Sampling from Social Networks with Attributes

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ABSTRACT

Sampling from large networks represents a fundamental challenge for social network research. In this paper, we explore the sensitivity of different sampling techniques (node sampling, edge sampling, random walk sampling, and snowball sampling) on social networks with attributes. We consider the special case of networks (i) where we have one attribute with two values (e.g., male and female in the case of gender), (ii) where the size of the two groups is unequal (e.g., a male majority and a female minority), and (iii) where nodes with the same or different attribute value attract or repel each other (i.e., homophilic or heterophilic behavior). We evaluate the different sampling techniques with respect to conserving the position of nodes and the visibility of groups in such networks. Experiments are conducted both on synthetic and empirical social networks. Our results provide evidence that different network sampling techniques are highly sensitive with regard to capturing the expected centrality of nodes, and that their accuracy depends on relative group size differences and on the level of homophily that can be observed in the network. We conclude that uninformed sampling from social networks with attributes thus can significantly impair the ability of researchers to draw valid conclusions about the centrality of nodes and the visibility or invisibility of groups in social networks.

Keywords: social networks; sampling methods; sampling bias; homophily

1. INTRODUCTION

Sampling from large networks represents a fundamental problem for social network research. In order to draw valid conclusions from network samples, understanding how accurately samples reflect the position of nodes in the original network is essential. Previous research has studied robustness of network samples from different angles, for example by examining the accuracy of network measures such as degree or betweenness centrality. A range of network properties has been found to be sensitive to the choice of sampling methods [4, 6, 11, 13, 15, 16, 18, 30].

Motivation and problem. In this paper, we focus on the specific problem of sampling nodes and edges from a social network with attributes, i.e., a network where nodes are colored. For example, the color of nodes might be determined by gender, ethnicity, or age. We consider the special case of networks (i) where one binary attribute can be observed (e.g., a male and a female group of nodes), (ii) where the size of the two groups is unequal (e.g., a male majority and a female minority), and (iii) where nodes with the same or different attribute value attract or repel each other, i.e., homophilic [26] or heterophilic networks [3]. While the general impact of sampling on network characteristics has been studied thoroughly in the past [4, 6, 11, 13, 15, 16, 30], the role of attributes in combination with fundamental social mechanisms such as homophily [21, 27] has only received little attention so far [19]. In fact little is known about whether or how different sampling techniques are able to conserve the ranking of nodes or the visibility of groups from the original network. Accurately capturing network characteristics of groups of nodes in sampled data, however, is crucial not only for researchers interested in directly studying these groups (e.g., gender or sociological studies), but also for researchers interested in analyzing the structure of the complete network since attributes of actors can impact the overall network structure [5, 21, 27].

Research questions. In this paper, we thus ask: How sensitive are different sampling techniques with respect to conserving the ranking of nodes and the visibility of groups in synthetic and empirical social networks with (i) different minority and majority group proportions, and (ii) various levels of homophily?

Methods and materials. We evaluate different sampling techniques (node sampling, edge sampling, random walk sampling, and snowball sampling) with respect to reflecting the ranking of nodes and the visibility of groups in network samples (see Figure 1). Instead of putting the focus on the whole population as in previous work, we specifically focus on sub-populations (or groups); we call the larger group majority and the smaller group minority. Our work is guided by the intuition that an ideal sample would allow to accurately preserve the original degree centrality ranking of nodes, and therefore preserve the relative importance between nodes and groups. That means, an ideal sample would not systematically rank nodes of one group higher and nodes of the other group lower than expected. This would be considered a biased sample or sampling error.
We additionally showcase observed artifacts on empirical networks. Based on the obtained insights, we provide indicators very popular minority) does not only fail to capture the sensitivity of different network measures with respect to miss-

We construct synthetic social networks and vary the structural mechanisms guiding the growth of the network (i.e., homophily, preferential attachment, and group sizes), to study the extent to which they impact the accuracy of samples. We additionally showcase observed artifacts on empirical networks. Based on the obtained insights, we provide indicators of why samples might have issues with capturing expected group characteristics.

Contributions and Findings. (i) We propose a method to measure the robustness of samples from networks with two attributes. (ii) Using synthetic and empirical networks, we provide evidence that different network sampling techniques have issues with capturing the expected centrality of nodes and the visibility of minority / majority groups in social networks. (iii) We discuss network characteristics that lead to observed discrepancies and quantify the impact of relative group size differences and homophily on sampling errors.

2. BACKGROUND AND RELATED WORK

Network analysis has long been plagued by issues of measurement error, usually in the form of missing data. Understanding the robustness of basic network measures is extremely important in order to assess the validity of network research. Prior research explored the impact of missing data on various network measures, but mainly focused on small sociometric networks [6, 11], small bipartite collaboration graphs [15], and random networks [4, 15].

Smith and Moody [28] extended this line of research and analyzed four classes of network measures on 12 relatively small (< 1000 nodes) empirical networks. They found that larger, more centralized networks, are in general more robust to missing nodes at random, especially for centrality and centralization measures. This is plausible since random node deletion in a centralized network (with skewed degree distribution) is less likely to remove hubs since few of them exist. In our work, we do not explore the effect of random node deletion, but, compare different sampling methods. Node sampling is the opposite of random node deletion, since the randomly selected nodes are included in the sample. Our results throughout this paper show that random node sampling from centralized networks (heterophilic networks with very popular minority) does not only fail to capture the centralization of the network well (since we miss the hubs), but also fails to accurately capture the relative importance of groups.

Wang et al [30] presented the first work that explores the sensitivity of different network measures with respect to missing data in two large online social networks and one random graph. They defined six different types of measurement errors (missing nodes, spurious nodes, missing edges, spurious edges, falsely aggregated nodes, and falsely disaggregated nodes) and simulated their effect on the complete network. Using Spearman rank correlation, the authors compared the list of nodes that is ranked based on the network measure in the original network with the one that is computed on the sample. The work finds support for Borgatti's findings [4], highlighting that different centrality measures are similarly robust to measurement errors. Interestingly, results show that more local network measures like clustering are more prone to missing data than more global measures such as centralities. Thus, the authors revised the general claim from past research that the more “global” a measure, the less resistant it is to measurement error.

Lee et al. [17] analyzed scale-free networks and three empirical networks suggesting that network properties such as betweenness centrality or clustering are sensitive to the choice of sampling method. Lee and Pfeffer [16] explored the quality of sampling by comparing the node-level network scores induced from the sample and the original network. They used edge-sampling and focused on degree and betweenness centrality for two empirical communication networks. Their results show that larger samples lead to high sampling accuracy and that centralized graphs in which fewer nodes enjoy higher attention offer more accurate samples when edge sampling is used. Our work extends their work, since we compare various sampling techniques and introduce groups and homophily.

Furthermore, Leskovec and Faloutsos [18] showed that network properties are sensitive to the choice of the sampling method. However, they assessed the quality of a sample by comparing the shape of the distribution of a network measure (e.g., degree) in the sample with the original one using the Kolmogorov-Smirnov Distance. This evaluation criterion is very different from what has been used in previous work and what we use in this work, since it does not take the accuracy of the ranking of nodes into account.

Most prior work shows that network estimates become more inaccurate with lower sample coverage, but there is a wide variability of these effects across different measures, network topologies and sampling errors. To our best knowledge, most previous work neglected the existence of heterogeneous attributes in networks and did not analyze the interplay between mechanisms that impact the topology of a social network and the accuracy of sampling techniques. A mention-
able exception is the work by Li and Ye [19] who explored the ratio of intra- and inter-group links in samples drawn from a sample of the follow-network of Twitter users. Our work extends their work by systematically exploring the effect of group sizes and homophily on the visibility of individual nodes. Our work focuses on undirected networks, but the work by Huisman [13] provides a comparison of sample bias in directed and undirected versions of the same network.

3. METHODS

In this work, we are interested in studying the accuracy of samples drawn from networks with unequally sized groups and various levels of homophily. We (i) describe used sampling techniques and (ii) explain how we assess the accuracy of a sample.

3.1 Sampling techniques

Our goal is to sample $K$ nodes from the overall set of $N$ nodes in a network. As pointed out in [18], we can split sampling algorithms into three groups: methods based on randomly selecting nodes, randomly selecting edges, and exploration techniques simulating random walks or virus propagation to find a representative sample of nodes. We focus on one sampling technique from each group:

Random node sampling. This is the most basic sampling technique where a random subset of $K$ nodes is selected. The sampled network then contains these $K$ nodes and all links between them. Random node sampling is e.g., used when a sample of individuals is first selected and then their contact behavior is observed. Numerous surveys and data collections use this method, e.g., measuring contact pattern among high school students using wearable sensors [20].

Random edge sampling. This strategy randomly samples edges from the network and filters the complete network by sampled edges. To be consistent with the other sampling strategies, we successively sample edges until $K$ nodes are selected. The sampled network then contains these $K$ nodes and sampled links, but not those links between selected nodes that have not been sampled. Random edge sampling is commonly used to construct a social graph by using information about contacts—e.g., phone calls are sampled and a graph of callers and receivers is constructed [12].

Snowball sampling. In snowball sampling, we randomly sample one starting node and add all its neighbors as well as the neighbors’ neighbors to the set of sampled nodes—i.e., two step snowball sampling. We repeat this until we have gathered $K$ nodes for the sample. If a full iteration does not catch $K$ nodes, we repeat the process again with a new randomly selected starting node. The sampled network then contains these $K$ nodes and all the links connecting them. Traditionally, snowball sampling is used when the population under study is not easily accessible (e.g., to study homeless people or illegal immigrants). Indeed, the promise of the snowball sampling is to access hard-to-reach population [1].

Random walk (RW) sampling. This strategy samples nodes by walking through the network. The walker starts at a random node in the network and chooses in each step one out-going link randomly and traverses it. All visited nodes are then added to the sample until $K$ nodes have been added. A teleport probability can be set for teleporting to another random node in the network instead of traversing a link in this iteration; we use 0.15 throughout this work. The sampled network then contains these $K$ nodes and all links between them. This technique of sampling is usually used in online social networks such as Facebook or Twitter, in which retrieving information about the whole population is overwhelming and computationally costly, but we can access and navigate the original network.

3.2 Evaluation measures

The ubiquity of sampled network data makes the understanding of the robustness of network measures crucial. Here, we focus on the most basic and widely used centrality measure: degree centrality [10]. The degree centrality of a node is defined as the fraction of nodes it is connected to.

Previous work explored the robustness of centrality measures in samples of networks without taking heterogeneous attributes of nodes into account. Therefore, simple rank correlation (see e.g., [6, 16, 28, 30]) and overlap measures (see e.g., [4]) have been used to assess how well a sample captures the ranking of nodes according to various network measures. In this work, we are interested in assessing how well a sample captures, on average, the overall position of nodes in the original network for each group of nodes separately. That means, we aim to reveal if the positions of nodes in both groups are equally well captured in a way that the relative group and node importance are preserved.

If we would compute the overall rank correlation (or overlap) between the two lists and ignore the group memberships, then the ranking of majority nodes would contribute more to the correlation coefficient (or overlap). A naive group-specific measure would be to compute a separate rank correlation (or overlap) for each group. However, this measure would only allow us to assess how well the relative importance of nodes within each group in the original network is preserved in the sample, but the relation between nodes across groups would be neglected. Therefore, simple rank correlation or overlap measures cannot be used to assess whether the relevance of nodes and groups is accurately captured in a sample.

In this work we define an ideal sample as a sample that allows to accurately reconstruct the original degree centrality ranking of nodes and therefore preserves the relative importance between nodes and groups. That means, an ideal sample does not systematically rank nodes of one group higher and nodes of the other group lower than expected. To assess the accuracy of the relative importance of nodes and groups, we propose the following two evaluation measures. Both evaluation measures focus on the top $k$ or top $k$ percent of the data, since (i) users focus on the first few results in ranked lists and (ii) the distribution of degree centralities are usually heavy tail distributions. Therefore, the contribution of disorders in the long tail (unpopular nodes) would dominate disorders in the head (popular nodes) if we would not limit our analysis to the head [32].

Top $k$ bias. To assess the accuracy of group visibility in a sample, we compare the fraction of minority nodes in the top $k$ nodes of a sample with its fraction in the top $k$ nodes of the complete network.

$$\text{bias}_{topk} = \text{expected}_{topk} - \text{observed}_{topk}$$

Expected refers to the fraction of minority nodes that we observe in the top $k$ nodes of the sample, while observed refers to the fraction of minority nodes in the top $k$ nodes of the original network. As sample size grows, the observed fraction in the sample approaches the expected fraction.
We analyze the log of the normalized cumulative relevance.

We cannot measure how much lower the ranking of a node is in the sample compared to its ranking in the complete network. To overcome this limitation, we first compute the relevance for each node $i$ by ranking nodes based on their centrality in the original network. The relevance of node $i$ is defined as the inverse rank that belongs to node $i$ normalized by the rank sum of all nodes ($N$) in the original network:

$$\text{rel}_i = \frac{\text{inv_rank}_i}{\sum_{j=1}^{N} \text{rank}_j} \tag{2}$$

The relevance shrinks linearly with the position of nodes in the list, but different weighting is possible. We compute for each group $g$ its cumulative group relevance ($\text{CGR}$) at rank $k$ in the original ranked list and compare it with the cumulative relevance at rank $k$ in the sample:

$$\text{CGR}_{topk}^{	ext{sample}} = \sum_{j=1}^{k} \text{rel}_{j \in g} \tag{3}$$

$$\text{nCGR}_{topk} = \frac{\text{CGR}_{topk}^{	ext{sample}} + \epsilon}{\text{CGR}_{topk}^{	ext{original}} + \epsilon} \tag{4}$$

The $\text{nCGR}_{topk}$ measures the extent to which the relevance of a group in the sample is above or below what we would expect from the original network with respect to the top $k$ nodes. If e.g., this normalized cumulative group relevance for the minority is $2$, then that means that the minority is twice as relevant in the sample than in the original network (for some top $k$). If it is $0.5$ then the group is half as relevant in the sample than in the original network. If it is $1$ then the group has equal relevance in the original network and the sample. We analyze the log of the normalized cumulative relevance since otherwise the measure is bound by zero; thus, the ideal nCGR is zero. To avoid division by zero and logarithm of zero, we add a small $\epsilon = 0.001$.

4. SIMULATION EXPERIMENTS

We construct synthetic networks and explore the effect of homophily and group size on the accuracy of samples in a controlled environment. First, we describe the network model which we use to create synthetic network data and second, we discuss the accuracy of centrality measures in samples drawn from these networks using different sampling methods.

4.1 Synthetic network generators

Preferential attachment (the tendency of nodes to connect to popular nodes) [2, 33] and homophily (the tendency of nodes to connect to similar nodes) [21, 27] have been extensively observed in many real-world social networks [7, 9, 23, 31] and information networks [22, 24]. Homophily implies the existence of at least one fixed or mutable attribute (e.g., gender, ethnicity, education status). Based on these attributes similarities between nodes can be defined.

We use an existing preferential attachment growth model with a homophily parameter that can be tuned and thus allows us to create networks with different levels of homophily and heterophily (see [8, 14] for details). The homophily parameter $h$ ranges between $0$ to $1$, $h \in [0,1]$, where $0$ means that nodes are only attracted by nodes that are dissimilar to them (heterophily), $1$ means nodes prefer to connect with similar nodes (homophily), and $0.5$ means that the link formation behavior is not driven by attributes. All nodes of the same group share the same homophily parameter $h$, because they share the same attribute value and thus have the same distance to other groups with different attribute values.

We generate all synthetic networks with $10,000$ nodes and a fixed minority ratio of $20\%$ (except when noted otherwise). An incoming node connects to $10$ nodes based on a specific homophily parameter and popularity (see [14]).

Figure 2 shows the degree distribution of both groups of nodes in networks that only vary in their degree of homophily. One can see that if we have two groups of unequal size and the network is heterophilic ($h < 0.5$), the degree distributions of majority and minority differ the most.

Figure 2: Degree distribution of synthetic networks. The average degree distribution of majority ($80\%$ of nodes) and minority ($20\%$ of nodes) in a synthetically generated preferential attachment network with various levels of homophily. One can see that the degree distributions are almost equal if homophily does not play a role ($h = 0.5$). In heterophilic networks ($h < 0.5$) the group-specific differences are much more pronounced than in homophilic networks ($h > 0.5$).
If we compare the degree distribution of the two groups in a moderate heterophilic network with $h = 0.25$ and a moderate homophilic network with $h = 0.75$, we see that the differences between the degree distributions are more pronounced in the heterophilic case. This asymmetric effect can be explained by the interplay between group size differences and homophily. The majority benefits from moderate homophily (e.g., $h = 0.75$) more than from high homophily (e.g., $h = 0.9$), because in high homophily conditions, their maximum degree is bound to the size of their group, while in moderate homophily conditions, sometimes also minority nodes will be attracted by the high degree of majority nodes. Unlike the majority in the homophilic case, the minority in the heterophilic case benefits more from extreme heterophily (i.e., $h = 0$) than from moderate heterophily (e.g., $h = 0.25$). That is because in the extreme heterophily condition, all majority nodes are attracted by the minority, but in moderate heterophily condition sometimes the majority is attracted by high degree nodes which can also be part of their group. So for the minority to gain popularity, it is better if they do not have to compete with the majority while the majority benefits from a competitive environment. In the next section, we will analyze how these group-specific differences in the degree distributions relate to sample biases.

### 4.2 Sample bias in synthetic networks

To assess sample bias, we generate synthetic networks, draw samples of varying size from them using different sampling techniques and assess the average visibility and relevance of different groups in samples. We repeat the random network generation process 10 times and draw 10 samples from each network; thus, in our evaluation, we report mean and standard error over 100 samples.

Figure 3 shows the visibility of the minority group in the top 100 nodes in samples of different size which have been created via different sampling methods. For example, in Figure 3 (a), the point for the green line at an $x$-value of 0.10 indicates that the top 100 ranked nodes based on degree centrality in a 10% sample from a moderate heterophilic network with $h = 0.25$, contains on average around 40% minority nodes. We can compare this observed percentage with the expected percentage from the original network (100% sample). In this case, we would expect to see close to 80% of minority nodes in the top 100 nodes indicating that the minority is underrepresented in small samples drawn from moderate heterophilic networks with unbalanced group sizes using node sampling.

Results show that especially node and snowball sampling reduce the visibility of minority groups in the top $k$ list if samples are drawn from extreme and moderate heterophilic networks. For node sampling, this is not surprising since all nodes have equal probability to be picked and therefore, a node’s sampling probability is proportional to its group size. Snowball samples aggregate the 2-hop neighbourhood of randomly selected seed nodes which likely are majority nodes. Since most majority nodes are unpopular (skewed degree distribution), the probability for picking a majority node that has only a few minority nodes as neighbours is high. Thus, we underestimate the visibility of the minority group in the top $k$. Figure 4 shows that in the heterophilic network, the bias of node and snowball samples decreases linearly with decreasing group size difference. Note that group sizes are balanced if the minority ratio is 0.5. We further find that RW samples are very robust against relative size differences between groups in homophilic and heterophilic networks.

In Figure 5 we show to what extent the original relevance of each group is preserved in the sample. We find that in most cases the relevance of the minority is underestimated. Only in moderate homophilic networks, minority is overrepresented. However, one needs to note that the extent with which the relevance of the minority is overestimated in moderate homophilic networks ($h = 0.75$, 4th row) is lower than the extent with which the relevance of the majority is overestimated in moderate heterophilic networks ($h = 0.25$, 2nd row).

Overall, we see that (i) the most accurate samples can be drawn from networks where homophily does not play a role, (ii) RW sampling performs best independent of the homophily conditions (see Figure 3 and 5) and relative group size differences (see Figure 4), (iii) all sampling methods perform similar if group size differences are small, and (iv)
the sampling error is always higher in heterophilic networks than in homophilic networks if the same sampling technique and group size differences are considered.

**Regression analysis.** To compare the impact of different factors on the sampling bias, we fit eight simple linear regression models, one model for each sampling technique and each error measure (top $k$ minority bias $\text{bias}_{topk}$ and the absolute sum of the normalized cumulative group relevance $\text{nCGR}_{topk}$ of the minority and the majority group). Each model was fitted to 3,200 observations (samples drawn from synthetically generated networks). Table 1 shows that across all sampling methods—perhaps not surprisingly—smaller samples lead to higher sampling errors and larger top $k$ lists lead to higher errors because the size of the network is constant. Interestingly, we see that only for node and snowball samples, the sampling error increases, if group size differences and the influence of the attribute on the edge formation behavior (i.e., the homophily parameter is closer to 0 or 1) increase. If only one of these factors changes, no significant effects on the sampling error can be observed, except for snowball samples. The bias of snowball samples also increases significantly if only homophily increases, because in extreme homophilic networks a snowball sample can only contain nodes of one group also if groups are of equal size. One can see that the sampling error of RW and edge samples cannot be explained by group size differences and homophily, which confirms our observation that these methods are rather robust against these factors.

5. **EMPIRICAL EXPERIMENTS**

Next, we analyse two empirical networks and explore the accuracy of samples drawn from these networks. We describe the statistical properties of these networks and contrast empirical findings with the findings obtained from simulation.

5.1 **Pokec social network**

**Dataset.** We study publicly available data\(^1\) obtained from the most popular Slovakian social network “Pokec” [29]. We added all friendship relations as undirected edges. The network contains 1,632,640 nodes (users) and 22,301,602 edges (friendship relations). The average degree of nodes is 27.32, the global clustering coefficient is 0.0069, and the graph diameter is 14.

For our experiments, we focus on the age of actors in the social network. Eliminating all nodes without age information results in a network with 1,138,314 nodes connected by 14,975,771 edges. For coloring nodes as minority and majority, we take the 80% percentile of the overall age distribution, and color all nodes with an age higher than this percentile as belonging to the minority (old users), and all below as belonging to the majority. This results in an age cut-off of 31 years, meaning that the minority—18.8% of all nodes—captures the oldest users in the network. Overall, around 92% of all edges in the network are between nodes of the same color—i.e., between two minority or two majority nodes. This exceeds the expectation of around 81.3% if edges would form totally at random. From that we can assert that the Pokec social network is moderately homophilic with respect to the defined age groups. Figure 6 shows the degree distribution of young and old users. One can see that the most popular users are part of the majority.

**Results.** Figure 7 shows that the visibility of the minority and the relevance of both groups is very well preserved in all samples. This is in line with what our model suggests for very homophilic networks (see Figure 7). Interestingly, random walk sampling produces the most accurate sample, which is also suggested by our model, especially for large relative groups size differences (see Figure 4(c)).

5.2 **Sexual contact network**

**Dataset.** We use a network of claimed sexual contacts between Brazilian escorts (prostitutes) and sex buyers [25].

1[https://snap.stanford.edu/data/soc-pokec.html](https://snap.stanford.edu/data/soc-pokec.html)
Figure 5: Normalized Cumulative Group Relevance. Each column depicts a different sampling technique, while each row refers to a different world for which the homophily level of the original network varies. The axis are aligned within each row, but not within each column, since the extent of error varies depending on the world. Again, each point refers to an average evaluation over 100 total iterations. One can see that in extreme heterophilic networks (first row) the relevance of the majority is overestimated in small and also in larger sized samples, while the relevance of the minority is slightly underestimated especially in small sized samples. In extreme homophilic networks (last row), it is the other way around, however the extent to which the relevance of the minority is overestimated is smaller than the extent to which the relevance of the majority is overestimated in the extreme heterophilic case. Overall, random walk sampling produces the most accurate samples, followed by edge sampling. In samples based on node and snowball sampling, the relevance of the minority is usually underestimated, except in moderate homophilic networks (4th row).
The network consists of 16,730 nodes (6,624 sex workers and 10,106 sex buyers) and 50,632 edges between them. The minority of nodes with a share of around 40% are sex workers, while the majority are sex buyers. The network is fully bi-partite, meaning that sex workers only connect with sex buyers to capture sexual contacts. Consequently, all edges within the networks are between nodes of different color and thus, the network is 100% heterophilic. The degree distributions of minorities and majorities show that minorities are more popular than majorities (see Figure 6). This is not surprising because the network is an example of an extreme heterophilic network since the majority nodes are attracted by the minority nodes and the other way around.

**Results.** Figure 7 shows that the minority (escorts) are very visible in the top 100 nodes ranked by degree centrality also in samples of small size. Node-based samples are the most inaccurate samples, since they underestimate the visibility and relevance of the minority most. Edge-based samples capture the visibility of the minority in the original network best if the original network is extremely heterophilic. Our model suggests that no large differences in the performance of different sampling techniques (as suggested by Figure 4) will exist because group size differences are rather small (40:60); but, edge and RW sampling will produce more accurate samples than node and snowball sampling. Further, we can expect that all samples will underestimate the relevance of the minority. These expectations are confirmed empirically (cf. Figure 7, bottom row).

### Table 1: Coefficients of eight linear regression models, one for each sampling technique and sampling error measure.

Each model was fitted to 3,200 observations (samples drawn from synthetically generated networks). The interaction term between group size difference and attribute influence is significant in node and snowball samples, but not in RW and edge samples. This indicates, the sampling error increases in node and snowball samples if the group size difference and the influence of attributes on the edge formation behavior are both increased. Edge and RW samples are rather robust against these factors. We compute the sampling error for lists of different length $k$ and control for the effect of $k$ in the model. The larger $k$, the higher the error. We also observe on average larger sampling errors on smaller sample sizes. Note: \( ^* p < 0.01; ^{**} p < 0.001 \).

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R2 0.281 0.135 0.418 0.428 0.225 0.152 0.275 0.135
overestimated while in homophilic networks, it is slightly underestimated.

One limitation of our network generation model is that we limit it to two groups and that it assumes that all nodes in a group are equally active and behave equally homophilic or heterophilic. In real world social networks, more groups and group-specific and individual behavioral differences can be present. Future research is necessary to study the effect of group-specific activity difference and asymmetric homophilic behavior and needs to explore the presence of multiple groups. Furthermore, we focus on one specific network measure and undirected networks warranting further explorations about the accuracy of various network measures in samples drawn from directed networks. Our work can be extended to more than one binary attribute by simply defining a similarity function that takes several attributes into account.

7. CONCLUSIONS

In summary, our work shows that the combination of two factors leads to sampling error in social networks with attributes: (i) group size differences and (ii) homophily. If unequal sized groups are present, random walk sampling always leads to the most accurate samples—indeed of the level of homophily. The sampling error is always larger if samples are drawn from heterophilic networks with unequally sized groups compared to homophilic ones. In heterophilic networks with unbalanced groups, random walk and edge sampling perform similar well, while in homophilic networks edge sampling produces more biased samples than random walk sampling. This can be explained by the fact that in homophilic networks edge sampling overestimates the importance of minority nodes, since minority nodes with high degree are more likely to be selected. Edge samples only include sampled edges, but not all other edges between selected node. Therefore, the difference in degree between minority and majority nodes can be skewed. Most sampling techniques produce accurate samples if the groups are of equal size. Only snowball samples can also be biased if homophily is a driving force behind the edge formation of nodes that belong to two equally sized groups.

Since researchers often do not have information about group size differences and homophily in the original network, random walk sampling is a robust choice. However, researchers cannot always choose their sampling method freely. Therefore, our results provide important guidance in estimating which groups will be over- or underestimated in samples drawn from social networks with unequally sized groups and various level of homophily. It is our hope that the research presented in this paper motivates more research into sampling from social networks with attributes.

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References


