even though the extremum distribution in the preferential attachment model is different from that of the configuration model their ranges seem to be quite close [12].) This in turn suggests that for the PA network $m$ should be chosen in the range 6000-18000 if $\alpha = 2$. As we can see from Figure 2 this is indeed a good range for the number of random walk steps. In the UK network $\gamma = 1.7$ and $C = 90$, which gives $D(1) \approx 329820$ and suggests a range of 20000-30000 for $m$ if $\alpha = 28.6$. Figure 3 confirms that this is a good choice. The degree distribution of the DBLP network does not follow a power law so we cannot apply the above reasoning to it.

We conclude this section with a remark that from equation (20), bound (14) and Proposition 1, it follows that we can find a $\beta$ fraction of top-$k$ largest degree nodes in sublinear expected time in the configuration model. That is, we have

$$E_u[\tilde{H}] \leq \frac{1}{1 - \beta} \left( \frac{2E[|D|]n}{C^{1/\gamma}(k - 1)^{-1/\gamma}n^{1/\gamma} + E[|D|]} \right) \sim \tilde{C}n^{\frac{2-\beta}{\gamma}}.$$ 

In particular, if $\gamma$ is close to one (which is often the case in complex networks), the computational savings with respect to the deterministic approach can be very significant. For instance, for the UK network with $k = 10$ and $\beta = 0.8$ the bound (15) gives

$$E_u[\tilde{H}] \leq 72531,$$

which means at least 255 fold computational savings.

4 Stopping criteria

Suppose now that we do not have any information about the range for the largest $k$ degrees. In this section we design stopping criteria that do not require knowledge about the structure of the network. As we shall see, knowledge of the order of magnitude of the average degree might help, but this knowledge is not imperative for a practical implementation of the algorithm.

Let us now assume that node $j$ can be sampled independently with probability $\pi_j(\alpha)$ as in (2). There are at least two ways to achieve this practically. The first approach is to run the random walk for a significant number of steps until it reaches the stationary distribution. If one chooses $\alpha$ reasonably large, say the same order of magnitude as the average degree, then the mixing time becomes quite small [1] and we can be sure to reach the stationary distribution in a small number of steps. Then, the last step of a run of the random walk will produce an i.i.d. sample from a distribution very
close to (2). The second approach is to run the random walk uninterruptedly, also with a significant value of $\alpha$, and then perform Bernoulli sampling with probability $q$ after a small initial transient phase. If $q$ is not too large, we shall have nearly independent samples following the stationary distribution (2). In our experiment, $q \in [0.2, 0.5]$ gives good results when $\alpha$ has the same order of magnitude as the average degree.

We now estimate the probability of detecting correctly the top $k$ list of nodes after $m$ i.i.d. samples from (2). Denote by $X_i$ the number of hits at node $i$ after $m$ i.i.d. samples. We note that if we use the second approach to generate i.i.d. samples, we spend approximately $m/q$ steps of the random walk. We correctly detect the top $k$ list with the probability given by the multinomial distribution

$$P[X_1 \geq 1, \ldots, X_k \geq 1] = \sum_{i_1 \geq 1, \ldots, i_k \geq 1} \frac{m!}{i_1! \cdots i_k!(m - i_1 - \ldots - i_k)!} \pi_1^{i_1} \cdots \pi_k^{i_k} (1 - \sum_{i=1}^k \pi_i)^{m - i_1 - \ldots - i_k}$$

but it is not feasible for any realistic computations. Therefore, we propose to use the Poisson approximation. Let $Y_j$, $j = 1, \ldots, n$ be independent Poisson random variables with means $\pi_j m$. That is, the random variable $Y_j$ has the following probability mass function $P[Y_j = r] = e^{-m\pi_j} (m\pi_j)^r / r!$. It is convenient to work with the complementary event of not detecting correctly the top $k$ list. Then, we have

$$P[\{X_1 = 0\} \cup \ldots \cup \{X_k = 0\}] \leq 2P[\{Y_1 = 0\} \cup \ldots \cup \{Y_k = 0\}]$$

$$= 2(1 - \prod_{j=1}^k P[\{Y_j \geq 1\}]) = 2(1 - \prod_{j=1}^k \left(1 - e^{-m\pi_j}\right)) =: a,$$  

where the first inequality follows from [14, Thm 5.10]. In fact, in our numerical experiments we observed that the factor 2 in the first inequality is very conservative. For large values of $m$, the Poisson bound without 2 works very well as proper approximation.

For example, if we would like to obtain the top 10 list with at most 10% probability of error, we need to have on average 4.5 hits per each top element. This can be used to design the stopping criteria for our random walk algorithm. Let $\bar{a} \in (0, 1)$ be the admissible probability of an error in the top $k$ list. Now the idea is to stop the algorithm after $m$ steps when the estimated value of $a$ for the first time is lower than the critical number $\bar{a}$. Clearly,

$$\hat{a}_m = 2(1 - \prod_{j=1}^k (1 - e^{-X_j}))$$
is the maximum likelihood estimator for $a$, so we would like to choose $m$ such that $\hat{a}_m \leq \bar{a}$. The problem, however, is that we do not know which $X_j$’s are the realisations of the number of visits to the top $k$ nodes. Then let $X_{j_1}, ..., X_{j_k}$ be the number of hits to the current elements in the top $k$ candidate list and consider the estimator

$$\hat{a}_{m,0} = 2(1 - \prod_{i=1}^{k}(1 - e^{-X_{ji}})),$$

which is the maximum likelihood estimator of the quantity

$$2(1 - \prod_{i=1}^{k}(1 - e^{-m\pi_{ji}})) \geq a.$$

(Here $\pi_{ji}$ is a stationary probability of the node with the score $X_{ji}$, $i = 1, \ldots, k$). The estimator $\hat{a}_{m,0}$ is computed without knowledge of the top $k$ nodes or their degrees, and it is an estimator of an upper bound of the estimated probability that there are errors in the top $k$ list. This leads to the following stopping rule.

**Stopping rule 0.** Stop at $m = m_0$, where

$$m_0 = \arg\min\{m : \hat{a}_{m,0} \leq \bar{a}\}.$$

The above stopping criterion can be simplified even further to avoid computation of $\hat{a}_{m,0}$. Since

$$\hat{a}_{m,1} := 2(1 - (1 - e^{-X_{jk}})^k) \geq \hat{a}_{m,0} \geq \bar{a},$$

where $X_{jk}$ is the number of hits of the worst element in the candidate list. The inequality $\hat{a}_m \leq \bar{a}$ is guaranteed if $\hat{a}_{m,1} \leq \bar{a}$. This leads to the following stopping rule for the random walk algorithm.

**Stopping rule 1.** Compute $x_0 = \arg\min\{x \in \mathbb{N} : (1 - e^{-x})^k \geq 1 - \bar{a}/2\}$. Stop at

$$m_1 = \arg\min\{m : X_{jk} = x_0\}.$$

We have observed in our numerical experiments that we obtain the best trade off between the number of steps of the random walk and the accuracy if we take $\alpha$ around the average degree and the sampling probability $q$ around $0.5$. Specifically, if we take $\bar{a}/2 = 0.15$ ($x_0 = 4$) in Stopping rule 1 for top 10 list, we obtain 87% accuracy for an average of 47 000 random walk steps for the PA network; 92% accuracy for an average of 174 468 random walk steps for the DBLP network; and 94% accuracy for an average of 247 168 random walk steps for the UK network. We have averaged over 1000 experiments to obtain tight confidence intervals.
5 Relaxation of top \( k \) lists

In the stopping criteria of the previous section we have strived to detect all nodes in the top \( k \) list. This costs us a lot of steps of the random walk. We can significantly gain in performance by relaxing this strict requirement. For instance, we could just ask for list of \( k \) nodes that contains 80% of top \( k \) nodes [3]. This way we can take an advantage of a generic 80/20 rule that 80% of result can be achieved with 20% of effort.

Let us calculate the expected number of top \( k \) elements observed in the candidate list up to trial \( m \). Define by \( X_j \) the number of times we have observed node \( j \) after \( m \) trials and

\[
H_j = \begin{cases} 
1, & \text{node } j \text{ has been observed at least once}, \\
0, & \text{node } j \text{ has not been observed}.
\end{cases}
\]

Assuming we sample in i.i.d. fashion from the distribution (2), we can write

\[
E[\sum_{j=1}^{k} H_j] = \sum_{j=1}^{k} E[H_j] = \sum_{j=1}^{k} P[X_j \geq 1] = \sum_{j=1}^{k} (1 - (1 - \pi_j)^m).
\]

(23)

In Figure 2 we plot \( E[\sum_{j=1}^{k} H_j] \) (the curve “I.I.D. sample”) as a function of \( m \) for \( k = 10 \) for the PA network with \( \alpha = 0 \) and \( \alpha = 2 \). In Figure 3 we plot \( E[\sum_{j=1}^{k} H_j] \) as a function of \( m \) for \( k = 10 \) for the UK network with \( \alpha = 0.001 \) and \( \alpha = 28.6 \). The results for the UK and DBLP networks are similar in spirit.

![Figure 2: Average number of correctly detected elements in top-10 for PA.](image)

Here again we can use the Poisson approximation

\[
E[\sum_{j=1}^{k} H_j] \approx \sum_{j=1}^{k} (1 - e^{-m\pi_j}).
\]
In fact, the Poisson approximation is so good that if we plot it on Figures 2 and 3, it nearly covers exactly the curves labeled “I.I.D. sample”, which correspond to the exact formula (23). Similarly to the previous section, we can propose stopping criteria based on the Poisson approximation. Denote

\[ b_m = \sum_{i=1}^{k} (1 - e^{-X_{j_i}}). \]

**Stopping rule 2.** Stop at \( m = m_2 \), where

\[ m_2 = \arg \min \{ m : b_m \geq \bar{b} \}. \]

Now if we take \( \bar{b} = 7 \) in Stopping rule 2 for top-10 list, we obtain on average 8.89 correct elements for an average of 16 725 random walk steps for the PA network; we obtain on average 9.28 correct elements for an average of 66 860 random walk steps for the DBLP network; and we obtain on average 9.22 correct elements for an average of 65 802 random walk steps for the UK network. (We have averaged over 1000 experiments for each network.) This makes for the UK network the gain of more than two orders of magnitude in computational complexity with respect to the deterministic algorithm.

6 Conclusions and future research

We have proposed the random walk method with the candidate list for quick detection of largest degree nodes and analyzed the complexity of the method by means of random walk hitting times. We have also supplied stopping criteria which do not require knowledge of the graph structure. In the case of large networks, our algorithm finds top-\( k \) list of largest degree nodes with
few mistakes with the running time orders of magnitude faster than the
deterministic algorithms. In future research we plan to obtain estimates
for the required number of steps for various types of complex networks and
to design methods for directed networks. In particular, it is interesting to
analyze in more detail how assortativity and clustering of networks affects
the performance of the method.

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