Learning Data Dependency with Communication Cost

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ABSTRACT
In this paper, we consider the problem of recovering a graph that represents the statistical data dependency among nodes for a set of data samples generated by nodes, which provides the basic structure to perform an inference task, such as MAP (maximum a posteriori). This problem is referred to as structure learning. When nodes are spatially separated in different locations, running an inference algorithm requires a non-negligible amount of message passing, incurring some communication cost. We inevitably have the trade-off between the accuracy of structure learning and the cost we need to pay to perform a given message-passing based inference task because the learnt edge structures of data dependency and physical connectivity graphs are often highly different. In this paper, we formalize this trade-off in an optimization problem which outputs the data dependency graph that jointly considers learning accuracy and message-passing cost. We focus on a distributed MAP as the target inference task due to its popularity, and consider two different implementations, ASYNC-MAP and SYNC-MAP that have different message-passing mechanisms and thus different cost structures. In ASYNC-MAP, we propose a polynomial time learning algorithm that is optimal, motivated by the problem of finding a maximum weight spanning tree. In SYNC-MAP, we first prove that it is NP-hard and propose a greedy heuristic. For both implementations, we then quantify how the probability that the resulting data graphs from those learning algorithms differ from the ideal data graph decays as the number of data samples grows, using the large deviation principle, where the decaying rate is characterized by some topological structures of both original data dependency and physical connectivity graphs as well as the degree of the trade-off, which provides some guideline on how many samples are necessary to obtain a certain learning accuracy. We validate our theoretical findings through extensive simulations, which confirm that it has a good match.

CCS CONCEPTS
• Computing methodologies → Learning in probabilistic graphical models; • Mathematics of computing → Probabilistic inference problems; • Theory of computation → Sample complexity and generalization bounds;

KEYWORDS
Graph structure learning, Distributed inference, Sample complexity

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1 INTRODUCTION
In many online/offline systems with spatially-separated agents (or nodes), a variety of applications involve distributed in-network statistical inference tasks, which have been widely studied, exploiting given knowledge of statistical dependencies among agents. As one example, in sensor networks with multiple targets, each sensor node measures the target-specific information in its coverage area (e.g., position, direction, distance), which further has a correlation among sensors. One well-recognized inference problem is a data association which determines the correct match between measurements of sensors and target tracks by maximum a posteriori (MAP) estimation that is executed in a distributed fashion by exchanging some information messages. Other examples include target tracking, and detection/estimation in sensor networks [23, 27, 29, 34] and de-anonymization, rumor/infection propagation in social networks [2, 11, 14, 19].

To solve these distributed in-network inference problems, it is of crucial importance to understand how data from nodes are inter-dependent. To that end, a notion of the graphical model has been one of the powerful frameworks in machine learning for a succinct modeling of the statistical uncertainty, where each node in the graphical models corresponds to a random variable and each edge specifies the statistical dependency between random variables. A wide variety of scalable inference algorithms on graphical models via message-passing have been developed, of which examples include belief propagation (BP) or max-product with a certain degree of convergence and accuracy guarantees [15, 24, 31, 32]. This graphical model, which we also call data dependency graph or simply data graph throughout this paper, is not given a priori, and it should be learnt only by using a given set of data samples from nodes. This problem, referred to as graph learning or structure learning [1, 13, 21, 25], has been an active research topic in statistical machine learning.

In this paper, for a collection of n data sample vectors generated by nodes, we study a problem of graph learning, which also considers the communication cost incurred by the distributed in-network inference algorithm being applied to the learnt data graph. Physical communication cost often becomes a critical issue, for example, exerting a significant impact on the lifetime of networked sensors. Clearly, there exists a trade-off between the amount of incurred cost...
and the learning accuracy of the data graph. Figure 1(a) illustrates the physical connection among 7 sensors, which differs from the exact data dependency graph in Figure 1(b). The sensor nodes s1 and s6 have non-negligible data dependency, requiring message-passing when performing inference, but they are three hops away from each other, incurring a large amount of communication cost. In this case, one may want to sacrifice the estimation accuracy a little bit and reduce communication cost by utilizing the data graph as shown in Figure 1(c). As done in many prior works on graph learning [3, 8, 9, 21], we restrict our attention to tree-structured data graphs due to its simplicity, yet a large degree of expressive powers and other benefits, e.g., some inference algorithms such as BP over tree-structured data graphs become optimal.

We now summarize our contributions in what follows:

(a) We first formulate an optimization problem of learning data graph, having as an objective function the weighted sum of learning accuracy and the amount of cost that will be incurred by a distributed inference algorithm. Out of many possible inference algorithms, we consider the maximum a posteriori (MAP) estimator that is popular for many inference tasks, and two versions for the MAP implementation: (i) asynchronous and (ii) synchronous, which we call \textsc{async-map} and \textsc{sync-map}. These implementations have different patterns of passing messages, thus leading to different forms of communication costs, being useful to understand how distributed algorithms’ cost affect the resulting data dependency graph.

(b) Next, for \textsc{async-map} we develop a polynomial-time algorithm to find an optimal (cost-efficient) data graph that corresponds to simply finding a maximum weight spanning tree. This simplicity stems from the cost structure of \textsc{async-map} that is characterized only by the sum of all ‘localized’ edge costs. Being in sharp contrast to \textsc{sync-map}, for \textsc{sync-map} we first prove that it is computationally intractable (i.e., NP-hard) in terms of the number of nodes, by reducing it to the problem of the Exact Cover by 3-sets. The hardness is due to the fact that the cost structure of \textsc{sync-map} depends on the diameter of the resulting tree which is the ‘global’ information involving the entire topology. As a practical solution, we propose a polynomial-time greedy heuristic to recover a sub-optimal, but cost-efficient data graph.

(c) Finally, for both \textsc{async-map} and \textsc{sync-map}, we quantify how the probability that the resulting (cost-efficient) data graph for a finite number of samples differs from the ideal data graph decays as n increases, using the large deviation principle (LDP), as a form of \(\exp(-n \cdot K)\). The error exponent \(K\) is characterized for each of \textsc{async-map} and \textsc{sync-map} by some topological information of physical/data graphs, cost structure for both inference mechanisms, and the degree of the trade-off.

To validate our theoretical results, we perform numerical simulations a pair of physical and data graphs with 20 nodes, where we quantitatively analyze (i) how estimating a data graph considering communication cost affects the resulting estimation for various values of trade-off parameters between inference accuracy and cost, (ii) how the estimation error decays as the same size increases.

1.1 Related Work

A variety of applications which involve distributed in-network statistical inference tasks among spatially inter-connected agents or sensors have been widely studied in many online/offline systems. In sensor networks, where the knowledge of statistical dependencies among sensed data is given, the tasks of target tracking [7, 22, 33], detection [6], parameter estimation [16, 27] are the examples, see [29] for a survey. In social networks, where the underlying social phenomenon of interest such as voting models, rumor/opinion propagation [2] evolves over a given social interaction graph, the inference tasks of distributed consensus-based estimation [19], de-anonymization of community-structured social network [14] and distributed observability [11] are studied. Message-passing has manifested as an efficient procedure for inference over graphical models that provide the framework of succinct model of the statistical uncertainty of multi-agents. Examples include belief propagation (BP) [24], max-product [15, 32] and references therein. They are known to be exact and efficient when the underlying graphical model is a tree [24, 31]. Recent research progress has been made for scalable message-passing for general graphs, e.g., junction tree [30] and graphs with loops [17].

In the area of structure learning, several algorithms have been proposed in the literatures to recover the statistical dependencies from a set of data samples [1, 13, 21, 25]. It is known that the exact structure learning for general graphical models is NP-hard. The research of structure learning for special graphical models includes: maximum likelihood estimation (MLE) [8, 21] for tree graphs, \(\ell_1\) regularized MLE for binary undirected graphs [25], convexified MLE for Gaussian graphical models, known as Lasso [13]. Theoretical guarantees for the learning accuracy have been established as the number of data samples, e.g., on tree graph [28], on binary undirected graphs [25], on a class of Ising model [4], or on Bayesian network [36]. Our work differs from all of the above works in that we consider physical communication cost incurred by some target inference algorithms when learning the data dependency graph.

There exists an array of work that addresses the trade-off between inference quality and cost in running distributed in-network inference on the known data graph, which are summarized as two directions: (i) developing novel inference algorithms with less communication of messages or (ii) constructing a new graphical model upon which the existing distributed in-network inference algorithms are performed with less communication resources.
the need of conserving resources requires to propose new message-passing schemes where the messages are compressed by allowing some approximation error in message values [10, 17, 20, 22], and/or some messages are censored (i.e., not to be transmitted) [7]. In (ii), most of the related works focused on constructing a junction tree that minimizes the inference cost [23], building a data dependency structure upon which message-passing is run energy-efficiently, where the communications among all agents are assumed to be done in one-hop [34], or optimizing the data dependency structure formulated by a multi-objective problem of inference quality and energy, assuming that the exact statistical dependencies are given as a complete graph [35]. While the main interest of this area has been focused on characterizing the desirable dependency structure for given complete knowledge of accurate data dependencies, our work is motivated by the practical situation where one can just be able to observe a finite number of data sample vectors of nodes, which do not provide such a complete knowledge. Therefore, our interest lies in learning the desirable data dependencies from a finite number of data samples.

2 MODEL AND PRELIMINARY

2.1 Model

Physical graph. We consider a (connected) physical network \(G = (V, E_p)\) with a set of \(d\) nodes \(V\) and links \(E_p\), where each node corresponds to an agent such as a sensor or an individual, and each link corresponds to a physical connectivity between two nodes. For example, in sensor networks, when nodes have wireless radios, then each link is established when two corresponding nodes over the link reach each other within each radio’s communication range.

Data samples. Each node \(i \in V\) generates a binary data, denoted by \(x_i \in X := \{0, 1\}\)\(^1\), where we denote by \(x = [x_i]_{i \in V}\) the data vector of all nodes, or simply a sample, e.g., target locations measured by all sensors. The underlying statistical uncertainty of samples can be represented by a joint distribution \(P(x)\) of a random vector \(X := [x_i]_{i \in V} \in X^d\), called data distribution, where each random variable \(X_i\) is associated to each node \(i \in V\). We often collect \(n\) multiple samples \((x^{1:n} = (x^1, x^2, \ldots, x^n))\) in order to infer what happens in the network by understanding the inter-dependence of data generated by nodes. For instance, when sensors measure the target location, then we infer the underlying statistical correlation among sensors from the observed samples, to estimate the true target location.

Data graph via graphical model. The underlying statistical dependency is often understood by the framework of graphical model, which has been a popular tool for modeling uncertainty by a graph structure, where each node corresponds to a random variable and each edge captures the probabilistic interaction between nodes. In particular, we model the data distribution \(P(x)\) as an undirected graph \(T = (V, E_p)\), which we call data graph, which consists of the same set \(V\) of nodes as that in the physical graph and nodes’ statistical dependencies captured by an edge structure \(E_p\) as: any two non-adjacent random variables are conditionally independent given all other variables, i.e., for any \((i, j) \notin E_p\),

\[
P(x_i, x_j | x_{V \setminus \{i,j\}}) = P(x_i | x_{V \setminus \{i,j\}}) \cdot P(x_j | x_{V \setminus \{i,j\}}).
\]

In this paper, we limit our focus on the tree-structured data graph (thus simply data tree), for which let \(T\) and \(P(X^d)\) be set of all spanning trees and set of all tree data distributions over \(V\), respectively, i.e., we assume \(T \in T\) and \(P \in P(X^d)\). Tree data graph is a class of graphical models that has received considerable attention in literatures [3, 8], since it possesses the following factorization property:

\[
P(x) = \prod_{i \in V} P_i(x_i) \prod_{(i, j) \in E} \frac{P_{i, j}(x_i, x_j)}{P_i(x_i)P_j(x_j)}.
\]

where \(P_i\) and \(P_{i, j}\) are the marginals on node \(i \in V\) and edge \((i, j) \in E_p\), respectively. Tree-structured data graph is known to strike a good balance between the expressive power and the computational tractability. In particular, the distribution \(P\) in (2) is completely specified only by the set of edges \(E_p\) and their pairwise marginals. Thus, if \(P\) has the factorization property as in (2), in other words, if \(P \in P(X^d)\), there exists a unique tree \(T = T(P)\) corresponding to \(P\). To abuse the notation, we henceforth denote by \(T(P)\) the unique data tree of a tree distribution \(P\). Figure 1 shows an example of the physical graph and two data graphs with 7 nodes.

2.2 Goal: Cost-efficient Learning of Data Graph

Learning data graph: What and why? To understand the underlying data dependency (2), it is enough to learn the structure of data graph \(E_D\) from the observed samples, which is known as the problem of (data graph) structure learning. Formally, when we are given a set of i.i.d. \(n\) samples \(x^{1:n}\) generated from an unknown (tree) data distribution \(P \in P(X^d)\) on a data tree \(T\), a structure learning algorithm is a (possibly randomized) map \(\phi\) defined by:

\[
\phi : (X^d)^n \rightarrow T.
\]

The quality of this algorithm \(\hat{T} = \phi(x^{1:n})\) is evaluated by how “close” \(\hat{T}\) is to the original data graph \(T\).

Distributed inference on data graph. One of the practical goals of estimating the data tree given a set of data samples is to perform an inference task based on \(T\). Thus, in many applications, primary interests are not focused on data itself but rather on how to exploit the data dependency for reliable decision making, such as target tracking, detection, estimation in sensor networks and/or social networks, which involves statistical inference about the networks described by a data graph. One example of inference tasks is the MAP (maximum a posteriori) based estimation. Distributed network inference has been widely studied with the help of various distributed algorithms on graphical models using message-passing. In particular, for a specific inference problem, a message between two nodes contains the information on influence that one node exerts on another, which is obtained based on the value contained in neighboring messages over an estimated data graph \(\hat{T}\). One critical issue of message-passing based inference algorithm is that messages are often passed along the multi-hop path on the physical graph \(G\), which incurs some amount of communication cost. Then, assuming that some inference algorithm would be run for the estimated data graph \(\hat{T}\), such a data graph learning must have the trade-off

\(^1\)We assume a binary data for simplicity, and our results are readily extended to any finite set \(X\).
between the accuracy of the learnt graph (i.e., how close the learnt graph is to the original data graph) and the communication cost generated by performing the distributed inference.

**Goal: Cost-efficient data graph learning.** Given an observed samples \( x^{1:n} \) from the unknown data distribution \( P \), our objective is to estimate a cost-efficient data tree, which captures the trade-off between (i) inference accuracy and (ii) communication cost for inference. For tree distributions, finding a distribution naturally gives rise to the corresponding data tree, as mentioned earlier. Thus, it is natural to find the tree distribution \( \hat{Q}^u(n) = \hat{Q}^u(x^{1:n}, G_i, \Pi, \gamma) \) that is the solution of the following optimization problem: for a constant parameter \( \gamma \geq 0 \) and a fixed inference algorithm \( \Pi \),

\[
\text{CDG}(n) \quad \hat{Q}^u(n) = \arg \min_{Q \in \mathcal{P}(X^d)} D(\hat{P}(x^{1:n}) || Q) + \gamma C(T(Q);G_i,\Pi),
\]

where \( \hat{P}(x^{1:n}) := \frac{1}{n} \sum_{k=1}^n \mathbb{1}\{x^k = x\} \) is the empirical distribution of \( x^{1:n} \), \( D(\cdot || \cdot) \) is some distance metric between two distributions, and \( C(T(Q);G_i,\Pi) \) is the communication cost paid by running an inference algorithm \( \Pi \) with respect to the data tree \( T(Q) \) over the physical graph \( G_i \). Recall that \( T(Q) \) is the data tree for the tree distribution \( Q \). The value of \( \gamma \) parameterizes how much we prioritize the communication cost compared to the inference accuracy \( D(\hat{P}(x^{1:n}) || Q) \). Note that as \( n \to \infty \), \( \hat{P}(x^{1:n}) \) converges to the original data distribution \( P \), which requires to solve \( \text{CDG}(\infty) \).

Then, this paper aims at answering the following two questions:

(a) What are good data-tree learning algorithms that compute \( T(\hat{Q}^u(n)) \) by solving \( \text{CDG}(n) \)? In Section 3, we consider the MAP estimator as an applied inference algorithm, and their two implementations having different cost functions, for which we propose two cost-efficient learning algorithms.

(b) How fast does \( \hat{Q}^u(n) \) converge to \( \hat{Q}^u(\infty) \) as the number of samples \( n \) grows? We use the large deviation principle (LDP) to characterize the decaying rate of the probability that \( T(\hat{Q}^u(n)) \neq T(\hat{Q}^u(\infty)) \) for two different learning algorithms in Section 3.

In this paper, we use the popular KL divergence as a distance metric \( D(\cdot || \cdot) \) for inference accuracy, denoted by \( D_{KL} \), where for two distributions \( P \) and \( Q \), \( D_{KL}(P || Q) := \sum_{x \in X^d} P(x) \log \frac{P(x)}{Q(x)} \).

For notational simplicity, we simply denote by \( \hat{Q}^u := \hat{Q}^u(\infty) \) the solution of \( \text{CDG}(\infty) \) throughout this paper.

### 3 COST-EFFICIENT DATA GRAPH LEARNING ALGORITHMS

In this paper, out of many possible inference tasks, we consider the maximum a posteriori (MAP) estimation, which is popularly applied in many applications such as data association for a multitarget tracking problem in sensor networks, community-structured social network de-anonymization problem in social networks [14].

#### 3.1 Distributed MAP and Cost

**Distributed MAP on tree-structured data graph.** The MAP estimator of some tree distribution \( Q \in \mathcal{P}(X^d) \) on its associated data tree \( T(Q) = (V,E_Q) \) is given by:

\[
\hat{x}^{\text{MAP}} := \arg \max_{x \in X^d} T(Q;G_i,\Pi) = \prod_{i \in V} \psi_i(x_i) \prod_{(i,j) \in E_Q} \psi_{ij}(x_i,x_j),
\]

where we use \( \psi_i(x_i) = Q_i(x_i) \) and \( \psi_{ij}(x_i,x_j) = \frac{Q(x_i|x_j,x_j)}{Q_i(x_i)Q_j(x_j)} \) for simplicity. A standard message-passing algorithm for the distributed MAP is a max-product algorithm, which defines a message \( m_{i \rightarrow j}^{(t)}(\cdot) \) from node \( i \) to \( j \) at \( t \)-th iteration with \( (i,j) \in E_Q \). Each node exchanges messages with its neighbors on the data tree \( T(Q) \), and these messages are updated over time in an iterative fashion by the following rule: at \( t \)-th iteration,

\[
m_{i \rightarrow j}^{(t+1)}(x_j) := k \cdot \max_{x_i \in X} \left[ \psi_i(x_i) \psi_{ij}(x_i,x_j) \prod_{k \in N(i) \setminus \{j\}} m_{k \rightarrow i}^{(t)}(x_j) \right],
\]

with the normalizing constant \( k \) to make the sum of all message values be 1, and \( N(i) \) denotes the neighboring nodes of \( i \).

**Communication cost of distributed MAP.** The communication cost of MAP is paid, depending on the actual protocol that specifies how to schedule message-passing procedures. Two natural message-passing protocols studied in literatures are: (a) asynchronous depth-first (unicast) update [12] and (b) synchronous (broadcast) parallel update [24]. Both protocols for a tree distribution \( Q \) with its data tree \( T(Q) \) have been shown to be consistent in that the message update (5) converges to a unique fixed point \( \{m_{i \rightarrow j}^{(t)}\}_{(i,j) \in E_Q} \) for two different learning algorithms for \( \text{CDG}(n) \) in (3) to estimate two cost-efficient data trees.

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\(^2\)For a tree \( T \) with \( d \) nodes, \( 2 \leq \text{diam}(T) \leq d - 1 \).
Algorithm 1: ASYNC-ALGO

Input: $x^{1:n}$, a set of $n$ samples, $\gamma$: the trade-off parameter, a physical graph $G = (V, E_p)$
Output: Estimated tree $T = (V, E)$

S0. $E = \emptyset$ and for each possible edge $e = (i, j) \in V \times V$, we initialize its weight by:

$$w_e = I_e(\hat{P}) - 2\gamma \cdot c_e,$$

(9)

where $I_e(\mu)$ is the mutual information between two end-points of edge $e$ with respect to a given joint distribution $\mu$.

S1. Run a maximum weight spanning tree algorithm for $H$, and save its resulting spanning tree at $T = (V, E)$.

S2. Return $T$.

3.2 Algorithm for Asynchronous MAP

Using the cost function for the asynchronous MAP in (6), the original optimization problem CDG(n) is re-cast into:

$$\text{CDG-A}(n) : \hat{Q}^*(n) = \arg \min_{Q \in \mathcal{P}(X^n)} D_{KL} (\hat{P}(x^{1:n}) \parallel Q) + \gamma \sum_{e \in E_{Q}} 2c_e.$$  

(8)

We now describe ASYNC-ALGO that computes $\hat{Q}^*(n)$ in (8) and thus estimates the cost-efficient data tree $T(\hat{Q}^*(n))$ in Algorithm 1. As we see, the algorithm is remarkably simple. Using given $n$ data samples, we construct a weighted complete graph, where the weight for each edge is assigned some combination of the mutual information of nodes $i$ and $j$ with respect to the empirical distribution $\hat{P}$ obtained from the data samples and the per-message cost, as in (9). Then, we run an algorithm that computes the maximum weight spanning tree, e.g., Prim’s algorithm or Kruskal’s algorithm, and the resulting spanning tree is the output of this algorithm.

Correctness of ASYNC-ALGO. We now present the correctness of the above algorithm in the sense that we can obtain the data tree corresponding to the optimal distribution formulated in (8), as explained in what follows: For some tree distribution $Q$ (thus, satisfying the factorization property in (2)), we have:

$$D_{KL}(\hat{P} \parallel Q) = -H(\hat{P}) - \sum_{x \in X^d} \hat{P}(x) \log Q(x)$$

$$\geq -H(\hat{P}) + \sum_{i \in V} H(\hat{P}_i) - \sum_{(i,j) \in E_Q} I(\hat{P}_{i,j}),$$

(10)

where $H(\cdot)$ is the entropy, and the inequality holds when the pairwise marginals over the edges of a fixed $E_Q$ are set to that of $\hat{P}$, i.e., $Q_{i,j}(x_i, x_j) = \hat{P}_{i,j}(x_i, x_j)$ for all $(i, j) \in E_Q$. Since the entropy terms are constant w.r.t. $Q$, it is straightforward that the structure of the estimator $\hat{Q}^*(n)$ of CDG-A(n) in (8) is given by:

$$\hat{E}^*(n) := \arg \max_{E_Q \subseteq \mathcal{P}(X^d)} \sum_{e \in E_Q} I_e(\hat{P}) - 2\gamma \cdot c_e,$$

(11)

$$\hat{Q}^*_{i,j}(n) = \hat{P}_{i,j}, \quad \forall (i,j) \in \hat{E}^*(n).$$

(12)

Then, it is easy to see that (11) requires us to find the maximum weight spanning tree using $I_e(\hat{P}) - 2\gamma \cdot c_e$ as the edge $e$’s weight, where the standard maximum weight spanning tree (MWST) computation algorithm runs in $O(d^2 \log d)$ time, where $|V| = d$.

3.3 Algorithm for Synchronous MAP

Similarly to ASYNC-MAP, using the cost in (7), the original optimization problem CDG(n) is re-cast into:

$$\text{CDG-S}(n) : \hat{Q}^*(n) = \min_{Q \in \mathcal{P}(X^n)} D_{KL} (\hat{P}(x^{1:n}) \parallel Q) + \gamma \cdot \text{diam}(T(Q)) \sum_{e \in E_Q} 2c_e.$$  

(13)

Following the similar arguments in Section 3.2, the structure of the above estimator of CDG-S(n) in (13) is given by

$$\hat{E}^*(n) := \arg \max_{E_Q \subseteq \mathcal{P}(X^d)} \sum_{e \in E_Q} I_e(\hat{P}) - 2\gamma \cdot \text{diam}(T(Q)) \cdot c_e,$$

(14)

$$\hat{Q}^*_{i,j}(n) = \hat{P}_{i,j}, \quad \forall (i,j) \in \hat{E}^*(n).$$

(15)

We comment that this optimization is non-trivial in that the objective function contains the diameter of the tree, which can be computed only when the solution is fully characterized.

Hardness. The key difference in the cost function of SYNC-MAP from ASYNC-MAP is simply the existence of $\text{diam}(T(Q))$. However, this simple difference completely changes the hardness of learning the optimal data tree in SYNC-MAP, as formally stated in the next theorem.

Theorem 3.1 (Hardness of CDG-S(n)). For any parameter $\gamma \geq 0$, obtaining the optimal distribution $\hat{Q}^*(n)$ in CDG-S(n) and thus its associated data tree $T(\hat{Q}^*(n))$ is NP-hard with respect to the number of nodes.

Due to space limitation, we present the full proof of Theorem 3.1 in our technical report [18]. The key step in proof is to reduce the CDG-S(n) in (13) to Exact Cover by 3-sets problem.

Greedy algorithm. Due to the above-mentioned hardness, we propose a greedy heuristic algorithm that outputs the tree structure denoted by $\hat{E}^*(n)$, called SYNC-ALGO($\beta$), as we describe in Algorithm 2, where $\beta$ is the algorithm parameter. The overall algorithm operates as follows:

S0. Initialize the weight of each possible edge with some initial value.

S1. Sequentially select the edge that has the maximum weight and add it to the temporary resulting tree.

S2. Update the weight of each edge whose one end-point is in the current resulting tree $V_S$ and another end-point is not, and go to S1 until we handle all nodes.

One of the central steps here is: first, we dynamically update the weight of the candidate edges (i.e., the set $E'$) that we will add and, second, which value is chosen as the weight is different from the “one-shot” weight assignment as done in ASYNC-ALGO. To see this intuition, we first note that from (14) it is easy to see that the degree of contribution in terms of weight by adding an edge $e \in E'$ to the existing resulting tree would be re-expressed as:

$$I_e(\hat{P}) - 2\gamma \cdot \text{diam}(T \cup \{e\}) \cdot c_e - K_e(T),$$

(20)

where $K_e(T)$ is defined in (18). Here, $K_e(T)$ corresponds to the change of the communication cost over the existing edges in $T$, ...
under the grown tree $T \cup \{e\}$. For example, $K_e(T) = 0$ if the diameter of the grown tree does not change by adding the edge $e$, or $K_e(T) = \sum_{e' \in E} 2\gamma \cdot c_{e'}$ if the diameter of the grown tree increases by 1.

In dynamically assigning the weight of the candidate edges in $E'$, we do not use the value of (20). Instead, as seen in (17), (i) we use the expected diameter growth of the tree, denoted by $D(T)$ in (19), and (ii) we use a tunable parameter $\beta > 0$ to compensate for the impact of the expected diameter growth of the tree and the tunable parameter $\beta$.

To summarize, these two modified choices of the weight are for handling a probable sacrifice of the performance when using a vanilla greedy method as in (18), since the edge weight should be modified suitably for the changed diameter on the way of tree construction. We expect that these two engineerings play an important role when the cost-efficient data graph is attained with a large diameter, where the edges chosen in the begging phase of the procedure (i.e., with a small diameter value) could exert much impact of communication cost at the end of the procedure. Our greedy algorithm runs in $O(d^2)$ times.

4 ESTIMATION ERROR FOR INCREASING SAMPLE SIZE

In this section, we provide the analysis of how the estimation error probability decays with the growing number of samples $n$, using the large deviation principle (LDP).

4.1 Estimation Error of ASYNC-ALGO

Clearly, when we use more and more data samples, $\hat{E}(n)$ approaches to $\hat{E}(\infty)$ that is the optimal edge structure solving CDG-A($\infty$). We are interested in characterizing the following error probability of the event $\mathcal{A}_n$:

$$P \left[ \mathcal{A}_n(x^{1:n}) : \{ \hat{E}(n) \neq \hat{E}(\infty) \} \right].$$

To characterize the probability in (21) that is one of the rare events, we use LDP that rare events occurs in the most probable way.

To this end, we aim at studying the following rate function $K = K(y)$:

$$K(y) := \lim_{n \to \infty} \frac{1}{n} \log P(\mathcal{A}_n(x^{1:n})).$$

whenever the limit exists.

We now consider a simple event, called crossover event, as defined in what follows: Recall that ASYNC-ALGO uses, for each edge $e$, the weight $\beta$ of $w_e(\hat{P}) = I_e(\hat{P}) - 2\gamma c_e$ based on the empirical distribution $\hat{P}$. Then, consider two edges $e$ and $e'$ such that the weight of $e$ exceeds that of $e'$ with respect to the true distribution $P$, i.e., $w_e(P) > w_{e'}(P)$. We now define the crossover event for two edges $e$ and $e'$ as:

$$C_n(e, e') := \{ w_e(\hat{P}) \leq w_{e'}(\hat{P}) \}.$$  

As the number of samples $n \to \infty$, the empirical distributions approach to the true distribution, thus the probability of the crossover event decays to zero, whose decaying rate which we call crossover rate is defined as $J_{e, e'} := \lim_{n \to \infty} -\frac{1}{n} \log P(C_n(e, e'))$. Using this definition of the crossover event, we present Theorem 4.1 that states the decaying rate of the estimation error probability as the number of data samples $n$ grows.

THEOREM 4.1 (DECAYING RATE OF ASYNC-ALGO). For any fixed parameter $\gamma \geq 0$,

$$\lim_{n \to \infty} -\frac{1}{n} \log P(\mathcal{A}_n(x^{1:n})) = K(\gamma),$$

where

$$K(\gamma) := \min_{e \in \hat{E}(\infty)} \min_{e' \notin \hat{E}(\infty)} J_{e, e'}.$$

where $\Psi(e') = (i, j); \hat{E}(\infty) := \{ v_l = (i), v_{l+1}, \cdots, v_{l+k} = (j) \}$ is the unique path between nodes $i$ and $j$, such that $(v_k, v_{k+1}) \in \hat{E}(\infty)$ for $1 \leq k \leq l - 1$, and

$$J_{e, e'} := \begin{cases} \inf_{Q \in \mathcal{P}(X^4)} \left\{ D_{KL}(Q || P_{e, e'}) : w_e(Q) = w_{e'}(Q) \right\}, & \text{if } \{Q \in \mathcal{P}(X^4) : w_e(Q) = w_{e'}(Q) \neq 0, } \\ \infty, & \text{otherwise.} \end{cases}$$

We interchangeably use $w_e(\hat{P})$ to denote the assigned weight of an edge $e$ in algorithms, with respect to the empirical distribution $\hat{P}$ from the given samples.
Moreover, we have the following (finite-sample) upper-bound on the error probability: for all \( n = 1, 2, \ldots \),
\[
\Pr \left[ \mathcal{A}_n(x^{1:n}) \right] \leq \frac{(d-1)^2(2d-2)}{2} \left( n - 1 + |X|^4 \right) \frac{1}{|X|^4 - 1} \exp(-n \cdot K(y)).
\] (27)

In Theorem 4.1, we observe that the decaying rate of error probability is specified by some topological information of physical/data graphs and the trade-off parameter \( \gamma \). In particular, the crossover event and its rate \( J_{e,e'} \) depend on how difficult it is to differentiate two edge weights under the true data distribution with a consideration of the trade-off parameter \( \gamma \) as well as per-message cost on edges. As interpreted from (26), when \( w_0(P) = I_0(P) - 2\gamma c_0 \) and \( w_1(P) = I_1(P) - 2\gamma c_1 \) are close, the confusion between edge and \( e' \) is present, leading to higher error probability, and we can show the existence of the infimum \( Q \) satisfying \( w_0(Q) = w_1(Q) \) as by slightly adjusting the true distribution \( P \). Moreover, we remark that the decaying rate \( J_{e,e'} \) (and thus \( K(y) \)) is characterized by a trade-off parameter \( \gamma \). The error rate becomes smaller (i.e., higher error probability) when \( \gamma \) nearly meets the condition \( w_0(P) = w_1(P) \), and the weights becomes deterministic with respect to the samples as \( \gamma \) increases since the portion of the cost in weights grows, resulting to \( J_{e,e'} \approx \infty \) in (26). These interpretations are well-matched to our numerical results in Section 5.

**Proof sketch.** The proof of Theorem 4.1 is presented in our technical report [18], and we describe the proof sketch for readers’ convenience. Our proof largely follows that of the related work in [28] that analyzes an error exponent of a standard tree structure learning (i.e., known as Chow-Liu algorithm [8]), whose goal is to solely estimate the true data distribution with no consideration of communication cost. Simply, the proof idea follows LDP in the following way. The error event \( \mathcal{A}_n(x^{1:n}) \) is expressed as a union of small events that \textit{ASYNC-ALGO} estimates only one wrong edge (see the definition of the crossover event in (23)), two wrong edges, and three, etc. Following LDP, the decaying rate of the error probability equals to the decaying rate of the most probable crossover event, which corresponds to the case of only one wrong edge. In more detail, two minimums in (25) specify the most-probably error, whose edge set differs from the optimal data tree structure \( \hat{E}^*(\infty) \) exactly in one edge, i.e., \( \hat{E}^*(\infty) \) \( \setminus \{e\} \cup \{e'\} \), where it contains the non-neighbor node pair \( e' \) (as selected in the first minimization) instead of the most probable replacement edge \( e \) in the unique path along \( \hat{E}^*(\infty) \) (as in the second minimization). To obtain the minimum crossover rate \( J_{e,e'} \), we apply the Sanov’s theorem [5], which provides an expression of the probabilistic relationship between \( \hat{P} \) and \( P \) via their KL divergence. Finally, in addition to the asymptotic decaying rate of the estimation error probability, we also establish its upper bound of the error probability in terms of the number \( n \) of data samples, where the first term \((d-1)^2(2d-2)/2\) of the bound in (27) implies the number of possible crossover events, and the second term \((n-1+|X|^4)/|X|^4-1\) represents the number of possible empirical distributions \( \hat{P}_{e,e'} \).

### 4.2 Estimation Error of \textit{ASYNC-ALGO}

We conduct a similar analysis here for \textit{ASYNC-ALGO} to what we did for \textit{ASYNC-ALGO}, which has more complicated issues for the following reasons: We first denote by \( w_0(\hat{P}, T) \) in (17) the assigned weight for edge \( e \) to stress its dependence on the corresponding resulting tree structure \( T \) and its associated empirical distribution \( \hat{P} \). Then, we need to investigate the most probable pattern in the rare event through a certain tree \( T \) at some iteration. Simply, the crossover event for two edges \( e \) and \( e' \) occurs if the order of edge weights from the given finite number of samples becomes reversed to the order of weights from the true data distribution. Among all possible crossover events, we are interested in the crossover event under every tree structure that is obtained on the way of constructing the ideal data structure, denoted by \( \hat{E}^*(\infty) \). Let \( e_{true}^t \) and \( T_{true}^t \) be the selected edge and constructed tree at \( t \)-th iteration obtained by running \textit{ASYNC-ALGO} w.r.t. the true data distribution \( P \), which would finally find \( \hat{E}^*(\infty) \). Then, it is obvious that \( e_{true}^t \) has the unique highest edge weight for \( P \), and the crossover event of our interest is defined as:

\[
C_n(e_{true}^t, e'; T_{true}^t) := \left\{ w_{e_{true}^t}(\hat{P}; T_{true}^t) \leq w_{e'}(\hat{P}; T_{true}^t) \right\}.
\] (28)

We now state Theorem 4.2 that establishes the decaying rate of the estimation error probability as the number of data samples grows.

**Theorem 4.2 (Decaying rate of \textit{ASYNC-ALGO}).** For any fixed parameter \( \gamma \geq 0 \),
\[
\lim_{n \to \infty} \frac{1}{n} \log \Pr(\mathcal{A}_n(x^{1:n})) \geq K(y),
\] (29)

where
\[
K(y) := \min_{t \in \{1, \ldots, |V|-1\}} \min_{e' \in T_{true}^t} J_{e_{true}^t, e'}(T_{true}^t).
\] (30)

where \( e_{true}^t \) and \( T_{true}^t \) are the selected edge and constructed tree at \( t \)-th iteration by running \textit{ASYNC-ALGO} w.r.t. the true data distribution \( P \), i.e., \( w_{e_{true}^t}(P) \) has the maximum edge weight under the tree \( T_{true}^t \), and it is given by: under some tree \( T \), for any \( e, e' \),
\[
J_{e,e'}(T) = \begin{cases} 
\inf_{Q \in \mathcal{P}(X^t)} \left| D_{KL}(Q \parallel P_{e,e'}) \right| : w_e(Q) = w_{e'}(Q), \\
|Q \in \mathcal{P}(X^t) : w_e(Q) = w_{e'}(Q) \neq 0, \\
\infty, \\
\end{cases}
\] (31)

Moreover, we have the following (finite-sample) upper-bound on the error probability: for all \( n = 1, 2, \ldots \),
\[
\Pr \left[ \mathcal{A}_n(x^{1:n}) \right] \leq \frac{(d-1)(d+1)}{6} \left( n - 1 + |X|^4 \right) \frac{1}{|X|^4 - 1} \exp(-n \cdot K(y)).
\] (32)

In Theorem 4.2, as seen in (29), the error rate function \( K(y) \) in (30) indeed provides a lower-bound of the actual decaying rate of the error event \( \mathcal{A}_n(x^{1:n}) \), since the crossover event \( C_n(e_{true}^t, e'; T_{true}^t) \) which estimates an edge \( e' \) \( \notin T_{true}^t \) rather than \( e_{true}^t \) at any \( t \)-th iteration does not guarantee that \( e' \) is a wrong edge. Intuitively, the edge weights of \textit{ASYNC-ALGO} dynamically change according to a diameter of \( T_{true}^t \) as iteration \( t \) proceeds, which makes the characterization of the exact error rate of \textit{ASYNC-ALGO} be non-trivial.

**Proof sketch.** Due to space limitation, we present the complete proof in our technical report [18], and we provide a brief proof sketch. The basic idea is similar to the proof of Theorem 4.1. As mentioned there, the crossover event \( C_n(e_{true}^t, e'; T_{true}^t) \) is not a...
subset of the error event \( \mathcal{A}_n(x^{1/n}) \), and as a result, we provide a lower-bound of the decaying error rate in the proof, as established by two minimizations in (31). In particular, the first minimization is taken over all iterations \( 1 \leq t \leq |V| - 1 \) so that it selects the iteration where the error occurs in the most probable way, and the second minimization specifies the non-neighbor node pair \( e' \), which can be estimated instead of \( e^{\text{true}} \), having the minimum \( J_{e^{\text{true}}e'}(T_t^{\text{true}}) \). In other words, the most probable pattern in the error event of SYNC-ALGO is to estimate \( T_t^{\text{true}} \setminus \{ e^{\text{true}} \} \cup \{ e' \} \) attained in two minimizations in (30). For the crossover rate \( J_{e^{\text{true}}, e'}(T_t^{\text{true}}) \) in (31), when two edges \( e^{\text{true}} \) and \( e' \) can be clearly differentiated via their edge weights, since the difference of the cost between two edges dominantly determines the order of the edge weights, \( i.e., \) the condition in (31) does not hold, the crossover event does not happen, \( i.e., \) \( J_{e^{\text{true}}, e'}(T_t^{\text{true}}) = \infty \). This mostly corresponds to the situation of a large value of the trade-off parameter \( \gamma \), where the communication cost plays an important role of the error event, which do not depend on the number of samples \( n \). Otherwise, the crossover rate is attained in a similar way to (26). Finally, we establish the upper bound of the error probability in terms of the first minimization of two algorithms estimates a different structure for data tree, since the goal is to achieve the highest inference accuracy. However, as \( \gamma \) grows, each of two algorithms estimates a different structure for data tree, since ASYNC-MAP and SYNC-MAP have different forms of communication costs. In particular, in ASYNC-ALGO, as \( \gamma \) grows, we observe that the algorithm produces the smallest data tree with more resemblance to the physical graph, and finally it estimates the data tree that is the same as the physical graph with \( \gamma = 4 \), see Figures 3(b) and 3(c). We note that for a large value of \( \gamma \), the goal of ASYNC-ALGO is to find an MWST of minimum total cost, which accords with the physical graph of line-topology. However, the communication cost of SYNC-ALGO increases in proportion to the diameter of the estimated tree, thus it estimates a tree that is of a star-like topology, \( i.e., \) a tree with a small diameter as seen in Figures 4(b) and 4(c), to significantly reduce the cost, as \( \gamma \) grows.

### 5 NUMERICAL RESULTS

In this section, we provide a set of numerical experiments to validate our analytical results of ASYNC-ALGO and SYNC-ALGO under various numbers of data samples, communication costs, and trade-off parameters.

#### 5.1 Setup

**Physical graph.** We use a physical network \( G = (V, E_P) \) consisting of 20 nodes forming a line topology, where node \( i \) can directly communicate only with nodes \( i-1 \) and \( i+1 \), see Figure 2(a). We assign some constant cost of single message-passing for each edge \( e = (i, i+1) \) : \( c_{i,i+1} = \kappa \times 1.1^i \), except for \( c_{1,2} = 4 \kappa, c_{3,4} = 2 \kappa, c_{6,7} = 0.1 \kappa \), where we appropriately choose \( \kappa \) to adjust the scale of total communication cost of two learning algorithms in the same range, for clear comparison with the same values of \( \gamma \). In the message-passing between non-neighboring (w.r.t. the physical graph) node pairs \( (i, j) \), we simply assume that it expenses the sum of the costs when it is passed along the unique shortest multi-hop path \( \Psi((i, j); G) \) on \( G \), \( i.e., \) \( c_{i,j} = \sum_{e \in \Psi((i, j); G)} c_e \). For example, \( c_{1,4} = c_{1,2} + c_{2,3} + c_{3,4} \). We use this line topology for an exemplar physical graph to clearly observe the difference between ASYNC-ALGO and SYNC-ALGO, where it leads to a significantly huge amount of communication cost for SYNC-MAP, due to large diameter value \( \text{diam}(G) = 19 \).

**Data graph.** As an underlying statistical dependences among 20 nodes in the data graph, we consider a 3-regular tree \( T = (V, E_D) \), except for boundary nodes, where the node 1 is a root node and every node has a degree of 3 or less, as depicted in Figure 2(b). Each random variable \( X_i \) associated to a node \( i \) is set to follow a Bernoulli distribution. For a root node 1, it has \( P(X_1 = 0) = 0.7 \) and \( P(X_1 = 1) = 0.3 \), and for other neighboring node pairs \( i \) and \( j \), we

![Figure 2: Physical and data graphs used for our simulations.](image-url)

set the conditional distribution between \( X_i \) and \( X_j \) by

\[
P(X_i = 0|X_j = 0) = 0.7, \quad P(X_i = 0|X_j = 1) = 0.3
\]

whenever \( i < j \). With this setting of per-node distribution, it turns out that neighboring node pairs have high correlations, and thus have distinct values of the mutual information.

Under this choice of physical and data graphs, we obtain numerical examples to show the performance of ASYNC-ALGO and SYNC-ALGO for various values of trade-off parameter \( \gamma \), ranging from 0 to 4, and a fixed \( \beta = 1 \) in our results. For a fixed \( n \in \mathbb{N} \), we first generate \( n \) i.i.d. samples \( x^{1/n} \) from \( P(x) \) in (33). Then, we compute the empirical distribution \( \hat{P}(x^{1/n}) \) and the empirical mutual information of all possible node pairs \( \{I_{e}(\hat{P})\}_{e \in V \times V} \). Then, we learn the cost-efficient data tree by running ASYNC-ALGO or SYNC-ALGO, and estimate how well the proposed algorithms recover the ideal data graph by investigating the estimation error probability as \( n \) grows.
parameterized by $\gamma$ in the optimization problem in (3), we vary $\gamma$ from 0 to 4 and plot the accuracy of MAP estimator and the total cost on the learnt data dependency graph as the red and blue lines, respectively, in Figures 3(d) and 4(d). In particular, we run the ASYNC-ALGO and SYNC-ALGO with $n = 200$ samples, respectively, and run the max-product algorithm on the learnt data tree to obtain the MAP estimator. We repeatedly run for 200 times, and measure the error probability that the MAP estimator on the learnt data tree differs from the MAP estimator on the true data graph, as a metric of inference accuracy. The average (over the 200 results) of the communication cost on the learnt data tree is measured by the form of (6) and (7) for each algorithm. In Figure 4(d), we observe that the MAP estimation error and cost with $\gamma = 0.5$ is 0.33 and 0.083, respectively, while those with $\gamma = 2.5$ is 0.91 and 0.034, respectively. The impact of $\gamma$ on the trade-off for two algorithms seems similar, as seen in Figures 3(d) and 4(d).

(iii) Impact of data sample size on graph estimation accuracy.
Finally, we demonstrate the theoretical findings in Theorems 4.1 and 4.2 on the decaying rate of the error probability w.r.t. the number of samples $n$ for various values of $\gamma$. In both ASYNC-ALGO and SYNC-ALGO, for a fixed $\gamma$, we run both algorithms for 200 times each, and measure their error probabilities. In Figures 5(a) and 6(a), we observe that the error probability $\mathbb{P}(Q)\varepsilon$ for every $\gamma$ decays exponentially as the sample size $n$ increases, as established in (27) and (32). It is interesting to see that a different choice of $\gamma$ leads to a different decaying rate, which can be understood by our analytical findings of the crossover rate in (26) and (31), simply given by:

$$J_{\epsilon,\gamma}(T) = \inf_{Q\epsilon\mathbb{P}(X^\epsilon)} \{D_{\text{KL}}(Q\parallel P_{\epsilon,\gamma}) : w_{\epsilon}(Q) = w_{\gamma}(Q)\},$$

where the edge weights for ASYNC-ALGO and SYNC-ALGO are assigned in different forms, yet depending on the value of $\gamma$, as
seen in (9) and (17). Some choice of $\gamma$ makes a difference of the corresponding edge weights highly small, so that it becomes easier to estimate wrong edges with an insufficient number of samples. In our simulation, ASYNC-ALGO with $\gamma = 2$ shows higher error probability of 0.2 with $n = 10^4$ samples, while that with $\gamma = 0$ achieves almost 0 error probability with less than 3000 samples, see Figure 5(a). This impact of $\gamma$ on the error probability is presented in Figures 5(b) and 6(b) for both algorithms, where for large $\gamma$, we observe the error probability decays at a higher rate in general, since the priority to the inference accuracy is insignificant, leading to less chance of experiencing the crossover event.

6 CONCLUSION

In many multi-agent networked systems, a variety of applications involve distributed in-network statistical inference tasks, such as MAP (maximum a posteriori), exploiting a given knowledge of statistical dependencies among agents. When agents are spatially-separated, running an inference algorithm leads to a non-negligible amount of communication cost due to inevitable message-passing, coming from the difference between data dependency and physical connectivity. In this paper, we consider a structure learning problem which recovers the statistical dependency from a set of data samples, which also considers the communication cost incurred by the applied distributed inference algorithms to the learnt data graph. To this end, we first formulate an optimization problem formalizing the trade-off between inference accuracy and cost, whose solution chooses a tunable point in-between them. As an inference task, we studied the distributed MAP and their two implementations ASYNC-MAP and SYNC-MAP that have different cost generation structures. In ASYNC-MAP, we developed a polynomial time, optimal algorithm, inspired by the problem of finding a maximum weight spanning tree, while we proved that the optimal learning in SYNC-MAP is NP-hard, thus proposed a greedy heuristic. For both algorithms, we then established how the error probability that the learnt data graph differs from the ideal one decays as the number of samples grows, using the large deviation principle.

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