Framework and Algorithms for Network Bucket Testing

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ABSTRACT

Bucket testing, also known as split testing, A/B testing, or 0/1 testing, is a widely used method for evaluating users' interaction with new features, products, or services. Usually, a small set of uniformly randomly chosen users are given the new service and the overall satisfaction rate is evaluated from the sample. In a recent work, Backstrom and Kleinberg, defined the notion of network bucket testing. Here the services are social in nature and users' satisfaction is only valid for measurement if some minimal number of their friends are also given the service (core set users). The goal is to estimate the mean user satisfaction rate while providing the service to the least number of users. This challenging problem is becoming increasingly relevant with the growing popularity of social networks.

In this paper we introduce a simple general framework for evaluating network bucket testing algorithms. The framework is constructed in a way that testing algorithms are only required to produce core sets of users. Given an algorithm, the framework produces an unbiased user satisfaction rate estimator and a corresponding variance bound for any network and any user satisfaction function. Furthermore, we present several simple testing algorithms which are evaluated using both synthetic and real social networks. Our experiments corroborate the theoretical results, and demonstrate the effectiveness of the proposed framework and algorithms.

Keywords

Social networks, Bucket testing, A/B testing, Network Bucket Testing, Unbiased estimation

1. INTRODUCTION

Bucket testing, a.k.a. split testing, A/B testing, or 0/1 testing, is a well know and widely used method for evaluating user engagement or satisfaction from a new service, feature, product etc. Before releasing a new service, providers often choose a small subset of users to which the service is

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given. Based on these users' behavior, the overall satisfaction of users can be estimated. If the satisfaction is high enough, the service is released to the entire user population. For example, web page layouts significantly impacts users' engagement. If a new page layout is considered, bucket testing is used to verify that indeed the new layout is better than the existing one. This example also explains why the number of users exposed to the new layout should is an issue. On one hand, it should be large enough so that the measurements are statistically valid. On the other hand, it should be as small as possible. If the new layout is indeed worse, these users 'suffered' from the experiment and the quality of service provided to them was reduced. Each bucket test is therefore given a budget, \mathcal{B} , which is the maximal number of users it is allowed to effect.

In social networks and services, however, the situation is more complex. Users' satisfaction might depend on whether the service is also available to their friends. For example, a messaging service might inherently be very useful but no user can enjoy it if none of his friends have it too. Thus, to measure users' engagement the service must be given to at least some of their friends. Other examples include, content tagging, games, certain kinds of adds, event invitations, etc. In [1] the authors suggest to set a parameter d > 0 such that users' interactions can be measured only if at least d of their friends have also received the service. We adopt this model as well.

Introducing some notations, let the graph G(V, E) represent the network in the standard way. Each node corresponds to a user and $\{i, j\} \in E$ iff users i and j are connected (or 'friends'). Also denote |V| = n and |E| = m. As in [1] we assume for simplicity matters that all the nodes in G have at least d neighbors. Let $f: V \to [0, 1]$ be an arbitrary function over the users. The function f can be thought of as measuring the user's engagement or satisfaction. The aim of the test is to estimate the mean value $\mu = \frac{1}{n} \sum_{i \in V} f_i$. If the algorithm chooses a set of users B (to give the service to) it pays |B| from the total budget \mathcal{B} . For the algorithm to evaluate f on a core set of nodes A it must pays |B| form the budget. The set B is the d-closure (or fringe) set of A which is the minimal set containing A such that all nodes in A have at least d neighbors in B.

An important difference between our work and [1] is the restrictions put on the satisfactions function. In [1] a random biased coin model generated the function $f: V \to \{0, 1\}$. While this model makes sense and indeed is very useful for analysis, we argue that it is still restrictive. The first generalization is that our function receives real values $f: V \to [0, 1]$.

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The reasoning for the latter is that in many cases, measures of satisfaction take scalar values. These include: the amount of time spent in an application, number of times it was launched, increase in activity, etc. Moreover, a random model for f is not always justified. Here, we consider *any* fixed or even adversarial satisfaction functions.

A fixed (or almost fixed) function f is a realistic scenario but it is also mathematically justified. In fact, one can easily see that the variance of estimating the mean of random function f is always dominated by variance obtained (over the random bits of the algorithm) for an adversarial choice of f.

Second, the authors of [1] suggest to choose the core set A according to different random walk procedures. While this gives good results we show that, in fact, there are virtually an unlimited number of valid algorithms. (it is worth mentioning that they also consider core sets including single nodes and pairs of nodes). We argue that *any* distribution over core subsets of nodes can be used. If the probability of any node being in the sample is strictly positive, the obtained estimator Z for μ is also *unbiased*. The only difference between different distributions is the variance this estimator exhibits. Thus, for every network, one should choose the distribution which minimizes the estimator variance. Since one must choose an algorithm before starting the bucket test, we give bounds on the variance from above for any f.

The rest of this work is organized as follows. In Section 2 we present the meta algorithm which applies the specific algorithms presented in Section 3. Connections of our concept to the random walk approach of [1] is considered in Section 4. A gradient decent optimization procedure aimed to reduce the estimator variance is briefly described in Section 5. Experimental setup and corresponding results are described in Section 6. We discuss the results and conclude the work in Section 7.

2. META ALGORITHM

We start by describing the meta-algorithm which is identical to all the distributions over core sets. The meta algorithm is straight forward and is identical to the one in [1]. Accordingly, a given algorithm produces core sets from some distribution over the graph's set of core sets. Then, for each selected core set it produces an unbiased estimator. It continues to do so until it exhausts the budget \mathcal{B} . The output estimator is the mean of all estimators obtained during the process. A formal description of the meta-algorithm is elaborated in Algorithm 1.

The final estimator Y is unbiased since each of the Z estimators are:

$$\mathbb{E}(Z) = \sum_{A \in supp(Q)} Q(A) \sum_{i \in A} \frac{1}{nq(i)} f(i)$$
$$= \frac{1}{n} \sum_{i \in V} \frac{f(i)}{q(i)} \left(\sum_{A \in supp(Q)} Q(A) \mathbb{1}_{\{i \in A\}} \right) = \frac{1}{n} \sum_{i \in V} f_i = \mu,$$

where Q is the core set distribution over 2^V , and q(i) gives the probability that the *i* is in the chosen core set (see more details in Algorithm 1). Note that if q(i) = 0 for some *i* this expression is not valid.

Since the overall estimator Y is unbiased for any distribution, the goal is to reduce its variance. Given the fact that core sets A_{ℓ} are chosen i.i.d. we have that $\operatorname{Var}(Y) =$

Algorithm 1 Network Bucket Testing Meta-Algorithm
Input: \mathcal{B} , budget
input: d , minimal degree threshold
Input: $G(V, E)$, input graph
Input: Q , core set distribution over 2^V
Input: q, such that $q(i) \leftarrow \sum_{A \in supp(Q)} Q(A) \mathbb{1}_{\{i \in A\}}$
Output: Y, estimation of $\mu = \frac{1}{n} \sum_{i \in V} f(i)$
$b \leftarrow 0 \ ; \ \ell \leftarrow 1 \ ; \ L \leftarrow 1$
while True do
$A_{\ell} \leftarrow \text{drawn according to core set distribution } Q$
$B_{\ell} \leftarrow d$ -closure (fringe) set of A
$b \leftarrow b + B $
$\mathbf{if} \ b > \mathcal{B} \ \mathbf{then}$
break
end if
$Z_{\ell} \leftarrow \sum_{i \in A} \frac{1}{nq(i)} f(i) ; L \leftarrow \ell ; \ell \leftarrow \ell + 1$
end while
return: $Y \leftarrow \frac{1}{L} \sum_{\ell=1}^{L} Z_{\ell}$

 $\frac{1}{L^2} \sum_{\ell=1}^{L} \operatorname{Var} (Z_\ell) = \frac{1}{L} \operatorname{Var} (Z).$ To approximate the size of L we compute the cost of this experiment which is $\sum_{\ell=1}^{L} |B_\ell|$. Assuming $|B_\ell| \ll \mathcal{B}$ and enforcing that the cost is less than the budget we get that $L \approx \mathcal{B}/\mathbb{E}(|B_1|)$. Note that since $|B_\ell|$ are also i.i.d. random variables, applying Chernoff's inequality shows that the value of $L = (1 \pm o(1))\mathcal{B}/\mathbb{E}(|B_1|)$ with high probability, the details are omitted. Finally, we have that

$$\operatorname{Var}(Y) \approx \frac{1}{\mathcal{B}} \mathbb{E}(|B|) \operatorname{Var}(Z)$$
 . (1)

where we omit the subscript and use B and Z instead of B_1 and Z_1 . Since the budget is fixed, the correct choice of distribution over core subsets and estimators is the one minimizing $\mathbb{E}(|B|) \operatorname{Var}(Z)$. This quantity is highly related to the efficiency defined in [1] as |B|/|A|. If the engagement function f is itself random, as assumed in [1], we can expect $\operatorname{Var}(Z)$ to be proportional to 1/|A|.

It remains to compute the variance $\operatorname{Var}(Z) = \mathbb{E}(Z^2) - (\mathbb{E}(Z))^2$. We start by calculating the second moment of the estimate Z,

$$\mathbb{E}(Z^{2}) = \sum_{A \in 2^{V}} Q(A) \sum_{i \in A} \sum_{j \in A} \frac{1}{n^{2}q(i)q(j)} f(i)f(j)$$

$$= \sum_{i \in V} \sum_{j \in V} \frac{1}{n^{2}} \frac{1}{q(i)q(j)} f(i)f(j) \sum_{A \in 2^{V}} Q(A) \mathbb{1}_{\{\{i,j\} \subset A_{\ell}\}}$$

$$= \sum_{i \in V} \sum_{j \in V} \frac{1}{n^{2}} \frac{q(i,j)}{q(i)q(j)} f(i)f(j)$$
(2)

where $q(i,j) \equiv \sum_{A \in 2^V} Q(A) \mathbb{1}_{\{\{i,j\} \subset A_\ell\}}$ is the probability that both nodes *i* and *j* are simultaneously included in a core set *A*. Maximizing this expression over functions *f* such that $\sum f(i) = \mu n$ gives the worst variance possible. It is easily verified that the maximal obtainable value is $\mathbb{E}(Z^2) = \frac{\mu^2}{q(1)}$, assuming w.l.o.g. that $\max_i \frac{1}{q(i)} = \frac{1}{q(1)}$. This bound, however, is overly pessimistic since it is obtained in the unrealistic case where $f(1) = \mu n$ and all other values are f(i) = 0. We therefore need to enforce that the values of *f* are distributed over many values. A natural way to achieve this is to limit ourselves to functions *f* such that $f(i) \in [0, 1]$. **Proposition 1** Let $W(i,j) = \frac{q(i,j)}{q(i)q(j)}$. Moreover, let $U_{\mu} = \arg \max_{|U| \le \mu n} = \sum_{\{i,j\} \subset U} W(i,j)$.

$$\operatorname{Var}(Z) \le \sum_{\{i,j\} \subset U_{\mu}} W(i,j)/n^2 - \mu^2.$$
(3)

PROOF. First, note that W is a positive semidefinite matrix. This is because $f^T W f = n^2 \mathbb{E} (Z^2) \ge 0$ for all f. Therefore, $f^T W f$ is a convex function of f defined over the convex set $f(i) \in [0,1]$ and $\sum f(i) = \mu n$. The maximal value of such functions is obtained in an extreme point of the body. Let u_{μ} be this extreme point $u_{\mu}(i) \in \{0,1\}$ and $\sum_i u_{\mu}(i) = \mu n$. That is, for μn nodes we have $u_{\mu}(i) = 1$ and for $(1 - \mu)n$ nodes $u_{\mu}(i) = 0$. Setting $U_{\mu} = \{i | u_{\mu}(i) = 1\}$ completes the claim. \Box

Computing this quantity amounts to finding the heaviest subgraph of size μn of G when the weights of the edges are gives by W. The heaviest subgraph problem is notoriously hard [4][2]. It does, however admit scalable approximation algorithms that work well in practice [3][7]. Regardless, in our scenario, it is natural to assume that μ is at least a small constant. Therefore, a random choice of U_{μ} is expected to yields a μ^2 approximation factor to the optimal. Moreover, if any algorithms improves on the random choice by a factor of t then the solution is guaranteed to by a $t\mu^2$ approximation to the optimal. This discussion, unfortunately, goes beyond the scope of this paper.

For the sake of simplicity, we use a more relaxed bound which uses the spectral norm of W. Applying the Cauchy-Schwarz inequality yields $\operatorname{Var}(Z) \leq \frac{1}{n^2} \lambda_1(W) \|u_{\mu}\|^2 - \mu^2$. Here $\lambda_1(W)$ denotes the spectral norm of W (its largest eigenvalue). Substituting $\|u_{\mu}\|^2 = \mu n$ we get that:

$$\operatorname{Var}\left(Z\right) \le \left(\frac{1}{n}\lambda_1(W) - \mu\right)\mu , \qquad (4)$$

The largest eigenvalue $\lambda_1(W)$ can be easily calculated using *power iteration* method. Except for this bound being significantly easier to compute, we shall see in the results section that it is also tight enough to give valuable information.

As seen above, the overall variance of the step estimator Z is proportional to $\mathbb{E}(|B|) \operatorname{Var}(Z)$, where both $\mathbb{E}(|B|)$ and $\operatorname{Var}(Z)$ are complex function of the distribution over core sets Q. In what follows we describe specific algorithms. Those enable us to efficiently draw core subsets from a distribution Q and produce the probability vector q for every graph. The goal, of course, is to reduce $\mathbb{E}(|B|) \operatorname{Var}(Z)$ as much as possible.

3. SPECIFIC ALGORITHMS

In order to describe the algorithm we require some additional notations. Let $N_i = \{j \in V : (i, j) \in E\}$ indicate the set of neighbors of node i and $N_i^+ = \{\{i\} \cup N_i\}$. We denote by $N_{i,j} = N_i \cap N_j$ (similarly $N_{i,j}^+ = N_i^+ \cap N_j^+$) and $M(i, j) = \min\{|N_{i,j}|, d-1\}$. We denote Q the distribution over subsets of nodes and Q(A) the probability of core set being chosen. Also, let supp(Q) be the support of Q, i.e., $A \in supp(Q)$ iff Q(A) > 0.

3.1 Naïve Algorithm

Here the core sets are simply the nodes of the graph, $supp(Q) = \{\{i\} | i \in V\}$. In addition, the core set distribution Q is simply Pr(A) = p(i), where p is some distribution defined over the nodes of G. Hence, $q(i) = Pr(i \in A) = p(i)$. Since the core set contains only one node and we randomly pick d of its neighbors to form the closure set, we clearly have $|B| \leq d + 1$.¹ To compute the variance we note that $W(i, j) = \frac{1}{q(i)}$ for i = j and zero otherwise. Since W is a diagonal matrix in this case, its top eigenvalue equals its maximal diagonal entry, we have that the spectral bound (4) reduces to

$$\operatorname{Var}\left(Z\right) \le \left(\frac{1}{n}\frac{1}{\min_{i} p(i)} - \mu\right)\mu.$$

This is minimized using the uniform distribution p(i) = 1/nand gives $\operatorname{Var}(Z) \leq (1-\mu)\mu$. In this case, it turns out, that the naïve spectral bound of 4 is tight.

The overall variance achieved by the naïve algorithm for any f and uniform node distribution is therefore

$$\operatorname{Var}(Y_{naive}) = \frac{1}{\mathcal{B}}(d+1)(1-\mu)\mu .$$
 (5)

3.2 Edge Algorithm

The first non trivial core set distribution can include any two nodes which are connected in the graph. Namely $\{i, j\} \in$ supp(Q) if $\{i, j\} \in E$. Setting the probability of choosing edge $\{i, j\}$ to be p(i, j) we have

$$\begin{split} \mathbb{E}\left(|B|\right) &\leq 2d - \sum_{\{i,j\}\in E} p(i,j)M(i,j) \\ W(i,j) &= \frac{p(i,j)}{\left(\sum_{k\in N_i} p(i,k)\right) \left(\sum_{k\in N_j} p(j,k)\right)} \end{split}$$

A possible good assignment for p could be achieved by producing a maximal weighted matching on the graph G, where the weight of edge $\{i, j\}$ is set to M(i, j). Assigning probability 2/n for all edges in the matching and probability zero to all other edges. Admittedly, not all graphs yield good maximal weighted matching, or even any matching which includes all nodes. We experimented with a simpler edge selection algorithm which applies to any graph.

3.3 Neighborhood Algorithm

Here the core sets are the graph nodes and their neighbors $supp(Q) = \{N_i^+ | i \in V\}$. In addition, we assign different probabilities to each core set according $Q(N_i^+) = p(i)$ where p is an arbitrary distribution defined over the nodes of G. The node i will be referred to as the center of N_i^+ . Hence, a node i belongs to the core set A if one of its neighbors (or itself) is the center node of A. Therefore we have

$$\mathbb{E}(|B|) \leq 1 + \sum_{\{i,j\} \subset E} (p(i) + p(j)) (d - M_{i,j})$$

$$W(i,j) = \frac{\sum_{k \in N_{i,j}^+} p(k)}{\left(\sum_{k \in N_i^+} p(k)\right) \left(\sum_{k \in N_j^+} p(k)\right)} .$$

4. CONNECTION TO RANDOM WALKS

In [1] the authors suggested generating the core sets according to a random walk. That is, start at any node and at each step move to one of the neighboring nodes uniformly at

 $^{^1\}mathrm{Recall}$ our assumption that all nodes have at least d neighbors.

random. One can theoretically consider a core set distribution Q which includes all length t paths in the graph. The probability of a core set A is the probability of it being the set of nodes produced by the random walk. Although it is computationally impossible to compute Q it is quite easy to sample from it simply by simulating the random walk. In order to execute the meta algorithm one must also be able to compute q(i). It is well known [6] that after a certain number of such steps, one reaches the stationary distribution. In the stationary distribution the probability of being at node *i* is proportional to its degree, denoted by deg(i). Therefore, the expected number of times a node is included in a length t random walk is $t \cdot deg(i) / \sum_{j \in V} deg(i).$ Setting $q(i) = t \cdot deg(i) / \sum_{j \in V} deg(i)$ completes the description of the algorithm. It is worth noting that there is a slight difference in notation between the ones in [1] and those used here. There, a node can appear multiple times in the core set, so in a sense, it behaves more like a list than a set. This is the reason the authors of [1] introduced the multiplicity of nodes in core sets into their estimator.

A similar view is also possible for the other variants of random walks proposed in [1]. The authors use random walks which try to balance the probabilities q(i). This is possible using a Metropolis-Hastings random walk as used, for example, in [5]. Another option is to assign weights to edges and transition with probabilities proportional to edge weights. It turns out that it is possible, in most cases, to assign such weights that the probability of visiting each node is roughly the same. Again, computing or storing Q is computationally impossible but sampling from it is easy. Setting q(i) = t/n and applying the meta algorithm is identical to the algorithms in [1].

One problematic aspect of using random walks is that it is impossible to compute the matrix W and hence impossible to analytically bound the variance for arbitrary unknown functions f. Surprisingly, there is a way to overcome this problem. In particular, one can simulate a very large number of random walks and produce an empirical matrix W'. It is not hard to see that after sufficiently many simulations, we would have $W' \sim W$ at least in the spectral sense. It can be shown using sampling argumentation, that $O(n^2 \log(n))$ simulations would suffice for W' to be close enough to W to give similar bounds.

5. GRADIENT DECENT OPTIMIZATION

In cases where the support of the core set distribution is small we can directly minimize the overall variance of the estimator. That is, find values for Q which minimize $\mathbb{E}(|B|) \operatorname{Var}(Z)$. One obstacle in doing so is that an exact expression for the variance $\operatorname{Var}(Z)$ is not available. From Proposition 1 we have that $\operatorname{Var}(Z) \leq \sum_{\{i,j\} \subset U_{\mu}} W(i,j)/n^2$. Alas, computing this value requires solving an NP-hard problem. We therefore replace it with a simple bound which sums over all elements of W, namely, $\operatorname{Var}(Z) \leq \sum_{\{i,j\} \subset V} W(i,j)/n^2$. Although this bound is extremely naïve, it serves well as a surrogate to the actual value. The second obstacle is that computing the closure set for a core set is also computationally hard. For this problem we also use a simple bound which is the size of closure set achieved by a greedy algorithm, B_g . Finally we are faced with minimizing $\psi(Q) = \mathbb{E}(|B_g|) \sum_{\{i,j\} \subset V} W(i,j)/n^2$. Since $\psi(Q)$ is a complex function of the core set distribution we cannot hope to minimize it exactly. We resort to minimizing $\psi(Q)$ heuristically using Gradient Decent. The results of using Gradient Decent on Neighborhood algorithm core sets are presented in the experimental section.

6. EXPERIMENTAL SETUP AND RESULTS

Evaluating algorithms' efficiency or preferring one algorithm to another is impossible in general. The correct choice of algorithm heavily depends on a wide range of parameters. While the authors of [1] report good results of their algorithms applied to portions of the Facebook network, we observe that it does less well for others. Likewise, our algorithms mostly outperform the naïve implementation but for some values of f and some graph the naïve algorithms beats them. The number of variations possible in the graphs, satisfaction functions, algorithms, and measures of success, is practically endless. Nevertheless, we tried to be as thorough as possible. Our choices are described below.

6.1 Graphs

The first crucial factor in the success of an algorithm is the Network it operates on. In this work we examine 3 different graphs, one real and two synthetic.

DBLP: we used the Digital Bibliography and Library Projects (DBLP) entire database. It contains data about authors of manuscripts. We associated each node in the graph with an author and an edge corresponds to coauthorship of at least one paper. The graph we obtained contains 845, 211 nodes, each with at least one edge (authors with no co-authors were discarded).

BA: a synthetic graph constructed according to the model of Albert and Barabási [8]. We start with a ring graph of size 10. Then we add nodes one at a time. Each new node is connected by edges to 10 other nodes already in the graph with probabilities proportional to their degrees.

WS: a synthetic graph constructed according to the model of Watts and Strogatz [9]. Here, to construct a network of n nodes we start with an n node ring graph. Then, we connect each node to 10 nodes to its right along the ring. Finally, we reroute each edge to a random node with probability 1/2.

One immediate problem we encounter is that, due to our model, nodes with degree lower than d cannot be measured. That is simply because, even if all their neighbors are chosen, they would not have d chosen neighbors. One can think of several solutions for this issue. For example, change the model by deciding that such nodes are still measurable if all their neighbors are chosen. Or, define the mean to not include those and never measure their satisfaction. However, since this is not the main point in of the paper we chose (as in [1]) to simply remove those nodes. Note that after removing some small degree nodes, other nodes might become removable too. Here we simply continued removing those until all degrees in the graph were at least d. This process will be referred to as trimming.

6.2 Satisfaction functions

As explained throughout the paper, one of the most crucial factors governing the variance of our estimation is the satisfaction function f. While our proofs bound the variance for all functions f simultaneously, we experimented with only three kinds. These were taken to represent extreme cases of network biases.

Uniform: samples uniformly, without replacement, exactly

 μn nodes from the graphs. For chosen nodes f(i) = 1 and f(i) = 0 otherwise. This serves mostly as a sanity check and as a baseline. We cannot expect any social feature to truly have such a satisfaction function.

BFS: starts a Breadth First Search algorithm in an arbitrary node in the graph and assigns a value of f(i) = 1 to all nodes it encounters until it encounters μn nodes. The rest are give value zero. This function gives an extreme graph topological bias.

Degree Percentile: assigns the value of f(i) = 1 to all nodes in the top μ percentile in terms of degree. In other words, f(i) = 1 for the μn nodes whose degree in the graph is the highest. Here we simulate another extreme case of degree bias. This is an important case for two reasons. First, our algorithms are heavily influenced by node degrees and so this choice of f might be 'difficult'. Second, in reality, social features are not independent of degree biases. This is because the node degree usually relates to the user's activity or 'socialness' in some sense. This function is the extreme case of all satisfaction distributions which are positively correlated with the degree.

Degree Bias: proposed in [1] and gives a less extreme degree bias. Nodes are picked randomly with probability proportional to $\log(deg(i))$. The process terminates when we have picked μn nodes. The function f assigns the value 1 to all picked nodes and 0 otherwise.

DBLP	Uniform	BFS	Degree Perce- ntile	Degree Bias
Naïve	0.41	0.41	0.42	0.41
Edge	0.38	0.37	0.20	0.36
Edge-Matching	0.32	0.38	0.34	0.32
Neighborhood	0.34	0.85	0.63	0.34
Neighborhood-20	0.32	0.53	0.39	0.32
Neighborhood-40	0.28	0.57	0.46	0.28
Neighborhood-GD	0.24	0.48	0.43	0.24
Simple-RW	0.30	0.47	0.32	0.29
Metropolis-H-RW	0.29	0.47	0.42	0.28
Matrix-Scaling-RW	0.27	0.49	0.43	0.27
Triangle-Closing-RW	0.26	0.49	0.44	0.26

Table 1: The table gives the Normalized RMSE scores for the various algorithms and satisfactions functions. The graph here is the DBLP graph and the satisfactions functions mean in is $\mu = 0.1$.

6.3 Algorithms

The algorithms we examined all produce their estimates according to meta algorithm (see Algorithm 1). Here we describe algorithms by the manner in which they choose core sets. All Random Walk (RW) based algorithms are described in detail in [1]. This serves mostly as a baseline but also serves to validate their results on graphs other than Facebook. For completeness we describe those shortly here. These algorithms perform random walks on the network and collect nodes they encounter into the core set. The difference between them is the transition probability between neighboring nodes. Below, we shortly recap each of the tested algorithms.

Naïve: denotes the algorithm in which each core set includes one single node chosen uniformly at random (see Section 3.1).

Barabási	Uniform	BFS	Degree Perce- ntile	Degree Bias
Naïve	0.31	0.31	0.31	0.31
Edge	0.34	0.30	0.17	0.33
Edge-Matching	0.31	0.30	0.29	0.30
Neighborhood	0.54	0.60	0.41	0.54
Neighborhood-20	0.40	0.38	0.23	0.38
Neighborhood-40	0.40	0.40	0.23	0.39
Neighborhood-GD	0.36	0.34	0.19	0.35
Simple-RW	0.33	0.31	0.17	0.32
Metropolis-H-RW	0.36	0.35	0.28	0.35
Matrix-Scaling-RW	0.30	0.31	0.28	0.30
Triangle-Closing-RW	0.31	0.31	0.28	0.30

Table 2: The table gives the Normalized RMSE scores for the various algorithms and satisfactions functions. The graph contains $n = 10^5$ nodes and is generated according to the model of Albert and Barabási [8]. As before, the satisfactions functions mean in is $\mu = 0.1$.

Watts- Strogatz	Uniform	BFS	Degree Perce- ntile	Degree Bias
Naïve	0.31	0.31	0.31	0.31
Edge	0.29	0.31	0.25	0.29
Edge-Matching	0.28	0.31	0.28	0.28
Neighborhood	0.27	0.44	0.24	0.27
Neighborhood-20	0.30	0.44	0.25	0.30
Neighborhood-40	0.27	0.44	0.25	0.27
Neighborhood-GD	0.28	0.41	0.26	0.28
Simple-RW	0.28	0.34	0.24	0.28
Metropolis-H-RW	0.29	0.35	0.27	0.29
Matrix-Scaling-RW	0.28	0.34	0.27	0.28
Triangle-Closing-RW	0.27	0.34	0.27	0.27

Table 3: The table gives the Normalized RMSE scores for the various algorithms and satisfactions functions. The graph contains $n = 10^5$ nodes and is generated according to the model of Watts and Strogatz [9]. As before, the satisfactions functions mean in is $\mu = 0.1$.

Edge: refers to core sets of size two. An edge is chosen uniformly at random and the core set contains its supporting nodes (see Section 3.2).

Edge-Matching: a variant to the former Edge algorithm. Here, the core sets are also pairs of nodes supported by edges. The idea is to create a set of edges which behaves like a matching but is simpler to obtain. The process proceeds as follows. Start with an empty edge set E_m . For every node, pick the edge connecting it to its neighbor (in G) with the least degree with respect to E_m . In case of ties, pick the one maximizing $M_{i,j}$ (for definition see Section 3). Add the picked edge to E_m and continue. The core sets are pairs of nodes supports of edges in E_m chosen uniform at random.

Neighborhood: corresponds to a uniform distribution over the sets N_i^+ . This, by selecting a node uniformly at random and selecting it and its neighbors (see Section 3.3).

Neighborhood-*k*: this algorithm, which is a variant of the former Neighborhood algorithm, also chooses nodes and their neighborhoods but avoids doing so for nodes of very

high degree. More accurately, Neighborhood-k chooses node i uniformly at random, if $|N_i^+| \leq k$ it returns N_i^+ otherwise it returns a singleton core set $\{i\}$.

Neighborhood-DG: here the core sets are still N_i^+ . However, the distribution Q over them is optimized using gradient decent (see Section 5) to reduce the overall estimation variance.

Simple-RW: a random walk algorithm which gives the standard transition probability. Move from node i to j with probability 1/deg(i).

Metropolis-Hastings-RW: moves from node *i* to *j* with probability $\min(1/deg(i), 1/deg(j))$, and stays in node *i* with the remainder probability. This produces a uniform stationary distribution but tends to visit the same nodes many time.

Matrix-Scaling-RW: transitions from i to j with probability w(i, j). These are computed by an iterative process to try to make the stationary distribution as uniform as possible.

Triangle-Closing-RW: the transition probability between i and j is depends on the node h visited before i. If $\{h, j\} \notin E$ the transition probability is w'(i, j). If $\{h, j\} \in E$ this probability is increased by a factor $\alpha \geq 1$ to be $\alpha w'(i, j)$. The weights w' are chosen to produce a node distribution which is as close to uniform as possible.

6.4 Measure of Success

Our main measure of success for an algorithm is the Root Mean Square Error (RMSE) of its outputs. Assume we execute an algorithm t times and produce outcomes Y_1, \ldots, Y_t . Since the satisfaction function f has mean μ the normalized RMSE is given by $\frac{1}{\mu} (\frac{1}{t} \sum_{i=1}^{t} (Y_i - \mu)^2)^{1/2}$. We chose RMSE as our measure of success since it embodies both accuracy and reliability. Note that since RMSE (squared) estimates $\mathbb{E}[(Y_i - \mu)^2]/\mu^2$, by Markov's inequality we also get confidence intervals.

6.5 Experimental Results

Tables 1–3 give the RMSE values achieved by the different algorithms for different choices of f. Each combination of algorithm and satisfaction function was run 1000 times and the RMSE value is calculated according to Section 6.4. For all tables we set the budget \mathcal{B} to 1% of the network size. This is a reasonable budget for actual bucket tests.

In Figure 1 the estimate normalized RMSE for the DBLP graph and the Degree Bias satisfaction function, is plotted vs. the budget \mathcal{B} . It is clearly visible that the RMSE decreases with the increase in budget.

6.6 Gradient Decent Experiments

To demonstrate the benefits of the Gradient Decent optimization (see Section 5) we applied the Neighborhood algorithm to the DBLP graph. As before, we iteratively trimmed that the minimal degree in the graph is 10. This resulted in a graph containing n = 57285 nodes. In Table 4 we provide several statistics for three different distributions over Neighborhood algorithm core sets, N_i^+ for all $i \in V$. The first is uniform (Uniform), the second is relative to $1/|N_i^+|$ (Degree) and the third is the probability $Q(N_i^+)$ assigned by the Gradient Decent procedure (GD).

Examining the table it is observed that the Gradient Decent optimization reduces $\psi(Q)$, mainly by reducing the average closure set size. In parallel, it also increases the effi-

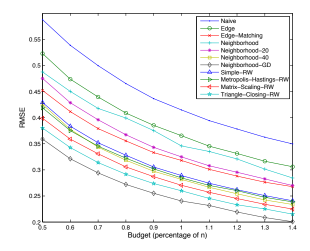


Figure 1: Estimate normalized RMSE for the DBLP graph and the Degree Bias satisfaction function vs. the budget \mathcal{B} .

ciency of the algorithm. On the other hand it increases the largest eigenvector when compared to that calculated for the Uniform distribution.

While it is hard to foresee the exact strategy that Gradient Decent optimization follows to reduce $\psi(Q)$, Figure 2 may provide some insights. In Figure 2, the DBLP graph degree PDF is plotted for the three core set distributions (Uniform, Degree, and GD). It is apparent that the GD optimization causes the graph degree PDF to drop much faster than those of the Uniform and Degree distributions. It turns out that the GD optimization reduces the probabilities of higher degree nodes. In fact, for the DBLP graph, it assigned zero probably to any Neighborhood core set of size greater than 114. This is more than half of the core sets!

	Uniform	Degree	GD
Average core set size $\mathbb{E}(A)$	22.92	17.096	15.31
Average closure set size $\mathbb{E}\left(B_g \right)$	72.56	54.77	26.29
Efficiency bound $\mathbb{E}(A) / \mathbb{E}(B_g)$	0.316	0.312	0.58
$\frac{1}{n^2} \Sigma_{i,j} W(i,j)$	1.360	1.315	1.20
$\psi(Q)$	98.71	72.00	31.66
$\frac{1}{n}\lambda_1(W)$	1.718	4.621	2.13

Table 4: Statistics for different distributions over Neighborhood algorithm core sets applied to the DBLP graph.

6.7 Spectral Bounds

Being able to analytically bound the accuracy (RMSE) of a network bucket test is crucial for two main reasons. Before the test, the administrator must choose the best algorithm to use. After the test, he/she must supply error bounds on the resulting estimate. Given the discussion following Proposition 1, this is a hard computational task. However, using the spectral bound of Equation (4), designers can get a rough bound for this quantity. To demonstrate the benefits of this bound we use values from Table 4, derived for

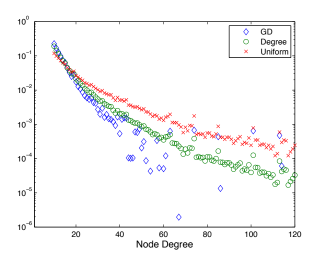


Figure 2: Probability distributions of the size of core sets for the DBLP graph. The three distributions represent different distributions over the core set N_{i+} . Uniform, selects each with constant probability, 1/n. Degree, selected N_{i+} w.p. proportional to $1/|N_{i+}|$, and GD selects N_{i+} w.p. according to the output of the Gradient Decent optimization.

the aforementioned Neighborhood algorithm core set distributions, to calculate bounds for the DBLP graph. The calculated bounds along with their corresponding simulation results are plotted in Figure 3 for $\mu = 0.1$.

7. DISCUSSION AND CONCLUDING RE-MARKS

In this paper we proposed and analyzed several algorithms for network bucket testing. The achieved results are comparable or better than previous algorithms depending on the setup. However, we argue that the contribution goes beyond that. First, our algorithms are simple to program, provide unbiased estimates, efficient to execute, and analyzable. Moreover, we can efficiently produce good error bounds for their performance. This gives us the ability to choose the best algorithm for a network well before running the test.

In addition, the framework lets algorithm designers analyze a very large variety of algorithms. For example, one can consider core sets of triangles in the graph. Or, cover the graph with small tightly connected subgraphs and consider that as core sets. The possibilities are endless. We hope the derivations also provide walk-through examples on how to analyze those.

An additional benefit which is not mentioned in the paper but is an immediate outcome of the Gradient Decent approach. That is, one can combine any number of different algorithms and consider the superset of their core sets. Applying the Gradient Decent process to the core superset can automatically mix the different algorithms. The resulting mixed algorithm is guaranteed to be better (no worse) than the best single algorithm.

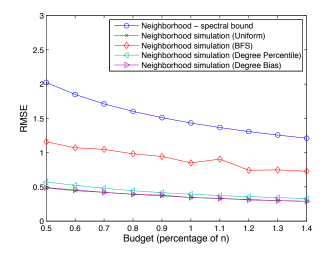


Figure 3: RMSE upper bound for the DBLP graph using the Neighborhood Algorithm. Along with the bounds we give simulation results for different functions, *f*. Namely. Uniform, BFS, Degree Percentile, and Degree Bias. As expected, all simulation results are lower than the theoretical bound.

8. **REFERENCES**

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