Identifying Spatial Invasion of Pandemics on Metapopulation Networks Via Anatomizing Arrival History

Jian-Bo Wang, Student Member, IEEE, Lin Wang, Member, IEEE, and Xiang Li, Senior Member, IEEE

Abstract—Spatial spread of infectious diseases among populations via the mobility of humans is highly stochastic and heterogeneous. Accurate forecast/mining of the spread process is often hard to be achieved by using statistical or mechanical models. Here we propose a new reverse problem, which aims to identify the stochastically spatial spread process itself from observable information regarding the arrival history of infectious cases in each subpopulation. We solved the problem by developing an efficient optimization algorithm based on dynamical programming, which comprises three procedures: 1) anatomizing the whole spread process among all subpopulations into disjoint componental patches; 2) inferring the most probable invasion pathways underlying each patch via maximum likelihood estimation; and 3) recovering the whole process by assembling the invasion pathways in each patch iteratively, without burdens in parameter calibrations and computer simulations. Based on the entropy theory, we introduced an identifiability measure to assess the difficulty level that an invasion pathway can be identified. Results on both artificial and empirical metapopulation networks show the robust performance in identifying actual invasion pathways driving pandemic spread.

Index Terms—Identifiability, infectious diseases, metapopulation, networks, process identification, spatial spread.

I. INTRODUCTION

THE FREQUENT outbreaks of emerging infectious diseases in recent decades lead to great social, economic, and public health burdens [1]–[3]. This trend is partially due to the urbanization process and, in particular, the establishment of long-distance traffic networks, which facilitate the dissemination of pathogens accompanied with passengers [4], [5]. Real-world examples include the transnational spread of severe acute respiratory syndrome (SARS) coronaviruses in 2003 [6], the global outbreak of A (H1N1) pandemic flu in 2009 [7], [8], avian influenza in Southeast Asia [9], [10], the spark of Ebola infections in western countries in 2014 [11], and recent potential outbreak of middle east respiratory syndrome [12].

During almost the same epoch, the theory of complex networks has been developed as a valuable tool for modeling the structure and dynamics of/on complex systems [13]–[16]. In the study of network epidemiology, networks are often used to describe the epidemic spreading from human to human via contacts, where nodes represent persons and edges represent interpersonal contacts [17]–[22]. To characterize the spatial spread between different geo-locations, simple network models are generalized with metapopulation framework, in which each node represents a population of individuals that reside at the same geo-region (e.g., a city), and the edge describes the traffic route that drives the individual mobility between populations [18], [19]. The networked metapopulation models have been applied to study the real-world cases such as SARS [6], A (H1N1) pandemic flu [23], and Ebola [11], which can capture some key dynamic features including peak times, basic epidemic curves, and epidemic sizes. Quantitative model results can be used to evaluate the effectiveness of control strategies [24]–[28], such as optimizing the vaccine allocation.

The numerical computing of large-scale metapopulation models is time-consuming, because of the requirement of high-level computer power. The model calibrations need high-resolution data for incidence cases, which may not be available or accurate during the early weeks of initial outbreaks [4]. Hence, continuous model training with data collected in real-time is essential in achieving a reliable model prediction [29]. Generally, model results are the ensemble average over numerous simulation realizations, which aims to predict the mean and variance of epidemic curves, while in reality there is no such thing described by the average over different realizations [30]. To extract more meaningful information from epidemic data generated by surveillance systems, recent studies (particularly in engineering fields) start paying attention to reverse problems, such as source detection and network reconstruction, which are briefly summarized here.

© IEEE 2016. This article is free to access and download, along with rights for full text and data mining, re-use and analysis.
A. Related Works

The theory of system identification has been established in engineering fields, usually used to infer system parameters. The use of system identification in epidemiology mainly focuses on inferring epidemic parameters, such as the transmission rate and generation time [31], which relies on constructing dynamical systems of ordinary differential equations. The methodology of system identification is not helpful in solving high-dimensional stochastic many-body systems, such as metapopulation models.

Source detection for rumor spreading on complex networks is becoming a popular topic, attracting extensive discussions in recent years. The target is to figure out the causality that can trigger the explosive dissemination across social networks, such as Facebook, Twitter, and Weibo. For example, using maximum likelihood (ML) estimators, Shah and Zaman [32] proposed the concept of rumor centrality that quantifies the role of nodes in network spreading. Luo et al. [33] designed new estimators to infer infection sources and regions in large networks. Wang et al. [34], [36] and Dong et al. [35] extended the scope by using multiple observations, which largely improves the detection accuracy. Another interesting topic is the network inference, which engages in revealing the topology structure of a network from the hint underlying the dynamics on a network [37]. Some useful algorithms (e.g., NetInf) have been proposed in [38]–[42]. Note that the algorithms for source detection and network inference are not feasible in identifying the spreading processes on metapopulation networks.

Using metapopulation networks models, some heuristic measures have been proposed to understand the spatial spread of infectious diseases, which are most related to this paper. Gautreau et al. [30] developed an approximation for the mean first arrival time between populations that have direct connection, which can be used to construct the shortest path tree (SPT) that characterizes the average transmission pathways among populations. Brockmann and Helbing [4] proposed a measure called “effective distance,” which can also be used to build the SPT. Using a different method based on the ML, Balcan et al. [43] generated the transmission pathways by extracting the minimum spanning tree from extensive Monte Carlo simulation results. Details about these measures will be given in Section IV, which compares the algorithmic performance.

B. Motivation

Current algorithms to inferring pandemic spatial spread generally make use of the topology features of metapopulation networks or extensive epidemic simulations. The resulting outcome is an ensemble average over all possible transmission pathways, which may fail in capturing those indeed transmitting the disease between populations, because of the high-level stochasticity and heterogeneity in the spreading process.

Good news comes from the development of modern sentinel and Internet-based surveillance systems, which becomes increasingly popular in guiding public health control strategies. Such systems can or will provide high-resolution, location-specific data on human and poultry cases [44]. Human mobility data are also available from mass transportation systems or GPS-based mobile Apps [3]. Integrating these data often used in different fields, a natural reverse problem poses itself, which is the central interest of this paper: is it probable to design an efficient algorithm to identify or retrospect the stochastic pandemic spatial spread process among populations by linking epidemic data and models.

C. Our Contributions

Main contributions of this paper are as follows.

1) A novel reverse problem of identifying the stochastic pandemic spatial spread process on metapopulation networks is proposed, which cannot be solved by existing techniques.

2) An efficient algorithm based on dynamical programming is proposed to solve the problem, which comprises three procedures. First, the whole spread process among all populations will be decomposed into disjoint component patches, which can be categorized into four types of invasion cases (INCs). Then, since two types of INCs contain hidden pathways, an optimization approach based on the ML estimation is developed to infer the most probable invasion pathways underlying each path. Finally, the whole spread process will be recovered by assembling the invasion pathways of each patch chronologically, without burdens in parameter calibrations and computer simulations.

3) An entropy-based measure called identifiability is introduced to depict the difficulty level an INC can be identified. Comparisons on both artificial and empirical networks show that our algorithm outperforms the existing methods in accuracy and robustness.

The remaining sections are organized as follows. Section II provides the preliminary definitions and problem formulation. Section III describes the procedures of our identification algorithm, and introduces the identifiability measure. Section IV performs computer experiments to compare the performance of algorithms. Section V gives the conclusion.

II. Preliminary and Problem Formulation

This section first elucidates the structure of networked metapopulation model, and then provides the preliminary definitions and problem formulation.

A. Networked Metapopulation Model

In the networked metapopulation model, individuals are organized into social units such as counties and cities, defined as subpopulations, which are interconnected by traffic networks of transportation routes. The disease prevails in each subpopulation due to interpersonal contacts, and spreads between subpopulations via the mobility of infected persons. Fig. 1 illustrates the model structure.

Within each subpopulation, individuals mix homogeneously. This assumption is partially supported by recent empirical findings on intraurban human mobility patterns [19], [45]–[48]. The intrapopulation epidemic
dynamics are characterized by compartment models. Considering the wide applications in describing the spread of pathogens, species, rumors, emotion, behavior, crisis, etc. [32], [33], [35], [49], we used the susceptible-infected (SI) model in this paper. Define \( N_i \) as the population size of each subpopulation \( i \), \( I_i(t) \) the number of infected cases in subpopulation \( i \) at time \( t \), \( \beta \) the transmission rate that an infected host infects a susceptible individual shared the same location in unit time. As such, the risk of infection within subpopulation \( i \) at time \( t \) is characterized by \( \lambda_i(t) = \beta I_i(t)/N_i \). Per unit time, the number of individuals newly infected in subpopulation \( i \) can be calculated from a binomial distribution with probability \( \lambda_i(t) \) and trails equalling the number of susceptible persons \( S_i(t) \).

The mobility of individuals among subpopulations is conceptually described by diffusion dynamics, \( \partial_t X_i = \sum_{j \in v(i)} p_{ij} X_j(t) - p_{ij} X_i(t) \), where \( X_i(t) \) is a placeholder for \( S_i(t) \) or \( I_i(t) \), \( v(i) \) is the set of subpopulations directly connected with subpopulation \( i \), and \( p_{ij} \) is the per capita mobility rate from subpopulation \( i \) to \( j \), which equals the ratio between the daily flux of passengers from subpopulation \( i \) to \( j \) and the population size of departure subpopulation \( i \). The ensemble of mobility rates \( 0 \leq p_{ij} < 1 \) defines a transition matrix \( P \), determined by the topology structure and traffic fluxes of the mobility network. The interpolation mobility of individuals is simulated with binomial or multinomial process (Appendix A). For more details in modeling rules, refer to [19].

B. Basic Definitions

The epidemic arrival time (EAT) is the first arrival time of infectious hosts traveling to a susceptible subpopulation. At a given EAT, at least an unaffected (susceptible) subpopulation will be contaminated, characterizing the occurrence of invasion events. Herein, \( S \) (I) denotes a (an) susceptible (infected) subpopulation.

For an invasion event, organizing newly contaminated subpopulations (remaining unaffected prior to that invasion event) into set \( S \), and infected subpopulations into set \( I \), we define the four types of INC as follows.

1) \( I \mapsto S \): \( I \) and \( S \) both are composed of a single subpopulation respectively, which represents that a previously unaffected subpopulation is infected by the new arrival of infectious host(s) from its unique neighboring infected subpopulation.

2) \( I \mapsto nS(n > 1) \): In this case, \( I \) only consists of a single subpopulation, while \( S \) contains \( n(n > 1) \) subpopulations. This represents that \( n \) previously unaffected subpopulations are contaminated due to the new arrival of infectious hosts from their common infected subpopulation in \( I \).

3) \( mI \mapsto S(m > 1) \): \( S \) only consists of a single subpopulation, and \( I \) contains \( m(m > 1) \) subpopulations. This means that the newly infected subpopulation in \( S \) is infected by the arrival of infected host(s) from \( m \) potential upstream subpopulations in \( I \) through the invasion edges.

4) \( mI \mapsto nS(m,n > 1) \): In this case, \( S \) and \( I \) both are composed of no less than two subpopulations, and they constitute a connected subgraph. Each previously unaffected subpopulation in \( S \) is contaminated due to the simultaneous arrival of infected hosts from \( m \) potential source subpopulations in \( I \). Each subpopulation in \( I \) may lead to the contamination of at least one but no more than \( n \) neighboring downstream subpopulations in \( S \) through the invasion edges. Multiple edges between any pair of subpopulations are forbidden.

Fig. 2(a) and (b) illustrates the two scenarios of \( mI \mapsto S(m > 1) \) and \( mI \mapsto nS(m,n > 1) \). A decomposition procedure of invasion partition (INP) is used to generate the components of INCs in each invasion event. The heuristic search algorithm to proceed the INP is given in Algorithm 1 if an invasion event occurs.

C. Problem Formulation

Suppose that the spread starts at an infected subpopulation. It forms the invasion pathways when this source invades many susceptible subpopulations and the cascading invasion goes on. We record the infected individuals of each subpopulation per unit time. From the data, we should know when a subpopulation is infected and how many infected individuals...
in this subpopulation, but we may not know which infected subpopulations invade this subpopulation if it has \( (m, m \geq 2) \) infected neighbor subpopulations through the corresponding edge(s) [see Fig. 2(a)] at that time step. The question of interest is how to identify the instantaneous spatial invasion process just according to the surveillance data. Herein, we know the network topology including subpopulation size and travel flows, such as the city populations of airports and travelers by an airline of the real network of American airports network (ANN).

Define an invasion pathway which are the directed edges that infected individuals invade to susceptible subpopulations at EAT. To identify it, we proceed the following invasion pathways identification (IPI) algorithm.

1) Decompose the whole pathways as four types of INCs by the IPI at each EAT; suppose the whole invasion pathways \( T \) are anatomized into \( \Lambda \) of four INCs. Let \( \hat{a}_i \) denote the identified invasion pathways based on the surveillance data \( G \) of that INC \( i \) and the given graph \( G \). According to the (stochastic) dynamic programming, we have the following equation to optimally solve this problem:

\[
T_{\text{whole invasion pathways}} = \text{opt} \sum_{i=1}^{\Lambda} \hat{a}_i. \tag{1}
\]

2) For each INC, we first judge whether it has a unique set of invasion pathways or more than one potential invasion pathways. When an INC has more than one possible invasion pathway, each set of which is called potential invasion pathway. If it has more than one potential invasion pathway, we estimate the true invasion pathways \( a^*_i \), denoted by \( \hat{a}_i \), based on the surveillance data \( G \) of that INC and the given graph \( G \). A potential pathway belonged to that INC is denoted by \( \forall a_i \in G_{\text{INC}} \). To make this estimation, we shall compute the likelihood of a potential invasion pathway \( a_i \). With respect to this setting, the ML estimator of \( a^*_i \) with respect to the networked metapopulation model given by that INC maximizes the correct identification probability. Therefore, we define the ML estimator

\[
\hat{a}_i = \text{arg max}_{a_i \in G_{\text{INC}}} P(a_i | G_{\text{INC}}) \tag{2}
\]

where \( P(a_i | G_{\text{INC}}) \) is the likelihood of observing the potential pathway \( a_i \) assuming it is the true pathway \( a^*_i \).

Thus, we would like to evaluate \( P(a_i | G_{\text{INC}}) \) for all \( a_i \in G_{\text{INC}} \), and then choose the maximal one.

III. IDENTIFICATION ALGORITHM TO INVASION PATHWAY

According to our above IPI decompose algorithm, it is easy to identify the invasion pathways for the INC scenario \( I \mapsto nS(n \geq 1) \) (they have the only invasion pathway from their neighbor infected subpopulation). Thus our invasion pathways’ identification algorithm mainly deals with the other two kinds of INCs \( ml \mapsto S(m > 1) \) and \( ml \mapsto nS(m, n > 1) \). To make the description clear, we restate the term \( I_i \) denotes subpopulation \( i \) which is infected, and its number of infected individuals of \( I_i \) at time \( t \) is denoted by \( I_i(t) \).

As time evolves, infected hosts travel among subpopulations, inducing the spatial pandemic dispersal. For each INC, by analyzing the variance of infected hosts in each subpopulation \( i \), we define three levels of extent of subpopulations observability to reflect the information held for the inference of relevant invasion pathway.

1) Observable Subpopulation: Subpopulation \( i \) is observable during an INC, given the occurrence of the three most evident (subpopulation’s) status transitions. The first refers to the transition \( S_i \mapsto I_i \), accounting that the previously unaffected subpopulation \( i \) is contaminated during that INC due to the arrival of infected hosts. The second concerns the transition \( I_i \mapsto S_i \), in which the previously infected subpopulation \( i \) becomes susceptible again during that INC, since the infected hosts do not trigger a local outbreak and leave \( i \). In the third transition \( S_i \mapsto S_i \), despite of having infected subpopulations in the neighborhood, subpopulation \( i \) remains unaffected during that INC due to no arrival of infected hosts. Fig. 3(a) illustrates such observable transitions.

2) Partially Observable Subpopulation: Subpopulation \( i \) is partially observable during an INC occurring at time \( t \), if its number of infected hosts is decreased, i.e., \( I_i(t) < I_i(t-1) \) and \( I_i(t) > 0 \), which implies that at least \( \Delta I_i(t) = |I_i(t) - I_i(t-1)| \) infected hosts leave \( i \) during that INC. It is impossible to distinguish their mobility destinations unless the INC \( I \mapsto S \) or \( I \mapsto nS \) occurs. Fig. 3(b) illustrates the partially observable subpopulation.
infected subpopulation in set \( \mathcal{I} \). Suppose that subpopulation \( S_j \) is contaminated at time \( t \) due to the appearance of \( \mathcal{H} \) infected hosts (\( \mathcal{H} \) is a positive integer number) that come from the potential sources in \( \mathcal{I} \). If the actual number of infected hosts from subpopulation \( I_i \) is \( \mathcal{H}_{i1} \), we have

\[
\sum_{i=1}^{m} \mathcal{H}_{i1} = \mathcal{H} \tag{3}
\]

with the conditions \( 0 \leq \mathcal{H}_{i1} \leq \mathcal{H} \) and \( \mathcal{H}_{i1} \leq I_i(t-1) \).

1) **Accurate Identification of Invasion Pathway:** Given a few satisfied prerequisites, (3) can have a unique solution, which implies that the invasion pathways of that INC can be identified accurately. Theorem 1 elucidates this scenario.

**Theorem 1 (Accurate Identification of Invasion Pathway):** With the following conditions: 1) among \( m \) possible sources illustrated in set \( \mathcal{I} \), there are only \( m'(m' \leq m) \) partially observable subpopulations \( \mathcal{I}' \), whose neighboring subpopulations (excluding the invasion destination \( S_j \)) only experience the transition \( S \rightarrow S \) or \( S \rightarrow \mathcal{I} \) at that EAT and 2) \( \sum_{i \in \mathcal{I}} [I_i(t-1) - I_i(t)] = \mathcal{H} \), the invasion pathway of an INC \( mI \rightarrow S(m > 1) \) can be identified exactly.

**Proof:** According to the definition of observability, in the INC, the number of local infected hosts in an involved partially observable source \( i \) will be decreased by \( [I_i(t-1) - I_i(t)] \) due to their departure. If the subpopulations in the neighborhood of \( i \) only experience the transition of \( S_j \) to \( S_i \) or \( I_j \) to \( S_i \) from \( t_{EAT-1} \) to \( t_{EAT} \), they are impossible to receive the infected hosts from subpopulation \( i \). Therefore, the newly contaminated subpopulation \( S_j \) is the only destination for those infected travelers departing from the partially observable sources. Since \( m' \leq m \), the second condition guarantees that (3) only has a unique solution, which corresponds to the accurate identification of invasion pathways of this INC.

2) **Potential Invasion Pathway:** If the conditions of Theorem 1 are unsatisfied, (3) has multiple solutions, each solution corresponds to a set of potential invasion pathways that can result in the related INC. Due to the heterogeneity in the traffic flow on each edge and the number of infected hosts within each contaminated source, each set of potential pathways is associated with a unique likelihood, which also identifies the occurrence probability of the corresponding solution of (3). Therefore, the identification of invasion pathway that induce an INC can be transformed to searching the most probable solution of (3).

We define the solution space \( \Phi \) of (3) of the INC \( mI \rightarrow S(m > 1) \), which subjects to two conditions: 1) \( \sum_{i=1}^{m} \mathcal{H}_{i1} = \mathcal{H} \) and 2) \( \forall \mathcal{I}_{i1}, \mathcal{H}_{i1} \leq I_i(t-1) \). The second condition is obvious, since the number of infected travelers departing from the source \( I_i \) cannot exceed \( I_i(t-1) \). Let us assume that \( \Phi \) contains \( M \) solutions, and a typical solution is formulated as \( \sigma_j = [\mathcal{H}_{i1}^{(j)}, i = [1, \ldots, m]] \). Obviously, each solution \( \sigma_j \) corresponds to a potential invasion pathway \( a_j \).

Through the INC \( mI \rightarrow S(m > 1) \), the observed event \( E \) shows that the destination \( S_j \) is contaminated due to the arrival of totally \( \mathcal{H} \) infected hosts from the potential sources \( I_i, i = [1, \ldots, m] \). With this posterior information,

\[
\sigma_j = [\mathcal{H}_{i1}^{(j)}, i = [1, \ldots, m]]
\]

This scenario illustrates that the accurate identification of invasion pathway is equivalent to finding the posterior probability of each invasion pathway. Therefore, the posterior probability \( P(E|\sigma_j) \) of the invasion pathway \( \sigma_j \) can be accurately identified by the following formula:

\[
P(E|\sigma_j) = \frac{P(\sigma_j|E)P(E)}{P(E)}
\]

where \( P(\sigma_j|E) \) is the occurrence probability of the corresponding solution, which corresponds to the accurate identification of invasion pathways of this INC.

**Fig. 3.** Illustration of neighbors classification in terms of status transitions: (a) Observable \( i \). (b) Partially observable \( i \). (c) Unobservable \( i \).
we first measure the likelihood of each possible solution \( \sigma_i \), which corresponds to the reasoning event that for each source \( I_i \), \( i \in \{1, m\} \), \( H_{t_{1}} \) infected hosts are transferred to \( S_1 \). It is evident that \( \forall j, \ P(E_{\text{mIS}}|\sigma_j) = 1 \), since \( \sigma_j \) will lead to the occurrence of event \( E_{\text{mIS}} \), which corresponds to \( G_{\text{mIS}} \).

According to Bayes’ theorem, the likelihood of the solution \( \sigma_j \) is characterized by

\[
P(\sigma_j|E_{\text{mIS}}) = \frac{P(E_{\text{mIS}}|\sigma_j)P(\sigma_j)}{P(E_{\text{mIS}})} = \frac{P(E_{\text{mIS}}|\sigma_j)P(\sigma_j)}{\sum_{j=1}^{M} P(E_{\text{mIS}}|\sigma_j)P(\sigma_j)}
\]

where \( M \) represents the number of potential solution \( \sigma_j \), and the last item \( \Omega(\mathcal{H}_{k_1}^{(i)}) \) represents the mobility likelihood transferring estimator of infected subpopulation \( I_i \) in \( \mathbb{I} \).

One linchpin of our algorithm in handling the scenario \( mI \rightarrow S(m > 1) \) is to estimate the probability of transferring \( H_{t_{1}} \) infected hosts from each infected subpopulation \( I_i \), \( i \in \mathbb{I} \), to the destination subpopulation \( S_1 \). Based on the independence between the intrasubpopulation epidemic reactions and the intersubpopulation personal diffusion, we introduce a transferring estimator to analyze the individual mobility of each source \( I_i \), which is in particular useful if there are partially observable and unobservable edges emanated from the focal infected subpopulation.

The specific formalisms of the transferring estimator are defined according to the three types of infected subpopulation \( I_i \) consisted of set \( \mathbb{I} \) which are unobservable subpopulation, partially observable subpopulation, and observable subpopulation with transition of \( I \) to \( S \).

a) Unobservable subpopulation \( I_i \): Due to the occurrence of \( \mathcal{H}_{t_{1}} \rightarrow S \), among all \( k_i \) edges emanated from subpopulation \( I_i \), there is only one invasion edge in that INC, labeled as \( k_i \), along which the traveling rate is \( p_{k_i} \) and \( H_{t_{1}} \) infected hosts are transferred to the destination \( S_1 \). Assume that there are \( \ell_i \) (1 ≤ \( \ell_i \) < \( k_i \)) unobservable and partially observable edges, labeled as 1, 2, ..., \( \ell_i \), respectively. Along each unobservable or partially observable edge, the traveling rate is \( p_{\ell_i} \), \( \ell \in \{1, \ell_i\} \), and \( x_{\ell} \) infected hosts leave \( I_i \). Accordingly, in total \( \eta_i = \sum_{\ell} x_{\ell} \) infected hosts leave \( I_i \) through the unobservable and partially observable edges. There remain \( k_i - \ell_i - 1 \) observable edges, labeled as \( \ell_i + 1, \ldots, k_i - 1 \), respectively. Along each observable edge, the traveling rate is \( p_{\ell_s} \), \( s \in \{\ell_i + 1, k_i - 1\} \), and \( x_{\ell_s} \) infected hosts leave \( I_i \). With probability \( \overline{p_{\ell_s}} = 1 - p_{\ell_s} - \sum_{\ell \neq s} p_{\ell} - \sum_{s \neq i} p_{\ell_s} \), an infected host keeps staying at source \( I_i \).

Since the infected hosts transferred by unobservable and partially observable edges are untraceable, it is unable to reveal the actual invasion pathways resulting in that INC accurately. Fortunately, the message of traveling rates on each edge is available by collecting and analyzing the human mobility transportation networks. Therefore, the mobility multinomial distribution \( \{31\} \) in Appendix A can be used to obtain the conditional probability that \( H_{t_{1}} \) infected hosts are transferred from infected source \( I_i \) to destination \( S_1 \), which is measured by the following transferring estimator:

\[
\Omega_{\text{a}}(\mathcal{H}_{t_{1}}) = P(H_{t_{1}}, p_{k_i}; I(t - 1); x_{\ell}, p_{\ell}, \ell = [1, \ldots, \ell_i]; x_{k_8}, p_{k_8}, \mathcal{R} = [\ell_i + 1, \ldots, k_i - 1]; \overline{x}_{\ell}, \overline{p_{\ell_i}})
\]

where \( \overline{x}_i \) accounts for the number of infected hosts that do not leave source \( I_i \) after the INC. Here, the observed number of infected persons in source \( I_i \) before the INC, i.e., \( I(t - 1) \), is used for the estimation, since the probability that a newly infected host also experiencing the mobility process is very low. Considering the conservation of infected hosts, and the implication of observable edges (i.e., \( x_{k_8} = 0, \forall \mathcal{R} \)), we have

\[
I(t - 1) = H_{t_{1}} + \sum_{\ell} x_{\ell} = H_{t_{1}} + \eta_i + \overline{x}_i
\]

Taking into account all scenarios that fulfill the condition \( \eta_i = \eta_i + \overline{x}_i = I(t - 1) - H_{t_{1}} \), the transferring estimator is simplified by the marginal distribution of (4), that is

\[
\sum_{\eta_i = I(t - 1) - H_{t_{1}}} P(\overline{x}_i, x_{\ell}, \ell = [1, \ldots, \ell_i])
\]

With independence, the transferring estimator becomes

\[
\Omega_{\text{a}} = \frac{I(t - 1)! \mathcal{H}_{t_{1}}!}{\eta_i \pi! \overline{p_{k_i}}} \left( \sum_{\ell} p_{\ell} + \overline{p_{\ell_i}} \right)^{\eta_i}.
\]

b) Observable subpopulation \( I_i \) (\( I_i \) to \( S_1 \)): If the infected hosts of source \( I_i \) all leave to travel from \( t_{E\text{AT}} - 1 \) to \( t_{E\text{AT}} \), the subpopulation \( I_i \) is observable at that INC. In this case, we have additional posterior messages, i.e., \( I(t) = 0 \) and \( \Delta I(t) = I(t - 1) \). Here, the number of infected hosts transferred to \( S_1 \) cannot exceed the total number of infected travelers departing from source \( I_i \), i.e., \( H_{t_{1}} \leq \Delta I(t) \). In this regard, the probability that \( H_{t_{1}} \) infected hosts arrive in destination \( S_1 \) is measured by the following transferring estimator:

\[
\left( \frac{\Delta I(t)}{\mathcal{H}_{t_{1}}} \right) \left[ \frac{\sum_{\ell} p_{\ell_i}}{p_{k_i}} \right]^{\mathcal{H}_{t_{1}}} \left[ \frac{1 - \sum_{\ell} p_{\ell_i} - \sum p_{k_i}}{\Delta I(t) - \mathcal{H}_{t_{1}}} \right].
\]

c) Partially observable subpopulation \( I_i \): If source \( I_i \) is partially observable, we can develop the inference algorithm with an additional posterior message, which reveals that at least \( \Delta I(t) = I(t) - I(t - 1) \geq 1 \) infected hosts leave the focal source \( I_i \) after the occurrence of that INC. In order to measure the conditional probability that \( H_{t_{1}} \) infected hosts are transferred from source \( I_i \) to destination \( S_1 \), we inspect all possible scenarios in detail, as follows.

Type 1: \( \Delta I(t) \leq \mathcal{H}_{t_{1}} \), i.e., the observed reduction in the number of infected hosts \( \Delta I(t) \) is less than those transferred from \( I_i \) to \( S_1 \). Here, we consider all cases that are in accordance with this condition.

If all \( \Delta I(t) \) confirmed infected travelers are transferred from \( I_i \) to \( S_1 \), the transferring estimator can be used to quantify the
conditional probability that the remaining $\mathcal{H}_{i1} - \Delta I_i(t)$ infected hosts concerned also visit $S_1$, that is

$$
\Omega_{p}(\mathcal{H}_{i1} - \Delta I_i(t)|I_i(t-1) - \Delta I_i(t)) = \frac{[I_i(t-1) - \Delta I_i(t)]!}{[\mathcal{H}_{i1} - \Delta I_i(t)]!}\left[\sum p_\ell + \bar{p}_i\right]^{\eta} \left[1 - \sum_{\ell} p_\ell + p_k \right]^{\Delta I_i(t)} \times \left[\sum p_\ell + \bar{p}_i\right]^{\Delta I_i(t)}
$$

where $p_{ki}/\left[\sum p_\ell + p_k \right]$ represents the relative traveling rate that any person from source $I_i$ is transferred to $S_1$, thus the last item on the right-hand side (RHS) accounts for the probability that $\Delta I_i(t)$ confirmed infected travelers all visit $S_1$.

If only a fraction of $\Delta I_i(t)$ confirmed infected travelers are transferred from $I_i$ to $S_1$, the situation is more complicated. Assume that $\Delta I_i(t) - \phi$ ($1 \leq \phi < \Delta I_i(t)$) confirmed travelers successfully come to $S_1$, the corresponding transferring estimator becomes

$$
\Omega_{p}(\mathcal{H}_{i1} - \Delta I_i(t) + \phi|I_i(t-1) - \Delta I_i(t)) = \left[\Delta I_i(t) - \phi\right] \left[\sum p_\ell + \bar{p}_i\right]^{\eta} \left[1 - \sum_{\ell} p_\ell + p_k \right]^{\phi} \frac{[I_i(t-1) - \Delta I_i(t)]!}{[\mathcal{H}_{i1} - \Delta I_i(t) + \phi]!}\left[\sum p_\ell + \bar{p}_i\right]^{\Delta I_i(t)}
$$

where the first item on the RHS accounts for the probability that $\Delta I_i(t) - \phi$ confirmed visitors visit $S_1$.

If all $\Delta I_i(t)$ confirmed infected travelers from $I_i$ are not transferred to $S_1$, the conditional probability that among the remaining $I_i(t-1) - \Delta I_i(t)$ infected hosts, $\mathcal{H}_{i1}$ infected travelers are transferred to $S_1$, which is measured by the following transferring estimator:

$$
\Omega_{p}(\mathcal{H}_{i1}|I_i(t-1) - \Delta I_i(t)) = \frac{[I_i(t-1) - \Delta I_i(t)]!}{[\mathcal{H}_{i1}]!}\left[\sum p_\ell + \bar{p}_i\right]^{\eta} \left[1 - \sum_{\ell} p_\ell + p_k \right]^{\Delta I_i(t)}
$$

where the last item on the RHS accounts for the probability that $\Delta I_i(t)$ confirmed infected travelers all do not visit $S_1$.

Taking into account all the above cases, the probability that $\mathcal{H}_{i1}$ infected hosts arrive at destination $S_1$ is measured by the following transferring estimator:

$$
\Omega_{p}(\mathcal{H}_{i1}) = \sum_{\phi=0}^{\Delta I_i(t)} \left[\Delta I_i(t)\right] [\sum p_\ell + \bar{p}_i\right]^{\eta} \left[1 - \sum_{\ell} p_\ell + p_k \right]^{\phi} \frac{[I_i(t-1) - \Delta I_i(t)]!}{[\mathcal{H}_{i1} - \Delta I_i(t) + \phi]!}\left[\sum p_\ell + \bar{p}_i\right]^{\Delta I_i(t)}
$$

Type 2: $\Delta I_i(t) > \mathcal{H}_{i1}$, i.e., the observed reduction in the number of infected hosts $\Delta I_i(t)$ exceeds the number of infected hosts transferred to $S_1$. Similar to the above analysis, we develop the transferring estimator by considering all possible cases that are in accordance with this condition.

If $\mathcal{H}_{i1}$ infected hosts transferred to $S_1$ are all from the observable travelers $\Delta I_i(t)$, the transferring estimator becomes

$$
\Omega_{p}(\mathcal{H}_{i1}) = \left[\Delta I_i(t)\right] [\sum p_\ell + \bar{p}_i\right]^{\eta} \left[1 - \sum_{\ell} p_\ell + p_k \right]^{\phi} \frac{[I_i(t-1) - \Delta I_i(t)]!}{[\mathcal{H}_{i1} - \Delta I_i(t) + \phi]!}\left[\sum p_\ell + \bar{p}_i\right]^{\Delta I_i(t)}
$$

where the last item accounts for the constraint that the remaining $I_i(t) - \Delta I_i(t)$ infected hosts will not be transferred to $S_1$.

Similar to type 1, the other two cases are: only a fraction of $\mathcal{H}_{i1}$ infected hosts transferred to $S_1$ are from the observable travelers $\Delta I_i(t)$, and $\mathcal{H}_{i1}$ infected hosts transferred to $S_1$ are all not from the observable travelers $\Delta I_i(t)$, we can also derive the transferring estimators.

Taking into account all these cases, the probability that $\mathcal{H}_{i1}$ infected hosts move to the destination subpopulation $S_1$ is measured by the following transferring estimator:

$$
\sum_{\mathcal{H}_{i1}=0}^{\mathcal{H}_{i1}} \left[\Delta I_i(t)\right] [\sum p_\ell + \bar{p}_i\right]^{\eta} \left[1 - \sum_{\ell} p_\ell + p_k \right]^{\phi} \frac{[I_i(t-1) - \Delta I_i(t)]!}{[\mathcal{H}_{i1} - \Delta I_i(t) + \phi]!}\left[\sum p_\ell + \bar{p}_i\right]^{\Delta I_i(t)}
$$

Generally, set $I$ consists of the three classes of subpopulations $I_i (1 \leq i \leq m)$ discussed above: unobservable subpopulation, partially observable subpopulation, and observable subpopulation of $I \rightarrow S$. According to (4), generally each potential pathway $a_i$ corresponds to a potential solution $\sigma_i$, the most-likely invasion pathway for a $m|l \rightarrow S$ can be identified as

$$
\hat{\sigma}_{m|l} = \text{arg max } P(\sigma_i|E_{m|l})
$$

B. Case of $m|l \rightarrow nS (m > 1, n > 1)$

Finally, we consider the case of $m|l \rightarrow nS (m > 1, n > 1)$, which is more complicated than $m|l \rightarrow S$, because some infectious populations in set $\mathbb{I}$ may have more than one invasion edge to the corresponding susceptible subpopulations in set $\mathbb{S}$, and the number of elements in set $\mathbb{S}$ is more than one, which obey a joint probability distribution of transferring likelihood. As shown in Fig. 2, an INC $m|l \rightarrow nS$ includes set $\mathbb{I} = \{I_i|i = 1, 2, \ldots, m\}$ and $\mathbb{S} = \{S_j|j = 1, 2, \ldots, n\}$. The first arrival infected individuals invaded each susceptible subpopulation in set $\mathbb{S}$ are $\{H_{i1}|i = 1, 2, \ldots, n\}$, respectively. Here, denote $U_i(i = 1, 2, \ldots, m)$ the subset of susceptible neighbor subpopulations in set $\mathbb{S}$ of infected subpopulation $I_i$, and $Y_j(j = 1, 2, \ldots, n)$ the subset of infected neighbor subpopulations in set $\mathbb{S}$ of susceptible subpopulation $S_j$. 
We define \( \sigma = \{ (H_{i1},i \in Y_1),\ldots, (H_{in},i \in Y_n) \} \) a potential solution for the \( ml \mapsto nS \), if subject to two conditions:

1) \[ \sum_{i \in Y_k} H_{ik} = H_k \quad (16) \]

\( H_{ik} \geq 0; \)

2) for any \( H_{ik} \) which denotes the number of infected hosts travel to subpopulation \( S_i \) from \( i \) at \( t_{\text{EAT}} \), we have

\[ \sum_{k \in U_i} H_{ik} \leq I_i(t-1), \quad 1 \leq i \leq m, \quad 1 \leq k \leq n. \]

If a \( ml \mapsto nS \) has \( M \) potential solutions, let \( \sigma = \{ H_{i1}^j^n | i \in Y_1, \ldots, H_{in}^j^n | i \in Y_n \} \), \( 1 \leq j \leq M \).

Similarly, we first discuss the accurately identifiable pathway for a given \( ml \mapsto nS \), then estimate the most-likely numbers of each \( H_{ik} \) as accurate as possible by designing our identification algorithm, since one solution of (16) corresponds to one invasion pathway of an INC \( ml \mapsto nS \).

1) Accurate Identification of Invasion Pathway: Given a few satisfied prerequisites, for all \( k \in U_i, i \in Y_k \) of the equations constituted by (16) can has a unique solution, which implies that the invasion pathway of that INC can be identified accurately. Theorem 2 elucidates this scenario.

**Theorem 2 (Accurate Identification of Invasion Pathway):** With the following conditions: 1) the number of invasion edges \( E_{in} \leq n + m; \) 2) the neighbor subpopulations of each subpopulation in set \( \mathbb{I} \) are with the transition \( S \) to \( S \) or \( I \) to \( S \) except their neighbor subpopulations in set \( \mathbb{S} \) during \( t_{\text{EAT}} \) to \( t_{\text{EAT}} \); and 3) \[ \sum_{i=1}^{n} \Delta I_i(t) = \sum_{k=1}^{n} \Delta H_k, \] the invasion pathway of an INC \( ml \mapsto nS(m,n > 1) \) can be identified accurately.

**Proof:** Since the number of infected individuals in the partially observable subpopulation \( i \) reduces at time \( t \), i.e., \( I_i(t) < I_i(t-1) \) and \( I_i(t) > 0 \), it is inevitable that a few infected carries diffuse away from subpopulation \( i \). Occurring the state transitions of \( S \to I, I \to S \) at time \( t \), subpopulations in the neighborhood of \( i \) (excluding the new contaminated subpopulation \( j \) cannot receive infected travelers. Therefore, the only possible destination for those infected travelers is subpopulation \( j \).

The conditions \( E_{in} \leq n + m \) and \[ \sum_{i=1}^{n} \Delta I_i(t) = \sum_{k=1}^{n} \Delta H_k \] make the equations \[ \sum_{i \in Y_k} H_{ik} = H_k \quad \text{and} \quad \sum_{k \in U_i} H_{ik} = \Delta I_i(t) \] only has the unique solution \( \sigma = \{ (H_{i1},i \in Y_1),\ldots, (H_{in},i \in Y_n) \} \). The reason is that rank(\( A_{\text{coef}} \)) = \( E_{in} \), where \( A_{\text{coef}} \) is the coefficient matrix of equations \[ \sum_{i \in Y_k} H_{ik} = H_k \quad \text{and} \quad \sum_{k \in U_i} H_{ik} = \Delta I_i(t). \] Thus the invasion pathway of this \( ml \mapsto nS(m,n > 1) \) can be identified accurately.

2) Potential Invasion Pathway: If the conditions of Theorem 2 are unsatisfied, the equations constituted by (16) has multiple solutions, each solution corresponds to a set of potential invasion pathways that can result in the related \( ml \mapsto nS(m,n > 1) \). We derive the transferring likelihood of each potential solution similar to case of \( ml \mapsto S \). Therefore, the likelihood of solution \( \sigma_j \) is characterized by

\[ P(\sigma_j | \varepsilon_{mns}) = \prod_{k=1}^{m} \Omega(H_{ik}^{j_k}) \left/ \sum_{j=1}^{M} \prod_{k=1}^{m} \Omega(H_{ik}^{j_k}) \right. \quad (17) \]

where \( M \) represents the number of solution \( \sigma_j \), and the last item \( \Omega(H_{ik}^{j_k}) \) represents the transferring estimator of infected subpopulation \( I_k \) in \( \mathbb{I} \), \( k_h \in Y_k \). Note that \( \sigma_j \) and \( \varepsilon_{mns} \) correspond to a potential invasion pathway \( a_j \) of \( ml \mapsto nS \) and \( G_{mns} \), respectively.

Now we discuss the transferring estimator of subpopulation \( I_i \) according to its extent of subpopulation observability.

1) Subpopulation \( I_i \) has only one neighbor (invasion edge) in set \( \mathbb{S} \). In this case, the transferring estimator is the same as the depicted one in \( ml \mapsto S \).

2) Subpopulation \( I_i \) has \( \rho \) \( (\rho \geq 2) \) neighbors (invasion edges) in set \( \mathbb{S} \).

Suppose there are totally \( k_i \) edges emanate from \( I_i \) which consist of the following three kinds.

1) There are \( \rho_i \) invasion edges \( 2 \leq \rho_i \leq n \), labeled \( 1, 2, \ldots, \rho_i \), along which the traveling rates are \( p_{h}, h \in [1, \rho_i] \), and \( H_{ih} \) invade the subpopulations in the subset \( \{ Y_i = i_h \} \), respectively.

2) There are \( \ell_i \) unobservable and partially observable labeled \( 1 + \rho_1, \ldots, \ell_i + \rho_i \), respectively. Along each unobservable or partially observable edge, the traveling rate is \( p_{\ell, \ell} \), \( \ell \in [1, \ell_i] \), and \( X_i \) infected hosts leave \( I_i \). Accordingly, in total \( n_i = \sum \ell_i x_i \) infected hosts leave \( I_i \) through the unobservable and partially observable edges.

3) There remain \( k_i - \ell_i - \rho_i \) observable edges, labeled as \( \ell_i + \rho_1 + 1, \ldots, k_i \), respectively. Along each observable edge, the traveling rate is \( p_{\rho_{i+1}, \rho_{i+1}} \in [1, \rho_{i+1}], x_{\ell_i+1} \) infected hosts stay at the source \( I_i \). There are \( x_{\ell_i+1} \) infected hosts staying in subpopulation \( I_i \) with the probability \( \pi_{i}^\ell \). Because \( I_i \) connects the unobservable and partially observable infected subpopulations, we only know the sum \( \sum \ell_i x_i + \pi_i = \gamma \).

Now, we employ the following estimators to evaluate the transferring likelihood of the three categories of \( I_i \).

\( a) \) Unobservable subpopulation \( I_i \): Because \( \Delta I_i(t) = I_i(t-1) - I_i(t) \leq 0 \), we do not know whether and how many infected individuals travel to which destinations. Similar to the INC \( ml \mapsto S \), the transferring likelihood estimator of \( I_i \) is

\[ \Omega_a(H_{ih}) = P(H_{ih}, p_h, h = [1, \ldots, \rho], x_\ell, p_\ell) \quad \ell = [1 + \rho, \ldots, l + \rho]; x_\ell, p_\ell \quad \rho = [l + \rho, \ldots, k]; \pi_i, \pi_i^\ell. \quad (18) \]

By means of the observable edges, the transferring estimator can be simplified as

\[ \sum_{\ell_i = l(t-1)} \cdot H_{ih} = P(H_{ih}, p_h, h = [1, \ldots, \rho]; x_\ell, p_\ell) \]

\[ = \sum_{\ell_i = l(t-1)} \prod_{l=1}^{t} \frac{I_i(t-1)!}{l \cdot x_i \cdot \pi_i^\ell} \cdot \prod_{\ell} p_{\ell}^{\ell - \rho_i} = \sum_{\ell_i} \frac{I_i(t-1)!}{\prod_{l=1}^{t} l \cdot x_i \cdot \pi_i^\ell} \cdot \prod_{\ell} p_{\ell}^{\ell - \rho_i}. \quad (19) \]
Then the transferring estimator becomes by the marginal distribution as

$$
\Omega_d = \frac{I_1(t - 1)! \prod_{h} \frac{H_{i_h}}{h} \left[ \sum_{\ell} p_{\ell} + \bar{p} \right]}{\prod_{h} H_{i_h}! (\Delta I(t) - \sum_{n} H_{i_n})! \prod_{h} \left( \frac{p_h}{\sum_{k=1}^{l_{p}} p_k} \right)^{H_{i_h}} \Delta I(t) - \sum_{n} H_{i_n}}}.
$$

(20)

b) Observable subpopulation $I_i$ ($I_i$ to $S_i$): For this situation, $H_i = \{H_{i_h}|h = 1, \ldots, \rho\}$ all come from $\Delta I_i(t)$. The transferring likelihood estimator of a $I \rightarrow S$ observable subpopulation $I_i$ is

$$
\Omega_{ob} = \frac{\Delta I_i(t)! \prod_{h} \frac{H_{i_h}}{h}! (\Delta I_i(t) - \sum_{n} H_{i_n})! \prod_{h} \left( \frac{p_h}{\sum_{k=1}^{l_{p}} p_k} \right)^{H_{i_h}} \Delta I_i(t) - \sum_{n} H_{i_n}}{\prod_{h} H_{i_h}! (\Delta I_i(t) - \sum_{n} H_{i_n})! \prod_{h} \left( \frac{p_h}{\sum_{k=1}^{l_{p}} p_k} \right)^{H_{i_h}} \Delta I_i(t) - \sum_{n} H_{i_n}}} \times \left( \sum_{i_{l_{p}}} p_{l} \right)^{H_{i_l}}. \tag{21}
$$

where $\Delta I_i(t) = I_i(t - 1) - I_i(t) = I_i(t - 1)$.

c) Partially observable subpopulation $I_i$: Due to $\Delta I_i(t) = I_i(t - 1) - I_i(t) > 0$, at least $\Delta I_i(t)$ infected hosts leave source $I_i$ from $\text{EAT}_1$ to $\text{EAT}_2$.

We first decompose $H_i = \{H_{i_h}|h = 1, \ldots, \rho\}$ as two subsets: $H_i' = \{H_{i_h}'|h = 1, \ldots, \rho\}$ and $H_i'' = \{H_{i_h}''|h = 1, \ldots, \rho\}$, $H_{i_h}' + H_{i_h}'' = H_{i_h}$, where $H_{i_h}' \geq 0, H_{i_h}'' \geq 0$. Denote $H_i' = \{H_{i_h}'|h = 1, \ldots, \rho\}$ the infected hosts coming from $I_i(t - 1) - \Delta I_i(t)$, and $H_i'' = \{H_{i_h}''|h = 1, \ldots, \rho\}$ the infected hosts coming from $\Delta I_i(t)$. Then we analyze the transferring estimator on the following two types.

Type 1 ($\sum_{h} H_{i_h}'' \geq \Delta I_i(t)$): Suppose $\phi = \sum_{h} H_{i_h}'' (0 \leq \phi \leq \Delta I_i(t))$, which represents the number of infected hosts coming from $\Delta I_i(t)$. Given a fixed $\phi$, there may be more than one permutation $H_i'' = \{H_{i_h}''|j = 1, \ldots, \rho\}$ for $H_i''$. The transferring likelihood estimator is

$$
\Omega_{pu} = \frac{\sum_{\phi=0}^{\Delta I_i(t)} \sum_{\sum H_{i_h}'' = \phi} P_1 P_2}{\sum_{\phi=0}^{\Delta I_i(t)} \sum_{\sum H_{i_h}'' = \phi}} \tag{22}
$$

where

$$
P_1 = \frac{\Delta I_i(t)! \prod_{h} \frac{H_{i_h}'}{h}! (\Delta I_i(t) - \phi)! \prod_{h} \left( \frac{p_h}{\sum_{k=1}^{l_{p}} p_k} \right)^{H_{i_h}'} \Delta I_i(t) - \phi}}{\prod_{h} H_{i_h}! (\Delta I_i(t) - \sum_{n} H_{i_n})! \prod_{h} \left( \frac{p_h}{\sum_{k=1}^{l_{p}} p_k} \right)^{H_{i_h}} \Delta I_i(t) - \sum_{n} H_{i_n}}} \times \left( \sum_{i_{l_{p}}} p_{l} \right)^{H_{i_l}} \tag{23}
$$

and

$$
P_2 = \frac{\sum_{\phi=0}^{\Delta I_i(t)} \sum_{\sum H_{i_h}'' = \phi} P_1 P_2}{\sum_{\phi=0}^{\Delta I_i(t)} \sum_{\sum H_{i_h}'' = \phi}} \tag{24}
$$

Type 2 ($\sum_{h} H_{i_h}'' < \Delta I_i(t)$): Suppose $\phi = \sum_{h} H_{i_h}'' (0 \leq \phi \leq \sum_{h} H_{i_h}'')$, which represents the number of infectious hosts coming from $\Delta I_i(t)$. Given a fixed $\phi$, there may be more than one solution for $H_i''$. The transferring likelihood estimator is

$$
\Omega_{pu} = \frac{\sum_{h} H_{i_h}'' \prod_{h} \frac{H_{i_h}'}{h}! (\Delta I_i(t) - \sum_{n} H_{i_n})! \prod_{h} \left( \frac{p_h}{\sum_{k=1}^{l_{p}} p_k} \right)^{H_{i_h}'} \Delta I_i(t) - \sum_{n} H_{i_n}}{\prod_{h} H_{i_h}! (\Delta I_i(t) - \sum_{n} H_{i_n})! \prod_{h} \left( \frac{p_h}{\sum_{k=1}^{l_{p}} p_k} \right)^{H_{i_h}} \Delta I_i(t) - \sum_{n} H_{i_n}}} \times \left( \sum_{i_{l_{p}}} p_{l} \right)^{H_{i_l}}. \tag{25}
$$

According to (17), the most-likely invasion pathways for an INC $mI \rightarrow nS$ can be identified as

$$
\hat{d}^{mI_{nS}} = \arg \max_{d} P(\sigma_i|E_{mI_{nS}}) = \arg \max_{d} P(\sigma_i|G_{mI_{nS}}). \tag{26}
$$

Note that if the first arrival infectious individuals $H \geq 3$, there may be multiple potential solutions corresponding to one potential pathway. For example, a $mI \rightarrow S$ is illustrated in Fig. 4. In this situation, we merge the transferring likelihood of potential solutions of $mI \rightarrow S$ or $mI \rightarrow nS$ if they belong to the same invasion pathways, then find out the most-likely invasion pathways, which are corresponding to the maximum transferring likelihood.

According to (1) and (2), the whole invasion pathway $T$ can be reconstructed chronologically by assembling each identified invasion pathway of each INC after identification of four classes of INCs. To depict the IPI algorithm explicitly, the pseudocode for our algorithm is given in Algorithm 2.
C. Analysis of IPI Algorithm

Science IPI algorithm is based on hierarchical-iteration-like decomposition technique, which reduce the temporal-spatial complexity of spreading, it can handle large-scale spatial pandemic. Note that the invasion infected hosts \( \mathcal{H}_t \) at EAT always are very small (generally \( \leq 3 \)). Therefore, the computation cost of our IPI algorithm is small, and we employ the enumeration algorithm to compute each of \( M \) potential permutations. In this section, we only discuss the simplest situation that one pathway only corresponds to one permitted solution in an INC. The situation of one pathway corresponds to multiplex potential solutions can be extended.

Denote \( \pi \) the probability corresponding to the most likely pathways for a given INC. Thus we have

\[
\pi(\sigma) = \sup_{\pi_i} \{P(\sigma_i | \mathcal{E})\}.
\]

(25)

**Property 1:** Given an INC “ml \( \mapsto S \)” or “ml \( \mapsto nS \),”

\[
P(\sigma | \mathcal{E}) = \left(\prod_{i=1}^{M} \Omega / \sum_{i=1}^{M} \prod_{j=1}^{M} \Omega\right),
\]

where\( \Omega \) is the number of potential solutions. Thus \( P_{\text{max}} = P(\sigma_1 | \mathcal{E}) + \ldots + P(\sigma_M | \mathcal{E}) \); Because \( \pi(\sigma) \geq 1/M \), let \( P_{\min} = 1/M, P(\sigma_1 | \mathcal{E})/P(\sigma_1 | \mathcal{E}) = \sum_{i=1}^{M} P(\sigma_i | \mathcal{E}) \).

We have \( P_{\min} \leq \pi(\sigma) \leq P_{\max}. \)

(26)

**Proof:** Suppose that \( P(\sigma_1 | \mathcal{E}) \leq \ldots \leq P(\sigma_M | \mathcal{E}) \), where \( M \) is the number of potential solutions. Thus \( P_{\text{max}} = P(\sigma_1 | \mathcal{E}) + \ldots + P(\sigma_M | \mathcal{E}) \); Because \( \pi(\sigma) \geq 1/M \), let \( P_{\min} = 1/M, P(\sigma_1 | \mathcal{E})/P(\sigma_1 | \mathcal{E}) = \sum_{i=1}^{M} P(\sigma_i | \mathcal{E}) \).

We have \( P_{\min} \leq \pi(\sigma) \leq P_{\max}. \)

D. Identifiability of Invasion Pathway

Accordingly, our IPI algorithm first decomposes the whole invasion pathways into four classes of INCs. Some INCs are easy to identify, but some are difficult. Therefore, it is important to describe how possible an INC can be wrongly identified. The identification extent of an INC relates with the absolute value of \( \pi(\sigma) \) and information given by the probability vector of all potential invasion pathways. We employ the entropy to describe the information of likelihood vector, which contains the all likelihood of \( M \) potential solutions/pathways of an INC.

**Definition 1 (Entropy of Transferring Likelihoods of M Potential Solutions):** According to Shannon entropy, we define the normalized entropy of transferring likelihood \( P(\sigma | \mathcal{E}) \), \( P(\sigma_M | \mathcal{E}) \) as

\[
S = -\frac{1}{\log M} \sum_{i=1}^{M} P(\sigma_i | \mathcal{E}) \log P(\sigma_i | \mathcal{E}).
\]

(27)

This likelihood entropy \( S \) tells the information embedded in the likelihood vector of the potential solutions of a given INC.

The bigger of \( \pi(\sigma) \) and the smaller of entropy \( S \), the easier to identify the epidemic pathways for an INC. Define identifiability of invasion pathways to characterize the feasibility an INC can be identified

\[
\Pi = \pi(\sigma)(1 - S).
\]

(28)

Although the likelihood entropies of some INCs are small (less than 0.5), they are still difficult to identify, because their \( \pi(\sigma) \) are much less than 0.5. Therefore, identifiability \( \Pi \) describes the practicability of a given \( ml \mapsto S \) or \( ml \mapsto nS \) better than only using \( \pi(\sigma) \) or likelihoods entropy \( S \). The identifiability statistically tells us why some INCs are easy to identify, whose \( \Pi \) are more than 0.5, and why some INCs are difficult to identify, whose \( \Pi \) are much less than 0.5.

Next we show that there exist the upper and lower boundaries of identifiability \( \Pi \) for a given INC.

**Theorem 3:** Given an INC “ml \( \mapsto S \)” or “ml \( \mapsto nS \),” \( \Pi = \pi(1 - S) \in (1/M(1 - S)) \) is the identifiability computed by the IPI algorithm. There exist a lower boundary \( \Pi_{\min} = 1/M (1 - S) \) and an upper boundary \( \Pi_{\max} = \pi - S(\pi(\sigma)) \) that

\[
\Pi_{\min} \leq \Pi \leq \Pi_{\max}
\]

(29)

where \( S' = -(1/\log M)(\pi \log \pi) + \sum ((1 - \pi)/(M - 1)) \log((1 - \pi)/(M - 1)). \)

**Proof:** \( \Pi = \pi(1 - S) \geq (1/M)(1 - S) \geq (1/M)(1 - S') \), where \( S' = -(1/\log M)(\pi \log \pi) + \sum ((1 - \pi)/(M - 1)) \log((1 - \pi)/(M - 1)) \) \( = -(1/\log M)(\pi \log \pi) + (1 - \pi) \log((1 - \pi)/(M - 1)) \) \( = -(1/\log M)(\pi \log \pi) + (1 - \pi) \log((1 - \pi)/(M - 1)). \) According to Fano’s inequality, the entropy \( S' \leq S' \).

On the other hand, we note that function \( f(y) = y \log(y) \) is strictly convex. According to Jensen’s inequality, \( \pi S(\pi) = \pi \times (-\pi \log \pi + \sum_{i=1}^{M} P(\sigma_i | \mathcal{E}) \log P(\sigma_i | \mathcal{E}) \) \( \geq -(\pi(\sigma)) \) \( \geq -\sum_{i=1}^{M} P(\sigma_i | \mathcal{E}) \log P(\sigma_i | \mathcal{E}) \).

Therefore, \( \Pi_{\min} \leq \Pi \leq \Pi_{\max} \), where \( \Pi_{\min} = (1/M(1 - S') \) and \( \Pi_{\max} = \pi - S(\pi(\sigma)) \). That completes the proof of Theorem 3.

IV. COMPUTATIONAL EXPERIMENTS

To verify the performance of our algorithm, we proceed networked metapopulation-based Monte Carlo simulation method to simulate stochastic epidemic process on the AAN and the Barabasi–Albert (BA) networked metapopulation.

The AAN is a highly heterogeneous network. Each node of the AAN represents an airport, the population size of which is the serving area’s population of this airport. The directed traffic flow is the number of passengers through this edge/airline. The data of the AAN we are used to simulate is based on the true demography and traffic statistics [50]. We take the maximal component consisted of 404 nodes (airports/subpopulations) of all American airports as the network size of the ANN. The average degree of the AAN is nearly \( \langle k \rangle = 16 \). The total population of the AAN is the \( N_{\text{total}} \approx 0.243 \times 10^9 \), which covers most of the population of the USA.

The BA network obeys heterogeneous degree distribution [51], which holds two properties of growth and preference attachment. For a BA networked metapopulation, each node is a subpopulation containing many individuals. The details of how to generate a BA networked metapopulation including travel rates setting is presented in Appendix B. To test the performance of our algorithm to handle large-scale network, the subpopulations number of the BA networked metapopulation is fixed as 3000. This is nearly equal to the number of the world airports network [52]. We fix \( \langle k \rangle = 16 \) as the average degree of the BA networked metapopulation.
The initial population size of each subpopulation is \( N_1 = N_2 = \cdots = N_N = 6 \times 10^5 \), and the total population is \( N_{\text{total}} = 6 \times 10^5 \times 3000 = 1.8 \times 10^9 \), which covers most of the active travelers of the world.

A. Networked Metapopulation-Based Monte Carlo Simulation Method to Simulate Stochastic Epidemic Process

At the beginning, we assume only one subpopulation is seeded as infected and others are susceptible. Thus \( I_i(0) = 5 \), \( I_i(0) = 0 (i = 2, \ldots, N) \). We record and update each individual’s state (i.e., susceptible or infected) at each time step. At each \( \Delta t \) (\( \Delta t \) is defined as the unit time from \( t-1 \) to \( t \)), the transmission rate \( \beta \) and diffusion rate \( p_{ij} \) are converted into probabilities. The rules of individuals reaction and diffusion process in \( \Delta t \) are as follows.

1) Reaction Process: Individuals which are in the same subpopulation are homogeneously mixing. Each susceptible individual (in subpopulation \( i \)) becomes infected with probability \( \beta(I_i(t)/N_i) \). Therefore, the average number of newly added infected individuals is \( \beta((N_i(t)−I_i(t))/N_i) \), but the simulation results fluctuate from one realization to another. The reaction process is simulated by binomial distribution.

2) Diffusion Process: After reaction, the diffusion process of individuals between different subpopulation posterior to the reaction process is taken into account. Each individual from subpopulation \( i \) migrates to the neighboring subpopulation \( j \) with probability \( p_{ij} \). The average number of new infectious travelers from subpopulation \( i \) to \( j \) is \( p_{ij}I_i(t) \). The diffusion process is simulated by binomial distribution or multinomial distribution.

B. Numerical Results

We compared our IPI algorithm with three heuristic algorithms that generate the SPT or minimum spanning tree of the metapopulation networks.

1) Average-Arrival-Time (ARR)-Based SPT [30]: The minimum distance path from subpopulation \( i \) to subpopulation \( j \) over all possible paths is generated in terms of mean first arrival time. Thus the average-arrival-time-based SPT is constructed by assembling all shortest paths from the seed subpopulation to other subpopulations of the whole network.

2) Effective (EFF)-Distance-Based Most Probable Paths Tree [4] Methods: From subpopulation \( i \) to subpopulation \( j \), the effective distance \( D_{ij} \) is defined as the minimum of the sum of effective lengths along the arbitrary legs of the path. The set of shortest paths from seed subpopulation to other subpopulations constitutes an SPT.

3) Monte Carlo-Maximum-Likelihood (MCML)-Based Most Likely Epidemic Invasion Tree [43]: To produce a most likely infection tree, they constructed the minimum spanning tree from the seed subpopulation to minimize the distance. Some recent works [53]–[55] uses machine learning or genetic algorithms to infer transmission networks from surveillance data. Because of the distinction in model assumptions and conditions, we do not perform comparison with them.

We consider to access the identification accuracy for the inferred invasion pathways. This accuracy is defined by the ratio between the number of corrected identified invasion pathways by each method and the number of true invasion pathways, respectively. We also compute the accuracy of accumulative INCs of \( ml \rightarrow S \) and \( ml \rightarrow nS \). This accuracy is defined by the ratio between the number of corrected identified invasion pathways by each method in this class of INC and the number of true invasion pathways in this classes of INC. Additionally, we investigate the identification accuracy of early stage of a global pandemic spreading, which is important to help understand how to predict and control the prevalence of epidemics.

In Fig. 5 (top and middle), we observe the whole identification accuracy and the early-stage identification accuracy. Fig. 5 (bottom) shows the early and whole accumulative identification accuracy of \( ml \rightarrow S \) and \( ml \rightarrow nS \) through 20 independent realizations on the AAN for each algorithm, respectively. The simulation results show our algorithm is outperformance, which indicates heterogeneity of structure of the AAN plays an important role.

Fig. 6 shows the results of the BA networked metapopulation with 3000 subpopulations, the top of which presents the identification accuracy of whole invasion pathway for each realization of the four algorithms, while Fig. 6 (middle) shows the identification accuracy of early stage invasion pathway for each realization. Fig. 6 (bottom) shows accumulative identified accuracy of \( ml \rightarrow S \) and \( ml \rightarrow nS \) of 20 realizations for four algorithms respectively. The simulation results indicate that our algorithm can handle a large scale networked metapopulation with robust performance. Note that the performance of the ARR for the BA networked metapopulation is the same as that of the EFF, because our parameter \( C \) is a constant in the diffusion model (see Appendix B). Fig. 7 shows the comparison of the actual invasion pathways and the most likely identified invasion pathways for a given realization, during the early stage on the AAN.
The numerical results suggest that networks with different topologies yield different identification performances, which indicate an identification algorithm should embed in both effects of spreading and topology. Our algorithm takes into account both heterogeneity of epidemics (the number of infected individuals) and network topology (diffusion flows).

We finally test the identifiability of an INC. Fig. 8 shows the entropy and identifiability of wrongly identified $ml \mapsto S$ of 20 realizations of epidemic spreading on the AAN. The identifiability depicts the wrongly identified $ml \mapsto S$ more reasonable than the likelihoods entropy. It indicates that identifiability $\Pi$ has a better performance to distinguish whether an INC is difficult to identify or not than using the likelihoods entropy.

**V. Conclusion**

To conclude, we have proposed an identification framework as the so called IPI algorithm to explore the problem of inferring invasion pathway for a pandemic outbreak. We first anatomize the whole invasion pathway into four classes of INCs at each EAT. Then we identify four classes of INCs, and reconstruct the whole invasion pathway from the source subpopulation of a spreading process. We introduce the concept of identifiability to quantitatively analyze the difficulty level that an INC can be identified. The simulation results on the AAN and large-scale BA networked metapopulation have demonstrated our algorithm held a robust performance to identify the spatial invasion pathway, especially for the early stage of an epidemic. We conjecture the proposed IPI algorithm framework can extend to the problems of virus diffusion in computer network, human to human’s epidemic contact network, and the reaction dynamics may extend to the susceptible-infected-removed or susceptible-infected-susceptible dynamics.

**APPENDIX A**

**MOBILITY OPERATOR**

We discuss the individual mobility operator. Due to the presence of stochasticity and independence of individual mobility, the number of successful transform of individuals between or among adjacent subpopulations is quantified by a binomial or a multinomial process, respectively. If the focal subpopulation $i$ only has one neighboring subpopulation $j$, the number of individuals in a given compartment $X \in \{S, I\} and \sum X = N_i$ transferred from $i$ to $j$ per unit time, $T_{ij}(U_i)$, is generated from a binomial distribution with probability $p$ representing the diffusion rate and the number of trials $U_i$, that is

$$\text{Binomial}(T_{ij}, U_i, p) = \frac{U_i!}{T_{ij}!(U_i-T_{ij})!}p^{T_{ij}}(1-p)^{(U_i-T_{ij})}.$$  

(30)
If the focal subpopulation $i$ has multiple neighboring subpopulations $j_1, j_2, \ldots, j_k$, with $k$ representing $i$’s degree, the numbers of individuals in a given compartment $l$ transferred from $i$ to $j_1, j_2, \ldots, j_k$ are generated from a multinomial distribution with probabilities $p_{ij_1}, p_{ij_2}, \ldots, p_{ij_k}$ ($p_{ij_1} + p_{ij_2} + \cdots + p_{ij_k} = p$) representing the diffusion rates on the edges emanated from subpopulation $i$ and the number of trails $U_i$, that is

$$Multinomial\left(\left\{T_{ij}\right\}, U_i, \left\{p_{ij}\right\}\right) = \frac{U_i!}{\prod_{l} T_{lj}!\left(U_i - \sum_{l} T_{lj}\right)!}\left(\prod_{l} p_{ij}^{T_{lj}}\right) \times \left(1 - \sum_{l} p_{ij}\right)^{U_i - \sum_{l} T_{lj}}$$

where $l \in [1, k]$.

**APPENDIX B**

**GENERIC DIFFUSION MODEL TO GENERATE BARABASI–ALBERT METAPOPULATION NETWORK**

We develop a general diffusion model to generate a BA metapopulation network in Section V, which characterizes the human mobility pattern on the empirical statistical rules of air transportation networks.

The diffusion rate from subpopulation $i$ to $j$ is $p_{ij} = (w_{ij}/N_i)$, where $w_{ij}$ denotes the traffic flow from subpopulation $i$ to $j$. These empirical statistical rules are verified in the air transportation network [52]: $p_{ij} \sim \left(\frac{k_i}{k_i + k_j}\right)^{\theta}$, $\theta' = 0.5 \pm 0.1$; $T \sim k^\theta$, $\beta' \simeq 1.5 \pm 0.1$; $N \sim T^\delta$ ($T = \sum_l w_{lj}$), $\lambda' \simeq 0.5$.

All the above empirical formulas relate to node’s degree $k$. To generate an artificial transportation network, we introduce a generic diffusion model to determine the diffusion rate

$$p_{ij} = \frac{b_{ij} k^\theta}{\sum_l b_{ij} k^\theta} C$$

where $b_{ij}$ stands for the elements of the adjacency matrix ($b_{ij} = 1$ if $i$ connects to $j$, and $b_{ij} = 0$ otherwise), $C$ is a constant, and $\theta$ is a variable parameter. We assume that parameter $\theta$ follows the Gaussian distribution $\theta \sim N(\hat{\theta}, \delta^2) = (1/\sqrt{2\pi}\delta)\exp(-(\hat{\theta} - \theta)^2/2\delta^2))$. Based on the empirical rule of $T \sim k^\theta$, $\beta' \simeq 1.5 \pm 0.1$, where $\beta$ is approximately linear with $\theta$, the least squares estimation is employed to evaluate parameters $\hat{\theta}$ and $\delta^2$ if we set the initial population of each node and constant C. Then, for a given BA network, we get a BA networked metapopulation network where the real statistic information is embedded by using the above method.

**REFERENCES**

[1] M. J. Keeling and P. Rohani, Modeling Infectious Diseases in Humans and Animals. Princeton, NJ, USA: Princeton Univ. Press, 2008.

[2] M. E. J. Newman et al., “Modeling infectious disease dynamics in the complex landscape of global health,” Science, vol. 347, no. 6227, pp. 1–10, Mar. 2015.

[3] J. P. Fitch, “Engineering a global response to infectious diseases,” Proc. IEEE, vol. 103, no. 2, pp. 263–272, Feb. 2015.

[4] D. Brockmann and D. Helbing, “The hidden geometry of complex, network-driven contagion phenomena,” Science, vol. 342, no. 6164, pp. 1337–1342, Dec. 2013.

[5] A. J. McMichael, “Globalization, climate change, and human health,” New England J. Med., vol. 368, no. 14, pp. 1335–1343, Apr. 2013.

[6] A. R. Mclean, R. M. May, J. Patterson, and R. A. Weiss, Eds., SARS: A Case Study in Emerging Infections. New York, NY, USA: Oxford Univ. Press, 2005.

[7] C. Fraser et al., “Pandemic potential of a strain of influenza A (H1N1): Early findings,” Science, vol. 324, no. 5934, pp. 1557–1561, Jun. 2009.

[8] Y. Yang et al., “The transmissibility and control of pandemic influenza A/H1N1 virus,” Science, vol. 326, no. 5953, pp. 729–733, Oct. 2009.

[9] H. Yu et al., “Human infection with avian influenza A H7N9 virus: An assessment of clinical severity,” Lancet, vol. 382, no. 9887, pp. 138–145, Jun. 2013.

[10] T.-Y. Lam et al., “Dissemination, divergence and establishment of H7N9 influenza viruses in China,” Nature, vol. 522, pp. 102–105, Jun. 2015.

[11] M. F. C. Gomes et al., “Assessing the international spreading risk associated with the 2014 West African Ebola outbreak,” PLoS Currents Outbreaks, vol. 6, Sep. 2014. Doi: 10.1371/currents.outbreaks.cd81f863d4e2a4ef7694d7d9f1e9d5a.

[12] B. J. Cowling et al., “Preliminary epidemiological assessment of MERS-CoV outbreak in South Korea, May to June,” Eurosurveillance, vol. 20, no. 25, pp. 7–13, Jun. 2015.

[13] X. Wang and G. Chen, “Complex network: Small-world, scale-free, and beyond,” IEEE Circuits Syst. Mag., vol. 3, no. 1, pp. 6–20, Jan./Mar. 2003.

[14] M. E. J. Newman, Networks: An Introduction. New York, NY, USA: Oxford Univ. Press, 2010.

[15] C. W. Tan, M. Chiang, and R. Srikan, “Fast algorithms and performance bounds for sum rate maximization in wireless networks,” IEEE/ACM Trans. Netw., vol. 21, no. 3, pp. 706–719, Jun. 2013.

[16] B. Yang, J. Liu, and D. Liu, “Characterizing and extracting multiplex patterns in complex networks,” IEEE Trans. Syst., Man, Cybern. B, Cybern., vol. 42, no. 2, pp. 469–481, Apr. 2012.

[17] X. Fu, M. Small, and G. Chen, Propagation Dynamics on Complex Networks: Models, Methods and Stability Analysis. New York, NY, USA: Wiley, 2013.

[18] R. Pastor-Satorras, C. Castellano, P. Van Mieghem, and A. Vespignani, “Epidemic processes in complex networks,” Rev. Mod. Phys., vol. 87, no. 3, pp. 925–979, Jul./Sep. 2015.

[19] L. Wang and X. Li, “Spatial epidemiology of networked metapopulation: An overview,” Chin. Sci. Bull., vol. 59, no. 28, pp. 3511–3522, Jul. 2014.

[20] L.-C. Chen and K.-M. Carley, “The impact of countermeasure propagation on the prevalence of computer viruses,” IEEE Trans. Syst., Man, Cybern. B, Cybern., vol. 36, no. 1, pp. 198–202, Feb. 2006.

[21] L.-C. Chen and K. M. Carley, “The impact of countermeasure propagation on the prevalence of computer viruses,” IEEE Trans. Syst., Man, Cybern. B, Cybern., vol. 36, no. 1, pp. 198–202, Feb. 2006.

[22] Y. Yang et al., “On identifiability of arrival times,” Proc. Nat. Acad. Sci. USA, vol. 111, no. 5, pp. 18095–18096, Dec. 2014.
[32] D. Shah and T. Zaman, “Rumors in a network: Who’s the culprit?” IEEE Trans. Inf. Theory, vol. 57, no. 8, pp. 5163–5181, Aug. 2011.

[33] W. Luo, W. P. Tay, and M. Leng, “Identifying infection sources and regions in large networks,” IEEE Trans. Signal Process., vol. 61, no. 11, pp. 2850–2865, Jun. 2013.

[34] Z. Wang, W. Dong, W. Zhang, and C. W. Tan, “Rumor source detection with multiple observations: Fundamental limits and algorithms,” in Proc. ACM SIGMETRICS, Austin, TX, USA, Jun. 2014, pp. 1–13.

[35] W. Dong, W. Zhang, and C. W. Tan, “Rooting out the rumor culprit from suspects,” in Proc. IEEE Int. Symp. Inf. Theory (ISIT), Istanbul, Turkey, Jul. 2013, pp. 2671–2675.

[36] Z. Wang, W. Dong, W. Zhang, and C. W. Tan, “Rooting out our rumor sources in online social networks: The value of diversity from multiple observations.” IEEE J. Sel. Topics Signal Process., vol. 9, no. 4, pp. 663–677, Jun. 2015.

[37] X. Han, Z. Shen, W.-X. Wang, and Z. Di, “Robust reconstruction of complex networks from sparse data,” Phys. Rev. Lett., vol. 114, no. 2, Jan. 2015, Art. ID 028701.

[38] M. Gomez-Rodriguez, J. Leskovec, and A. Krause, “Inferring networks of diffusion and influence,” in Proc. 16th ACM SIGKDD Conf. Knowl. Disc. Data Mining (KDD), Washington, DC, USA, Jul. 2010, pp. 1019–1028.

[39] M. Gomez-Rodriguez, J. Leskovec, and A. Krause, “Inferring networks of diffusion and influence,” ACM Trans. Knowl. Disc. Data, vol. 5, no. 4, pp. 1–37, Feb. 2012.

[40] M. Gomez-Rodriguez, D. Baldazzi, and B. Schölkopf, “Uncovering the temporal dynamics of diffusion networks,” in Proc. 28th Int. Conf. Mach. Learn. (ICML), Bellevue, WA, USA, Jul. 2011, pp. 561–568.

[41] M. Gomez-Rodriguez, J. Leskovec, D. Baldazzi, and B. Schölkopf, “Uncovering the structure and temporal dynamics of information propagation,” Netw. Sci., vol. 2, no. 1, pp. 26–65, Apr. 2014.

[42] H. Dunehm, M. Gomez-Rodriguez, L. Song, and B. Schölkopf, “Estimating diffusion network structures: Recovery conditions, sample complexity & soft-thresholding algorithm,” in Proc. 31st Int. Conf. Mach. Learn. (ICML), vol. 32, Beijing, China, Jun. 2014, pp. 793–801.

[43] D. Balcan et al., “Multiscale mobility networks and the spatial spreading of infectious diseases,” Proc. Nat. Acad. Sci. USA, vol. 106, no. 51, pp. 21484–21489, Dec. 2009.

[44] K.-L. Tsui, Z. S.-Y. Wong, D. Goldsmid, and M. Edessees, “Tracking infectious disease spread for global pandemic containment,” IEEE Intell. Syst., vol. 28, no. 6, pp. 60–64, Nov./Dec. 2013.

[45] M. Barthélémy, “Spatial networks,” Phys. Rep., vol. 499, nos. 1–3, pp. 1–101, Feb. 2010.

[46] A. Bazzani, B. Giorgini, S. Rambaldi, R. Gallotti, and L. Giovannini, “Statistical laws in urban mobility from microscopic GPS data in the area of Florence,” J. Stat. Mech., vol. 2010, no. 5, May 2010, Art. ID P05001.

[47] C. Peng, X. Jin, K.-C. Wong, M. Shi, and P. Lio, “Collective human mobility pattern from taxi trips in urban area,” PLoS ONE, vol. 7, no. 4, Apr. 2012, Art. ID e34487.

[48] X. Liang, J. Zhao, L. Dong, and K. Xu, “Unraveling the origin of exponential law in intra-urban human mobility,” Sci. Rep., vol. 3, Oct. 2013, Art. ID 2983.

[49] E. Brooks-Pollock, G. O. Roberts, and M. J. Keeling, “A dynamic model of bovine tuberculosis spread and control in Great Britain,” Nature, vol. 511, no. 7508, pp. 228–231, Jul. 2014.

[50] L. Wang, X. Li, Y.-Q. Zhang, Y. Zhang, and K. Zhang, “Evolution of scaling emergence in large-scale spatial epidemic spreading,” PLoS ONE, vol. 6, no. 7, Jul. 2011, Art. ID e21197.

[51] R. Albert and A.-L. Barabási, “Statistical mechanics of complex networks,” Rev. Mod. Phys., vol. 74, no. 1, pp. 47–97, Jan. 2002.

[52] A. Barrat, M. Barthélémy, R. Pastor-Satorras, and A. Vespignani, “The architecture of complex weighted networks,” Proc. Nat. Acad. Sci. USA, vol. 101, no. 11, pp. 3747–3752, Mar. 2004.

[53] X. Wan, J. Liu, W.-K. Cheung, and T. Tong, “Inferring epidemic network topology from surveillance data,” PLoS ONE, vol. 9, no. 6, Jun. 2014, Art. ID e100661.

[54] B. Shi, J. Liu, X.-N. Zhou, and G.-J. Yang, “Inferring plasmodium vivax transmission networks from tempo-spatial surveillance data,” PLoS Negl. Trop. Dis., vol. 8, no. 2, Feb. 2014, Art. ID e2682.

[55] X. Yang, J. Liu, X.-N. Zhou, and W.-K. Cheung, “Inferring disease transmission networks at a metapopulation level,” Health Inf. Sci. Syst., vol. 2, no. 8, Nov. 2014, Art. ID PMC4375841.