Information Source Detection in the SIR Model: A Sample Path Based Approach

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Abstract—This paper studies the problem of detecting the information source in a network in which the spread of information follows the popular Susceptible-Infected-Recovered (SIR) model. We assume all nodes in the network are in the susceptible state initially except the information source which is in the infected state. Susceptible nodes may then be infected by infected nodes, and infected nodes may recover and will not be infected again after recovery. Given a snapshot of the network, from which we know all infected nodes but cannot distinguish susceptible nodes and recovered nodes, the problem is to find the information source based on the snapshot and the network topology. We develop a sample path based approach where the estimator of the information source is chosen to be the node associated with the sample path that most likely leads to the observed snapshot. We prove for infinite-trees, the estimator is a node that minimizes the maximum distance to the infected nodes. A reverse-infection algorithm is proposed to find such an estimator in general graphs. We prove that for g-regular trees such that \( gq > 1 \), where \( g \) is the node degree and \( q \) is the infection probability, the estimator is within a constant distance from the actual source with a high probability, independent of the number of infected nodes and the time the snapshot is taken. Our simulation results show that for tree networks, the estimator produced by the reverse-infection algorithm is closer to the actual source than the one identified by the closeness centrality heuristic. We then further evaluate the performance of the reverse infection algorithm on several real world networks.

I. INTRODUCTION

Diffusion processes in networks refer to the spread of information throughout the networks, and have been widely used to model many real-world phenomena such as the outbreak of epidemics, the spreading of gossips over online social networks, the spreading of computer virus over the Internet, and the adoption of innovations. Important properties of diffusion processes such as the outbreak thresholds \([1]\) and the impact of network topologies \([2]\) have been intensively studied.

In this paper, we are interested in the reverse of the diffusion problem: given a snapshot of the diffusion process at time \( t \), can we tell which node is the source of the diffusion? The answer to this problem has many important applications, and can help us answer the following questions: who is the rumor source in online social networks? which computer is the first one infected by a computer virus? who is the one who uploaded contraband materials to the Internet? and where is the source of an epidemic?

We call this problem information source detection problem. This information source detection problem has been studied in \([3]-[5]\) under the Susceptible-Infected (SI) model, in which susceptible nodes may be infected but infected nodes cannot recover. The authors formulated the problem as a maximum likelihood estimation (MLE) problem, and developed novel algorithms to detect the source.

In this paper, we adopt the Susceptible-Infected-Recovered (SIR) model, a standard model of epidemics \([6], [7]\). The network is assumed to be an undirected graph and each node in the network has three possible states: susceptible (\( S \)), infected (\( I \)), and recovered (\( R \)). Nodes in state \( S \) can be infected and change to state \( I \), and nodes in state \( I \) can recover and change to state \( R \). Recovered nodes cannot be infected again. We assume that initially all nodes are in the susceptible state except one infected node (called the information source). The information source then infects its neighbors, and the information starts to spread in the network. Now given a snapshot of the network, in which we can identify infected nodes and healthy (susceptible and recovered) nodes (we assume susceptible nodes and recovered nodes are indistinguishable), the question is which node is the information source.

We remark that it is very important to take recovery into consideration since recovery can happen due to various reasons in practice. For example, a contraband material uploader may delete the file, a computer may recover from a virus attack after anti-virus software removes the virus, and a user may delete the rumor from her/his blog. In order to solve the information source detection problem in these scenarios, we study the SIR model in this paper, which makes the problem significantly more challenging than that in the SI model as we will explain in the related work section.

A. Main Results

The main results of this paper are summarized below.

- Similar to the SI model, the information source detection problem can be formalized as an MLE problem. Unfortunately, to solve the MLE problem, we need to consider all possible infection sample paths, and for each sample path, we need to specify the infection time and recovery time for each healthy node and the infection time for each infected node, so the number of possible sample paths is at the order of \( \Omega(t^N) \), where \( N \) is the network size.
and \( t \) is the time the snapshot is obtained. Therefore, the MLE problem is difficult to solve even when \( t \) is known. The problem becomes much harder when \( t \) is unknown, which is the assumption of this paper. To overcome this difficulty, we propose a sample path based approach. We propose to find the sample path which most likely leads to the observed snapshot and view the source associated with that sample path as the information source. We call this problem optimal sample path detection problem.

We investigate the structure properties of the optimal sample path in trees. Defining the infection eccentricity of a node to be the maximum distance from the node to infected nodes, we prove that the source node of the optimal sample path is the node with the minimum infection eccentricity. Since a node with the minimum eccentricity in a graph is called the Jordan center, we call the nodes with the minimum infection eccentricity the Jordan infection centers. Therefore, the sample path based estimator is one of the Jordan infection centers.

- We propose a low complexity algorithm, called reverse infection algorithm, to find the sample path based estimator in general graphs. In the algorithm, each infected node broadcasts its identity in the network, the node who first collect all identities of infected nodes declares itself as the information source, breaking ties based on the sum of distances to infected nodes. The running time of this algorithm is equal to the minimum infection eccentricity, and the number of messages each node receives/sends at each iteration is bounded by the degree of the node.

- We analyze the performance of the reverse infection algorithm on \( g \)-regular trees, and show that the algorithm can output a node within a constant distance from the actual source with a high probability, independent of the number of infected nodes and the time the snapshot is taken.

- We conduct extensive simulations over various networks to verify the performance of the reverse infection algorithm. The detection rate over regular trees is found to be around 60\%, and is higher than that of the infection closeness centrality (or called distance centrality) heuristic. The infection closeness of a node is defined to be the inverse of the sum of distances to infected nodes and the infection closeness centrality heuristic is to claim the node with the maximum infection closeness as the source. Note that in [3]–[5], the authors proved the node with the maximum infection closeness is the MLE on regular trees. For real world networks, our experiments also show that the reverse infection algorithm outperforms random guesses significantly. We then further evaluate the performance of the reverse infection algorithm on several real world networks.

### Related Work

There have been extensive studies on the spread of epidemics in networks based on the SIR model (see [1], [2], [3], [4] and references within). The work most related to this paper is [3]–[5], in which the information source detection problem was studied under the SI model. [10], [11] considers the problem of detecting multiple information sources under the SI model. This paper considers the SIR model, where infection nodes may recover, which can occur in many practical scenarios as we have explained. Because of node recovery, the information source detection problem under the SIR model differs significantly from that under the SI model. The differences are summarized below.

- The set of possible sources in the SI model [3]–[5] is restricted to the set of infected nodes. In the SIR model, all nodes are possible information sources because we assume susceptible nodes and recovered nodes are indistinguishable and a healthy node may be a recovered node so can be the information source. Therefore, the number of candidate sources is much larger in the SIR model than that in the SI model.

- A key observation in [3]–[5] is that on regular trees, all permitted permutations of infection sequences (an infection sequence specifies the order at which nodes are infected) are equally likely under the SI model. The number of possible permutations from a fixed root node, therefore, decides the likelihood of the root node being the source. However, under the SIR model, different infection sequences are associated with different probabilities, so counting the number of permutations are not sufficient.

- [3]–[5] proved that the node with the maximum closeness centrality is the an MLE on regular-trees. We define the infection closeness centrality to be the inverse of the sum of distances to infected nodes. Our simulations show that the sample path based estimator is closer to the actual source than the nodes with the maximum infection closeness.

Other related works include: (1) detecting the first adopter of innovations based on a game theoretical model [12] in which the authors derived the MLE but the computational complexity is exponential in the number of nodes, (2) network forensics under the SI model [13], where the goal is to distinguish an epidemic infection from a random infection, and (3) geospatial abduction problems (see [14], [15] and references within).

### II. Problem Formulation

#### A. The SIR Model for Information Propagation

Consider an undirected graph \( G = \{V, E\} \), where \( V \) is the set of nodes and \( E \) is the set of (undirected) edges. Each node \( v \in V \) has three possible states: susceptible (S), infected (I), and recovered (R). We assume a time slotted system. Nodes change their states at the beginning of each time slot, and the state of node \( v \) in time slot \( t \) is denoted by \( X_v(t) \). Initially, all nodes are in state S except node \( v^* \) which is in state I and is the information source. At the beginning of each time slot, each infected node infects each of its susceptible neighbors with probability \( q \), independent of other nodes, i.e., a susceptible node is infected with probability \( q \) independently.
The information source detection problem is to identify the information source can be formulated as a maximum likelihood detection problem as follows:

\[ v^* \in \arg \max_{v \in \mathcal{V}} \sum_{X[0,t]:F(X(t))=Y} \Pr(X[0,t]\mid v^* = v), \]

where \( \Pr(X[0,t]\mid v^* = v) \) is the probability to obtain sample path \( X[0,t] \) given the information source is node \( v \).

We note the difficulty of solving this maximum likelihood problem is the curse of dimensionality. For each \( v \) such that \( Y_v = 0 \), we need to decide its infection time and recovery time (the node is in susceptible state if the infection time is \( > t \)), i.e., \( O(t^2) \) possible choices; for each \( v \) such that \( Y_v = 1 \), we need to decide the infection time, i.e., \( O(t) \) possible choices. Therefore, even for a fixed \( t \), the number of possible sample paths is at least at the order of \( t^N \), where \( N \) is the number of nodes in the network. This curse of dimensionality makes it computationally expensive, if not impossible, to solve the maximum likelihood problem. To overcome this difficulty, we propose a sample path based approach which is discussed below.

### D. Sample Path Based Detection

Instead of using the MLE, we propose to identify the sample path \( X^*[0,t^*] \) that most likely leads to \( Y \), i.e.,

\[ X^*[0,t^*] = \arg \max_{t, X[0,t]} \Pr(X[0,t]\mid X(t) = Y), \tag{1} \]

where \( X(t) = \{X[0,t]\mid F(X(t)) = Y\} \). The source node associated with \( X^*[0,t^*] \) is then viewed as the information source.

## III. SAMPLE PATH BASED DETECTION ON TREE NETWORKS

The optimal sample paths for general graphs are still difficult to obtain. In this section, we focus on tree networks and derive structure properties of the optimal sample paths.

First, we introduce the definition of eccentricity in graph theory [16]. The eccentricity \( e(v) \) of a vertex \( v \) is the maximum distance between \( v \) and any other vertex in the graph. The Jordan centers of a graph are the nodes which have the minimum eccentricity. For example, in Figure 2, the eccentricity of node \( v_1 \) is 4 and the Jordan center is \( v_2 \), whose eccentricity is 3.

Following a similar terminology, we define the infection eccentricity \( \bar{e}(v) \) given \( Y \) as the maximum distance between \( v \) and any infected nodes in the graph. Define the Jordan infection centers of a graph to be the nodes with the minimum infection eccentricity given \( Y \). In Figure 2, nodes \( v_3 \), \( v_{10} \), \( v_{13} \) and \( v_{14} \) are observed to be infected. The infection eccentricities of \( v_1, v_2, v_3, v_4 \) are 2, 3, 4, 5, respectively, and the Jordan infection center is \( v_1 \).

We will show that the source associated with the optimal sample path is a node with the minimum infection eccentricity. We derive this result using three steps: first, assuming the information source is \( v_r \), we analyze \( t_{v_r}^* \) such that

\[ t_{v_r}^* = \arg \max_{t, X[0,t]} \Pr(X[0,t]\mid v^* = v_r), \]

We say \( F(X[t]) = Y \) if \( F(X_v(t)) = Y_v \) for all \( v \). Identifying the information source can be formulated as a maximum likelihood detection problem as follows:

\[ v^* \in \arg \max_{v \in \mathcal{V}} \sum_{X[0,t]:F(X(t))=Y} \Pr(X[0,t]\mid v^* = v), \]
The largest distance from \( v_0 \) to an infected node is \( k \). \( \mathcal{Y}^0 \) is the topology where there is only one infected node—the root node \( v_r \). Note that if no infected node is observed, no algorithm performs better than a random guess. To prove \( \mathcal{Y} \), we use induction over \( k \).

**Step 1:** First, we consider the case \( k = 0 \). All the sample paths considered in step 1 lead to observation \( \mathcal{Y} \in \mathcal{Y}^0 \). We denote by \( T_{v_r} \), the tree rooted in \( v_r \) and \( T_{v_r}^{-v_r} \) the tree rooted at \( u \) but without the branch from \( v_r \). For example, Figure 3 shows \( T_{v_1}^{-v_1} \), \( T_{v_2}^{-v_2} \), \( T_{v_3}^{-v_3} \) and \( T_{v_4}^{-v_4} \). The sample path from time slot 0 to \( t \) restricted to \( T_u^{-v_r} \) is denoted by \( \mathfrak{X}(\mathcal{Y}[0,t], T_u^{-v_r}) \).

Furthermore, denote by \( C(v) \) the set of children of \( v \). We have

\[
\Pr(\mathfrak{X}(\mathcal{Y}[0,t], T_u^{-v_r})|X_{v_r}(t) = I) = (1 - p)^t \prod_{u \in C(v_r)} \Pr(\mathfrak{X}(\mathcal{Y}[0,t], T_u^{-v_r})|X_{v_r}(t) = I),
\]

where the last equality holds since \( v_r \) is the only infected node in the network at time \( t \), which requires \( X_{v_r}(s) = I \) for \( 0 \leq s \leq t \). Node \( u \in C(v_r) \) has two possible states \( S \) or \( R \).

**Step 1.a** \( u \) is susceptible if it was not infected within time slots. In each time slot, \( v_r \) tries to infect \( u \) with probability \( q \). The probability that \( u \) is susceptible at time slot \( t \) is

\[
(1 - q)^t,
\]

which implies that

\[
\Pr(\mathfrak{X}(\mathcal{Y}[0,t], T_u^{-v_r})|X_{v_r}(t) = I) = (1 - q)^t
\]

if \( X_u(t) = S \).

**Step 1.b** If \( u \) is in the recovered state, we denote by \( t_u^R \) and \( t_u^R(u) \) its infection and recovery times, respectively. Then, we have if \( X_u(t) = R \),

\[
\Pr(\mathfrak{X}(\mathcal{Y}[0,t], T_u^{-v_r})|X_{v_r}(t) = I) = (1 - q)^{t_u^R(u)} q(1 - p)^{t_u^R(u) - t_u^R(u) - 1} p \prod_{u \in C(v_r)} \Pr(\mathfrak{X}(\mathcal{Y}[0,t], T_u^{-v_r})|t_u^R(u), t_u^R),
\]

where \( (1 - q)^{t_u^R(u)} q(1 - p)^{t_u^R(u) - t_u^R(u) - 1} p \) is the probability that node \( u \) was infected at time \( t_u^R(u) \) and recovered at time \( t_u^R(u) \). Since \( T_u^{-v_r} \) is also an infinite tree, there exists at least one node \( \xi \in T_u^{-v_r} \) such that the node is in the susceptible state but its parent node (say node \( \gamma \)) is in the recovered state. We denote by \( T_u^{-v_r} \setminus T_{\xi^{-v_r}} \) the set of nodes that are on subtree \( T_u^{-v_r} \) but not on subtree \( T_{\xi^{-v_r}} \). Then,

\[
\Pr(\mathfrak{X}(\mathcal{Y}[0,t], T_u^{-v_r})|t_u^R(u), t_u^R) \\
= \Pr(X(\mathcal{Y}[0,t], T_u^{-v_r}(\xi^{-v_r}))|t_u^R(u), t_u^R) \times \Pr(X(\mathcal{Y}[0,t], T_{\xi^{-v_r}}^{-v_r})|t_u^R(u), t_u^R) \leq (1 - q)^{t_u^R(u)} \leq (1 - q)^t,
\]

\[
(2)
\]

We divide all possible infection topologies \( \mathcal{Y} \) into countable subsets \( \{\mathcal{Y}^k\} \) where \( \mathcal{Y}^k \) is the set of infection topologies where i.e., \( t_{v_r}^* \) is the time duration of the optimal sample path in which \( v_r \) is the information source. It turns out that \( t_{v_r}^* \) equals to the infection eccentricity of node \( v_r \). Considering Figure 2 if the source is \( v_1 \), then the time duration of the optimal sample path starting from \( v_1 \) is 2.

In the second step, we consider two neighboring nodes, say nodes \( v_1 \) and \( v_2 \). We will prove that if \( \hat{e}(v_1) < \hat{e}(v_2) \), then the optimal sample path rooted at \( v_1 \) occurs with a higher probability than the optimal sample path rooted at \( v_2 \).

Finally, at the third step, we will show that given any two nodes \( u \) and \( v \), if \( v \) has the minimum infection eccentricity and \( u \) has a larger infection eccentricity, then there exists a path from \( u \) to \( v \) along which the infection eccentricity monotonically decreases, which implies that the source of the optimal sample path must be a Jordan infection center. For example, in Figure 2 node \( v_4 \) has a larger infection eccentricity than \( v_1 \) and \( v_4 \rightarrow v_3 \rightarrow v_2 \rightarrow v_1 \) is the path along which the infection eccentricity monotonically decreases from 5 to 2.

**A. The Optimal Time**

**Lemma 1.** Consider a tree network rooted at \( v_r \) and with infinitely many levels. Assume the information source is the root, and the observed infection topology is \( \mathcal{Y} \) which contains at least one infected node. If \( \hat{e}(v_r) \leq t_1 < t_2 \), then the following inequality holds

\[
\max_{X[0,t_1] \in \mathcal{X}(t_1)} \Pr(\mathfrak{X}(0,t_1)) \geq \max_{X[0,t_2] \in \mathcal{X}(t_2)} \Pr(\mathfrak{X}(0,t_2)),
\]

where \( \mathcal{X}(t) = \{X[0,t]|F(X(t)) = \mathcal{Y}\} \). In addition,

\[
t_{v_r} = \hat{e}(v_r) = \max_{u \in I} d(v_r, u),
\]

where \( d(v_r, u) \) is the length of the shortest path between \( v_r \) and \( u \) and also called the distance between \( v_r \) and \( u \), and \( I \) is the set of infected nodes.

**Proof:** We start from the case where the time difference of two sample paths is one, i.e., we will show that

\[
\max_{X[0,t] \in \mathcal{X}(t)} \Pr(\mathfrak{X}(0,t)) \geq \max_{X[0,t+1] \in \mathcal{X}(t+1)} \Pr(\mathfrak{X}(0,t+1)).
\]

\[
(2)
\]
where equation (7) holds because $\xi$ remained to be susceptible during the time slots at which $\gamma$ was in the infected state and (8) holds because $t_{u}^L - t_{u}^R \geq 1$. The maximum value of $\Pr(X([0, t], T_{u}^{-v_r})|t_{u}^L, t_{u}^R)$ can be achieved in the sample path in which $u$ was infected and then recovered in the next time slot so that $w$ was vulnerable to infection only in one time slot. Furthermore,

$$(1 - q) t_{u}^L - q(1 - p) t_{u}^R$$

is maximized when $t_{u}^L = 1, t_{u}^R = 2$ i.e., $u$ was infected at the first time slot and recovered in the second time slot. Therefore, if $X_u(t) = R$,

$$\Pr(X([0, t], T_{u}^{-v_r})|X_{v_r}(t) = I) \leq q(1 - q)^{|C(u)|}.$$  \hspace{1cm} (9) \hspace{1cm} \textbf{Step 1.c} Define $X^*([0, t])$ to be the optimal solution to

$$\max_{X([0, t]) \in X(t)} \Pr(X([0, t])).$$

For $t = 1$, since all $u \in C(v_r)$ are in the susceptible state,

$$\Pr(X^*([0, t]) = (1 - p)(1 - q)^{|C(v_r)|}.$$  \hspace{1cm} (10) \hspace{1cm} \text{For } t \geq 2, \text{ according to (5) and (9),}

$$\Pr(X^*([0, t]) = (1 - p)^t \prod_{u \in C(v_r)} \max \left\{ (1 - q)^t, q(1 - q)^{|C(u)|} \right\}.$$  \hspace{1cm} (11) \hspace{1cm} \text{(12)}

Note that $t$ is fixed in this optimization problem and (12) is a none-increasing function of $t$. Since $|C(u)| \geq 1$,

$$\Pr(X^*([0, 2]) \leq (1 - p)^2 \max \left\{ (1 - q)^2, (q(1 - q))^{|C(v_r)|} \right\} < (1 - p)(1 - q)^{|C(v_r)|}$$

$$= \Pr(X^*([0, 1])).$$

In a summary, $\Pr(X^*([0, t]))$ is a none-increasing function of $t \in [1, \infty)$ when $k = 0$.

\textbf{Step 2:} Assume (2) holds for $k \leq n$, and consider $k = n + 1$. Clearly $t \geq n + 1 \geq 1$ for each $X([0, t])$ such that

$$F(X([0, t]) \in \mathcal{Y}^{n+1}.$$

Furthermore, the set of subtrees $T = \{ T_{u}^{-v_r} | u \in C(v_r) \}$ are divided into two subsets: $T^h = \{ T_{u}^{-v_r} | u \in C(v_r), Y(T_{u}^{-v_r}) \cap \mathcal{I} = \emptyset \}$ and $T^v = T \setminus T^h$, where $Y(T_{u}^{-v_r})$ is the vector of $Y$ restricted to subtree $T_{u}^{-v_r}$. In Figure 3, $T^h = \{ T_{u}^{-v_r} \}$ and $T^v = \{ T_{v_1}^{-v_r}, T_{v_2}^{-v_r}, T_{v_3}^{-v_r} \}$. We note that given $t_{v_r}^R$, the infection processes on the sub-trees are mutually independent.

\textbf{Step 2.a} Recall that $T^h$ is the set of subtrees having no infected nodes. Following the argument for the $k = 0$ case, we can obtain that if $T_{u}^{-v_r} \in T^h$, then

$$\Pr(X^*([0, t], T_{u}^{-v_r})|t_{u}^R) = \max \left\{ (1 - q)^{t_{u}^R}, q(1 - q)^{|C(u)|} \right\}$$

when $t \geq t_{u}^R$ and

$$\Pr(X^*([0, t], T_{u}^{-v_r})|t_{u}^R) = \max \left\{ (1 - q)^t, q(1 - q)^{|C(u)|} \right\}$$

when $t < t_{u}^R$. So $\Pr(X([0, t], T_{u}^{-v_r})|t_{u}^R)$ is non-increasing in $t$ given any $t_{u}^R$.

\textbf{Step 2.b} For $T_{v_r}^{-v_r} \in T^v$, given the sample path $X([0, t + 1], T_{u}^{-v_r})$, we will construct a sample path $X([0, t], T_{u}^{-v_r})$ which occurs with a higher probability. Denote the infection time of $u$ in sample path $X([0, t], T_{u}^{-v_r})$ by $t_{u}^I$. We let $t_{u}^I$ denote the infection time in sample path $X([0, t + 1], T_{u}^{-v_r})$.

If $t_{u}^I > 1$, we choose $t_{u}^I = t_{u}^I - 1$, i.e., $u$ is infected one time slot later in $X([0, t + 1])$ than in $X([0, t])$. Assume the infection processes after $u$ was infected are the same in the two sample paths $X([0, t], T_{u}^{-v_r})$ and $X([0, t + 1], T_{u}^{-v_r})$. Therefore, we have

$$\Pr(X([0, t + 1], T_{u}^{-v_r}) = (1 - q)^{t_{u}^I - 1} q \Pr(X([0, t + 1], T_{u}^{-v_r})|t_{u}^I).$$

and

$$\Pr(X([0, t], T_{u}^{-v_r}) = (1 - q)^{t_{u}^I} q \Pr(X([0, t], T_{u}^{-v_r})|t_{u}^I).$$

where $\Pr(X([0, t], T_{u}^{-v_r})|t_{u}^I)$ is the probability of $X([0, t], T_{u}^{-v_r})$ after $u$ was infected. Since the sample paths $X([0, t], T_{u}^{-v_r})$ and $X([0, t + 1], T_{u}^{-v_r})$ are the same after $u$ was infected, we obtain

$$\Pr(X([0, t], T_{u}^{-v_r})|t_{u}^I) = \Pr(X([0, t + 1], T_{u}^{-v_r})|t_{u}^I).$$

Therefore, with $t_{u}^I = t_{u}^I - 1$, we get

$$\Pr(X([0, t + 1], T_{u}^{-v_r}) < \Pr(X([0, t], T_{u}^{-v_r})).$$

If $t_{u}^I = 1$, we set $t_{u}^I = t_{u}^I = 1$. Based on the induction assumption, for $k \leq n$ since $Y(T_{u}^{-v_r}) \in \mathcal{Y}^{n}, m \leq n$, we have

$$\max_{X([0, t], T_{u}^{-v_r}) \in \mathcal{X}(t, T_{u}^{-v_r})} \Pr(X([0, t], T_{u}^{-v_r})).$$

where $\mathcal{X}(t, T_{u}^{-v_r}) = \{ X([0, t], T_{u}^{-v_r}) : F(X([0, t], T_{u}^{-v_r}) = Y(T_{u}^{-v_r}) \}$. Therefore, given any $X([0, t + 1], T_{u}^{-v_r})$, we can always find a corresponding sample path $X([0, t], T_{u}^{-v_r})$, which occurs with a higher probability.

\textbf{Step 2.c} Now we consider the sample path $X^*([0, t + 1])$ and denote by $t_{v_r}^R$ the recovery time of node $v_r$ in $X^*([0, t + 1])$. We now construct a sample path $X([0, t])$ as follows:

\textbf{Note that we cannot apply the same argument to $t_{u}^I > 1$ because $t_{u}^I = t_{u}^I$ may not be feasible in a valid $X([0, t])$.
If $\hat{t}_u^R > t + 1$, i.e., $v_r$ is an infected node, then $\hat{t}_v^R > t$, where $\hat{t}_v^R$ is the recovery time of $v_r$ in $X[0, t]$.

- If $\hat{t}_u^R \leq t$, we choose $\hat{t}_v^R = \hat{t}_u^R$.
- If $\hat{t}_u^R = t + 1$, we choose $\hat{t}_v^R = t$.

We further complete $X(0, t)$ by having optimal ones on $T^h$ and constructing the ones in $T$ following step 2.b. According to steps 2.a and 2.b, it is easy to verify that $X(0, t)$ occurs with a higher probability than $X^*([0, t + 1])$. Therefore, we conclude that inequality (2) holds for $k = n + 1$, hence for any $k$ according to the principle of induction.

**Step 3** Repeatedly applying inequality (2), we obtain that $t_v^*$ is the minimum amount of time required to produce the observed infection topology. The minimum time required is equal to the maximum distance from $v_r$ to an infected node. Therefore, the lemma holds.

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**B. The Sample Path Based Estimator**

After deriving $t_v^*$, we have a unique $t_u^*$ for each $v \in V$. The next lemma states that the optimal sample path starting from a node with a smaller infection eccentricity is more likely to occur.

**Lemma 2.** Consider a tree network with infinitely many levels. Assume the information source is the root, and the observed infection topology is $Y$ which contains at least one infected node. For $u, v \in V$ such that $(u, v) \in E$, if $t_u^* > t_v^*$, then

$$\text{Pr}(X_u^*([0, t_u^*])) < \text{Pr}(X_v^*([0, t_v^*])),$$

where $X_u^*([0, t_u^*])$ is the optimal sample path starting from node $u$.

**Proof:** Recall that $T_v$ denotes the tree rooted at $v$ and $T_v^{-u}$ denotes the tree rooted at $u$ but without the branch from $v$. Furthermore, $C(v)$ is the set of children of $v$, and $X([0, t_v], T_v^{-u})$ is the sample path $X([0, t_v])$ restricted to $T_v^{-u}$.

**Step 1:** The first step is to show $t_u^* > t_v^* + 1$. First we claim $T_v^{-u} \cap I \neq \emptyset$. Otherwise, all infected nodes are on $T_v^{-u}$. Since on a tree, $v$ can only reach nodes in $T_v^{-u}$ through edge $(u, v)$, $t_u^* = t_v^* + 1$, which contradicts $t_u^* > t_v^*$. If $T_v^{-u} \cap I \neq \emptyset$, $\forall a \in T_v^{-u} \cap I$, we have

$$d(u, a) = d(v, a) + 1 \leq t_v^* + 1,$$

and $d(u, b) = d(v, b) + 1 \leq t_v^* + 1$.

Hence,

$$d(u, a) = d(u, b) + 1 \leq t_v^* + 1,$$

which implies that

$$t_u^* \leq t_v^* + 1,$$

i.e., $t_u^* = t_v^* + 1$.

If $T_v^{-u} \cap I = \emptyset$, all infected nodes are in $T_v^{-u}$, so it is obvious $t_u^* = t_v^* + 1$.

**Step 2:** In this step, we will prove that $t_u^* = 1$ on the sample path $X_u^*([0, t_u^*])$. If $t_u^* > 1$ on $X_u^*([0, t_u^*])$, then

$$t_u - t_v = t_v^* + 1 - t_u^* < t_v^*.$$

Note that according to the definition of $t_u^*$ and $t_v^*$, within $t_u^* - t_v^*$ time slots, node $v$ can infect all infected nodes on $T_v^{-u}$. Since $t_u^* = t_v^* + 1$, the infected node farthest from node $u$ must be on $T_v^{-u}$, which implies that there exists a node $a \in T_v^{-u}$ such that $d(u, a) = t_u^* = t_v^* + 1$ and $d(v, a) = t_v^*$. So node $v$ cannot reach $a$ within $t_v^* - t_u^*$ time slots, which contradicts the fact that the infection can spread from node $v$ to $a$ within $t_v^* - t_u^*$ time slots along the sample path $X_u^*([0, t_u^*])$. Therefore, $t_u^* = 1$.

**Step 3:** Now given sample path $X_u^*([0, t_u^*])$, we construct $X_v([0, t_v^*])$ which occurs with a higher probability. We divide the sample path $X_u^*([0, t_u^*])$ into two parts along subtrees $T_v^{-u}$ and $T_v^{-u}$. Since $t_u^* = 1$, we have

$$\text{Pr}(X_u^*([0, t_u^*])) = q \text{Pr}(X_u^*([0, t_u^*], T_v^{-u}) \mid t_v^* = 1) \text{Pr}(X_v^*([0, t_v^*], T_v^{-u}) \mid t_u^* = 1).$$

For the subtree $T_v^{-u}$, given $X_u^*([0, t_u^*], T_v^{-u})$, in which $t_u^* = 1$, we construct the partial sample path $X_v^*([0, t_v^*], T_v^{-u})$ to be identical to $X_v^*([0, t_v^*], T_v^{-u})$ except that all events occur one time slot earlier, i.e.,

$$X_v^*([0, t_v^*], T_v^{-u}) = X_v^*([1, t_v^*], T_v^{-u}).$$

This is feasible because $t_v^* = t_u^* - 1$. Then

$$\text{Pr}(X_u^*([0, t_u^*], T_v^{-u}) \mid t_v^* = 1) = \text{Pr}(X_v^*([0, t_v^*], T_v^{-u}) \mid t_u^* = 1).$$

For the subtree $T_v^{-u}$, we construct $X_v^*([0, t_v^*], T_v^{-u})$ such that

$$X_v^*([0, t_v^*], T_v^{-u}) \in \text{arg}\max_{X([0, t_v^*], T_v^{-u}) \in X([t_v^*], T_v^{-u})} \text{Pr}(X([0, t_v^*], T_v^{-u}) \mid t_v^* = 1).$$

Based on Lemma 2, we have

$$\text{max}_{X([0, t_v^*], T_v^{-u}) \in X([t_v^*], T_v^{-u})} \text{Pr}(X([0, t_v^*], T_v^{-u}) \mid t_v^* = 1) = \text{max}_{X([0, t_v^*], T_v^{-u}) \in X([t_v^*], T_v^{-u})} \text{Pr}(X([0, t_v^*], T_v^{-u}) \mid t_u^* = 1) > \text{max}_{X([0, t_v^*], T_v^{-u}) \in X([t_v^*], T_v^{-u})} \text{Pr}(X([0, t_v^*], T_v^{-u}) \mid t_v^* = 1).$$

Therefore, given the optimal sample path rooted at $u$, we have constructed a sample path rooted at $v$ which occurs with a higher probability. The lemma holds.

Next, we give a useful property of the Jordan infection centers in the following lemma.

**Lemma 3.** On a tree network with at least one infected node, there exist at most two Jordan infection centers. When the network has two Jordan infection centers, the two must be neighbors.

\[\square\]
Proof: First, we claim if there are more than one Jordan infection centers, they must be adjacent. Suppose \( v, u \in \mathcal{V} \) are two Jordan infection centers and \( \bar{e}(v) = \bar{e}(u) = \lambda \). Suppose \( v \) and \( u \) are not adjacent, i.e., \( d(v, u) > 1 \). Then, there exists \( w \in \mathcal{V} \) such that
\[
d(w, u) = 1,
\]
and
\[
d(w, v) = d(v, u) - 1,
\]
i.e., \( w \) is a neighbor of \( u \) and is on the shortest path between \( u \) and \( v \). Note in a tree structure \( w \) is unique.

If \( \mathcal{I} \cap \mathcal{T}_w = \emptyset \), then \( \forall a \in \mathcal{I} \),
\[
d(w, a) = d(u, a) - 1 < d(u, a),
\]
which contradicts the fact that \( u \) is a Jordan infection center.

If \( \mathcal{I} \cap \mathcal{T}_w \neq \emptyset \). Since \( \forall b \in \mathcal{I} \cap \mathcal{T}_w \),
\[
d(v, b) = d(v, w) + d(w, b),
\]
i.e.,
\[
d(w, b) = d(v, b) - d(v, w) \leq \lambda - 1.
\]
On the other hand, since \( \bar{e}(u) = \lambda \), \( \forall h \in T_w^{-u} \cap \mathcal{I} \),
\[
d(w, h) = d(u, h) - 1 \leq \lambda - 1.
\]
In a summary, \( \forall h \in \mathcal{I} \),
\[
d(w, h) \leq \lambda - 1,
\]
which contradicts the fact that the minimum infection eccentricity is \( \lambda \).

Therefore all Jordan infection centers must be adjacent to each other. However, suppose there exist \( n \) infection eccentric centers where \( n > 2 \), they would form a clique with \( n \) nodes which contradicts the fact that the graph is a tree. Therefore, there exist at most two adjacent Jordan infection centers.

Based on Lemma 2 and Lemma 3 we finish this section with the following theorem.

**Theorem 4.** Consider a tree network with infinitely many levels. Assume that the observed infection topology \( \mathbf{Y} \) contains at least one infected node. Then the source node associated with \( \mathbf{X}^*[0, t^*] \) (the solution to the optimization problem (7)) is a Jordan infection center, i.e.,
\[
v^* = \arg \min_{v \in \mathcal{V}} \bar{e}(v).
\]

Proof: We assume the network has two Jordan infection centers: \( w \) and \( u \), and assume \( \bar{e}(w) = \bar{e}(u) = \lambda \). The same argument works for the case where the network has only one Jordan infection center.

Based on Lemma 3, \( w \) and \( u \) must be adjacent. We will show for any \( a \in \mathcal{V} \setminus \{w, u\} \), there exists a path from \( a \) to \( u \) (or \( w \)) along which the infection eccentricity strictly decreases.

**Step 1:** First, it is easy to see from Figure 4 that \( d(\gamma, w) \leq \lambda - 1 \) \( \forall \gamma \in T_w^{-u} \cap \mathcal{I} \). We next show that there exists a node \( \xi \) such that the equality holds.

In a summary, \( \forall \gamma \in \mathcal{I} \),
\[
d(\gamma, u) \leq \lambda - 1.
\]
This contradicts the fact that \( \bar{e}(w) = \bar{e}(u) = \lambda \). Therefore, there exists \( \xi \in T_w^{-u} \cap \mathcal{I} \) such that
\[
d(\xi, w) = \lambda - 1.
\]

**Step 2:** Similarly, \( \forall \gamma \in T_w^{-u} \cap \mathcal{I} \),
\[
d(\gamma, u) \leq \lambda - 1,
\]
and there exists a node such that the equality holds.

**Step 3:** Next we consider \( a \in \mathcal{V} \setminus \{w, u\} \), and assume \( a \in T_w^{-u} \) and \( d(a, u) = \beta \). Then for any \( \gamma \in T_w^{-u} \cap \mathcal{I} \), we have
\[
d(a, \gamma) = d(a, u) + d(u, w) + d(w, \gamma) \leq \beta + 1 + \lambda - 1 = \lambda + \beta,
\]
and there exists \( \xi \in T_w^{-u} \cap \mathcal{I} \) such that the equality holds. On the other hand, \( \forall \gamma \in T_w^{-u} \cap \mathcal{I} \),
\[
d(a, \gamma) \leq d(a, u) + d(u, \gamma) \leq \beta + \lambda - 1.
\]
Therefore, we conclude that
\[
\bar{e}(a) = \lambda + \beta,
\]
so the infection eccentricity decreases along the path from \( a \) to \( u \).

**Step 4:** Repeatedly applying Lemma 2 along the path from node \( a \) to \( u \), we can conclude that the optimal sample path rooted at node \( u \) is more likely to occur than the optimal sample path rooted at node \( a \). Therefore, the root node associated with the optimal sample path \( \mathbf{X}^*[0, t^*] \) must be a Jordan infection center, and the theorem holds.
IV. REVERSE INFECTION ALGORITHM

Since in tree networks with infinitely many levels, the estimator based on the sample path approach is a Jordan infection center, we view the Jordan infection centers as possible candidates of the information source. We next present a simple algorithm to find the information source in general networks. The algorithm is to first identify the Jordan infection centers, and then break ties based on the sum of distances to infected nodes.

The key idea of the algorithm is to let every infected node broadcast a message containing its identity (ID) to its neighbors. Each node, after receiving messages from its neighbors, checks whether the ID in the message has been received. If not, the node records the ID (say \(v\)), the time at which the message is received (say \(t_v\)), and then broadcasts the ID to its neighbors. When a node receives the IDs of all infected nodes, it claims itself as the information source and the algorithm terminates. If there are multiple nodes receiving all IDs at the same time, the tie is broken by selecting the node with the smallest \(\sum t_v\).

The tie-breaking rule we proposed is to choose the node with the maximum infection closeness [17]. The closeness measures the efficiency of a node to spread information to all other nodes. The closeness of a node is the inverse of the sum of distances from the node to any other nodes. In our model, we define the infection closeness as the inverse of the sum of distances from a node to all infected nodes, which reflects the efficiency to spread information to infected nodes. We select a Jordan infection center with the largest infection closeness, breaking ties at random.

**Algorithm 1 Reverse Infection Algorithm**

```markdown
for \(i \in I\) do
    \(i\) sends its ID \(\omega_i\) to its neighbors.
end for

while \(t \geq 1\) and STOP == 0 do
    for \(u \in V\) do
        if \(u\) receives \(\omega_i\) for the first time then
            Set \(t_{ui} = t\) and then broadcast the message \(\omega_i\) to its neighbors.
            If there exists a node who received \(|I|\) distinct messages, then set \(STOP == 1\).
        end if
    end for
end while

return \(u^* = \arg\min_{u \in S} \sum_{i \in I} t_{ui}\), where \(S\) is the set of nodes who receive \(|I|\) distinct messages when the algorithm terminates. Ties are broken at random.
```

V. PERFORMANCE ANALYSIS

The reverse infection algorithm is based on the structure properties of the optimal sample paths on trees. While the MLE is the node that maximizes the likelihood of the snapshot among all possible nodes, the sample path based estimator does not have such a guarantee. To demonstrate the effectiveness of the sample path based approach, we next show that on \((g+1)\)-regular trees where each node has \(g+1\) neighbors, the information source generated by the reverse infection algorithm is within a constant distance from the actual source with a high probability, independent of the number of infected nodes and the time at which the snapshot \(Y\) was taken.

**Theorem 5.** Consider a \((g+1)\)-regular tree with infinitely many levels where \(g > 2\) and \(gq > 1\). Assume that the observed infection topology \(Y\) contains at least one infected node. Given \(\epsilon > 0\), there exists \(d_{\epsilon}\) such that the distance between the optimal sample path estimator and the actual source is \(d_{\epsilon}\), with probability \(1 - \epsilon\), where \(d_{\epsilon}\) is independent of the number of infected nodes and the time the snapshot \(Y\) was taken.

**Proof:** Consider the tree rooted at the information source \(v^*\). We say \(v^*\) is at level 0. We denote by \(Z_l\) the set of infected and recovered nodes at level \(l\). Furthermore, we define \(Z_l^\ast\) to be the set of infected and recovered nodes at level \(l\) whose parents are in set \(\mathcal{I}\) and who were infected within \(\tau\) time slots after their parents were infected. We assume \(Z_\ast = \{v^*\}\). In addition, let \(Z_l = |Z_l|\) and \(Z_l^\ast = |Z_l^\ast|\).

Note

\[
\lim_{\tau \to \infty} Z_l^\ast = Z_l,
\]

and given \(v\) and \(u \in Z_l^\ast\),

\[
|t_u^\ast - t_v^\ast| \leq l(\tau - 1),
\]

i.e., the infection times of nodes in \(Z_l^\ast\) differ by at most \(l(\tau - 1)\) (note that the difference is not \(\tau - 1\) since the parents of \(u\) and \(v\) may be infected at different times). Our proof is based on the Galton Watson (GW) branching process [18]. A GW branching process is a stochastic process \(B(l)\) which evolves according to the recurrence formula \(B(0) = 1\) and

\[
B(l) = \sum_{i=1}^{B(l-1)} \zeta_i,
\]

where \(\{\zeta_i\}\) is a set of random variables, taking values from nonnegative integers. The distribution of \(\zeta_i\) is called the offspring distribution of the branching process. In a \((g+1)\)-regular tree, the evolution of \(Z_l^\ast\) is a branching process, where the offspring distribution is a function of \(\tau\). We use \(B^\ast\) to denote the corresponding branching process, and \(B^\ast(l)\) to denote the number of offsprings at level \(l\), i.e., \(B^\ast(l) = Z_l^\ast\) (we use these two notations interchangeably). Given a node is in the infected state for \(l\) time slots, the number of infected offsprings follows a binomial distribution. Note the following two facts:
The number of survived processes on the subtree rooted at the node. We define $Y$ as the number of infected nodes in the distance between two events:

1. Event 1: The number of infected nodes at level $l$ occurs, then we have $Z = 0$.
2. Event 2: The number of infected nodes at level $l$ is at least $l$. Recall that at time $t$, the distance between any infected node and the actual source is no more than $t$, which implies the eccentricity of a Jordan infection center is at most $t$.

Figures 5. A pictorial description of the positions of $v', \tilde{v}, u_1$, and $w$.

- The number of time slots at which a node is in the infected state follows a geometric distribution with parameter $p$.
- A child remains to be susceptible with probability $(1-q)^\tau$ when the parent has been in the infected state for $\tau$ time slots.

Therefore, the offspring distribution of the branching process $B^\tau$ at level $\geq t$ is

$$\Pr(\gamma = i) = \sum_{t=1}^{\tau-1} (1-p)^{t-1}p \binom{g}{i} (1-(1-q)^t)^i (1-q)^{\tau-g-i},$$

where $\gamma$ is the number of offsprings of a node. The offspring distribution of branching process $B^\infty$ is

$$\Pr(\gamma' = i) = \sum_{t=1}^{\infty} (1-p)^{t-1}p \binom{g}{i} (1-(1-q)^t)^i (1-q)^{t(g-i)}.$$ 

Each infected node can be viewed as the source of branching processes on the subtree rooted at the node. We define $K_l$ to be the number of survived $B^1$ branching processes whose roots are in set $Z^*_l$, where a branching process survives if it never dies out.

Now given $L \geq 2$, we consider the following events:

- Event 1: $Z_L = 0$
- Event 2: $K_l \geq 2$ for some $l \leq L$. In other words, at least two $B^1$ branching processes starting from $Z^*_l$ survive for some $l \leq L$.

We note that these two are disjoint events.

When $Z_l = 0$, no node at level $L$ is infected and the infection process terminates at level $L - 1$. When there is at least one infected node in $\mathbf{Y}$, since $\widehat{e}(v^*) \leq L - 1$, the minimum infection eccentricity is at most $L - 1$. Therefore, the distance between $v^*$ and $v^1$ is no more than $2(L - 1)$.

The source node has $g + 1$ children while other nodes have $g$ children.

Given $K_l \geq 2$ for some $l \leq L$, we will argue that the distance between the sample path based estimator and the actual one is upper bounded by $(\tau + 1)L - 1$. Consider Figure 5 where the shaded nodes are infected and recovered nodes. We will show that if two $B^1$ branching processes starting from $l \leq L$ survive, a node at level $\geq (\tau + 1)L - 1$ cannot be a Jordan infection center. Recall that at time $t$, the distance between any infected node and the actual source is no more than $t$, which implies the eccentricity of a Jordan infection center is at most $t$.

Now consider a node $\tilde{v}$ at level $\geq (\tau + 1)l - 1$. Recall that at least two $B^1$ branching processes starting from level $l$ survive. Let $u_1 \in Z^*_l$ be the root of a survived $B^1$ branching process, and assume node $\tilde{v}$ is not on the subtree rooted at $u_1$. Further, assume $v'$ is an infected node at the lowest level on sub-tree $T^w_1$. Since the branching process $B^1_1$ survives, the infection process propagates one level lower at each time slot and node $v'$ is at level $l + t - t^l_{u_1}$.

From Figure 5, it is easy to see that the distance between $v'$ and $\tilde{v}$ is at least

$$t - t^l_{u_1} + 2 + (\tau + 1)(l - 1) = t - t^l_{u_1} + \tau l + 1,$$

which occurs when the first common predecessor of nodes $v'$ and $\tilde{v}$ is at level $\leq 1$. Note that the common predecessor cannot appear at level $\geq l$ since $v'$ is not on $T^w_1$. Since $u_1 \in Z^*_l$, the infection time of node $u_1$ is no later than $\tau l$, i.e., $t^l_{u_1} \leq \tau l$. Therefore, the distance between $v'$ and $\tilde{v}$ is at least $t + 1$, which is larger than $t$. Hence, $v'$ cannot be a Jordan infection center. Since $\leq L$, any node at or below level $(\tau + 1)L - 1$ cannot be a Jordan infection center. In a summary, if event 2 occurs, then we have

$$d(v^*, \tilde{v}) \leq (\tau + 1)L - 1.$$ 

We next show that given any $\epsilon$, we can find sufficiently large $\tau$ and $L$, independent of $t$ and the number of infected nodes, such that the probability that either event 1 or event 2 occurs is at least $1 - \epsilon$.

Given $n_0 > 0$ and $\tau > 0$, we define

$$l^1 = \min \{ l : Z^*_l > n_0 \},$$

i.e., $l^1$ is the first level at which $B^\tau$ has more than $n_0$ nodes. We first have

$$\Pr(Z_L = 0) + \Pr(K_l \geq 2 \text{ for some } l \leq L) \geq \Pr(Z_L = 0) + \Pr(K_l \geq 2 \text{ and } l^1 \leq L),$$

$$= \Pr(Z_L = 0) + \Pr(l^1 \leq L) \Pr(K_l \geq 2 \mid l^1 \leq L),$$

$$= \Pr(Z_L = 0) + \Pr \left( \bigcup_{i=1}^{L} \{ Z^*_i > n_0 \} \right) \Pr(K_l \geq 2 \mid l^1 \leq L),$$

$$\geq \left( 1 - \Pr \left( \bigcap_{i=1}^{L} \{ Z^*_i \leq n_0 \} \right) \right) - \Pr \left( \bigcup_{i=1}^{L} \{ Z^*_i = 0 \} \right),$$

$$\times \Pr \left( K_l \geq 2 \mid l^1 \leq L \right) + \Pr(Z_L = 0).$$
Note that we have
\[
\Pr(K_l \geq 2|t| \leq L) = \sum_{i=1}^{L} \Pr(K_{it} \geq 2|t| = l) \Pr(l|t| \leq L).
\] (13)
According to Lemma 6, given any \(\epsilon > 0\), we can find a sufficiently large \(n_0\) such that
\[
\Pr(K_l \geq 2|t| = l) \geq (1 - \epsilon_1),
\]
which implies that for sufficiently large \(n_0\),
\[
\Pr(K_l \geq 2|t| \leq L) \geq 1 - \epsilon_1.
\]
We can then conclude
\[
\Pr(Z_L = 0) + \Pr(K_l \geq 2 \text{ for some } l \leq L)
\geq \left(1 - \Pr \left( \bigcup_{i=1}^{L} \{0 < Z_i^L \leq n_0\} \right) \right) (1 - \epsilon_1)
- \Pr \left( \bigcup_{i=1}^{L} \{Z_i^L = 0\} \right) + \Pr(Z_L = 0)
= \left(1 - \Pr \left( \bigcap_{i=1}^{L} \{0 < Z_i^L \leq n_0\} \right) \right) (1 - \epsilon_1)
+ \Pr(Z_L = 0) - \Pr(Z_L^\ast = 0),
\]
where \(\Pr(\bigcup_{i=1}^{L} \{Z_i^L = 0\}) = \Pr(Z_L^\ast = 0)\) because \(Z_i^L = 0\) implies that \(Z_L^\ast = 0\) for \(l \leq L\).
According to Lemma 7 and Lemma 8, given any \(\epsilon_2 > 0\) and \(\epsilon_3 > 0\), there exist sufficiently large \(\tau\) and \(L\) such that
\[
1 - \Pr \left( \bigcap_{i=1}^{L} \{0 < Z_i^L \leq n_0\} \right) > 1 - \epsilon_2,
\]
and
\[
\Pr(Z_L = 0) - \Pr(Z_L^\ast = 0) \geq -\epsilon_3.
\]
Hence, we have
\[
\Pr(Z_L = 0) + \Pr(K_l \geq 2 \text{ for some } l \leq L)
\geq (1 - \epsilon_1)(1 - \epsilon_2) - \epsilon_3.
\]
Now choosing \(\epsilon_1 = \epsilon_2 = \epsilon_3 = \epsilon_4/3\) for some \(\epsilon_4 > 0\), we have
\[
\Pr(Z_L = 0) + \Pr(K_l \geq 2 \text{ for some } l \leq L) \geq 1 - \epsilon_4.
\]
Now let \(|Y|\) denote the number of infected nodes in the observation \(Y\). Define events \(E_1 = \{Z_L = 0\}\) and \(E_2 = \{K_l \geq 2 \text{ for some } l \leq L\}\). We have
\[
\Pr(E_1||Y| = 1) + \Pr(E_2||Y| = 1)
= \frac{1}{\Pr(|Y| = 1)} \left( \Pr(E_1 \cap \{|Y| = 1\}) + \Pr(E_2 \cap \{|Y| = 1\}) \right).
\]
Since \(E_2\) implies that \(|Y| = 1\), we have
\[
\Pr(E_1||Y| = 1) + \Pr(E_2||Y| = 1)
= \frac{1}{\Pr(|Y| = 1)} \left( \Pr(E_1 \cap \{|Y| = 1\}) + \Pr(E_2) \right)
= \frac{1}{\Pr(|Y| = 1)} (\Pr(E_1) - \Pr(E_1 \cap \{|Y| = 0\}) + \Pr(E_2))
\geq \frac{1}{\Pr(|Y| = 1)} (\Pr(E_1) - \Pr(|Y| = 0) + \Pr(E_2))
\geq \frac{1}{\Pr(|Y| = 1)} (\Pr(|Y| = 1) - \epsilon_4)
= 1 - \frac{\epsilon_4}{\Pr(|Y| = 1)}.
\]
Note that \(\Pr(|Y| = 1)\) is a positive constant since the \(B^1\) branching process starting from the information source survives with non-zero probability. The theorem holds by choosing \(\epsilon_4 = \epsilon\Pr(|Y| = 1)\).

**Lemma 6.** Consider \(n_0\) i.i.d GW branching processes with a binomial offspring distribution with parameters \(y\) and \(q\) such that \(yq > 1\). Denote by \(K\) the number of branching processes that survive. Given any \(\epsilon > 0\), if
\[
n_0 \geq \frac{8 \log 2}{1 - \rho} \geq \epsilon
\]
then
\[
\Pr(K \geq 2) \geq 1 - \epsilon,
\]
where \(\rho\) is the extinction probability of the GW branching process. In the binomial case, \(\rho\) is the smallest non-negative root of equation \(\rho = (1 - q + q\rho)^y\).

**Proof:** The extinction probability of a GW branching process is denoted by \(\rho\), which is the smallest non-negative root of equation \(\rho = G(\rho)\) according to [18], where \(G(\rho)\) is the moment generating function of offspring distribution. In the binomial case we have \(G(\rho) = (1 - q + q\rho)^y\). \(\rho < 1\) when \(yq > 1\).

We define a Bernoulli random variable \(H_i\), for the \(i^{th}\) branching process such that
\[
H_i = \begin{cases} 
1, & \text{if the } i^{th} \text{ branching process survives;}
0, & \text{otherwise.}
\end{cases}
\]
So \(K = \sum_{i=1}^{n_0} H_i\), and
\[
E[K] = n_0(1 - \rho).
\]
According to the Chernoff bound [19], we have
\[
\Pr(K \leq (1 - \delta)(1 - \rho)n_0) < e^{-\frac{(1-\rho)n_0\delta^2}{2}}.
\]
Choose \(\delta = 0.5\). The Lemma holds if
\[
(1 - \rho)n_0/2 \geq 2,
\]
and
\[
(1 - \rho)n_0/8 \geq \log 1/\epsilon.
\]

\[\square\]
Lemma 7. Given any $\epsilon > 0$, there exists a constant $L'$ such that for any $L \geq L'$,

$$
\Pr\left( \bigcap_{i=1}^{L} \{0 < Z_t^i \leq n_0\} \right) \leq \epsilon.
$$

Proof: Define $p_r$ to be the probability that a node infects at least one of its children if it is in the infection state for $\tau$ time slots. We have

$$
p_r = \sum_{\tau-1} p(1 - p)^{\tau-1} p(1 - q)^{\tau},
$$

and

$$
\Pr(0 < Z_t^i \leq n_0) \leq \Pr(Z_t^i > 0) \leq (1 - (1 - p_r)^n). 
$$

which implies that

$$
\Pr\left( \bigcap_{i=1}^{L} 0 < Z_t^i \leq n_0 \right) = \Pr(0 < Z^L_t \leq n_0) \leq \Pr(0 < Z^L_{t-1} \leq n_0) \cdots \leq \Pr(0 < Z^L_1 \leq n_0) \Pr(0 < Z^L_1 \leq n_0) \leq (1 - (1 - p_r)^n)^L.
$$

The lemma holds by choosing

$$
L' = \left\lceil \frac{\log \epsilon}{\log (1 - (1 - p_r)^n)} \right\rceil.
$$

Lemma 8. Given any $\epsilon > 0$, there exist $\tau'$ and $L'$ such that for any $\tau > \tau'$ and $L > L'$

$$
\Pr(Z_L = 0) - \Pr(Z^\tau_L = 0) \geq -\epsilon.
$$

Proof: Note the difference can be re-written as

$$
\Pr(Z_L = 0) - \Pr(Z^\tau_L = 0) = (\Pr(Z_L = 0) - \Pr(Z^\infty_L = 0)) + (\Pr(Z^\infty_L = 0) - \Pr(Z^\tau_L = 0)).
$$

Step 1 Since $\{Z^\infty_L = 0\} \subseteq \{Z^\tau_L = 0\}$,

$$
\Pr(Z^\infty_L = 0) - \Pr(Z^\tau_L = 0) \geq 0.
$$

Step 2 We know

$$
\lim_{L \to \infty} \Pr(Z_L = 0) = \Pr(Z^\infty_L = 0).
$$

Then for $\epsilon/2 > 0$, there exists $L'$ such that for any $L \geq L'$,

$$
\left| \Pr(Z_L = 0) - \Pr(Z^\infty_L = 0) \right| \leq \epsilon/2,
$$

which implies that

$$
\Pr(Z_L = 0) - \Pr(Z^\infty_L = 0) \geq -\epsilon/2.
$$

Step 3 In this step, we will show

$$
\lim_{\tau \to \infty} \Pr(Z^\tau_L = 0) = \Pr(Z^\infty_L = 0).
$$

Define the generating functions of the offspring distributions of $B^\tau$ and $B^\infty$ to be $G_r(s)$ and $G(s)$, respectively. We know that $G_r(s) - s$ and $G(s) - s$ are convex functions when $s \in [0, 1]$. Let $\rho = \Pr(Z^\infty_L = 0)$, i.e., the extinction probability, we know that $\rho$ is the smallest nonnegative root of $G(\rho) = \rho$ and $\rho < 1$.

Similarly, define $\rho'_r = \Pr(Z^\tau_L = 0)$, and $\rho' = \lim_{\tau \to \infty} \rho'_r$. Taking limit on both sides of $G_r(\rho') = \rho$, we have

$$
G(\rho') = \rho.
$$

Note that

$$
\rho \leq \rho_s \leq \rho_1 < 1
$$

for any $\tau$, so

$$
\rho \leq \rho' \leq \rho_1 < 1.
$$

Since $G(s) - s = 0$ has at most two solutions in $[0, 1]$ and $s = 1$ is one of them, we conclude $\rho' = \rho$. Therefore, for given $\epsilon/2 > 0$, there exists $\tau \geq \tau'$ such that

$$
\Pr(Z^\infty_L = 0) - \Pr(Z^\tau_L = 0) \geq -\epsilon/2.
$$

Hence, the lemma holds.

VI. SIMULATIONS

In this section, we evaluate the performance of the reverse infection algorithm on different networks, including different tree networks and some real world networks.

A. Tree Networks

In this section, we evaluate the performance of the reverse infection algorithm on tree networks. We compare the reverse infection algorithm with the closeness centrality heuristic, which selects the node with the maximum infection closeness as the information source. Note that the node with the maximum closeness is the maximum likelihood estimator of the information source on regular trees under the SI model.

1) Small-size tree networks: We first studied the performance on small-size trees. The infection probability $q$ was chosen uniformly from $(0, 1)$ and the recovery probability $p$ was chosen uniformly from $(0, q)$. The infection process propagates $t$ time slots where $t$ was uniformly chosen from $[3, 5]$. To keep the size of infection topology small, we restricted the total number of infected and recovered nodes to be no more than 100. For small-size trees, we first calculated the MLE using dynamic programming for fixed $t$ and then searching over $t \in [0, t_{\text{max}}]$ for a large value of $t_{\text{max}}$ to find the optimal estimator.

The detection rate is defined to be the fraction of experiments in which the estimator coincides with the actual source. We varied $q$ from 2 to 10 and the results are shown in Figure...
We can see that the detection rate of the reverse infection algorithm is almost the same as that of the MLE, and is higher than that of the closeness centrality heuristic by approximately 20% when the degree is small and by 10% when the degree is large.

2) General $g$-regular tree networks: We further conducted our simulations on large-size $g$-regular trees. The infection probability $q$ was chosen uniformly from $(0, 1)$ and the recovery probability $p$ was chosen uniformly from $(0, q)$. The infection process propagates $t$ time slots where $t$ was uniformly chosen from $[3, 20]$. We selected the networks in which the total number of infected and recovered nodes is no more than 500.

We varied $g$ from 2 to 10. Figure 7 shows the detection rate as a function of $g$. We can see the detection rates of both the reverse infection and closeness centrality algorithms increase as the degree increases and is higher than 60% when $g > 6$. However, he detection rate of the reverse infection algorithm is higher than that of the closeness centrality algorithm, and the average difference is 8.86%.

3) Binomial random trees: In addition, we evaluated the performance on binomial random trees where the number of children of each node follows a binomial distribution with number of trials $g'$ and success probability $\beta$. We fixed $g' = 10$ and varied $\beta$ from 0.1 to 0.9. The durations of the infection process and the observed infected networks were selected according to the same rules for the $g$-regular tree case. The results are shown in Figure 8. Similar to the regular tree case, as $\beta$ increases, the tree is more denser which increases the number of survived branching processes and the detection rate. The reverse infection algorithm outperforms the closeness centrality algorithm by 10.16% on average.

B. Real World Networks

We next conducted experiments on three real world networks — the Internet Autonomous Systems network (IAS), the Wikipedia who-votes-on-whom network (Wikipeida), and the power grid network (PG). We compare the reverse infection algorithm with random guessing, which randomly selects a node and declares it as the information source. In these networks, the infection probability $q$ was chosen uniformly from $(0, 0.05)$ and the recovery probability $p$ was chosen uniformly from $(0, q)$. Here we chose small infection probabilities since the network was of finite size so the infection process should be controlled to make sure that not all nodes were infected when the network was observed. The duration $t$ was an integer uniformly chosen from $[3, 200]$. We selected the networks in

3Available at http://snap.stanford.edu/data/index.html
4Available at http://www-personal.umich.edu/~mejn/netdata/
which the total number of infected and recovered nodes was in the range of [50, 500].

1) The Internet autonomous systems network: Figure 9 shows the results on the Internet autonomous systems network. An Internet autonomous system is a collection of connected routers who use a common routing policy. The Internet autonomous system network is obtained based on the recorded communication between the Internet autonomous systems inferred from Oregon route-views on March, 31st, 2001. The network consists of 10,670 nodes and 22,002 edges. According to Figure 9, more than 80% of the estimators identified by the reverse infection algorithm are no more than two hops away from the actual sources, comparing to 10% under the random guessing.

2) The Wikipedia who-votes-on-whom network: Figure 10 shows results on the Wikipedia who-votes-on-whom network, in which two nodes are connected if one user voted on the other in the administrator promotion elections. The network has 100,736 links and 7,066 nodes. We have similar observations as for the Internet autonomous systems network: the majority of the estimators produced by the reverse infection algorithm are no more than two hops away from the actual sources; and only less than 20% of the estimators of random guessing are within two hops from the actual sources.

3) The power grid network: Figure 11 shows the results on the power grid, which has 4,941 nodes and 6,594 edges. As we can see, the reverse infection algorithm performs better than the random guessing. The peak of the reverse infection algorithm appears at the third hop versus the seventeenth hop under random guessing.

VII. Conclusion

In this paper, we developed a sample path based approach to find the information source under the SIR model. We proved that the sample path based estimator is a node with the minimum infection eccentricity. Based on that, a reverse infection algorithm has been proposed. We analyzed the performance of the reverse infection algorithm on regular trees, and showed that with a high probability the distance between the estimator and actual source is a constant, independent of the number of infected nodes and the time the network was observed. We evaluated the performance of the proposed reverse infection algorithm on several different network topologies.

APPENDIX A

NOTATION TABLE

| Symbol | Description |
|--------|-------------|
| $q$    | the probability an infected node infects its neighbors |
| $p$    | the probability an infected node recovers |
| $v^*$  | the actual information source |
| $e(v)$ | the infection eccentricity of node $v$ |
| $d(v, u)$ | the length of shortest path between node $v$ and node $u$ |
| $C(v)$ | the set of children of node $v$ |
| $e(v)$ | the infection eccentricity of node $v$ |
| $t^v_u$ | the time duration associated of the optimal sample path in which node $v$ is the information source |
| $t^v_u$ | the infection time of node $v$ |
| $t^v_u$ | the recovery time of node $v$ |
| $T^{-v}_u$ | the snapshot of all nodes |
| $T^{-v}_u$ | the tree rooted at node $u$ but without the branch from $v$. |
| $X_v(t)$ | the state of node $v$ at time $t$ |
| $X(t)$ | the states of all nodes at time $t$ |
| $X[0, t]$ | the sample path from 0 to $t$ |
| $X[(0, t), T^{-v}_u]$ | the sample path from time slot 0 to $t$ restricted to $T^{-v}_u$ |
| $X(t)$ | the set of all valid sample path from time slot 0 to $t$ |
| $X(t, T^{-v}_u)$ | the set of all valid sample path from time slot 0 to $t$ restricted to $T^{-v}_u$ |
| $I$    | the set of the infected nodes |

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