Local Tomography of Large Networks Under the Low-Observability Regime

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Abstract—This article studies the problem of reconstructing the topology of a network of interacting agents via observations of the state-evolution of the agents. We focus on the large-scale network setting with the additional constraint of partial observations, where only a small fraction of the agents can be feasibly observed. The goal is to infer the underlying subnetwork of interactions and we refer to this problem as local tomography. In order to study the large-scale setting, we adopt a proper stochastic formulation where the unobserved part of the network is modeled as an Erdős-Rényi random graph, while the observable subnetwork is left arbitrary. The main result of this work is to establish that, under this setting, local tomography is actually possible with high probability, provided that certain conditions on the network model are met (such as stability and symmetry of the network combination matrix). Remarkably, such conclusion is established under the low-observability regime, where the cardinality of the observable subnetwork is fixed, while the size of the overall network scales to infinity.

Index Terms—Topology inference, network tomography, graph learning, low-observability, local tomography, large-scale networks, Erdős-Rényi model, random graphs, diffusion networks.

I. INTRODUCTION

IN NETWORKED dynamical systems [1]–[4] the state of the agents comprising the network evolves over time and is affected by peer-to-peer interactions. In general, information about the profile of interactions is unavailable. It is the goal of network tomography to infer network connectivity from observing the evolution of the graph nodes. Problems of this type arise in many domains where knowledge of the underlying topology linking the agents is critical for better inference and control mechanisms. For example, in distributed processing over networks, the underlying topology is critical for the performance delivered by the distributed strategies such as consensus [5]–[16] and diffusion [17]–[22]; in the context of epidemics, it is well-known that the network topology may foster or hinder the outbreak of diseases or opinions [23]; in the context of brain functionality, it is also known that the connectivity among brain regions impacts the efficiency and robustness of the brain dynamics [24] and can help explain brain functional disorders [25], [26]; in cyber-security applications it is important to determine and understand the underlying network structure to devise effective countermeasures [27]; and tomography is also a relevant problem in economics [28] (in the context of causal inference) and physics applications [29]. Depending on the particular application, an appropriate model for the underlying networking dynamical system must be selected. In this work, we focus on a linear stochastic dynamical system that will be detailed in Sec. I-A — see also Sec. I-C for an example.

This article focuses on the large-scale network setting, where one can typically observe and/or process limited portions of the network. More formally, we address a local tomography problem: a subset of the agents is observed and their subnetwork of interactions is inferred from these observations. Figure 1 depicts the local tomography paradigm. There are three main reasons that cause this observability limitation in the large-scale network setting:

- **Accessibility-limit.** Some portions of the network are not accessible and, hence, unobservable. Moreover, in many large-scale settings the existence of some sources of interactions (i.e., unobserved network links) might be unknown.
- **Probing-limit.** The acquisition of data and storage capacities can be smaller than the scale of the network.
- **Processing-limit.** The complexity of the data-mining further constrains the size of the data that can be processed.

For instance, one may probe the activity of a subset of nodes – as it is unfeasible to track the activity of all the nodes in a large-scale network – in order to reconstruct its underlying profile of interactions. This requires that we partially observe the system and extract information about its underlying subnetwork of interactions.

Under the aforementioned local tomography setting, the problem of inferring the subnetwork topology across the observed agents becomes exceedingly challenging or even
ill-posed. It is therefore important to devise nontrivial conditions (if any) under which the problem is still well-posed, i.e., the information about the topology can be effectively inferred from the observable samples. In this article, we establish that under an appropriate setting, the problem of local tomography becomes well-posed with high probability in the thermodynamic limit: when the number of interacting agents $N$ grows, the (fixed) subnetwork topology associated with the observed agents can be perfectly recovered. We refer to such framework as “low-observability” to emphasize that we are interested in studying the local tomography problem in the thermodynamic limit of large networks while the observed part is fixed and finite. Besides ascertaining conditions under which the problem is well-posed with high probability, we further derive a procedure on the space of observables to recover the subnetwork topology. Finally, as an application of these results, we propose a strategy to learn the topology sequentially, by partitioning the observable network into small patches, and launching successive instances of the local tomography algorithm on these patches.

### A. Preview of the Main Result

Network tomography is associated with retrieving the underlying network structure of a distributed dynamical system via observation of the output measurements of the constituent elements. The typical formulation of the network tomography problem involves two main objects: $i)$ the statistical model that governs the laws of evolution of the (stochastic) dynamical system of interest; $ii)$ a set of observables. In this article, we consider a stochastic dynamical system described by a first-order Vector Auto-Regressive (VAR) or diffusion model, which is a popular theoretical model arising across many disciplines. Under this model, $N$ entities corresponding to the network agents interact over time $n$ according to the following law:

$$ y_n = A y_{n-1} + \beta x_n $$

Here, $A$ is a stable $N \times N$ matrix with nonnegative entries, and

$$ y_n = \begin{bmatrix} y_1(n), y_2(n), \ldots, y_N(n) \end{bmatrix}^T, $$

$$ x_n = \begin{bmatrix} x_1(n), x_2(n), \ldots, x_N(n) \end{bmatrix}^T, $$

with the vector $y_n$ collecting the state (or output measurements) at time $n$ of the $N$ agents comprising the network; and $x_n$ representing a random input (e.g., a source of noise or streaming data) at time $n$. The ensemble $\{x_i(n)\}$ are independent and identically distributed (i.i.d.) both spatially (i.e., w.r.t. to index $i$) and temporally (i.e., w.r.t. to index $n$). Without loss of generality, we assume that the random variables $x_i(n)$ have zero mean and unit variance.

The support-graph of $A$ reflects the underlying connections among the agents. Indeed, we have from (1) that:

$$ y_i(n) = \sum_{\ell=1}^N a_{i\ell} y_\ell(n-1) + \beta x_i(n), $$

which shows that, in order to update its output at time $n$, agent $i$ combines the outputs of other agents from time $n-1$. In particular, agent $i$ scales the output of agent $\ell$ by using a combination weight $a_{i\ell}$. Note that the output of agent $\ell$ is employed by agent $i$ only if $a_{i\ell} \neq 0$. The uppermost panel in Figure 2 offers a pictorial view of the local tomography problem, whereas the lowermost panel illustrates the role of the combination weights in determining the mutual influences between nodes.

Model (4) further shows how the observation $y_i(n)$ is affected by the source value $x_i(n)$, which is available locally at agent $i$ at time $n$.

The problem of support-graph recovery addressed in this work is generally referred to as network tomography in the literature because only indirect observations are available. In our framework, only output measurements from a subset of the nodes are accessible, and no information is available about the unobserved nodes including their number or connectivity. We refer to this paradigm as local tomography. Under this challenging framework, the goal is identifiability of the topology linking the observable agents. That is, we consider the problem of inferring the topology associated with a subset $S$ of observable interacting agents, by measuring only the outputs produced by such agents.

Let us ignore for a while the restriction of partial observability. It is tempting (and actually not that uncommon in the literature) to estimate the connections between the network agents by measuring the correlation between their output measurements. There is, however, one critical issue related to the use of the correlation measure for topology inference, arising from the streaming nature of the data. In general, when an external observer starts collecting output measurements, the network would have been in operation since some time already. Therefore, after a transient phase, over a connected network abiding by a stable linear stochastic dynamical system as described in (1), all agent pairs will become correlated. In order to illustrate this point in greater detail, let us introduce the correlation matrix at time $n$, namely, $R_S(n) \triangleq E[y_n y_n^T]$. When $A$ is symmetric (which is the case considered in this work), using (1), and neglecting transient terms associated with the thermodynamic limit: when the number of interacting agents $N$ grows, the (fixed) subnetwork topology associated with the observed agents can be perfectly recovered.
with the initial state \( y_0 \), we have that:

\[
R_0(n) = \beta^2 \sum_{i=0}^{n-1} A^2 i \xrightarrow{n \to \infty} R_0 = \beta^2 (I - A^2)^{-1}
\]  

(5)

where the latter series is guaranteed to converge whenever \( A \) is a stable matrix – all its eigenvalues lie inside the unit disc. Assuming that the system is observed at steady-state (i.e., that the system is in operation since some time), we must focus on the limiting correlation matrix, \( R_0 \). However, we immediately see from (5) that, even if the correlation matrix were perfectly known, direct retrieval of the support graph of \( A \) from \( R_0 \) is obfuscated by the fact that the correlation matrix depends on (a superposition of) powers of \( A \), and not only on \( A \). Moreover, even with full observation, when inversion of the matrix \( R_0 \) can be performed, in view of equation (5), one would retrieve \( A^2 \), and not \( A \). Then, since the mapping from \( A^2 \) to (the support graph of) \( A \) is not bijective in general, one would be faced with the inverse problem of retrieving the support graph of \( A \) from \( A^2 \) – such inverse problem can still be explored by properly reinforcing some sparsity constraints, refer, e.g., to [30].

Tomography relies primarily in developing a scheme to properly process the observables – e.g., the state evolution of the interconnected agents – so as to infer the underlying network structure. The above naïve scheme, based purely on the correlation \( R_0 \), can be improved by introducing the one-lag correlation matrix, which, in view of (1), takes the form:

\[
R_1(n) \triangleq \mathbb{E}[y_n y_{n-1}^T] = A R_0(n - 1) \xrightarrow{n \to \infty} R_1 = AR_0.
\]  

(6)

Therefore, we obtain the following relationship:

\[
A = R_1 R_0^{-1}
\]  

(7)

In principle, since there exist many ways to estimate \( R_0 \) and \( R_1 \) consistently as \( n \to \infty \), expression (7) reveals one possible strategy to estimate \( A \) (and hence its support-graph) from the observations.

Topology estimation based on relation (7) is viable whenever full-observation of the system is permitted. Under a partial observability restriction, however, when only a subset \( S \) of the network is accessible, only the covariance submatrices associated with the observable agents, denoted by \( [R_0]_S \) and \( [R_1]_S \), are available. One is certainly free to introduce a truncated version of (7), say, as:

\[
\hat{A}_S = [R_1]_S ([R_0]_S)^{-1}
\]  

(8)

It is clear from basic linear algebra that (8) is in general distinct from the ground truth matrix \( A_S = [R_1 R_0^{-1}]_S \), namely, from the combination matrix corresponding to the subnetwork connecting the observable agents \( S \) whose support must be inferred.

Despite this difference, it has been shown in the recent work [31], [32] that the support of the observable network can still be recovered (consistent tomography) through the truncated estimator in (8), under certain conditions that can be summarized as follows: i) the overall network graph is drawn from a connected Erdős-Rényi random graph with vanishing connection probability; ii) the cardinality of the observed subnetwork grows linearly with the size of the overall network; iii) the matrix \( A \) is a symmetric combination matrix belonging to a certain class.

The work [31] leads to several insightful conclusions about network tomography for Erdős-Rényi models. In this work, we pursue the same network tomography problem albeit for a different more demanding network setting, explained below, which will require new arguments and lead to new results that extend the results from [31]. In particular, the proof techniques used here will rely on graph theoretical techniques and on special graph constructs to arrive at the important conclusion that the naïve truncated estimator (8) is still able to deliver consistent tomography under partial network observations and more relaxed requirements.

It is worth noting that in comparison with [31], this work establishes a strong consistency result (Theorem 1) that does not require an independence condition. The main features of the framework proposed in the current manuscript in comparison with [31] can be summarized as follows:

— Topology of the accessible network portion. We assume that the subnetwork connecting the observable agents has an arbitrary topology, which is modeled through a deterministic graph. This subgraph is the object of inference. The remaining unobserved part is assumed to be drawn from an Erdős-Rényi random graph. The overall network construction is therefore referred to as a partial Erdős-Rényi model. It is useful to interpret and motivate this model in the classical “signal plus noise” paradigm in the following sense. For what concerns the object of the inference (i.e., the support graph of the observable nodes), it is modeled as an arbitrary deterministic signal. For what concerns the undesired component (i.e., the unobserved subnet), it is modeled as a noisy component. To get insightful results, we must choose some model for this random component. In the absence of any prior information, it is meaningful to opt for a uniform model, namely, the Erdős-Rényi random graph, where the presence/absence of each edge is determined through a sequence of i.i.d. Bernoulli experiments. Accordingly, few connections (i.e., high sparsity in the unobserved portion) take on the meaning of a controlled noise level. In contrast, in [31] it was assumed that the overall network (observed portion + unobserved portion) is drawn from an Erdős-Rényi random graph. Such a construction poses limitations on the subgraph that we wish to identify, which cannot be selected in an arbitrary fashion any longer. Moreover, the network construction used in [31] assumes a vanishing fraction of connected nodes within the observable set; a condition that is removed in the current analysis.

— Cardinality of the accessible network portion. In [31], it was assumed that the cardinality \( S \) of the observed subset \( S \) scales linearly with \( N \), so that the ratio \( S/N \) converges to some positive fraction \( \xi \) as \( N \to \infty \). In contrast, we assume here that the subnetwork of observable nodes \( S \) is fixed. This means that, in our framework, we focus on retrieving the support of a subnetwork \( S \) that is embedded in a network that becomes infinitely larger as \( N \to \infty \), i.e., the size of the unobserved component becomes asymptotically
drawn from an Erdős-Rényi random graph with connection probability setting. We shall prove that, if the unobserved network is consistent tomography is achievable under the aforementioned work (Theorem 1 further ahead) is to establish that consistency is not formulated in terms of an entry-wise recovery. Instead, consistency there is formulated in a strong sense, meaning that the whole subgraph is exactly recovered as $N \to \infty$, and no independence approximation is required.

B. Related Work

The existing approaches to network reconstruction can be categorized based on two major features:

— $\mathcal{F}_1$: Class of networked dynamical systems governing the state-evolution of the agents, e.g., the diffusion model in (1), and related observables, e.g., the process $y_n$ in (1).

— $\mathcal{F}_2$: Topology-retrieval methods that should exploit the relation between the observables and the underlying support-topology. Such methods are sensitive to the dynamics and the observables arising from the model in $\mathcal{F}_1$.

Regarding $\mathcal{F}_1$, most works focus on linear systems. Non-linear dynamics are often dealt with by linearizing via considering variational characterizations of the dynamics (under small-noise regimes) [33]–[35] or by appropriately increasing the dimension of the observable space [36], [37]. In the context of linear (or linearized) systems, particular attention is paid to autoregressive diffusion models [28], [30], [38]–[40].

For what concerns $\mathcal{F}_2$, the majority of the literature considers methods aimed at identifying commonalities between correlation constructs and graph topologies. We now make a brief summary of the available results as regards the existing topology-retrieval methods that are more closely related to our setting. To get some ideal benchmark, it is useful to start with the full observation case, and then focus on the case of interest of partial observation.

— Tomography under full observations. In [41], the authors introduce directed information graphs, which are used to reveal the dependencies in networks of interacting processes linked by causal dynamics. Such a setting is enlarged in [42], where a metric is proposed to learn causal relationships by means of functional dependencies, over a (possibly nonlinear) dynamical network. Causal graph processes are exploited in [40], where an algorithm (with a detailed convergence analysis) is proposed for topology recovery. Recently, the inverse problem of recovering the topology via correlation structures has been addressed through optimization-based methods, by reinforcing some (application-dependent) structural constraints such as sparsity, stability, symmetry. For instance, in [30], [39], since the combination matrix and the correlation matrix share the same eigenvectors, the set of candidate topologies is reduced by computing these
eigenvectors, and the inverse problem is then tackled with optimization methods under sparsity constraints.

An account of topology inference from node measurements (still under the full observations regime) is offered in [43], where a general linear model is considered and an approach based on Wiener filtering is proposed to infer the topology.

However, as already noted in [43] a Wiener filtering approach is redundant, since exact topology recovery can be obtained (with full observations) through the estimator in (7). As it is well known, this solution admits the following useful interpretation: the combination weights \( \{a_j\}_{j=1}^N \) obtained through (7) are the coefficients of the best one-step linear predictor (a.k.a., in the context of causal analysis, as Granger estimator), i.e., they yield the minimum expected squared error in estimating \( y_i(n) \) from the past samples \( \{y_j(n-1)\}_{j=1}^N \) — see, e.g., [44]. We remark that the case where (7) is applied with correlation matrices estimated empirically from the measurements provides the best one-step linear predictor in a least-squares sense (i.e., when the expected squared error is replaced by the empirical squared error evaluated on the measurements collected over time). However, all the aforementioned results pertain to the case where node measurements from the whole network are available. It is instead necessary to consider the case when only partial observation of the network is permitted.

— Tomography under partial observations, identifiability. The case of partial observations is addressed in [45], [46], for cases when the network graph is a polytree.

The case of more general topologies is instead addressed in [38], [47], where technical conditions for exact or partial topology identification are provided. It is useful to contrast such identifiability conditions with the approach pursued in the present work. Basically, the identifiability conditions offered in [38], [47] act at a “microscopic” level, namely, they need a detailed knowledge of the topology and/or the statistical model (e.g., type of noise, joint distribution of the observable data).

For these reasons, the approach is not practical for large-scale network settings (which are the main focus of this work).

In contrast, in this work we pursue a statistical asymptotic approach that is genuinely tailored to the large-scale setting: the conditions on the network topology are described at a macroscopic level through average descriptive indicators, such as the connection probability between any two nodes. Under these conditions, we focus on establishing an achievability result that holds (in a statistical sense) as the size of the network scales to infinity.

— Tomography under partial observations, methods.

As already noted, the classic, exact solution to the topology problem under full observation is provided by (7), and arises from the solution of a one-step linear prediction problem [38], [43]. Under partial observations, we propose to keep the same approach, except that the best one-step linear prediction is enforced on the observable nodes only. As a matter of fact, the combination weights estimated through (8) provide the best one-step linear prediction of the observable measurement \( y_i(n) \) (for \( i \in S \)) from the past observable measurements \( \{y_j(n-1)\}_{j \in S} \). We remark that this solution, which can still be interpreted as a Granger estimator, is widely adopted in causal inference from time series, when one ignores and/or neglects the existence of latent components. However, there is in principle no guarantee that such an estimator can provide reliable tomography. Our main goal is to establish that it actually can, under the demanding setting illustrated in Sec. I-A.

— Connections with graphical models. In a nutshell, a graphical model can be described as a collection of random variables indexed by nodes in a network, whose pairwise relationships (which determine the topology, i.e., the undirected graph) are encoded in a Markov random field. One of the fundamental problems in graphical models is to retrieve the network topology by collecting measurements from the network nodes. It is useful to comment on some fundamental differences, as well as useful commonalities, between the graphical model setting and our problem.

In the standard graphical model formulation (and, hence, in the vast majority of the available related results) the network evolution over time (e.g., the dynamical system in (1)) is not taken into account. Rather, the samples \( \{y_n\}_{n \in \mathbb{N}} \) are assumed independent across the index \( n \). This difference has at least two relevant implications.

The first difference pertains to the type of estimators used for topology retrieval. For example, in a Gaussian graphical model, the inverse of the correlation matrix \( R_0^{-1} \), a.k.a. concentration matrix, contains full information of the graph topology: the \((i, j)\)-th entry of the concentration matrix is nonzero if, and only if, nodes \( i \) and \( j \) are connected. In contrast, we see from the Granger estimator in (7) that in our case an additional operation is needed (namely, multiplication with the one-lag correlation matrix \( R_1 \)) to obtain the matrix that contains the topology information (in our case, the combination matrix, \( A \)). This difference is an inherent consequence of the system dynamics described by the first-order VAR model in (1). Second, the dynamical system ruling the network evolution usually enforces some degree of dependence between subsequent measurements. For this reason, while in our case the observations collected over time are correlated, in the standard graphical model formulation the samples upon which the topology inference is based are usually assumed statistically independent. Keeping in mind these fundamental distinctions, we now list some recent works about topology recovery on graphical models.

The idea of studying the large-network behavior through an Erdős-Rényi model has been applied in [48], where the emergence of “large” paths over the random graph (a property that we will use in our treatment) has been exploited for topology inference. However, reference [48] addresses the case of full observations. Instead, for the case of partial observations, in [49] an efficient method is proposed, which is suited for the case of large-girth graphs, such as, e.g., the bipartite Ramanujan graphs and the random Cayley graphs. In [50], still for the case of partial observations, an inference method is proposed under the assumption that the connection matrix is sparse, whereas the error matrix associated to the latent-variables component exhibits a low-rank property.

In summary, contrasted with recent results about topology recovery on graphical models, the results obtained in the
present work constitute an advance because: i) we deal with a dynamical system, see (1); ii) the partial observations setting considered in the present work relies on assumptions different from those used in [49], [50]: in our case the unobserved component is Erdős-Rényi, but the subnetwork of observable nodes is deterministic and arbitrary, and the combination matrix obeys transparent conditions borrowed from the adaptive networks literature. We believe that the possibility of working with dynamical models, the arbitrariness of the monitored subnetwork, as well as the direct physical meaning of the conditions on the combination matrix, provide useful novel insights on the problem of topology inference under partial observations.

To sum up, our major contribution lies in establishing technical guarantees for graph structural consistency of the Granger estimator applied to the subset of observable nodes. This is formally stated in the main result of this paper, Theorem 1.

Short versions of this work were reported in [51], [52].

C. Motivating Example: Adaptive Diffusion Networks

A network of $N$ agents observes a spatially and temporally i.i.d. sequence of zero-mean and unit-variance streaming data $\{x_i(n)\}$, for $n = 1, 2, \ldots$, and $i = 1, 2, \ldots, N$. Here, the letter $n$ refers to the time index while the letter $i$ refers to the node index. In order to track drifts in the phenomenon we are monitoring, the network agents implement an adaptive combination step followed by an adaptation step.

This notation is crucial in our treatment, since it will allow us to identify nodes without cumbersome and redundant double-index notation.

Finally, $\mathbb{I}$ denotes a column vector with all its entries equal to one; $0_{N 	imes N}$ denotes an $N \times N$ matrix with all its entries equal to zero; $\mathbb{I}_C$ denotes the indicator function, which is equal to one if condition $E$ is true, and is equal to zero, otherwise; the $N \times N$ identity matrix is denoted by $I$; and $\log(\cdot)$ denotes the natural logarithm.

B. Graph Notation

The set of all undirected graphs that can be defined on a set of nodes (vertex set) $\mathcal{V}$ is denoted by $\mathcal{G}(\mathcal{V})$. When $N$ is the number of nodes, the notation $\mathcal{G}(N)$ implies that the vertex set is $\mathcal{V} = \{1, 2, \ldots, N\}$.

When dealing with a graph $G \in \mathcal{G}(N)$, its connection structure (i.e., the edges of the graph) can be described through its $N \times N$ adjacency matrix. The $(i, j)$-th entry of the adjacency matrix of the graph $G$ will be denoted by the lower-case symbol $g_{ij}$, with $g_{ij} = 1$ if the nodes $i$ and $j$ are connected, and $g_{ij} = 0$ otherwise. Henceforth, we assume that $g_{ii} = 1$, i.e., all nodes exhibit self-loops. This reflects
the fact that usually each agent uses information from its own output measurement to update its state.

Given \( G \in \mathcal{G}(N) \), and a subset \( S \subseteq \{1, 2, \ldots, N\} \), the subgraph corresponding to \( S \) is denoted by \( G_S \in \mathcal{G}(S) \). The support graph of a matrix \( A \) is denoted by \( G(A) \). The \((i, j)\)-th entry of its adjacency matrix is \( \mathbb{1}_{|a_{ij} > 0|} \), namely, nodes \( i \) and \( j \) are connected on \( G(A) \) if, and only if, \( a_{ij} \) is strictly positive.

A path from \( i \) to \( j \) is a sequence of edges where the first edge originates from \( i \) and the last edge terminates at \( j \). The existence of a path of length \( r \) can be expressed as:

\[
g_{i, n_1, n_2 \ldots, n_{r-1}, j} = 1,
\]

for a certain sequence of vertices \( n_1, n_2, \ldots, n_{r-1} \) belonging to \( V \). According to this definition, a path can also pass multiple times through the same node, or can linger for one or more steps at the same node when it has a self-loop.

The set of neighbors of the node \( i \) (including \( i \) itself) in the undirected graph \( G \) will be denoted by \( N_i(G) \). The degree of the node \( i \) is the cardinality of \( N_i(G) \), whereas \( \Delta_{\text{max}}(G) \) is the maximum degree in \( G \). Likewise, the \( r \)-th order neighborhood of the node \( i \) (including \( i \) itself) is denoted by \( N_i^{(r)}(G) \), and is formally given by:

\[
N_i^{(r)}(G) = \{ j \in V : \delta_{i,j}(G) \leq r \},
\]

where \( \delta_{i,j}(G) \) is the distance between the nodes \( i \) and \( j \) on the graph \( G \), i.e., the length of the shortest path linking \( i \) and \( j \).

A random graph \( G \) obeys the Erdős-Rényi model if each edge of \( G \) is drawn, independently from the other edges, with identical probability \( p_N \). Equivalently stated, the adjacency random variables \( g_{ij} \), for \( i = 1, 2, \ldots, N \) and \( i < j \), are independent and identically distributed (i.i.d.) Bernoulli variables. The notation \( G \sim \mathcal{G}^*(N, p_N) \) signifies that the graph \( G \) belongs to the Erdős-Rényi class with connection probability that vanishes as \( N \to \infty \), and that obeys the following scaling law:

\[
p_N = \frac{\log N + c_N}{N},
\]

where \( c_N \to \infty \) as \( N \to \infty \) (in an arbitrary way, provided that \( p_N \to 0 \)). It is a well-known result that random graphs belonging to the family \( \mathcal{G}^*(N, p_N) \) are connected with high probability [54].

**Remark 1.** As a note of clarity, in the forthcoming treatment, we assume that all random variables find domain in a common probability space \((\Omega, \mathcal{F}, \mathbb{P})\), where \( \Omega \) is the set of realizations, \( \mathcal{F} \) is the sigma-algebra of measurable sets and \( \mathbb{P} \) is the probability measure. For instance, the event

\[
\{ \omega \in \Omega : \delta_{i,j}(G(\omega)) \leq r \} \in \mathcal{F},
\]

represents the set of realizations \( \omega \in \Omega \) yielding a graph \( G(\omega) \) whose distance between the (fixed) nodes \( i \) and \( j \) does not exceed \( r \). To render a more compact notation, we henceforth omit the realization \( \omega \) in the characterization of the events. For instance, in the case of the event (19) we represent it rather as

\[
\{ \delta_{i,j}(G) \leq r \},
\]

where the random quantities are emphasized by the boldface letter – in the event in (20), the only random object is the graph \( G \) as it is the only boldface variable.

### C. Useful Graph Operations

In our exposition, we will be performing certain operations over graphs, as well as evaluate certain functions such as comparing distances between nodes over distinct graphs. Therefore, it is useful to introduce the following graph operations for later use (which are illustrated in Figure 3):

1. **Graph embedding.** Given a vertex set \( V \), and a subset thereof, \( S \subseteq V \), the embedding of a graph \( G^{(1)} \in \mathcal{G}(S) \) into the larger graph \( G^{(2)} \in \mathcal{G}(V) \) will be denoted by \(^1\):

\[
G = G^{(1)} \oplus G^{(2)}, \quad G \in \mathcal{G}(V),
\]

and results in a graph with the following properties: 

i) the connections between nodes in \( S \) that are present in \( G^{(2)} \) are cancelled; ii) the nodes in the vertex set \( S \) of graph \( G^{(1)} \) are mapped into the corresponding nodes of graph \( G^{(2)} \), and so are the pertinent connections. We stress that the connections from \( S' \) to \( S \) are determined by the graph \( G^{(2)} \). We notice that the operation in (21) is not commutative (because the first graph is embedded into the second graph, and not vice versa), and that the output graph \( G \) does not depend on the connections existing in \( G^{(2)} \) among nodes belonging to the set \( S \).

\(^1\)In order to avoid confusion, we remark that the symbol \( \oplus \) is also used, in the graph literature, to denote a different kind of operation called “ring sum”. However, we prefer to denote our embedding operation by the same summation symbol to emphasize the “signal-noise” structure that is relevant in our application.
2) **Local disconnection.** Given a graph $G \in \mathcal{G}(V)$, the notation:

$$G_{U_1 \rightarrow U_2} \in \mathcal{G}(V),$$

(22)

describes the graph that is obtained from $G$ by removing all the edges that connect nodes in $U_1$ to nodes in $U_2$, namely, all the connections between $U_1$ and $U_2$.

3) **Connections inheritance.** The notation:

$$G_{j \rightarrow U} \in \mathcal{G}(V),$$

(23)

describes the graph that is obtained from $G$ through the following chain of operations: i) all edges within $U$ are removed; ii) all edges connecting nodes in $U$ to the rest of the network are removed; iii) all connections from $U$ to the rest of the network are inherited by node $j$.

All the above graph operations preserve self-loops unless otherwise stated.

**III. PROBLEM FORMULATION**

Consider a graph $G(V)$ and assume we are able to observe data from a subset $S \subset V$ of the nodes. From these observations, we would like to devise a procedure that allows us to discover the connections among the nodes in $S$, under the assumption that the structure of the graph in the complement set, $S'$, as well as the connections between $S$ and $S'$, will be random, following i.i.d. drawing of the pertinent edges. The desired construction can be formally described as follows.

Let $G^{(\text{obs})} \in \mathcal{G}(S)$ be a deterministic graph on the observable set $S$, with some unknown topology (which is not restricted in any way), and let $G^{(\text{unobs})} \sim \mathcal{G}^*(N, p_N)$ be an Erdős-Rényi random graph on $N$ nodes. We assume that the overall network graph, $G$, is of the form:

$$G = G^{(\text{obs})} \oplus G^{(\text{unobs})}$$

(24)

Specifically, the connections within the observable set $S$ are described through the graph $G^{(\text{obs})}$, while the connections within $S'$, as well as the connections between $S'$ and $S$, are described through the graph $G^{(\text{unobs})}$. Note that $G^{(\text{unobs})}$ is an Erdős-Rényi random graph on $N$ nodes, but its subgraph $G^{(\text{unobs})}_S$ is replaced by $G^{(\text{obs})}$ in characterizing $G$, in view of (24) (refer also to Figure 3). Therefore, the structure of $G^{(\text{unobs})}$ within the observable subnet becomes immaterial. Equation (24) highlights the “signal-noise” construction, with the boldface notation emphasizing the random (i.e., noisy) component that corresponds to the unobserved network portion, and with the normal font emphasizing the deterministic component that corresponds to the arbitrary topology of the observed network portion.

The aforementioned construction will be referred to as a *partial* Erdős-Rényi graph. The class of partial Erdős-Rényi random graphs with a deterministic graph component $G^{(\text{obs})}$ placed on the set $S$, will be formally represented by the notation $\mathcal{G}^*(N, p_N, G^{(\text{obs})})$. We shall often refer to the observable graph over $S$ by the simpler notation $G_S$. As such, we can also write,

$$G \sim \mathcal{G}^*(N, p_N, G_S)$$

(25)

to denote partial Erdős-Rényi random graphs with deterministic component $G_S$. We assume that $G$ (and hence $G_S$) is unknown. In this context, the goal of local tomography is to estimate $G_S$ via observing the state evolution of the observable agents in $S$. Figure 4 illustrates the partial Erdős-Rényi construction just described.

Before formulating the tomography problem, we observe that, under condition (9), the partial Erdős-Rényi graph is asymptotically connected with high probability for any choice of the subgraph $G_S$, as stated in the following lemma.

**Lemma 1 (Connectivity of partial Erdős-Rényi graphs).** Given any graph $G_S \in \mathcal{G}(S)$, the partial Erdős-Rényi graph

$$G \sim \mathcal{G}^*(N, p_N, G_S)$$

(26)

is connected with high probability, i.e.,

$$\lim_{N \rightarrow \infty} \mathbb{P}[G \text{ is connected}] = 1.$$  

(27)

**Proof:** See Appendix A.

**A. Combination Assignment.**

A combination assignment is a rule that builds the combination matrix $A$ as a function of the underlying graph $G$. We assign (positive) weights to the edges of $G$ and denote the resulting matrix of weights by $A$. Some useful and popular choices are the Laplacian and the Metropolis rules, which arise naturally in the context of adaptive diffusion networks [53], and are defined as follows:

**Laplacian rule.**

$$a_{ij} = \rho \times \left\{ \begin{aligned} \frac{\lambda g_{ij}}{d_{\text{max}}} & , & i \neq j \\ 1 - \frac{\lambda}{d_{\text{max}}} \sum_{l \neq i} g_{il} & , & i = j \end{aligned} \right.$$ 

(28)

2Strictly speaking, in the network literature the Laplacian and Metropolis rules are defined with weights that add up to one, which would correspond to (28) and (29) without the multiplying factor $\rho$. The multiplying factor $\rho$, which provides the matrix stability, is usually left separate and not absorbed into the combination matrix. For instance, in the case of diffusion networks (14) we have $\rho = 1 - \mu$, where $\mu$ is the step-size. In our treatment, it is more convenient to include this scaling factor into the combination matrix, as done in (28) and (29).
**Metropolis rule.**

\[ a_{ij} = \rho \times \begin{cases} \frac{g_{ij}}{\max(d_i, d_j)}, & \text{for } i \neq j \\ 1 - \sum_{\ell \neq i} \frac{g_{i\ell}}{\max(d_i, d_\ell)}, & \text{for } i = j \end{cases}, \quad (29) \]

where \( d_i \) is the degree of agent \( i \) and \( d_{\max} \) is the maximum degree in the network. It is useful to remark that different combination rules may have different impact on the performance of the topology estimators, as we will see in the examples presented in Sec. V.

In this paper, we shall focus on the family of nonnegative symmetric combination policies introduced in [31], and whose characterizing properties we recall next.

**Property 1 (Bounded-norm).** The maximum row-sum norm,

\[ ||A||_{\infty} \triangleq \max_i \sum_{\ell = 1}^N |a_{i\ell}|, \quad (30) \]

is upper bounded by some \( \rho < 1 \). \( \square \)

For nonnegative symmetric matrices, Property 1 becomes:

\[ ||A||_{\infty} = \max_{i=1,2,...,N} \sum_{\ell = 1}^N a_{i\ell} = \max_{i=1,2,...,N} \sum_{\ell = 1}^N a_{i\ell} \leq \rho \]

From Property 1 we see that (most of) the combination weights \( a_{i\ell} \) typically vanish as \( N \) gets large, since a finite mass of value at most \( \rho \) must be allocated across an ever-increasing number of neighbors – on an Erdős-Rényi graph, the average number of neighbors scales as \( Np_N \), and in the regime considered in this paper we have \( Np_N \to \infty \) in view of (18).

The next property identifies a useful class of combination policies, for which degeneracy to zero of the combination weights is prevented by proper scaling. As highlighted below, such property is broad enough to encompass typical combination rules, such as the Laplacian (28) and the Metropolis (29) rules.

**Property 2 (Non-degeneracy under \((Np_N)\)-scaling).** Consider a combination policy applied to a partial Erdős-Rényi graph \( G \sim \mathcal{F}^*(N, p_N, G_S) \). The combination policy belongs to class \( \mathcal{C}_\epsilon \) for some \( \epsilon > 0 \), if for all \( i, j = 1, 2, ..., N \) with \( i \neq j \):

\[ P[Np_N a_{ij} > \tau | g_{ij} = 1] \geq 1 - \epsilon_N \]

where \( \epsilon_N \) goes to zero as \( N \to \infty \). In other words, if two nodes \( i \) and \( j \) are connected (corresponding to \( g_{ij} = 1 \)), then the scaled combination coefficient \( Np_N a_{ij} \) lies above a certain threshold value denoted by \( \tau \), with high probability, for large \( N \). \( \square \)

It is useful to remark that, since condition (32) is applied to a partial Erdős-Rényi construction, the nodes belonging to the observable set \( S \) are connected in a deterministic fashion. This means that, for \( i, j \in S \), the random variable \( g_{ij} = g_{ij} \) is in fact deterministic. In this case, the condition in (32) should be rewritten, for any connected pair \((i, j)\) in \( G_S \), as:

\[ P[Np_N a_{ij} > \tau | g_{ij} = 1] = P[Np_N a_{ij} > \tau] \geq 1 - \epsilon_N, \quad (33) \]

because conditioning on a deterministic event becomes immaterial.

We now introduce a sufficient condition under which a combination rule fulfills Property 2. The relevance of this condition is that it can be readily verified for certain typical combination assignments such as Laplacian and Metropolis rules, and it automatically provides one value of \( \tau \) to identify the class \( \mathcal{C}_\epsilon \).

**Lemma 2 (Useful policies belonging to \( \mathcal{C}_\epsilon \)).** Any policy obeying the inequality

\[ a_{ij} \geq \frac{\gamma}{d_{\max}(G)} g_{ij} \]

for all \( i \neq j \) and for some \( \gamma > 0 \), satisfies Property 2 with the choice \( \tau = \gamma/e \).

**Proof:** See Appendix A. \( \square \)

We denote by \( \mathcal{C}_{\rho, \tau} \) the class of weight-assignment policies for which Properties 1 and 2 hold simultaneously.

Using the definition of the Laplacian rule in (28), it is readily verified that this rule possesses Property 1 and fulfills (34) with the choice \( \gamma = \rho \lambda_2 \). Likewise, using (29), it is readily verified that the Metropolis rule possesses Property 1 and fulfills (34) with the choice \( \gamma = \rho \). As a result, both policies belong to the class \( \mathcal{C}_{\rho, \tau} \), with the following choices of \( \tau \) (the meaning of the subscripts should be obvious):

\[ \tau_L = \frac{\rho \lambda_2}{e}, \quad \tau_M = \frac{\rho}{e}. \]

Before proving the main result of this work, it is useful to illustrate the physical meaning of Property 2 in connection with the network tomography problem. We introduce the \( S \times S \) error matrix that quantifies how much the truncated estimator in (8) differs from the true sub-matrix \( A_S \), namely,

\[ E_S \triangleq \hat{A}_S - A_S \]

The magnified \((i, j)\)-th entry of the truncated estimator in (8), \( Np_N[A_S]_{ij} \), can be written as:

\[ \begin{cases} Np_N a_{ij} + Np_N e_{ij}, & \text{if } i \text{ and } j \text{ are connected,} \\ \text{not vanishing} \end{cases} \]

(37)

where \( e_{ij} \) is an error quantity, and the qualification of being “not vanishing” is a consequence of Property 2. According to (37), if we want the nonzero entries \( Np_N a_{ij} \) to stand out from the error floor, when \( i \) and \( j \) are interacting, or to be bounded above (by \( \tau \)), when \( i \) and \( j \) are non-interacting, as \( N \) grows large, we must be able to control the impact of the error term \( Np_N e_{ij} \).

We are now ready to summarize the main problem treated in this article.

**Local Tomography.** Let \( A \) be an \( N \times N \) matrix obtained from any combination assignment belonging to the class \( \mathcal{C}_{\rho, \tau} \),
over a graph $G \sim \mathcal{G}(N, p_N, G_S)$ on $N$ nodes with a given (arbitrary) subgraph $G_S$. Let $\{[y_n]_S\}_{n \in \mathbb{N}}$ be the state-evolution associated with the observable subset of agents $S$ and obeying the stochastic dynamical law

$$y_n = Ay_{n-1} + \beta x_n.$$  

(38)

**Problem:** given $\{[y_n]_S\}_{n \in \mathbb{N}}$, can we determine $G_S$?

**IV. MAIN RESULT**

The main result of this work is to establish that the truncated estimator $\hat{A}_S = [R_1]_S (\{R_0\}_S)^{-1}$ introduced in (8), contains enough information to recover the true support graph, $G_S$, of the combination matrix $A_S$ that corresponds to the observable subset, $S$. More specifically, we establish that a positive threshold $\tau$ exists, such that the graph obtained by comparing the entries of $\hat{A}_S$ against this threshold matches, with high probability, the true support graph $G_S$. This implies that the topology of the subnetwork $G_S$ can be fully recovered, with high probability, via the output measurements $\{[y_n]_S\}_{n \in \mathbb{N}}$ used to construct $\{R_0\}_S$ and $\{R_1\}_S$, namely, via the observable nodes only. Even if broader observation is permitted, the result enables the possibility of surmounting the curse of dimensionality by processing smaller $S \times S$ matrices $\{R_1\}_S$ and $\{R_0\}_S$, instead of large-scale $N \times N$ matrices $R_0$ and $R_1$, wherein one of the operations involves the often expensive inversion of a large matrix $R_0$ and yet attaining exact recovery with high probability.

Before stating the main theorem, let us introduce a useful thresholding operator. We consider a matrix $M \in \mathbb{S}^{2 \times 2}$, whose $(i, j)$-th entry is $m_{ij}$, with $i \in S$ and $j \in S$. The thresholding operator compares the off-diagonal entries against some threshold $\tau > 0$, and produces as output a graph, $\Gamma_r(M) \in \mathcal{G}(S)$, whose adjacency matrix has $(i, j)$-th entry equal to

$$\begin{cases} 1, & i \neq j, \\ 0, & m_{ij} > \tau. \end{cases}$$  

(39)

In other words, the thresholding operator returns a graph whereby two nodes $i$ and $j$ are connected only if the $(i, j)$-th entry of the matrix $M$ exceeds the threshold. We assume all entries of the main diagonal of $G$ equal to one. We are now ready to state the main theorem.

**Theorem 1** (Exact recovery of $G_S$). Let $G_S$ be a given deterministic graph (with arbitrary topology) on $S$ nodes, and let $G \sim \mathcal{G}(N, p_N, G_S)$ be a partial Erdős-Rényi random graph where the sequence $c_N$ that determines the connection probability, $p_N = (1/N) \log N + c_N$, obeys the condition:

$$\frac{[\log (\log N + c_N)]^2}{\log N} \rightarrow 0.$$  

(40)

Let also $A$ be a combination matrix with support graph $G$. If there exists $\tau > 0$ and $0 < \rho < 1$ such that $A$ belongs to $\mathcal{G}_{p, \tau}$, then the following results hold.

i) If $i, j \in S$ are interacting, then the $(i, j)$-th magnified entry of the truncated estimator, $Np_N[\hat{A}_S]_{ij}$, exceeds the threshold $\tau$ with high probability as $N \rightarrow \infty$.

ii) If $i, j \in S$ are non-interacting, then the $(i, j)$-th magnified entry of the error matrix, $Np_N e_{ij}$, converges to zero in probability.

iii) The graph obtained by applying the thresholding operator in (39) to the magnified truncated estimator, $Np_N \hat{A}_S$, matches the true support graph, $G_S$, with high probability as $N \rightarrow \infty$, namely,

$$\lim_{N \rightarrow \infty} \mathbb{P}[\Gamma_r(Np_N \hat{A}_S) = G_S] = 1.$$  

(41)

**Proof:** See Appendix B.

---

**A. Outline of the Main Proof**

We offer here an outline of the proof of Theorem 1. The detailed proof is reported in Appendix B and related appendices C, D, and E.

First, we use the fact (proved in [31]) that the entries of the matrix $[\hat{A}_S]_{ij}$ converges to zero in probability over the active pairs) are desirable.

$$Np_N[\hat{A}_S]_{ij} = Np_N a_{ij} + Np_N e_{ij} \geq Np_N a_{ij}.$$  

(42)

In view of Property 2, Eq. (42) implies that, when $i$ and $j$ are interacting nodes, the quantity $Np_N[\hat{A}_S]_{ij}$ exceeds a positive threshold $\tau$ with high probability and, hence, part $i$ of Theorem 1 is proved. If, in addition, we show that the magnified error $Np_N e_{ij}$ converges to zero in probability over the non-interacting pairs, i.e., if we prove part $ii)$, then

$$Np_N [\hat{A}_S]_{ij} \rightarrow 0 \quad \text{and, hence,} \quad Np_N [\hat{A}_S]_{ij} \quad \text{stays below} \quad \tau \quad \text{as} \quad N \quad \text{goes to infinity}$$

over the non-interacting pairs. Now, in order to prove part $iii)$, we need to show that it is possible to classify correctly, as $N \rightarrow \infty$, each pair of nodes by comparing the truncated estimator $\hat{A}_S$ against the threshold $\tau$ (or any smaller value): if $Np_N[\hat{A}_S]_{ij} > \tau$, then classify $(i, j)$ as an interacting pair, otherwise classify it as non-interacting. Since the cardinality of the observable set is finite, parts $i)$ and $ii)$ imply part $iii)$ by direct application of the union bound. Accordingly, it remains to examine part $ii)$.

Using one result in [31], we rewrite the entries of the error matrix in (36) as:

$$e_{ij} = \sum_{\ell, m \in S} a_{ij} h_{\ell m} b_{mj}, \quad i, j \in S,$$  

(43)

where $B \triangleq A^2$ and $H \triangleq (I_B - B_S)^{-1}$. The error in (43) is determined by three main factors, namely: $i)$ $a_{ij}$, which is nonzero only if node $i$ (from subset $S$) and agent $\ell$ (from subset $S'$) are neighbors; $ii)$ $b_{mj}$, which is nonzero only if node $m$ (from subnet $S'$) and agent $j$ (from subset $S$) are second-order neighbors (i.e., connected in one or two steps); $iii)$ the term $h_{\ell m}$, which is the $(\ell, m)$-th entry of the matrix $H$. Clearly, in (43), the relevant entries $h_{\ell m}$ are those that are “activated” by nonzero values of $a_{ij}$ and $b_{mj}$. The $(\ell, m)$ pairs for which $a_{ij}$ and $b_{mj}$ are nonzero will be accordingly referred to as “active pairs”. Refer to Figure 5 for a graphical illustration of the active pairs.

It is also clear from (43) that, in order to get a small error, small values of $h_{\ell m}$ (over the active pairs) are desirable. In Theorem 2 – see Appendix C – we are able to show that, for large $N$, vanishing values of $h_{\ell m}$ are obtained when the distance between nodes $\ell$ and $m$ gets large. In particular, Theorem 2 states that the distance between $\ell$ and $m$ that plays an important role in the magnitude of $H$ is the one
inhomogeneous coupling is a useful formal tool that is used to reduce the
constraints graph procedure that we are going to introduce is able to solve this further result, namely, a result that holds carefully replacing the partial Erdős-Rényi random graph
by an Erdős-Rényi graph, are also rare over the large, the remaining part of the proof consists of showing that the distance on the transformed graph
is small if the distance on the transformed graph is non-interacting pairs, Theorem 1 ensures that, asymptotically, any interacting, otherwise classify them as non-interacting. The inverse dependence of η on NpN can be explained as follows. Since we compare the unnormalized matrix entries, [A_S]_ij, against the threshold η, and since in view of Theorem 1 these entries (over the connected pairs) scale essentially as 1/(NpN), a proper classification threshold should exhibit the same type of scaling.

From a practical perspective, it is necessary to select an appropriate value for η, in order to correctly classify the interacting and non-interacting pairs. In this connection, prior information on the dynamical system in (1) can be useful to set the detection threshold. Indeed, we see from (44) that knowledge is needed about: (i) the average degree NpN; and (ii) the parameter τ that characterizes the class ℋp,τ where the combination matrix stems from. Let us assume that such a knowledge is available. Now, using the values of τ reported in (35), for the Laplacian and Metropolis rules we have, respectively:

\[ \eta_L = \frac{\rho \ell}{eNpN}, \quad \eta_M = \frac{\rho \ell}{eNpN}. \]  

We observe that Theorem 1 is an asymptotic (in the size N) result. For a numerical practical application of this result, it is useful to make three remarks.

First, given a detection threshold η that guarantees exact asymptotic classification, any threshold smaller than η still guarantees exact asymptotic classification. In order to explain why, let us consider two thresholds η_1 and η_2, with η_1 < η_2, and assume that η_2 is known to provide exact asymptotic classification. Then we have that: (i) for interacting pairs, if [A_S]_ij is higher than η_2, then it is obviously higher than η_1, implying correct classification also with threshold η_1; (ii) for non-interacting pairs, Theorem 1 ensures that, asymptotically, [A_S]_ij will be smaller than any small value ε, implying correct existence of a threshold τ such that the entries of the naïve estimator ̂A_S provide correct reconstruction of the observable network with high-probability. In particular, introducing a detection threshold:

\[ \eta \triangleq \tau/(NpN), \]  

the topology of the observable network can be recovered for sufficiently large N as follows: If [A_S]_ij > η, then classify nodes i and j as interacting, otherwise classify them as non-interacting. The inverse dependence of η on NpN can be explained as follows. Since we compare the unnormalized matrix entries, [A_S]_ij, against the threshold η, and since in view of Theorem 1 these entries (over the connected pairs) scale essentially as 1/(NpN), a proper classification threshold should exhibit the same type of scaling.

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V. APPLYING THEOREM 1

Theorem 1 asserts the possibility of performing local tomography over large-scale diffusion networks as it asserts the

\[ \eta_L = \frac{\rho \ell}{eNpN}, \quad \eta_M = \frac{\rho \ell}{eNpN}. \]  

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classification also with threshold \( \eta_1 \). In other words, \( \eta_1 \) also provides exact classification.

Second, we observe that a combination rule can fulfill Property 2 for different values of \( \tau \). For example, assume that we proved that a combination rule fulfills Property 2 with a certain value \( \tau_1 \). Then, the same policy fulfills Property 2 with a higher value, e.g., \( \tau_2 > \tau_1 \).

Third, consider a pair \((i, j)\) of interacting nodes, and let us examine (42). According to Property 2, the selection of \( \tau \) relates only to the properties of the combination matrix, namely, to the behavior of \( N_{\rho N}a_{ij} \) for interacting nodes. On the other hand, for finite sizes of the network, the error \( e_{ij} \) is small, but not zero. As a result the quantity \( N_{\rho N}[\hat{A}_S]_{ij} \) will be greater than \( N_{\rho N}a_{ij} \), namely, the entries of the estimated combination matrix are, on average, shifted upward due to this additional (and positive) error. As a result, one expects that, for finite values of \( N \), increasing the values of \( \tau \) may be beneficial for classification purposes.

The aforementioned issues show that there is freedom in selecting the threshold parameter to attain exact topology recovery asymptotically (i.e., as \( N \) grows to infinity). On the other hand, we remark that different threshold choices are expected to behave differently for finite network sizes. In fact, the following trade-off arises: a higher threshold reduces the likelihood that a zero entry of the combination matrix gives rise to a (false) threshold crossing, while concurrently increasing the likelihood that a nonzero entry gives rise to a (correct) threshold crossing.

A. Nonparametric Strategies

From the analysis conducted in the previous section, we have learned the following facts about tomography based on the thresholding operator. First, a good threshold tuning requires some a-priori knowledge of the model (e.g., of the average degree, \( N_{\rho N} \), or of the class of combination matrices to set the constant \( \tau \)). Second, even with some good a-priori knowledge, it is not clear how the threshold should be optimized to maximize the performance, because a trade-off arises for finite network sizes, whose (nontrivial) solution would require an even more detailed knowledge of the underlying model.

For all these reasons, it is useful to verify the possibility of employing some nonparametric pattern recognition strategies, which work blindly (without any a-priori knowledge), and which are capable to automatically adapt the classification threshold to the empirical data. In particular, in the forthcoming experiments we will consider a \( k \)-means clustering algorithm (with \( k = 2 \)) that will be fed with the entries of the truncated estimator matrix in (8). The clustering algorithm will attempt to find some separation threshold empirically on the data, and to split accordingly the matrix entries into two clusters (connected and non-connected). The cluster with higher arithmetic mean will be then elected as the cluster of connected nodes.

B. Unknown Correlation Matrices

Until now, we have implicitly assumed that the correlation matrices, \( R_0 \) and \( R_1 \), are perfectly known, and, hence, that the truncated estimator \( \hat{A}_S \) in (8) could be evaluated exactly. However, in practice such correlation matrices are unknown, and must be estimated from the data. For this reason, we will consider numerical simulations where we empirically estimate the truncated correlation matrices \([R_0]_S\) and \([R_1]_S\) from the observed data through the sample-average estimator (boldface notation is omitted to emphasize that the observed \( y_n \) refers to a particular realization):

\[
\begin{align*}
([R_0])_S &= \frac{1}{n_{\max} + 1} \sum_{n=0}^{n_{\max}} [y_n]_S[y_n]_S^\top, \\
([R_1])_S &= \frac{1}{n_{\max}} \sum_{n=0}^{n_{\max}-1} [y_{n+1}]_S[y_n]_S^\top.
\end{align*}
\]

We remark that it is possible to optimize such estimates by exploiting prior information on the structural properties of the system, and such an optimization could boost the performance of the algorithm. This estimation-tuning is outside the scope of this paper, but showing that our strategy works with the (perhaps) simplest correlation estimators is definitely encouraging. In the next section, we will additionally display the performance of the algorithm under the ideal case of known correlations, where the exact computation of the truncated estimator in (8) can be accomplished. This ideal case provides a superior limit in performance also with respect to optimized correlation estimators.

VI. NUMERICAL EXPERIMENTS

We are now ready to present the results of the numerical experiments. In Figure 7, we display the (empirically-estimated) topology-recovery probability, with reference to an overall network with number of nodes \( N \) ranging from 10 to 200, and for the case of a Laplacian combination rule with \( \rho = 0.9 \) and \( \lambda = 0.95 \). The observable network is made up of \( S = 10 \) nodes and the connections within the latent part (and between the latent and observable nodes) are drawn according to an Erdős-Rényi model with connection probability \( p_N = 2 \log(N)/N \). We see that the probability of correct recovery gets close to 1 as \( N \) increases for all the considered scenarios: parametric versus \( k \)-means thresholding, and empirically estimated truncated correlation matrices (as in (46) and (47)) versus known truncated correlation matrices. We notice that the recovery probability curve is not monotonic. Such behavior matches perfectly our theoretical results, as we now explain. First, when \( N = S = 10 \) (first point in Figure 7), all the network is observed, and in view of (6) and the comments that follow this equation) the recovery probability must be equal to 1. Second, Theorem 1 ensures that a probability of correct recovery equal to 1 must be also attained asymptotically (in \( N \)). Accordingly, since the error probability curve starts from the value 1, and converges to 1 as \( N \) increases, the non-monotonic behavior exhibited in Figure 7 makes sense.

Let us now compare the performance of the different strategies shown in Figure 7. As one expects, the strategies that know the true correlation matrices outperform the strategies that do not know them. A separate comment is called for
diffusion recursion in (4) links the nodes through the overall nodes from all across with thus delivering a better performance. Nonparametric clustering algorithm might automatically adapt thresholding estimator is not optimized at all, whereas the theoretical considerations made in the previous section. Indeed, in the simulations the threshold employed by the algorithm. Perhaps unexpectedly, the latter strategy outperforms the former one. However, this behavior matches well with known correlations are also displayed as a superior limit in performance.

while comparing the thresholding estimator and the clustering algorithm. Perhaps unexpectedly, the latter strategy outperforms the former one. However, this behavior matches well with the theoretical considerations made in the previous section. Indeed, in the simulations the threshold employed by the thresholding estimator is not optimized at all, whereas the nonparametric clustering algorithm might automatically adapt thresholding estimator is not optimized at all, whereas the nonparametric clustering algorithm might automatically adapt to the empirical evidence arising from the data, thus delivering a better performance.

The above analysis is repeated for the case of a Metropolis combination rule with \( \rho = 0.9 \), and the result is shown in Figure 8. The same general conclusions that we draw for the Laplacian rule apply. However, we see here that the performance of the thresholding operator seems slightly worse. One explanation for this behavior is the following. The choice of the constant \( \tau \) in (35) is perhaps over-conservative. Indeed, such choice follows by estimating the asymptotic scaling law of the maximal degree (see Lemma 2), whereas the nonzero entries of the Metropolis matrix in (29) are determined only by the maximum over pairs of degrees. This means that, on average, the nonzero entries of the Metropolis matrix are higher than what is predicted by the chosen \( \tau \). It is expected that for the Metropolis rule, a larger threshold can be used without affecting the identification of connected pairs, while reducing the errors corresponding to the disconnected pairs.

A. Beyond Theorem 1

Theorem 1 establishes that, under certain technical conditions, it is possible to retrieve the topology of a subset \( S \), even when the majority of the network nodes are not observed. This appears to be a nontrivial result, since an observable measurement \( y_i(n) \), \( i \in S \), is subject to the influence of nodes from all across the network. This happens because the diffusion recursion in (4) links the nodes through the overall \( N \times N \) matrix \( A \), which takes into account also the influences that unobserved nodes have on observed nodes.

The possibility of inferring the topology of a subnet by taking measurements from this subnet only, and by ignoring the unobserved components, is of paramount importance, in view of the accessability, probing and processing limitations that arise unavoidably in practical applications. In particular, it is tempting to think about a sequential reconstruction strategy, where a larger network is reconstructed through local tomography experiments over smaller network portions, and where each local experiment obeys some probing/processing constraints. We start by illustrating the perhaps simplest sequential reconstruction strategy.

Assume that there is an observable subset of nodes \( S \), which is embedded in a larger network with many unobserved components. Due to probing and processing limitations, it is possible to probe and process at most \( M \) nodes per single experiment. Accordingly, the set \( S \) is divided into the “patches” \( S_1, S_2, \ldots, S_P \). For simplicity, we assume that the patches do not overlap each other and that they cover completely the set \( S \) (i.e., the patches form a partition of \( S \)). Each local tomography experiment will correspond to probing the union of two patches, \( S_i \cup S_j \). For this reason, each patch has cardinality \( S_i \leq M/2 \), for \( i = 1, 2, \ldots, P \), which allows coping with the probing-and-processing constraint. The process is pictorially illustrated in Figure 9. Clearly, if a particular pair of nodes does not belong to the union of patches under test, we cannot make inference on that pair. The maximum number of experiments that allows testing all pair of nodes is \( P(P - 1)/2 \). Per each experiment, we apply the local tomography strategy described in the previous section, namely: i) we compute the empirical correlation matrices, \( [R_0]_{S_i \cup S_j} \) and \( [R_1]_{S_i \cup S_j} \), from which ii) the truncated estimator \( \hat{A}_{S_i \cup S_j} \) is computed; and iii) we apply the k-means algorithm to obtain an estimated subgraph \( G_{S_i \cup S_j} \). As more and more pairs of patches are
tested, the connection profile of the network is reconstructed. A pseudo-code for the sequential reconstruction algorithm, nicknamed “Patch&Catch”, is shown in Algorithm 1.

Before seeing the Patch&Catch algorithm in operation, it is important to make a fundamental remark. Proving that the sequential reconstruction strategy retrieves the topology of \(G_S\) consistently (as \(N \to \infty\)) seems a nontrivial task. In particular, we now explain why consistency of the Patch&Catch algorithm does not come as a corollary of Theorem 1.

Assume first that \(G_S\) is an arbitrary deterministic network. In order to apply Theorem 1 to each local experiment, the unobserved component should be an Erdős–Rényi graph. However, given a union-of-patches under test, \(S_i \cup S_j\), the unobserved component is a mix of an Erdős–Rényi graph and of a portion of \(G_S\) (refer to Figure 9). Since the latter portion is not purely Erdős–Rényi (because \(G_S\) is arbitrary), Theorem 1 does not directly apply. On the other hand, if we assume that the whole graph (and, hence, also \(G_S\)) is Erdős–Rényi, then the network \(G_S\) would be not fixed as \(N \to \infty\). In particular, since \(S\) has finite cardinality, it will become asymptotically disconnected, with high probability as \(N \to \infty\).

In summary, we make no claim that the sequential reconstruction can grant consistent recovery. Therefore, the numerical results we are going to illustrate in this subsection must be intended as a preliminary test aimed at checking whether, in the finite network-size regime, a sequential reconstruction strategy might be successfully applicable.

We are now ready to see an application of the Patch&Catch algorithm. The overall network is made up of \(N = 300\) nodes, and is generated according to an Erdős–Rényi graph with probability of connection \(p_N = 5(\log N)/N\). The combination matrix \(A\) is obtained via the Metropolis rule, and the system is observed over a time scale of \(n_{\text{max}} = 10^5\) samples. We run the Patch&Catch algorithm in a sub-region \(S\), assuming a strict probing constraint of \(M = 10\) nodes per experiment.

In Figure 10 we consider a subset \(S\) of cardinality \(S = 20\), and we display the evolution of the algorithm for an increasing number of tested patches. Since \(M = 10\), and choosing equal-sized patches, we get \(P = S/(M/2) = 4\) patches. For each experiment, we depict the true graph of connections (blue edges), as well as the overall graph of connections estimated up to the current experiment (red edges). The network nodes that form the patches tested in the single experiment are highlighted in red. We see from Figure 10 that the network is faithfully reconstructed, sequentially as the number of experiments grows, until the complete subnetwork topology is correctly retrieved after \(P(P - 1)/2 = 6\) experiments.

In Figure 11, the same procedure is applied to a larger subset with \(S = 60\). For this case, we illustrate the performance delivered by the Patch&Catch algorithm in a more quantitative way. More precisely, we display the evolution, as more experiments are performed, of the normalized distance between the true graph \(G_S\) and the estimated graph \(\hat{G}_S\), namely:

\[
\text{dist}(G_S, \hat{G}_S) \triangleq \frac{2}{S(S-1)} \sum_{i,j} |g_{ij} - \hat{g}_{ij}|, \quad (48)
\]

and we assume that initially the estimated graph has no edges. We see from Figure 11 that the aforementioned distance exhibits a desirable decreasing behavior: the discrepancies between the true graph and the estimated graph diminish progressively as more experiments are conducted.

Before concluding this section, it is useful to comment on two important aspects. First, the algorithm can easily be generalized to account for overlapping patches. This would simply require to set a tie-break rule for managing the case where a particular pair, say \((h, k)\), is present in two distinct experiments. Since usually the connection probability is small, one meaningful rule could be an AND rule, where the pair \((h, k)\) is labeled as connected only if so they are in both experiments. The simplest tie-break rule might be retaining just the first classification of one pair of nodes, ignoring the results possibly arising from subsequent experiments. Second, in some applications, the reconstruction can be formed sequentially,
Sequential topology reconstruction

Probing experiment no. 1

Probing experiment no. 2

Probing experiment no. 3

Probing experiment no. 4

Probing experiment no. 5

Probing experiment no. 6

Fig. 10. Illustration of the sequential graph reconstruction. We consider \( S = 20 \) nodes with probing limit \( M = 10 \). Each patch \( S_i \) has cardinality equal to \( S_i = 5 \) nodes. At each experiment two patches are probed. The red nodes represent the nodes being probed at each experiment and the red edges represent the inferred edges up to the current experiment. All pairs were correctly classified.

by exploiting, at each experiment, the information coming from past experiments. For example, having ascertained the structure of a given subset of nodes might be informative of important network-level features of some nodes – e.g., high degree nodes – and hence, informative on their level of importance on the network. Likewise, some prior knowledge on a particular network structure (e.g., a tree structure), could help to optimize the formation of successive patches.

VII. Comparison with the Results in [31]

It is useful to contrast the results of this work with the results in [31]. As already explained in the introductory sections, the main differences can be summarized as follows. In [31], the network is homogeneous, since all the connections (also those in \( S \)) obey a classic Erdős-Rényi construction with connection probability \( p_N \). Also, the size of the network, \( S \), scales linearly with \( N \) as \( \zeta N \), meaning that the fraction of observable nodes, \( \zeta \), is constant (and greater than zero). Finally, the consistency result in [31] holds in a weak sense: it shows that the fraction of correctly identified edges converges to one in the limit and an independence approximation is used to control the error rate. In this work, consistency holds in a strong sense and the independence approximation is not required. The results obtained here generalize the above framework in several directions.

A. The Case of Fixed \( S \)

In this work, we prove that perfect recovery is achievable even in the extreme case that the number of observable nodes is fixed when \( N \) diverges, namely, when the observable network portion is embedded into an infinitely large number of unobservable nodes. We remark that the case of fixed \( S \) cannot be addressed with the tools used in [31]. Let us now explain why. The result proved in [31] relies essentially on the following result (Theorem 1 in [31]):

\[
\sum_{j=1}^{S} e_{ij} \leq \rho,
\]
which reveals that the (column-wise) sum of the errors is limited, irrespectively of the network size. This result is obtained by exploiting matrix algebra tools. It is shown in [31] how (49) leads to the useful conclusion that, on average, the off-diagonal entries of the error matrix scale as $1/S$, which further implies that:

$$\mathbb{P}[Np_Ne_{ij} > \epsilon] \lesssim \frac{N}{S} p_N^s, \quad \text{(Ref. [31])} \quad (50)$$

where the symbol “$\lesssim$” here means that the quantity appearing on the left-hand side is upper bounded by a quantity that scales, asymptotically with $N$, as the quantity appearing on the right-hand side. Equation (50) reveals two useful facts. First, when $S/N$ stays constant as $N$ grows, and since $p_N$ goes to zero, we see that the magnified error vanishes. This is one fundamental conclusion ascertained in [31]. At the same time, Eq. (50) highlights how, for fixed $S$, we are no longer in the position of establishing from (50) that the magnified error converges to zero, because the product $Np_N$ diverges with $N$. In summary, the matrix-algebra tools taken in [31] are not powerful enough to address the challenging case when $S$ is fixed, namely, when the fraction of observable nodes goes to zero as $N$ grows.

On the other hand, in this work we show how this more challenging scenario can be addressed, by exploiting matrix-graph tools, i.e., by evaluating paths and distances over graphs. One important benefit of the new approach is that the results now hold for an arbitrary topology of the observable network portion, while in [31] this latter component was constrained to be Erdős-Rényi.

B. The Case of $S \sim \xi N$

We notice that the results of this work can be applied to the case addressed in [31]. Indeed, when $G_S$ is Erdős-Rényi, we can repeat the proof of Theorem 1 by essentially skipping the homogenization-and-coupling step, because the overall graph is homogeneous ab initio. Then we would get, for any $i \neq j$ (also for the connected pairs, in this particular case):

$$\mathbb{P}[Np_Ne_{ij} > \epsilon] \lesssim N \xi p_N^s. \quad (51)$$

Therefore, both the matrix-algebra approach (used in [31]), and the matrix-graph approach (used here) lead to the result that the topology of the observable network portion can be reconstructed faithfully. However, it must be remarked that the matrix-graph approach requires some additional conditions on the connection probability, $p_N$, which translate into a slightly more restrictive requirement in terms of sparsity.

On the other hand, and interestingly, the matrix-algebra approach and the matrix-graph approach lead to different estimates on how the error probability in (51) converges to zero. Indeed, with the approach used in [31], one is able to see that the rate of decay is at least in the order of $p_N$ (see (50), and observe that $N/S \sim 1/\xi$). Moreover, in [31] it is shown that the decay rate is actually faster than $p_N$.

In contrast, with the approach adopted in the current work we get the upper bound in (51), which provides the (looser) asymptotic prediction that the decay rate is slower than $p_N$. In summary, we conclude that, under the regime $S \sim \xi N$, and for a full Erdős-Rényi construction, the results of [31] are more powerful in predicting the decay rate of the error probabilities. It could be interesting at this point to ask whether it is possible to combine the matrix-algebra approach with the matrix-graph approach to obtain refined estimates.

APPENDIX A

SOME USEFUL LEMMATA

Proof of Lemma 1: In order to prove that the partial Erdős-Rényi graph $G \sim \mathcal{G}_s(N, p_N, G_S)$ is connected, it suffices to consider the worst case where the embedded graph, $G_{S'}$, is internally disconnected, i.e., where no edges exist between nodes in $S$. We note that the nodes in $S$, even if disconnected, can still be connected to nodes belonging to the unobserved set, $S'$. The latter property enables the possibility that the overall graph, $G$, is connected, as we are going to show.

Since we are assuming that $G_S$ is internally disconnected, the overall graph is connected if both $G_{S'}$ is connected, and any node in $S$ connects to some node in $S'$. Refer to Figure 12 for a graphical illustration. We prove Lemma 1 via the contrapositive statement: the overall graph is not connected if either $G_{S'}$ is not connected, or if at least one node in $S$ is not connected to $S'$, namely, we have that:

$$\{G \text{ not connected} \} \subseteq \{G_{S'} \text{ not connected} \} \cup \exists \text{ an isolated node of } G \text{ in } S.$$  \quad (52)

Therefore, applying the union bound we get:

$$\mathbb{P}[G \text{ is not connected}] \leq \mathbb{P}[G_{S'} \text{ is not connected}] + \mathbb{P}[\exists \text{ an isolated node of } G \text{ in } S]$$  \quad (53)

$$= \mathbb{P}[G_{S'} \text{ is not connected}] + 1 - (1 - p_N)^{N-S}.$$
Since $G_S$ is a classic Erdős-Rényi $\mathcal{G}^*(N-S, p_N)$, we have that:
\[
\lim_{N \to \infty} P[G_S \text{ is not connected}] = 0.
\] (54)

Moreover, since $S$ is fixed, we have that:
\[
(1 - p_N)^{N-S} = \left(1 - \frac{\log N + c_N}{N}\right)^{N-S} \leq \left(1 - \frac{\log N}{N}\right)^{N-S} \to 0.
\] (55)

**Proof of Lemma 2:** In order to prove the claim of the lemma, we must show that (34) implies (32) with the choice $\tau = \gamma / e$. Let us observe preliminarily that (34) yields the following implication:
\[
d_{\max}(G) < eNp_N, g_{ij} = 1 \leq [Np_N a_{ij} > \tau, g_{ij} = 1].
\] (56)

Therefore, we can write:
\[
P[Np_N a_{ij} > \tau | g_{ij} = 1] \geq P[d_{\max}(G) < eNp_N | g_{ij} = 1].
\] (57)

Now, by trivial upper bounding techniques, we can obtain the following chain of inequalities:
\[
P \left[ d_{\max}(G) \geq eNp_N | g_{ij} = 1 \right]
\leq P \left[ \max_{n \in S} \sum_k g_{nk} > eNp_N | g_{ij} = 1 \right]
+ P \left[ \max_{n \in \bar{S}} \sum_k g_{nk} > eNp_N | g_{ij} = 1 \right]
= P \left[ \max_{n \in S} \sum_k g_{nk} > eNp_N | g_{ij} = 1 \right]
+ P \left[ \sum_{k \in S} g_{nk} + \sum_{k \notin S} g_{nk} > eNp_N | g_{ij} = 1 \right]
\leq P \left[ \max_{n \in S} \sum_k g_{nk} > eNp_N | g_{ij} = 1 \right]
+ P \left[ S + \max_{n \in \bar{S}} \sum_{k \notin S} g_{nk} > eNp_N | g_{ij} = 1 \right]
= P \left[ \max_{n \in S} \sum_k g_{nk} > eNp_N | g_{ij} = 1 \right]
+ P \left[ \max_{n \in \bar{S}} \sum_{k \notin S} g_{nk} > eNp_N | g_{ij} = 1 \right]
\leq \epsilon_N \rightarrow 0,
\] (62)

where the last inequality follows directly from Lemma 1 in [31], since $S$ is fixed and since the subgraph formed by the edges $g_{nk}$ with either $n \notin S$ or $k \notin S$ is Erdős-Rényi with parameter $p_N$ defined in equation (18). Actually, for $i, j \in S$ (i.e., when $g_{ij} = g_{ij}$ is deterministic) a simplified version of Lemma 1 in [31] does suffice, where the conditioning can be skipped.

**Appendix B**

**Proof of Theorem 1**

We first prove part i). It is shown in [31] that the entries of the error matrix defined in (36) are nonnegative, i.e., $E_S \geq 0_S \times S$, and, hence, we can write, for $i, j \in S$:
\[
P[Np_N[A_S]_{ij} = Np_Na_{ij} + Np_Ne_{ij} \geq Np_Na_{ij}.]
\] (63)

Therefore, from Property 2 we get immediately the claim in part i). If we further show that the magnified error $Np_Ne_{ij}$ converges to zero in probability over the non-interacting pairs (i.e., if we prove part ii) of the present theorem), then we can attain exact (with high probability) classification via inspection on the truncated estimator $A_S$; if $Np_N[A_S]_{ij} > \tau$, then classify $(i, j)$ as an interacting pair, otherwise classify it as non-interacting, where $\tau$ is the threshold characterizing the family $\mathcal{C}_{p, \tau}$ of weight assignments from where $A$ is obtained, in view of Property 2. As a result, and since the cardinality of the observable set is finite, part iii) would follow if we are able to prove part ii). The proof of part ii) is demanding and will be developed through a sequence of five steps.

**Step 1: Relating the error to the distance between nodes belonging to $S'$**. It is shown in [31] that the error matrix in (36) can be represented as:
\[
E_S = A_{SS}H B_{S'} S
\] (64)

where
\[
B \triangleq A^2, \quad H \triangleq (I_S - B_{S'})^{-1}.
\] (65)

From (64) we can write, for $i, j \in S'$:
\[
\epsilon_{ij} = \sum_{\ell, m \in S'} a_{ij} h_{\ell m} b_{mj}
\] (66)

where $\epsilon_{ij}$ is the error at the pair $(i, j)$. Therefore, in order to control the size of the error $\epsilon_{ij}$, small values of the factors $h_{\ell m}$, for $\ell, m \in S'$, would be desirable. In view of the definition for $H$ in (65), we have that:
\[
h_{\ell m} = [(I_S - B_{S'})^{-1}]_{\ell m} = \left[ \sum_{k=0}^{\infty} (B_{S'})^k \right]_{\ell m},
\] (67)

as the matrix $B_{S'} = [A^2]_{S'}$ is stable, since $\rho(B_{S'}) < \|B_{S'}\|_{\infty} < 1$, from Property 1, where $\rho(B_{S'})$ is the spectral radius of $B_{S'}$. It is useful at this point to recall the following known fact from matrix algebra that relates the entries associated with the powers of a matrix with the distances between nodes on its underlying support graph.

Let $M \in S_N^{+} \times N$ be a nonnegative symmetric matrix with positive diagonal entries, and let $G(M)$ be its underlying support graph. Consider the powers of the matrix $M$, namely, $M_k$, for $k = 1, 2, \ldots$. Then we have that [55]:
\[
\delta_{\ell, m}(G(M)) = r \Leftrightarrow \text{ the smallest } k \text{ with } [M^k]_{\ell m} > 0 \text{ is } r,
\] (68)

5During this first step the boldface notation will be skipped because we focus on properties that depend solely on the structure of the matrix, and not on the statistical model of the underlying graph.
where $\delta_{\ell,m}(G(M))$ represents the distance between the nodes $\ell$ and $m$ in the graph $G(M)$ as defined in Sec. II-B. In fact, note that, if $\ell$ is not connected to $m$ in the support graph $G(M)$, then $M_{\ell m} = 0$. If the smallest path connecting $\ell$ to $m$ has a length of two hops (in particular, $\ell$ is not connected to $m$, hence $M_{\ell m} = 0$), then there exists $k$ so that $M_{\ell k} > 0$ and $M_{km} > 0$. Thus, $[M^2]_{\ell m} = \sum_i [M_{\ell i}][M_{im}] > M_{\ell k}M_{km} > 0$. Reasoning by induction one can establish (68). The following observation follows: if $M$ is stable, i.e., $\rho(M) < 1$, and if the distance $\delta_{\ell,m}(G(M)) = r$ is large, then $[M^k]_{\ell m}$ is small for all $k$ as for $k < r$ we have $[M^k]_{\ell m} = 0$ and for $k \geq r$ the corresponding power $[M^k]_{\ell m}$ is small since $k$ is large and $M$ is stable.

Now, examining (67), and using (68) with $M = B_S$, one might be tempted to conclude that a small $h_{\ell m}$ would result if nodes $\ell$ and $m$ are distant from each other. The reasoning is correct, but note that, in general, the distance between $\ell$ and $m$ is dependent on the topology of the network $G_S$, which is arbitrary. In other words, by relying solely on the elementary observation in (68), one would not be able to draw useful conclusions about the magnitude of the entries $h_{\ell m}$ (and, hence, of the entries in the error matrix $E_S$) in our context where $G_S$ is arbitrary.

As a matter of fact, as stated in Theorem 2 (proved in Appendix C) the distance affecting $h_{\ell m}$ is the one between $\ell$ and $m$ on a transformed graph, $G_{S_{\ell,m}S}$, which is the graph obtained from $G$ by removing all the edges connecting nodes inside the observable subset $S$, introduced in Sec. II-C – refer to Figure 6 for a graphical illustration of the contrast between $G$ and $G_{S_{\ell,m}S}$. Note that the edges possibly connecting nodes from $S$ to nodes in $S'$ are not removed in the graph $G_{S_{\ell,m}S}$.

**Theorem 2.** Given two distinct nodes $\ell, m \in S'$, we have that:

$$\delta_{\ell,m}(G_{S_{\ell,m}S}) = r \Rightarrow h_{\ell m} \leq \frac{\rho^r}{1 - \rho^2}$$

(69)

where $h_{\ell m}$ is the ($\ell, m$)-th entry of the matrix

$$H = (I_S - B_S)^{-1},$$

(70)

$B_S = [A^2]_S$ and $0 < \rho < 1$ is an upper-bound for the maximum row-sum of the matrix $A$, in view of Property 1, remarking that $A$ is a combination matrix satisfying Properties 1 and 2, i.e., obtained from any weight assignment in the class $\mathcal{C}_{\rho,1}$, as e.g., the Metropolis and the Laplacian weight assignment rules.

In words, Theorem 2 relates the magnitude of the entries of $H$ with the distance between nodes in a manner that does not depend on the subnetwork $G_S$. We remark that we do not assume that the nodes in $G_S$ are not connected among each other. In fact, we impose no restrictions whatsoever on the topology of $G_S$ to prove the main theorem. Reference to the graph $G_{S_{\ell,m}S}$ is used only when devising universal bounds on the terms $h_{\ell m}$, in view of Theorem 2. In other words, we are able to rule out the role of the subnetwork topology $G_S$ in as much as computing upper bounds for $H$.

In the next step, we will show in detail how (69) is helpful to control the size of the error in (66).

**Step 2: Large distances vs. small distances.** The summation appearing in (66) can be restricted to nodes that obey the conditions:

$$\ell \in N_i(G), \quad m \in N_i^{(2)}(G),$$

(71)

namely, to nodes $\ell \in S'$ that are neighbors of the node $i \in S$ (so that $a_{i\ell} > 0$), and to nodes $m \in S'$ that are second-order neighbors of the node $j \in S$ (so that $b_{mj} > 0$). Henceforth, we refer to such pair $(\ell, m)$ as an active pair. Figure 5 depicts the possible configurations of the active pairs. In words, the summation characterizing the error in equation (66) runs only over the active pairs. In fact, the error in (66) can be represented as:

$$Np_Ne_{ij} = Np_N \sum_{\ell, m \in S'} a_{i\ell} h_{\ell m} b_{mj}$$

(72)

$$= Np_N \sum_{\ell, m \in S'} a_{i\ell} J_{\ell m} h_{\ell m} b_{mj},$$

(73)

where the randomness of the various quantities, arising from the randomness of the underlying random graph $G$, has been now emphasized through the boldface notation, and where we have introduced the variable:

$$J_{\ell m} \triangleq \{a_{i\ell} > 0, b_{mj} > 0\} \cap \{i \in N_i(G), m \in N_i^{(2)}(G)\}$$

(74)

as the indicator of an active pair $(\ell, m) \in S' \times S'$, i.e., $J_{\ell m} = 1$ if $(\ell, m)$ is an active pair and $J_{\ell m} = 0$ otherwise. Now, in order to prove part ii) of Theorem 1, we need to prove that, for two non-interacting nodes $i$ and $j$, and for any $\epsilon > 0$:

$$\mathbb{P}[Np_Ne_{ij} > \epsilon] \xrightarrow{N \to \infty} 0.$$  

(75)

As stated in Theorem 2, in Step 1, the distance between nodes $\ell, m \in S'$ on the aforementioned reference graph, $G_{S_{\ell,m}S}$, plays a role in the size of $h_{\ell m}$ and hence, in the magnitude of the error. In addition, we have seen that the relevant nodes are those obeying (71), i.e., the active pairs. It is therefore useful to introduce the following events. For $\ell, m \in S'$, with $\ell \neq m$, we define:

$$D_{\ell,m} \triangleq \{\delta_{\ell,m}(G_{S_{\ell,m}S}) \leq r_N, \ell \in N_i(G), m \in N_i^{(2)}(G)\},$$

(76)

where $r_N$ is a certain sequence of distances, with $r_N \to \infty$ as $N \to \infty$, in a way that will be specified later. The event in (76) certifies that the distance on the graph $G_{S_{\ell,m}S}$ between two distinct nodes, $\ell, m \in S'$, does not exceed a prescribed value $r_N$, and also certifies the membership of the nodes $\ell$ and $m$ to the pertinent neighborhoods defined on the graph $G$, i.e., it certifies that $(\ell, m)$ is an active pair. We remark that $D_{\ell,m}$ is, formally, a (measurable) set and that the only random object characterizing $D_{\ell,m}$ in equation (76) is the random graph $G$ – refer to Remark 1 in Sec. II-B. The observable subset $S$, the sequence $r_N$ and the indexes $i, j, \ell, m$ are fixed (or deterministic). Accordingly, $D_{\ell,m}$ represents the set of realizations of the partial Erdős-Rényi random graph $G$ where the constraints of distance and neighborhood among the fixed nodes $i, j, \ell, m$ in equation (76) are met. Refer to Figure 13 for an illustration.
Likewise, for \( \ell = m \) the set \( D_{\ell,m} \) reduces to:
\[
D_{\ell,\ell} = \{ \ell \in N_1(G) \cap N_2^{(2)}(G) \},
\]
where we see that the event \( D_{\ell,\ell} \) simply certifies the membership of the node \( \ell \) to the pertinent neighborhoods defined on the graph \( G \). We finally introduce the union event:
\[
D_{\text{small}} = \bigcup_{\ell, m \in S'} D_{\ell,m},
\]
where the event is small, i.e., \( \delta_{\ell,m}(G) \leq r_N \), for at least one active pair \((\ell, m)\) in \( S' \times S' \). We can write:
\[
P[N_P e_{ij} > \epsilon] = P[N_P e_{ij} > \epsilon, D_{\text{small}}] \tag{77}
+ P[N_P e_{ij} > \epsilon, D'_{\text{small}}] \leq P[D_{\text{small}}] + P[N_P e_{ij} > \epsilon, D'_{\text{small}}],
\]
where \( D'_{\text{small}} \) is the complement of the event (or measurable set) \( D_{\text{small}} \), i.e., \( D'_{\text{small}} \cap D_{\text{small}} = \emptyset \), and can be read as the event (or set of realizations of \( G \)) where the distances are large, i.e., \( \delta_{\ell,m}(G) > r_N \), for all active pairs \((\ell, m)\). For the sake of a compact notation, we write
\[
P[N_P e_{ij} > \epsilon, D'_{\text{small}}] \leq \frac{P[N_P e_{ij} > \epsilon]}{P[D_{\text{small}}]} \tag{80}
\]
and instead of
\[
P[N_P e_{ij} > \epsilon] \cap D'_{\text{small}} \leq \frac{P[N_P e_{ij} > \epsilon]}{P[D_{\text{small}}]} \tag{81}
\]
Then, the route that we follow to prove the claim in (75) goes as follows: i) (small-distance) we show that:
\[
P[D_{\text{small}}] \underset{N \to \infty}{\to} 0, \tag{82}
\]
i.e., the occurrence of a small distance in at least one active pair \((\ell, m)\) is rare, with high probability, and ii) (large-distance) we show that large distances\(^6\) imply small errors, formally:
\[
P[N_P e_{ij} > \epsilon, D'_{\text{small}}] = 0 \quad \text{for sufficiently large} \quad N. \tag{83}
\]
\(^6\)The terminology “small distances” and “large distances” will be often coined for simplicity to denote \( \min_{\ell,m} \text{is active } \delta_{\ell,m}(G) \leq r_N \) and \( \min_{\ell,m} \text{is active } \delta_{\ell,m}(G) > r_N \), respectively.

Equations (82) and (83) will imply the desired result in equation (75) in view of equation (80). Let us start by proving (83). From the definition in equation (78), we have that
\[
D'_{\text{small}} = \bigcap_{\ell, m \in S'} D'_{\ell,m}. \tag{84}
\]
Using (76) and (77), from (84) we conclude that the complementary event \( D'_{\text{small}} \) can be compactly expressed through the indicator variables in (74) as follows:
\[
D'_{\text{small}} = \left\{ J_{\ell,m}^{(i)} = 0 \text{ for all } \ell, m \in S' \right\}. \tag{85}
\]
where we have further introduced the indicator variable:
\[
J_{\ell,\ell}^{(i)} = 1, \quad J_{\ell,m}^{(i)} = I_{[\delta_{\ell,m}(G_{S'' \cap S'}) \leq r_N]} \forall \ell \neq m. \tag{86}
\]
That is, the event \( D'_{\text{small}} \) represents the set of all realizations of the random network \( G \), where each pair \((\ell, m)\) in \( S' \times S' \) is either non-active or, if active, then \( \ell \) is distant from \( m \), i.e., \( \delta_{\ell,m}(G) > r_N \). We now show that, in view of (69), the occurrence of \( D'_{\text{small}} \) implies an upper bound on the entries \( h_{\ell,m} \), namely,
\[
D'_{\text{small}} \subseteq \left\{ h_{\ell,m} J_{\ell,m} \leq \frac{\rho^{r_N+1}}{1 - \rho^2} J_{\ell,m} \text{ for all } \ell, m \in S' \right\} \tag{87}
\]
Indeed, we know from (85) that the occurrence of \( D'_{\text{small}} \) implies that the product \( J_{\ell,m}^{(i)} \) is equal to zero for all \( \ell, m \in S' \). Let us consider first the degenerate case \( \ell = m \). Since \( J_{\ell,\ell}^{(i)} = 1 \), the variable \( J_{\ell,m} \) must be equal to zero and (87) holds trivially. We switch to the case \( \ell \neq m \). If \( J_{\ell,\ell} = 0 \), i.e., \( (\ell, m) \) is not an active pair, then (87) holds trivially. If, instead, \( J_{\ell,\ell} = 1 \), then we must have \( J_{\ell,m}^{(i)} = 0 \), i.e.,
\[
\delta_{\ell,m}(G_{S'' \cap S'}) \geq r_N + 1. \tag{88}
\]
As a consequence, Eq. (87) holds true in view of (69) (proved in Theorem 2). Applying now (87) to (73), we conclude that, when \( D'_{\text{small}} \) occurs, we must have that:
\[
N_P e_{ij} \leq N_P \frac{\rho^{r_N+1}}{1 - \rho^2} \sum_{\ell \in S'} a_{i\ell} \sum_{m \in S'} b_{mj} J_{\ell,m} \tag{89}
\]
\[
\leq N_P \frac{\rho^{r_N+4}}{1 - \rho^2}, \tag{90}
\]
where the last inequality holds from the row-sum stability of \( A \in \mathbb{C}_{p,t} \) (Property 1):
\[
||A||_{\infty} \leq \rho, \quad ||B||_{\infty} \leq \rho^2. \tag{91}
\]
Accordingly, from (90) we have that:
\[
P[N_P e_{ij} > \epsilon, D'_{\text{small}}] \leq P\left[ N_P \frac{\rho^{r_N+4}}{1 - \rho^2} > \epsilon \right], \tag{91}
\]
where we remark that the event appearing in the latter probability is in fact a deterministic event. If we now find a sequence \( r_N \) that drives to zero the quantity \( N_P \rho^{r_N+4} \), then, for sufficiently large \( N \), the probabilities appearing in (91) are eventually zero. We will illustrate how to make
a proper selection of \( r_N \) in the final step (i.e., Step 5) of this proof. It suffices for now to assume that such a sequence exists, namely, that:

\[
N p_N N^{r_N+4} \xrightarrow{N \to \infty} 0
\]

(92)

which, in view of (91), yields the desired claim in (83). Therefore, we conclude that, if nodes in \( S' \) forming active pairs, i.e., obeying (71), lie sufficiently far apart on the graph \( G_{S' \to S} \), then the magnified error can be driven to zero as \( N \to \infty \). This observation corroborates the claim that large distances imply small values of the error. In light of (80), the claim of Theorem 1 will be proved if we show that the occurrence of a small distance on at least one active pair (\( \ell, m \)) is a rare event, namely, if we prove the claim in (82). We will address this challenge in the forthcoming steps.

**Step 3: Relating partial Erdős-Rényi to a standard Erdős-Rényi via homogenization.** Two sources of asymmetry make the proof of (82) challenging. First, we see that the events in (76) refer to different graphs, namely, \( G_{S' \to S} \) and \( G \). The proof of (82) challenging. Further, the graph \( \tilde{G} \) in (76) refers to different graphs, namely, \( G_{S' \to S} \) and \( G \) – the graph \( G \) for the neighborhood constraint and \( G_{S' \to S} \) for the distance constraint. Second, both the local disconnection implied by \( G_{S' \to S} \) and the partial Erdős-Rényi construction implied by \( G \) (recall Figure 6), introduce additional non-homogeneity across nodes that makes the estimation of the probability of the event \( D_{\ell m} \) in (76), and hence the estimation of the probability in (82), rather intricate.

In order to overcome this issue, Theorem 3 further ahead states that, without loss of generality, we can replace the events \( D_{\ell m} \) in (76), by the events

\[
\tilde{D}_{\ell m} \triangleq \{ \delta_{\ell m}(\tilde{G}) \leq r_N, \ell \in N_i(\tilde{G}), m \in N_j^2(\tilde{G}) \},
\]

(93)

where \( \tilde{G} \sim \mathscr{G}^*(N, \tilde{p}_N) \) is a standard Erdős-Rényi graph with \( \tilde{p}_N =Sp_N \) (for sufficiently large \( N \)), in that if we prove the convergence

\[
P[\tilde{D}_{\text{small}}] \to 0,
\]

(94)

then, the convergence in (82) holds, where we have defined

\[
D_{\text{small}} \triangleq \bigcup_{\ell, m \in S'} \tilde{D}_{\ell m}.
\]

(95)

This is accomplished, in the proof of Theorem 3, via constructing a graph \( \tilde{G} \) that is Erdős-Rényi and that is coupled with \( G \) in the sense that

\[
D_{\ell m} \subseteq \tilde{D}_{\ell m}
\]

(96)

for all \( \ell, m \in S' \) and hence,

\[
D_{\text{small}} \subseteq \tilde{D}_{\text{small}}.
\]

(97)

Therefore, the induced coupling yields:

\[
P[D_{\text{small}}] \leq P[\tilde{D}_{\text{small}}],
\]

(98)

implying that if one is able to prove that the probability for the homogeneous case vanishes, equation (94), so does the probability for the original (non-homogeneous) case. We refer to this coupling procedure as homogenization as the conditions characterizing the event \( D_{\text{small}} \) refer to the same graph \( \tilde{G} \) and, further the graph \( \tilde{G} \) is a standard Erdős-Rényi. That is, the inhomogeneity that characterizes the events \( D_{\text{small}} \) is not present in the new event \( \tilde{D}_{\text{small}} \).

We now state Theorem 3 and prove it in Appendix D.

**Theorem 3 (Coupling and homogenization).** Let \( G \sim \mathscr{G}^*(N, p_N; G_S) \) be a partial Erdős-Rényi random graph, and let \( \tilde{G} \) be a pure Erdős-Rényi random graph \( \tilde{G} \sim \mathscr{G}(N, \tilde{p}_N) \) where, for \( N \) sufficiently large:

\[
\tilde{p}_N =Sp_N = \frac{\log N + c_N}{N}.
\]

(99)

If \( i, j \in S \) are non-interacting (\( g_{ij} = 0 \)), then we have that:

\[
P[D_{\text{small}}] \leq P[\tilde{D}_{\text{small}}]
\]

(100)

**Step 4: Managing the small-distance pairs.** The final step to prove Theorem 1, in view of inequality (100), consists in proving that (82) holds true on the homogenized graph \( \tilde{G} \sim \mathscr{G}^*(N, \tilde{p}_N) \), namely, that:

\[
P[\tilde{D}_{\text{small}}] \xrightarrow{N \to \infty} 0.
\]

(101)

Using (93) and (95), we observe that:

\[
\tilde{D}_{\text{small}} \subseteq \{ \delta_{ij}(\tilde{G}) \leq r_N + 3 \},
\]

(102)

i.e., if the distance between \( \ell, m \) of an active pair (\( \ell, m \)) is bounded by \( r_N \), then, since \( \ell \) and \( m \) are neighbor and second order neighbor of \( i \) and \( j \), respectively, the distance between the nodes \( i \) and \( j \) cannot exceed \( r_N + 3 \). Refer to Figure 14 for an illustration. Therefore,

\[
P[\tilde{D}_{\text{small}}] \leq P[\delta_{ij}(\tilde{G}) \leq r_N + 3],
\]

(103)

and the estimation of \( P[\delta_{ij}(\tilde{G}) \leq r_N + 3] \) amounts to a standard analysis of distance scaling on Erdős-Rényi random graphs as the graph \( \tilde{G} \) is a pure Erdős-Rényi. In fact, Lemma 3 (included in Appendix E for completeness) asserts that

\[
P[\delta_{ij}(\tilde{G}) \leq r_N + 3] \leq \tilde{p}_N (N\tilde{p}_N)^{r_N+2} \left( \frac{1}{1 - 1/(N\tilde{p}_N)} \right).
\]

(104)
Now, since by assumption \( r_N \to \infty \) and in the Erdős-Rényi regime that we are assuming we have \( Np_N \to \infty \), as \( N \) grows large, we conclude from (104) that \( \mathbb{P}[D_{\text{small}}] \) vanishes if we are able to choose a sequence \( r_N \) yielding:

\[
\bar{p}_N(Np_N)^{\rho+2} N^{\to \infty} \to 0
\]  

(105)

Note that the requirement (92) implies that \( r_N \) cannot diverge too slow, whereas the requirement in (105) implies that \( r_N \) cannot diverge too fast. The next step illustrates how to choose a sequence \( r_N \), with \( r_N \to \infty \), yielding both (92) and (105).

**Remark 2** (More on homogenization). Ultimately, the homogenization in Theorem 3 reduces the estimation of \( \mathbb{P}[D_{\text{small}}] \) to a simple analysis of distance scaling between only one pair of nodes \((i, j)\) in a pure Erdős-Rényi random graph, in view of the subset inequalities (97) and (102). Note that, to prove the convergence (82), one may be tempted to directly apply the following inequality (instead of invoking the extra homogenization, inequality (97), granted by Theorem 3)

\[
\mathcal{D}_{\text{small}} \subseteq \{\delta_{ij}(G) \leq r_N + 3\},
\]

(106)

for the original heterogeneous event \( D_{\text{small}} \), and with \( G \) in the RHS instead of the pure Erdős-Rényi \( \mathcal{G} \). But the probability \( \mathbb{P}[\{\delta_{ij}(G) \leq r_N + 3\}] \) does not converge to zero as \( N \) grows large in this case as the distance \( \delta_{ij}(G) \) depends on \( G_S \) which is arbitrary. Therefore, the inequality (106) is not useful to establish the convergence (82). In this line of thought, one can attempt to find a reference graph \( G_{\text{ref}} \), if any, so that

\[
\mathcal{D}_{\text{small}} \subseteq \{\delta_{ij}(G_{\text{ref}}) \leq r_N + 3\},
\]

and for which \( \mathbb{P}[\{\delta_{ij}(G_{\text{ref}}) \leq r_N + 3\}] \) converges to zero. Another natural candidate is \( G_{\text{ref}} := G_{S_{\text{ref}}} \), but the referred inequality does not hold in this case, i.e.,

\[
\mathcal{D}_{\text{small}} \nsubseteq \{\delta_{ij}(G_{S_{\text{ref}}}) \leq r_N + 3\}.
\]

(108)

Refer to Figure 15 for a graphical counter-example on this. One can attempt to simply fill in the gap of \( G_{S_{\text{ref}}} \) by considering the reference graph

\[
G_{\text{ref}} := G_S \oplus G_{S_{\text{ref}}}
\]

(109)

where \( G_S \sim \mathcal{G}(S, \bar{p}_N) \), but this also does not work as the inequality (107) is not satisfied (for the same reason as \( G_{S_{\text{ref}}} \) fails to meet it).

The homogenization in Theorem 3 provides a careful construction of a reference graph that allows us to lip this difficulty and rigorously reduce the computation of \( \mathbb{P}[D_{\text{small}}] \) to a simple distance-scaling of a particular pair of nodes in an Erdős-Rényi random graph. Such construction may be of independent interest.

**Step 5: Choosing the sequence \( r_N \).** In the above steps (specifically, in Step 1 and Step 4), we have maintained that a certain sequence of distances, \( r_N \), exists that fulfills the two conditions in (92) and (105). We start by examining these conditions in more detail. Taking the logarithm of the functions appearing in (92), we get:

\[
\log \left( Np_N^\rho N^{\rho + 4} \right) = \log(\log N + c_N) + (r_N + 4) \log(\rho)
\]

where we have used the expression for \( p_N \) in (9). Observing that \( \log(\rho) < 0 \) since \( \rho < 1 \), and letting:

\[
\alpha \triangleq |\log(\rho)|, \quad \omega_N \triangleq \log(\log N + c_N),
\]

from (110) we can write:

\[
\omega_N - \alpha r_N - 4\alpha N^{\to \infty} \to -\infty
\]

(112)

Let us switch to the analysis of (105). Taking the logarithm of the functions appearing in (105) we can write:

\[
\log \left( \bar{p}_N(Np_N)^{\rho+2} K^{N+3} \right) = \log \left( \frac{(Np_N)^{\rho+3}}{N} \right) = (r_N + 3) \log(Np_N) - \log N
\]

which, since \( \bar{p}_N = Sp_N \), in view of (9) and (111), yields:

\[
(r_N + 3)[\log S + \omega_N] - \log N, N^{\to \infty} \to -\infty
\]

(114)

We first show why the assumption in (40) is related to (112) and (114). From (112) and (114) we conclude that, for sufficiently large \( N \), we must necessarily have:

\[
\frac{\omega_N}{\alpha} < r_N < \frac{\log N}{\omega_N}
\]

(115)

which in turn implies:

\[
\frac{\omega_N^2}{\log N} < \alpha \iff \frac{[\log(\log N + c_N)]^2}{\log N} < \log(\rho),
\]

(116)

where in the last step we used the definitions in (111). Now, if we want to guarantee the verification of (116) irrespectively of the particular value of \( 0 < \rho < 1 \), we need to enforce condition (40).

Next we illustrate how to choose a sequence \( r_N \) that, under assumption (40), fulfills simultaneously (112) and (114). We set:

\[
r_N = \left\lfloor \frac{1}{2} \log N \right\rfloor.
\]

(117)
Substituting (117) into (112), and observing that \(|x| > x - 1\), where \(|x|\) stands for the greatest integer smaller than or equal to \(x\), we can write:

\[
\omega_N - a \left( \frac{1}{2} \frac{\log N}{\omega_N} \right) - 4a < \omega_N - a \frac{\log N}{\omega_N} + a - 4a = \omega_N \left( 1 - \frac{a \log N}{2 \omega_N^2} - \frac{3a}{\omega_N} \right) \rightarrow \infty \text{ from (40)}
\]

\[
N \rightarrow \infty \rightarrow - \infty, \quad (118)
\]

which shows that the condition in (112) is met with the choice in (117).

Likewise, substituting (117) into (114), and observing that \(|x| \leq x\), we have:

\[
\left( \frac{1}{2} \frac{\log N}{\omega_N} \right) + 3 \left( \log S + \omega_N \right) - \log N \leq \left( \frac{1}{2} \frac{\log N}{\omega_N} + 3 \right) \left( \log S + \omega_N \right) - \log N = \frac{\log N}{\omega_N} \left[ \log S + 3 \log S + \frac{\omega_N}{2} \log N + 3 \omega_N - \log N \right] = \frac{\log N}{\omega_N} \left[ \log S + 3 \log S + \frac{\omega_N}{2} \log N - \frac{\omega_N}{2} + 3 \frac{\omega_N^2}{2} \log N \right]
\]

\[
N \rightarrow \infty \rightarrow - \infty, \quad (119)
\]

with the convergence holding true because \(\omega_N \rightarrow \infty\) as \(N \rightarrow \infty\), while \(\omega_N^2 / \log N\) and \(\omega_N / \log N\) vanish in view of (40).

We have in fact shown that the condition in (114) is met with the choice in (117).

Refer to Figure 16 for a summary of the proof of Theorem 1.

**APPENDIX C**

**PROOF OF THEOREM 2**

First, we start by observing that the matrix \(H\) in (65) is not sensitive to the submatrix \(A_S\). Note that this is not immediately clear via simple inspection, since computation of \(H\) involves the matrix \(B = A^2\). However, note that

\[
A = \begin{bmatrix} A_S & A_{SS} \\ A_{SS} & A_S \end{bmatrix} \quad (120)
\]

and from the rules for block-matrix multiplication, we can write:

\[
B_S = [A^2]_S = A_{SS}A_{SS} + (A_S)^2 \quad (121)
\]

which highlights that the matrix \(B_S\) does not depend on the submatrix \(A_S\).

As a corollary to this observation, it follows that the matrix \(H\) in (65) is not a function of the particular submatrix \(A_S\), and, hence, it is insensitive to the particular topology of the subnetwork connecting the observed agents. Since we are to devise bounds for the terms \(h_{\ell m}\), we can assume without loss of generality that all the entries of \(A_S\) are equal to zero,\(^7\)

\(^7\)Assuming that the diagonal entries are equal to zero contradicts our rules for constructing a combination policy. However, this is inessential, because setting the \(A_S\) block to zero is only a mathematical expedient to compute suitable bounds, without any physical meaning.

namely, that \(A_S = 0_{S \times S}\). In other words, we can remove the edges among the observable agents as far as computing bounds on \(H\) goes. This will imply that an appropriate distance between nodes in \(S\) (which will play a role in bounding \(h_{\ell m}\)) is given by \(\delta_{\ell m}(G_{S \cup S})\), namely, by the distance between \(\ell\) and \(m\) on the graph \(G_{S \cup S}\) where the edges among the agents in \(S\) have been removed.

Theorem 2, proved next, establishes an exponential bound on \(h_{\ell m}\), which holds if \(\ell\) and \(m\) are distant on \(G_{S \cup S}\) (not necessarily on \(G\), and hence not dependent on \(G_S\)).

**Proof of Theorem 2:** We remind that:

\[
H = (I_{G_S} - B_S)^{-1} = \sum_{k=0}^{\infty} (B_S)^k, \quad (122)
\]

since \(A\) is row-sum stable and \(B = A^2\) (refer, e.g., to [31], [53]). Let now \(\tilde{A}\) be the matrix obtained from \(A\) by replacing the submatrix \(A_S\) with the void matrix, \(0_{S \times S}\), and let accordingly \(\tilde{B} = \tilde{A}^2\). Since, in view of (121), modifying the submatrix \(A_S\) does not alter the submatrix \(B_S\), we can safely write:

\[
B_S = \tilde{B}_S \quad (123)
\]

Moreover, it is known that, for any two nonnegative matrices \(Q\) and \(R\)

\[
[QR]_S = Q_{SS}R_{SS} + Q_{S}R_s \geq Q_{S}R_s, \quad (124)
\]

with entry-wise inequality. Taking \(Q = R = \tilde{B}\), and reasoning by induction, we have then:

\[
(B_S)^0 = (\tilde{B}_S)^0 \leq [\tilde{B}^n]_S = [\tilde{A}^2]^n_S, \quad (125)
\]

The first equality follows from (123). Rephrasing (125) on an entry-wise basis, we get, for all \(\ell, m \in S\):

\[
[(B_S)^n]_{\ell m} \leq [\tilde{A}^2^n]_{\ell m}. \quad (126)
\]

We recall that the support graph of \(\tilde{A}\) is given by \(G_{S \cup S}\). Since by assumption \(\delta_{\ell m}(G_{S \cup S}) = r\), then in view of (68), the smallest \(k\) yielding \([A^k]_S > 0\) is \(k = r\). In view of (126), this property implies that one can consider only the terms \([B_S^n]_{\ell m}\) for \(n \geq r\), where \(r\) is not necessarily an even number, we could in general consider all the terms for which \(n \geq [r/2]\), where \([x]\) stands for the smallest integer that is greater than or equal to \(x\). With this choice, the series in (122) can be truncated as (the term \(n = 0\) is zero because \(\ell \neq m\)):

\[
h_{\ell m} = \sum_{n=1}^{\infty} [(B_S)^n]_{\ell m} = \sum_{n=1}^{\infty} [(B_S)^n]_{\ell m}
\]

\[
\leq \sum_{n=[r/2]}^{\infty} [A^2^n]_{\ell m}
\]

\[
\leq \sum_{n=[r/2]}^{\infty} A_{[r/2]}^n \leq \frac{\rho^r}{1 - \rho^2} \leq \frac{\rho^r}{1 - \rho^2}, \quad (127)
\]

where inequality (a) follows by using (126) with \(A\) in place of \(\tilde{A}\); inequality (b) follows due to (31) as we know that \(\sum_{n=1}^{\infty} [A^2^n]_{\ell m} \leq \rho^2\).
modify the structure of the partial Erdős-Rényi $G$ in such a way that the resulting transformed graph $\tilde{G}$ fulfills the following properties: i) $\tilde{G}$ is a homogeneous (i.e., classic) Erdős-Rényi graph; ii) the original event $D_{\ell,m}$ on $G$ implies its counterpart $D_{\ell,m}$ defined on the new graph $\tilde{G}$, i.e.,

$$D_{\ell,m} \subseteq \tilde{D}_{\ell,m} \quad (132)$$

for all $\ell, m \in S'$, and hence,

$$D_{\text{small}} \subseteq \tilde{D}_{\text{small}}. \quad (133)$$

This further yields the desired inequality

$$P[D_{\text{small}}] \leq P[\tilde{D}_{\text{small}}]. \quad (134)$$

As a result, showing the convergence in (128) for the homogeneous system implies the convergence in (129) for the original heterogeneous partial Erdős-Rényi. We refer to this coupling procedure simply as homogenization.

**Remark 3.** It is tempting at first glance to simply replace the graph $G_{S \mapsto S}$ in (131) with the standard Erdős-Rényi $G$ by reinforcing the coupling $\tilde{G}_{S'} = G_{S'}$. This, in fact, contracts the distance, i.e., $\tilde{\delta}_{\ell,m}(G) \leq \delta_{\ell,m}(G_{S \mapsto S})$ which yields the implication

$$\tilde{\delta}_{\ell,m}(G_{S \mapsto S}) \leq r_N \Rightarrow \tilde{\delta}_{\ell,m}(\tilde{G}) \leq r_N \quad (135)$$

at the same time as the new graph $\tilde{G}$ is homogeneous, but the neighborhood constraint $m \in N_j^2(G)$ is jeopardized as it is not implied by its counterpart on $G$, since the condition $m \in N_j^2(G)$ also depends on $G_S$, which is not Erdős-Rényi, but arbitrary (hence, the subset inclusion (133) does not follow from this simple homogenization). The structure modification on the original graph $G$ is carefully performed in the proof of Theorem 3 to both grant the contraction of the distances at the same time as preserving the neighborhood constraints.

**Proof of Theorem 3:** In Figure 17, middle panel, we display the graph $G_{S \mapsto S}$. Moreover, we denote by $\tilde{S}_{ij}$ the set $S$ deprived of the nodes $i$ and $j$, namely, $\tilde{S}_{ij} \triangleq S \setminus \{i, j\}$. The
basic trick that allows homogenization is defining a new graph
where all the connections from \( \tilde{S}_{ij} \) to \( S' \) are inherited by the
node \( j \). The transformed graph is denoted by (and is displayed
in the rightmost panel of Figure 17):

\[
\mathcal{G} \triangleq (G_{S \rightarrow S})_{j \rightarrow \tilde{S}_{ij}} \tag{136}
\]

This operation achieves the twofold goal of ensuring that i) the
distance \( \delta_{\ell,m}(G_{S \rightarrow S}) \) between any two nodes \( \ell \) and \( m \) in
\( S' \) is reduced, namely,

\[
\delta_{\ell,m}(G_{S \rightarrow S}) \leq \delta_{\ell,m}(G_{S \rightarrow S}) \tag{137}
\]

and ii) if node \( m \) is second-order neighbor of \( j \) on the original
graph \( G \), so is on the transformed graph, namely,

\[
m \in N_j^{(2)}(G) \Rightarrow m \in N_j^{(2)}(\mathcal{G}) \tag{138}
\]

Note that equations (137) and (138) induce the desired cou-
pling between the transformed graph, \( \mathcal{G} \), and the graphs
\( G_{S \rightarrow S}, \ G \), in that for all \( \ell, m \in S' \):

\[
\begin{cases}
\delta_{\ell,m}(G_{S \rightarrow S}) \leq r_N, \ell \in N_i(G), m \in N_j^{(2)}(G) \\
\delta_{\ell,m}(\mathcal{G}) \leq r_N, \ell \in N_i(G), m \in N_j^{(2)}(\mathcal{G})
\end{cases}
\tag{139}
\]

At this point, we observe that \( \mathcal{G} \) is still not homogeneous (in
particular, the nodes in \( \tilde{S}_{ij} \) on the graph \( \mathcal{G} \) are isolated) and
hence, the proof is not finished. Before proceeding on this
point, we first justify equations (137) and (138).

The inequality (137) stems from the following observation. The
only modification in \( G_{S \rightarrow S} \) to get \( \mathcal{G} \) is related to \( \tilde{S}_{ij} \).
Therefore, if there exists a path from \( \ell \in S' \) to \( m \in S' \)
on \( G_{S \rightarrow S} \), which flows through \( \tilde{S}_{ij} \), such path (or a shortened
version thereof) is also present in \( \mathcal{G} \), but now via \( j \). Refer to
Figure 18 for an illustration.

Indeed, each path on \( G_{S \rightarrow S} \) hopping across \( \tilde{S}_{ij} \) is mapped
into a path traversing node \( j \) (instead of traversing the corre-
spounding nodes in \( \tilde{S}_{ij} \)), since the node \( j \) has inherited all
connections between \( \tilde{S}_{ij} \) and \( S' \).

The neighborhood implication (138) results from the follow-
ing observation. If on the graph \( G \), the node \( m \) is connected
to \( j \) through an intermediate node belonging to \( \tilde{S}_{ij} \), then it
is connected to \( j \) in one step on the graph \( \mathcal{G} \). One difficulty
might arise if, on graph \( G \), node \( m \) is connected to \( j \) through
node \( i \), because on \( \mathcal{G} \) nodes \( i \) and \( j \) are disconnected. This is
not a problem, however, because to prove our result we need
to examine only the case that \( i \) and \( j \) are disconnected on the
original graph \( G \) (as stated in the theorem). We remark that,
for the case that \( \ell = m \), condition (137) is redundant and

\[
\left\{ \ell \in N_i(G) \cap N_j^{(2)}(G) \right\} \subseteq \left\{ \ell \in N_i(\mathcal{G}) \cap N_j^{(2)}(\mathcal{G}) \right\}. \tag{140}
\]

Now, we return to the observation that the transformed
graph, \( \mathcal{G} \), is still asymmetrical, because, apart from the fact
that the set \( S \) contains disconnected nodes, the probability that
the node \( j \) is connected to a node in \( S' \) is now augmented as \( j \)
inherited all the connections from \( \tilde{S}_{ij} \) to \( S' \). Since under the
partial Erdős-Rényi construction, these connections follow a
standard Bernoulli law, we conclude that the probability of \( j \)
being connected to a particular node in \( S' \), in the new random
graph \( \mathcal{G} \), is simply given by (recall that \( \tilde{S}_{ij} \) does not contain
node \( i \)):

\[
1 - (1 - p_N)^{S-1}. \tag{141}
\]

To see why, \( j \) is not connected to a particular node in \( S' \), say
\( k \in S' \), in the graph \( \mathcal{G} \), if and only if, \( j \) is not connected

\[
\begin{cases}
\delta_{\ell,m}(G_{S \rightarrow S}) \leq r_N, \ell \in N_i(G), m \in N_j^{(2)}(G) \\
\delta_{\ell,m}(\mathcal{G}) \leq r_N, \ell \in N_i(G), m \in N_j^{(2)}(\mathcal{G})
\end{cases}
\tag{139}
\]
define a new (random) graph \( G \). Simple coupling between Bernoulli random variables, we can the implications shown in (139) and (140) hold true with wise. Since fleshing out a graph with new connections can only \( \delta_{\ell,m}(\tilde{G}) \leq \delta_{\ell,m}(G_{S \leftrightarrow S}) \)

Fig. 18. Illustration of a particular path (blue color) connecting \( t \) to \( m \) in the graph \( G_{S \leftrightarrow S} \) on the LHS of the figure. The edges linking to nodes in \( \bar{S}_{ij} \) on the graph \( G_{S \leftrightarrow S} \), link only to \( j \) on the graph \( \tilde{G} \). A path that crosses \( M \geq 1 \) nodes in \( \bar{S}_{ij} \) in the graph \( G_{S \leftrightarrow S} \), only crosses \( j \) in the new graph \( \tilde{G} \).

to \( k \) in \( G \), i.e., \( g_{jk} = 0 \), and \( g_{ak} = 0 \) for all \( a \in \bar{S}_{ij} \). In other words,

\[
P[G_{jk} = 0] = P[g_{jk} = 0, g_{ak} = 0 \forall a \in \bar{S}_{ij}] = (1 - p_N)^{S-1}. \quad (142)
\]

Hence, \( G_{jk} = 1 \) with probability \( p_N^\ast = 1 - (1 - p_N)^{S-1} \). But now homogenizing the transformed graph \( \tilde{G} \) is an easy task. It suffices to augment the connection probabilities of all the remaining pairs (including those in \( S \)), in order to match the connection probability in (141). More formally, resorting to a simple coupling between Bernoulli random variables, we can define a new (random) graph \( G^\ast \) from \( \tilde{G} \) as

\[
g_{uv}^\ast = \max \{ \tilde{g}_{uv}, q_{uv} \}
\]

where \( q_{uv} \) are i.i.d. Bernoulli random variables with

\[
P[q_{uv} = 1] = \begin{cases} 
1 - (1 - p_N)^{S-1}, & \text{if } u \in \bar{S}_{ij} \\
1 - (1 - p_N)^{S-1}, & \text{if } (u, v) = (i, j) \\
1 - (1 - p_N)^{S-2}, & \text{if } u \in \{i\} \cup S' \text{ and } v \in S', \\
0, & \text{if } u = j \text{ and } v \in S'.
\end{cases} \quad (144)
\]

The resulting graph \( G^\ast \) is Erdős-Rényi with \( p_N^\ast = 1 - (1 - p_N)^{S-1} \). Figure 19 graphically summarizes the idea. Moreover, since \( 1 - (1 - p_N)^{S-1} \leq S_W N \), and in order to obtain a random graph whose connection probability is explicitly given by (9), we can further define a graph \( \tilde{G} \) with connection probability given by

\[
\tilde{p} = S_W N \frac{\log N + c_N}{N} = \frac{\log N + c_N}{N}, \quad (145)
\]

where \( c_N = (S - 1) \log N + a_N \) and with the coupling \( g_{uv}^\ast \leq \tilde{g}_{uv} \) (realization-wise) for all \( u, v \). This can be easily obtained via a standard coupling between Bernoulli random variables. Therefore \( \tilde{G} \subseteq G \), i.e., \( \tilde{G} \) is a subgraph of \( G \), realization-wise. Since fleshing out a graph with new connections can only decrease distances and favor membership to any neighborhood, the implications shown in (139) and (140) hold true with \( \tilde{G} \) replaced by \( G \). This implies, in view of (93) and (95), that:

\[
D_{\text{small}} \subseteq \tilde{D}_{\text{small}}, \quad (146)
\]

which in turn implies the claim of the theorem.

**APPENDIX E**

**MANAGING SMALL DISTANCES**

**Lemma 3.** Let \( G \) be a pure Erdős-Rényi random graph \( G(N, p_N) \). We have that

\[
P[\delta_{i,j}(G) \leq r] \subseteq p_N^\ast = \frac{1}{1 - 1/(N p_N^\ast)} \left( \frac{1}{1 - (1 - p_N^\ast)^{S-1}} \right) \quad (147)
\]

**Proof:**

Since the event \( \delta_{i,j}(G) = r \) signifies that the shortest path connecting \( i \) to \( j \) has length equal to \( r \), there must exist a path connecting \( i \) to \( j \) obeying the following conditions: \( i \) all intermediate nodes are visited only once through the path (otherwise the path itself could be squeezed to one of a shorter length); \( ii \) along the path, one cannot spend one or more steps lingering on the same node. Accordingly, we can write:

\[
\{ \delta_{i,j}(G) = r \} \subseteq \bigcup_{M} \{ g_{i,n_1} g_{n_1, n_2} \cdots g_{n_{r-1}, j} = 1 \}, \quad (148)
\]

where the set \( M \) is defined as \( M \equiv M_1 \cap M_2 \) with

\[
M_1 \equiv \{ n := (n_1, \ldots, n_{r-1}) \in \mathbb{N}^{r-1} : n_u \neq n_v \forall u,v \}, \quad (149)
\]

\[
M_2 \equiv \{ n := (n_1, \ldots, n_{r-1}) \in \mathbb{N}^{r-1} : n_k \neq i, j \forall k \}. \quad (150)
\]

It is useful to remark that the event \( \mathcal{E} \) in (148) does not coincide with the event that the shortest path has length equal to \( r \), because the possibility of having paths longer than \( r \) is not ruled out. The event \( \mathcal{E} \) in (148) simply underlies the existence of at least one path of length \( r \) with the necessary characteristics, which explains the one-sided implication in (148), and
yields $P[\delta_{t,m}(G) = r] \leq P[\mathcal{E}]$. We have
\[
P[\mathcal{E}] \leq \sum_M \prod_{i=1}^M P[g_{i1} g_{i2} \cdots g_{ir} = j] = M^r p_N^r.
\]
where we recall that $M$ stands for the cardinality of the set $\mathcal{M}$ in view of the notation in Sec. II-A, where sets are represented by calligraphic letters and the corresponding cardinalities are represented by normal font letters. Observe that
\[
M = (N - 2)(N - 3) \ldots (N - r) \leq (N - 2)^{r-1}.
\]
(152)
Therefore,
\[
P[\delta_{t,j}(G) = r] \leq (N - 2)^{r-1} p_N^r \leq p_N (N p_N)^{r-1}
\]
(153)
and as a result,
\[
P[\delta_{t,j}(G) \leq r] = \sum_{a=1}^r P[\delta_{t,j}(G) = a] \leq p_n \sum_{a=1}^r (N p_N)^{a-1}
\]
\[
= p_n \sum_{a=0}^{r-1} (N p_N)^a \leq p_n (N p_N)^{r-1} \cdot \frac{1}{1 - 1/(N p_N)}.
\]
(154)
\]
\]

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