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Frontiers of Statistical Network Analysis: Inference, Tensors and Beyond

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HIGHER-ORDER GRAPHON THEORY: FLUCTUATIONS, INFERENCE, AND DEGENERACIES

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Classification AMS 2020: 05C80, 60F05, 05C60

Keywords: Inhomogeneous random graphs, network analysis, generalized U-statistics, subgraph counts.

Exchangeable random graphs, which include some of the most widely studied network models, play a central role in statistical network analysis. Graphons, which are the central objects in graph limit theory, provide a natural way to sample exchangeable random graphs. It is well known that network moments (motif/subgraph counts) identify a graphon (up to an isomorphism), hence, understanding the sampling distribution of subgraph counts in random graphs sampled from a graphon is pivotal for nonparametric network inference. Although there are a few results regarding the asymptotic normality of subgraph counts in graphon models, for many commonly appearing graphons this distribution is degenerate. This degeneracy phenomenon was overlooked until very recently and its consequences in network inference have remained unexplored. Towards this, in joint works with Chatterjee and Janson [1] and Chatterjee and Dan [2] we obtain the following results:

- We derive the joint asymptotic distribution of any finite collection of network moments in random graphs sampled from a graphon, that includes both the non-degenerate case (where the distribution is Gaussian) as well as the degenerate case (where the distribution has both Gaussian or non-Gaussian components). This provides the higher-order fluctuation theory for subgraph counts in the graphon model.
- We develop a novel multiplier bootstrap for graphons that consistently approximates the limiting distribution of the network moments (both in the Gaussian and non-Gaussian regimes). Using this and a procedure for testing degeneracy, we construct joint confidence sets for any finite collection of motif densities. This provides a general framework for statistical inference based on network moments in the graphon model.

We also discuss various structure theorems and open questions about degeneracies of the limiting distribution and connections to quasirandom graphs.

References

- [1] B. B. Bhattacharya, A. Chatterjee, and S. Janson, Fluctuations of subgraph counts in graphon based random graphs, *Combinatorics, Probability, and Computing*, Vol. 32 (3), 428–464, 2023.
- [2] A. Chatterjee, S. Dan, B. B. Bhattacharya, Higher-order graphon theory: Fluctuations, degeneracies, and inference, *arXiv:2404.13822*, 2024.

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LOW-DIMENSIONAL ADAPTATION OF DIFFUSION MODELS

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Keywords: diffusion models, low-dimensional structure, acceleration

Motivated by the practical efficacy of diffusion models, the past few years have witnessed a flurry of activity towards establishing convergence theory for diffusion generative models, particularly the two mainstream algorithms: DDPM and DDIM. For a fairly general family of target distributions \mathbb{P}_{data} (without assuming smoothness and log-concavity), the state-of-the-art theory demonstrated that for both DDPM and DDIM, it takes at most the order of (modulo some log factor)

(0.1)
$$\frac{d}{\varepsilon}$$
 iterations

to yield a sample whose distribution is ε -close in total variation (TV) distance to the target distribution, provided that perfect score function estimates are available.

Nevertheless, even linear scaling in the ambient dimension d can still be prohibitively expensive for many contemporary applications. Take the ImageNet dataset for instance: each image might contain 150,528 pixels, while its intrinsic dimension is estimated to be 43 or less. As a result, applying the state-of-the-art theory (0.1) could suggest an iteration complexity that exceeds one million, even though practical implementations of DDIM and DDPM often produce high-quality samples in just a few hundred (or even a few ten) iterations. The discrepancy between theory and practice suggests that worst-case bounds, such as (0.1), may be overly conservative. To reconcile this discrepancy, it is crucial to bear in mind the intrinsic dimension of the target data distribution and explore whether and how diffusion models can harness this potentially low-dimensional structure.

Motivated by this, in this talk we would like to explore how diffusion models leverage low-dimensional structure to speed up the sampling process. Focusing on two mainstream samplers — the denoising diffusion implicit model (DDIM) and the denoising diffusion probabilistic model (DDPM) — and assuming accurate score estimates, we prove that their iteration complexities are no greater than the order of k/ε (up to some log factor), where ε is the precision in total variation distance and k is some intrinsic dimension of the target distribution. Our results are applicable to a broad family of target distributions without requiring smoothness or log-concavity assumptions. Further, we develop a lower bound that suggests the (near) necessity of the coefficients introduced by Ho et al. 2020 and Song et al. 2020 in facilitating low-dimensional adaptation. Our findings provide the first rigorous evidence for the adaptivity of the DDIM-type samplers to unknown low-dimensional structure, and improve over the state-of-the-art DDPM theory regarding total variation convergence.

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AGNOSTIC CHARACTERIZATION OF INTERFERENCE IN RANDOMIZED EXPERIMENTS (EXTENDED ABSTRACT)

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Classification AMS 2020: 62D99

Keywords: causal inference, spillovers, social networks, interference

In randomized experiments, it may be possible for the participants to affect each other, by mechanisms such as transmission of disease, sharing of information, peer influence, or economic competition. Such phenomena (termed "interference between units") violates assumptions that are commonly used for statistical inference.

Mechanisms for interference often play fundamental roles in our understanding of social outcomes. For this reason, the empirical characterization of interference (such as its nature, prevalence, or strength) may be of scientific interest. For experiments with binary-valued outcomes, we give a general approach for characterizing the prevalence of interference, which can be used to explore questions such as

- Q1. How many units are affected by any treatment (including their own)?
- Q2. How many units are affected by the treatment of others? of distant others?
- Q3. How many units are affected by the treatment of others, provided that their own treatment satisfies some condition?

For each of these questions, our approach gives conservative point estimates and one-sided confidence intervals, which both lower bound the true value. Under reasonable experiment designs, the point estimate will be consistent for a lower bound on the true value, while the one-sided interval will cover the true value at the desired level. These consistency and coverage properties hold without any additional assumptions or restrictions on the nature of the interference, requiring only a randomized experiment whose design is known. As a result, our estimates remain valid even if they use an observed social network that is only a crude proxy for the actual social mechanisms.

A previous attempt to answer such questions relied on inversion of a test statistic, and produced quite conservative (though valid) lower bounds. Our new approach is significantly tighter, and may be more practical as a result. Our point estimates are asymptotically equal to Hajek-normalized contrasts, such as comparisons of treated versus untreated, or comparisons of different levels of indirect exposure, or comparisons that combine measures of direct and indirect treatment. Under stronger assumptions, such contrasts arise naturally as estimates of treatment effects. Our results indicate that without assumptions on interference, these contrasts may be interpreted more weakly as lower bounds on the number of units who are affected by the treatments. We also find empirically that our interval estimates have efficiency (i.e., interval widths) which is competitive with, and often better than, that of the expected average treatment effect (EATE), an assumption-lean treatment effect.

0.1. Idea of Method. Consider an experiment on N units, with $X = (X_1, \ldots, X_N)$ denoting the binary-valued treatment of each unit, and $Y = (Y_1, \ldots, Y_N)$ denoting their binary outcomes. We allow for arbitrary interference, so that the outcome Y_i of unit i may potentially depend on all N treatments,

(0.1)
$$Y_i = f_i(X_1, ..., X_N), \quad i \in [N]$$

where the potential outcome mappings $\{f_i\}_{i=1}^N$ may be arbitrary and unknown. Suppose that we wish to estimate τ^{basic} , the number of units who are affected by any treatment, including their own treatment or the treatment of others. To define this estimand, let $\mathcal{I} \subset [N]$ denote the subset of units who are unaffected by treatment and have constant outcome mappings,

$$\mathcal{I} = \{i : f_i(X) \text{ is constant in } X\}$$

so that $\tau^{\text{basic}} = N - |\mathcal{I}|$.

Our high-level approach to estimating τ^{basic} is the following:

- (1) Propose idealized estimators $\hat{\tau}_1$ and $\hat{\tau}_2$ which will have good statistical properties, such as consistency and asymptotic normality, but require knowledge of \mathcal{I}
- (2) Show that Δ , the difference in average outcomes between treated and control (which can be computed without knowledge of \mathcal{I}) converges to a lower bound for $\max(\hat{\tau}_1, \hat{\tau}_2)$, so that if τ_1 and τ_2 are both consistent for τ^{basic} , then Δ is an asymptotic lower bound.
- (3) Lower bound the boundary of the lower 1-sided confidence intervals induced by $\hat{\tau}_1$ and $\hat{\tau}_2$ and their variance estimates, by minimizing the tighter of the two boundaries over all hypotheses for the unknown subset \mathcal{I} .

To define $\hat{\tau}_1$ and $\hat{\tau}_2$, let S_i denote the indicator of whether unit i's treatment and outcome have the same binary value,

(0.2)
$$S_i = 1\{(X_i, Y_i) = (1, 1) \text{ or } (0, 0)\},\$$

and let $\hat{\tau}_1$ and $\hat{\tau}_2$ denote sampling-based estimators of $\tau^{\text{basic}} = N - |\mathcal{I}|$, in which the unknown cardinality of \mathcal{I} is unbiasedly estimated by a probability-weighted (i.e., Horvitz-Thompson) sample:

(0.3)
$$\hat{\tau}_1 = N - \sum_{i \in \mathcal{I}} \frac{1\{S_i = 1\}}{P(S_i = 1)}$$
 and $\hat{\tau}_2 = N - \sum_{i \in \mathcal{I}} \frac{1\{S_i = 0\}}{P(S_i = 0)}$

Because $\hat{\tau}_1$ and $\hat{\tau}_2$ involve only units in \mathcal{I} whose outcomes are unaffected by treatment and hence are constant, they often will exhibit simple statistical behavior, even if strong interference exists between units who are not in \mathcal{I} . For example, if treatment is assigned by independent Bernoulli randomization, then $\hat{\tau}_1$ and $\hat{\tau}_2$ are sums of independent variables. Similarly, if the dependencies between the unit treatments are bounded, then $\hat{\tau}_1$ and $\hat{\tau}_2$ are sums of variables whose dependencies will be similarly bounded. For this reason, under a variety of designs we may expect the values of $\hat{\tau}_1$ and $\hat{\tau}_2$, while unknown due to \mathcal{I} being unknown, to concentrate at their expectation (which equals τ^{basic}) and to be asymptotically normal.

Our motivation for constructing $\hat{\tau}_1$ and $\hat{\tau}_2$ is the following: under mild conditions on the experiment design, the maximum of $\hat{\tau}_1$ and $\hat{\tau}_2$ is lower bounded by the magnitude of the propensity-weighted difference in outcomes between treated and control, given by

$$\Delta = \sum_{i=1}^{N} \left(\frac{X_i}{P(X_i = 1)} - \frac{1 - X_i}{P(X_i = 0)} \right) Y_i,$$

as stated by Theorem 0.1 below:

Theorem 0.1. Let the total weights of the treated and control converge to their expectations, so that

(0.4)
$$\sum_{i=1}^{N} \frac{X_i}{P(X_i=1)} = N + O_P(N^{1/2}) \text{ and } \sum_{i=1}^{N} \frac{1-X_i}{P(X_i=0)} = N + O_P(N^{1/2})$$

Then it holds that

(0.5)
$$\left| \sum_{i=1}^{N} \left(\frac{X_i}{P(X_i = 1)} - \frac{1 - X_i}{P(X_i = 0)} \right) Y_i \right| \le \max(\hat{\tau}_1, \hat{\tau}_2) + O_P(N^{1/2})$$

If $\hat{\tau}_1^{\text{Haj}}$ and $\hat{\tau}_2^{\text{Haj}}$ are asymptotically normal, with consistent variance estimators denoted by \hat{V}_1 and \hat{V}_2 , then by combining 1-sided confidence intervals it holds with probability converging to at least $1 - \alpha$ that

(0.6)
$$\tau^{\text{basic}} \ge \max\left\{\hat{\tau}_1^{\text{Haj}} - z_{1-\frac{\alpha}{2}}\sqrt{\hat{V}_1}, \, \hat{\tau}_2^{\text{Haj}} - z_{1-\frac{\alpha}{2}}\sqrt{\hat{V}_2}\right\}.$$

As the right hand side of (0.6) requires knowledge of \mathcal{I} , it cannot be computed.

To construct a computable one-sided confidence interval for τ^{basic} , we will lower bound (0.6) by minimizing over all possible hypotheses for the unknown \mathcal{I} . Doing so results in the confidence statement that with probability at least $1 - \alpha$,

(0.7)
$$\tau^{\text{basic}} \ge \max\left(\min_{\phi \in \{0,1\}^N} \hat{\tau}_1^{\text{Haj}}(\phi) - z_{1-\frac{\alpha}{2}} \sqrt{\hat{V}_1(\phi)}, \min_{\phi \in \{0,1\}^N} \hat{\tau}_2^{\text{Haj}}(\phi) - z_{1-\frac{\alpha}{2}} \sqrt{\hat{V}_2(\phi)}\right),$$

where $\hat{\tau}_k^{\text{Haj}}(\phi)$ and $\hat{V}_k(\phi)$ denote $\hat{\tau}_k^{\text{Haj}}$ and \hat{V}_k evaluated under the hypothesis that $\mathcal{I} = \{i : \phi_i = 1\}$ for $\phi \in \{0, 1\}^N$. (See paper for further details, such as the form of the variance estimators \hat{V}_1 and \hat{V}_2 .)

0.2. **Illustrative Example.** In an experiment described in [Cai et al., 2015], rural farmers in China were randomly assigned to information sessions where they would be given the opportunity to purchase weather insurance. The sessions were randomized to give either high or low levels of information about the insurance product. First round sessions were held three days before second round sessions, so that first round attendees would have opportunity for informal conversations with their second round friends, in which they might share their opinions about the insurance product. Social network information was elicited, with the farmers instructed to list 5 close friends with whom they specifically discussed rice production or financial issues.

The goal of the experiment was to broadly demonstrate the importance of information sharing, by measuring its effects in a randomized setting. One of the conclusions of [Cai et al., 2015] was that the decision to purchase insurance was affected not only by a farmer's own treatment assignment, but also by that of their friends; specifically, farmers assigned to a second round low-information session were

more likely to purchase insurance if more of their listed friends in the first round were assigned to a high-information session.

For this experiment, our point estimate is that at least 23% of second round farmers, if assigned to a low information session, would be affected by information given to the first round farmers (1-sided 95% CI: at least 9%). This point estimate of 23% is asymptotically equal to a Hajek-normalized comparison of second round units who received low information directly but had many first round friends with high information, versus those in the second round who received low information directly and had few or no first round friends with high information :

point estimate
$$\approx \sum_{i=1}^{N} \left(\frac{1}{\hat{N}_{1}} \frac{1\{X_{i}=0, W_{i}=1\}}{P(X_{i}=0, W_{i}=1)} - \frac{1}{\hat{N}_{0}} \frac{1\{X_{i}=0, W_{i}=0\}}{P(X_{i}=0, W_{i}=0)} \right) Y_{i}$$

Here $i \in [N]$ enumerates the second round units, $X_i = 0$ if unit *i* was assigned to a low information session, $W_i = 1$ if all of unit *i*'s first round friends received high information, Y_i denotes unit *i*'s decision of whether or not to purchase insurance, and \hat{N}_1 and \hat{N}_0 denote the Hajek normalization factors, where $\hat{N}_a = \sum_{i=1}^{N} (P(X_i = 0, W_i = a))^{-1}$ for a = 0, 1.

For comparison, we consider an EATE-type treatment effect that considers the relative effects of receiving (X_i, W_i) equal to (0, 1) versus (0, 0):

treatment effect =
$$\frac{1}{N} \sum_{i=1}^{N} (\mathbb{E}[Y_i | X_i = 0, W_i = 1] - \mathbb{E}[Y_i | X_i = 0, W_i = 0])$$
,

where the expectation is taken over the randomization of treatment under the experiment design. For this target parameter, the method of [Gao and Ding, 2023] gives a Hajek-normalized point estimate of 23%, and 95% CI of [2%, 45%]. This confidence interval requires an assumption of "approximate neighborhood interference", in which the interference between farmers in different villages is asymptotically negligible. Such an assumption might be debatable, as farmers listed cross-village friendships. In contrast, no assumptions on interference are required for our estimand. Thus for the purposes of demonstrating the presence of social influence (as opposed to policy recommendation), our estimand may be an appropriate target parameter, and has tighter, less questionable CIs when compared to an analogous treatment effect.

REFERENCES

[[]Cai et al., 2015] Cai, J., De Janvry, A., and Sadoulet, E. (2015). Social networks and the decision to insure. *American Economic Journal: Applied Economics*, 7(2):81–108.

[[]Gao and Ding, 2023] Gao, M. and Ding, P. (2023). Causal inference in network experiments: regression-based analysis and design-based properties. *arXiv preprint arXiv:2309.07476*.

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HIGH-DIMENSIONAL NETWORK CAUSAL INFERENCE

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Keywords: Network interference, Nonparametric interference effects, Average direct treatment effect on the treated, Confidence intervals, Neighborhood size confidence set

We propose a new method of high-dimensional network causal inference (HNCI) that provides both valid confidence intervals for the average direct treatment effect on the treated (ADET) and valid confidence sets for the neighborhood size affecting the interference effect. Consider a sample of n units indexed by $i \in [n] := \{1, 2, ..., n\},\$ connected through an interference network G, where each unit is randomly assigned a binary treatment Z_i Bernoulli (p_i) for some p_i \sim \in (0, 1).Let $\mathbf{z} = (z_1, z_2, \cdots, z_n)^T \in \{0, 1\}^n$ denote the treatment assignments, which serves as a realization of the random vector $\mathbf{Z} = (Z_1, Z_2, \cdots, Z_n)^T$. For example, z could indicate that a tax incentive is offered to a specific subset of businesses in a region. In the network setting, the units are referred to as nodes in G, which are rarely independent of each other. Hence, the effect of a tax incentive on a specific company may depend on whether its collaborators or competitors also receive the tax incentive. For the n nodes connected through G, the potential outcome of the *i*th node is defined as $\widetilde{Y}_i(\mathbf{z}) = \widetilde{Y}_i(z_i, \mathbf{z}_{-i})$, where $\widetilde{Y}_i(\cdot) : \{0, 1\}^n \to \mathbb{R}$, and z_i and \mathbf{z}_{-i} are the treatment assignments for the *i*th node and the remaining nodes, respectively. In practice, we may observe node covariates $\{\mathbf{C}_i\}_{i \in [n]}$.

We exploit the following potential outcome model framework introduced in [1], where the potential outcome of the *i*th node is defined as

(0.1)
$$Y_i(z_i, \mathbf{z}_{-i}) = z_i \tau_i + f\left(\gamma_0(G_i^{\mathbf{z}}(k_0))\right) + \epsilon_i$$

Here, $\tau_i := \mathbb{E}\{\widetilde{Y}_i(1, \mathbf{0}_{-i}) - \widetilde{Y}_i(0, \mathbf{0}_{-i}) | \mathbf{C}_i\}$ is the average direct effect of the treatment on the *i*th node, $i \in [n]$, $\gamma_0(\cdot)$ is a known mapping satisfying the *nested matching* property that $\gamma_0(G_i^{\mathbf{z}}(k)) = \gamma_0(G_j^{\mathbf{z}}(k))$ implies $\gamma_0(G_i^{\mathbf{z}}(k')) = \gamma_0(G_j^{\mathbf{z}}(k'))$ for all $k' \in [k]$, $f(\cdot)$ is an unknown interference function, k_0 is the smallest neighborhood size that satisfies (0.1), and ϵ_i 's are independent errors with $\mathbb{E}(\epsilon_i) = 0$, $\operatorname{Var}(\epsilon_i) = \sigma_0^2$.

We work under model (0.1) to estimate and infer the average direct treatment effect on the treated (ADET)

(0.2)
$$\tau := \frac{1}{\sum_{i=1}^{n} Z_i} \sum_{i=1}^{n} Z_i \tau_i,$$

which represents the average incremental response of treated units to their own treatments. We are also interested in estimating the neighborhood size k_0 with statistical uncertainty guarantee.

For untreated nodes $z_i = 0$, we have

(0.3)
$$\widetilde{Y}_i(0, \mathbf{z}_{-i}) = f(\gamma_0(G_i^{\mathbf{z}}(k_0))) + \epsilon_i.$$

Thanks to the nested matching property, for each pre-specified neighborhood size $k \ge k_0$, the set of untreated nodes can be partitioned into d(k) disjoint subsets, denoted as $S_k = \{S_{k,1}, S_{k,2}, \ldots, S_{k,d(k)}\}$, where each subset $S_{k,j}$ contains nodes with the same interference function value $\gamma_0(G_i^{\mathbf{z}}(k))$ for all $i \in S_{k,j}$. Define the vector of true interference function values over the node partition S_k as

(0.4)
$$\beta_k^0 = (\beta_{k,1}^0, \beta_{k,2}^0, \cdots, \beta_{k,d(k)}^0)^T$$

Based on this property, the response vector $\mathbf{y}_{obs} \in \mathbb{R}^{n_0}$ of untreated nodes can be rewritten in the form of a linear regression model

$$\mathbf{y}_{obs} = \mathbf{X}_k \boldsymbol{\beta}_k^0 + \boldsymbol{\varepsilon}_0,$$

where $\mathbf{X}_k \in \{0, 1\}^{n_0 \times d(k)}$ is the design matrix with each row indicating the corresponding unit's membership in S_k , and ε_0 is the error term. Since k can be larger than k_0 and the function f can be many-to-one, there exisits unknown homogeneity in the regression coefficient vector $\boldsymbol{\beta}_k^0$, and the true interference function values $\{f(\gamma_0(G_i^{\mathbf{z}}(k))) : z_i = 0, i = 1, \dots, n\}$ can be estimated by estimating the regression coefficients $\boldsymbol{\beta}_k^0$ in (0.5).

By considering this linear representation, we reformulate the original nonparametric model into a linear regression model where the regression coefficients, corresponding to the underlying true interference function values of nodes, exhibit a latent homogeneous structure. This formulation enables us to leverage existing literature on homogeneity pursuit [3] to conduct valid statistical inferences with theoretical guarantees for estimating the unknown β_k^0 . This gives us the estimates of the set of interference function values $\{f(\gamma_0(G_i^z(k))) : z_i = 0, i = 1, \dots, n\}$ and the confidence interval for these estimates.

By using the matching technique, the estimates of ADET can also be constructed and the confidence interval can be calculated. We theoretically justify the inference for the ADET through establishing asymptotic normality with estimable variances. By employing the repro samples approach [4], we further provide the confidence set for the interference of neighborhood size k_0 with theoretical guarantees. The practical utility of the newly suggested methods is demonstrated through simulations and real data examples.

REFERENCES

- [1] Alexandre Belloni, Fei Fang, and Alexander Volfovsky. Neighborhood adaptive estimators for causal inference under network interference. *arXiv preprint arXiv:2212.03683*, 2022.
- [2] Wenqin Du, Rundong Ding, Yinging Fan, and Jinchi Lv. HNCI: High-Dimensional Network Causal Inference *arXiv preprint arXiv:2412.18568*, 2024.
- [3] Xiaotong Shen and Hsin-Cheng Huang. Grouping pursuit through a regularization solution surface. *Journal of the American Statistical Association*, 105, 727–739, 2010.
- [4] Peng Wang, Min-Ge Xie, and Linjun Zhang. Finite- and large-sample inference for model and coefficients in high- dimensional linear regression with repro samples. *arXiv preprint arXiv:2209.09299*, 2022.

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LEARNING LATENT FEATURES FROM NETWORK DATA

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Keywords: seriation, latent model, recursive trees, Jordan ordering

1. LATENT VARIABLE MODEL

We describe a graph by its adjacency matrix $X \in \{0, 1\}^{n \times n}$.

Definition 1.1. Latent variable model. The graph is generated randomly as follows:

- each vertex *i* is characterized by a latent variable $z_i \in \mathcal{Z}$
- conditionally on z, the X_{ij} are independent, with

$$\mathbb{P}[X_{ij} = 1|z] = \mathbb{E}[X_{ij} = 1|z] = f(z_i, z_j)$$

where $f : \mathcal{Z} \times \mathcal{Z} \rightarrow [0, 1]$.

This model encompass the graphon model, random geometric graphs where $f(z_i, z_j) = g(d(z_i, z_j))$ with d a distance on \mathcal{Z} , the Robinson model where f decreases when moving away from the diagonal, the stochastic block model, ranking models and so on.

An ideal objective is to recover the latent values z_1, \ldots, z_n from the observation of X. Yet, it is an ill-posed problem, due to the lack of identifiability: while it is a minor issue in parametric models (estimation up to some "invariant" transformation), it is a much more severe issue in non-parametric models. For simplicity, we focus on the case where

- *f* belongs to some non-parametric class, with smoothness or shape assumptions;
- the latent positions are $z_i = \pi^*(i)$ for π^* a permutation of $\{1, \ldots, n\}$.

The goal is to recover π^* from $X = \pi^* F \pi^{*T} + E$, with $F_{ij} = f(i, j)$ unknown, π^* an unknown permutation matrix and E_{ij} independent sub-Gaussian random variables. What is the rate of estimation without computational constraints? What is the rate of estimation with poly-time algorithms? Is there a gap between the two?

Statistical-computational gaps exist in latent variable model [2], for example both in the geometric seriation model $F_{ij} = \lambda \mathbf{1}_{|i-j| \le \sqrt{n}}$, with $\lambda > 0$, and in the Hölder graphon model $f : [0, 1] \times [0, 1] \rightarrow [0, 1]$ with α -Hölder regularity, $0 < \alpha < 1$. We outline below two settings where estimation can be performed in poly-time at the optimal statistical rate.

2. BI-LIPSCHITZ SHAPE CONSTRAINT

Definition 2.1. Bi-Lipschitz $\mathcal{BL}(\alpha, \beta)$ **.** Assume that $F \in [0, 1]^{n \times n}$ is symmetric and

•
$$F_{ik} - F_{jk} \ge \alpha \frac{|i-j|}{n}$$
 for $k < i < j$, and $F_{jk} - F_{ik} \ge \alpha \frac{|i-j|}{n}$ for $i < j < k$;
• $|F_{ik} - F_{jk}| \le \beta \frac{|i-j|}{n}$.

The parameter α drives the signal strength. The parameter β is a smoothness parameter.

Example 2.2. Toeplitz matrix. The simple Toeplitz matrix $F_{ij} = 1 - \alpha \frac{|i-j|}{n}$ belongs to $\mathcal{BL}(\alpha, \alpha)$.

We define the max-error loss as

$$\ell_{\infty}(\hat{\pi}, \pi^*) = \min_{\pi^* admissible} \frac{1}{n} \max_{i \in [n]} |\hat{\pi}_i - \pi_i^*|$$

Theorem 2.3. [3] There exist some poly-time estimators $\hat{\pi}^{poly}$ such that, for any $F \in \mathcal{BL}(\alpha,\beta)$, any $n \geq C_{\alpha,\beta}$, and for some numerical constant c > 0, with probability at least $1 - n^{-2}$

$$\ell_{\infty}(\hat{\pi}^{poly}, \pi^*) \leq \frac{c}{\alpha} \sqrt{\frac{\log n}{n}} .$$

Furthermore, this rate is optimal for the Toeplitz matrix, up to a possible log factor.

This result proved in [3] ensures that the optimal statistical rate $n^{-1/2}$ for estimating Bi-Lipstchitz permuted matrices can be achieved by poly-time algorithms. Hence, there is no statistical-computational gap in this case. The optimal algorithm essentially

(1) first estimates the neighborhood distance

$$D_{ij}^* = \sqrt{n \sum_{k} (F_{ik}^{\pi^*} - F_{jk}^{\pi^*})^2}$$

at optimal rate $|\hat{D} - D^*|_{\infty} = O(n^{3/4});$

- (2) then perform a first partial ordering of points separated by $O(n^{3/4})$ based on this estimation;
- (3) then refine this partial ordering by comparing partial sums, providing a reliable partial ordering of points separated by $O(n^{1/2})$.

Furthermore, Theorem 2.3 remains valid for weak local Bi-Lipschitz functions.

3. Recursive trees

Going beyond conditional i.i.d. graphs, we can consider recursive trees such as

Definition 3.1. Random Recursive Tree (RRT) Build a tree recursively by connecting each new node to existing nodes uniformly at random

Definition 3.2. Preferential Attachement Tree (PA) Build a tree recursively by connecting each new node to existing nodes with a probability proportional to their degree

Let us define a Jordan centroid $\hat{\sigma}_J$ as a vertex such that, when removing it, it splits the tree into components of size smaller than $\lfloor n/2 \rfloor$. At least one, maximum two such vertices exist. For a node u, we can define $\hat{d}(u)$ as the number of descendants of u in the tree rooted at $\hat{\sigma}_J$. Then, we can order the nodes according to the number of descendant $\hat{d}(u)$, ties being broken randomly. We call Jordan ordering this ordering $\hat{\sigma}_J(u)$, which can be computed in $O(n \log n)$ time.

Theorem 3.3. [1] For any $\alpha \ge 1$, in the RTT model

$$R_{\alpha}(\hat{\sigma}_J) := \sum_{u \in V} \frac{|\hat{\sigma}_J(u) - \sigma(u)|}{\sigma(u)^{\alpha}} \le \kappa(\alpha) n^{2-\alpha} + C \log^4 n$$

with

$$\kappa(\alpha) = \frac{2}{2-\alpha} + \frac{2e^2}{(2-\alpha)^2} + \frac{2}{(2-\alpha)^3}$$

Furthermore, the rate $n^{2-\alpha}$ is optimal for $\alpha \in [1, 2)$.

Most of the story is the same for the PA model, except that Descendant ordering is optimal up to constant only for $1 \le \alpha < 5/4$.

References

- [1] Simon Briend, Gabor Lugosi, Christophe Giraud, and Deborah Sulem. Estimating the history of a random recursive tree. *Bernoulli*. To appear
- [2] Bertrand Even, Christophe Giraud, and Nicolas Verzelen. Computational lower bounds in latent models: clustering, sparse-clustering, biclustering *arXiv*, https://arxiv.org/abs/2506.13647
- [3] Yann Issartel, Christophe Giraud, Nicolas Verzelen. Minimax optimal seriation in polynomial time *arXiv*, https://arxiv.org/abs/2405.08747

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STATISTICAL ANALYSIS OF RECIPROCITY

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Classification AMS 2020: 62F10, 62F12, 62C20, 62R99

Keywords: Asymptotic normality; Effective sample size; Maximum likelihood estimator; Reciprocity; Sparse networks.

Consider a directed network with *n* nodes, denoted by $G_n = (V, E)$, where $V = \{1, \ldots, n\}$ is the set of nodes and $E \subseteq V \times V$ represents the edge set. We focus on simple graphs, so no self-loops are allowed, i.e., $(j, j) \notin E$ for any $j \in V$. Let $A_{ij} \in \{0, 1\}$ denote the random variable indicating the presence of a directed link from node *i* to node *j*. Assuming that dyads (A_{ij}, A_{ji}) and (A_{kl}, A_{lk}) are independent whenever $\{i, j\} \cap \{k, l\} = \emptyset$, the Bernoulli model with reciprocity (BR) specifies multinomial probabilities for each dyad as follows (Krivitsky and Kolaczyk, 2015):

(0.1) **BR model:** $p_{ij}(0,0) \propto 1$, $p_{ij}(1,0) = p_{ij}(0,1) \propto \exp(\mu_n)$, $p_{ij}(1,1) \propto \exp(2\mu_n + \rho_n)$,

where $p_{ij}(a, b) = p(A_{ij} = a, A_{ji} = b)$. In this model, μ_n represents the baseline tendency of nodes *i* and *j* to connect, while ρ_n captures *reciprocity*, the propensity for pairs of nodes to form mutual links. BR model serves as a natural extension of the Erdős–Rényi model (Erdős and Rényi, 1959, 1960) for undirected graphs, adapted to incorporate reciprocity for the analysis of directed networks. This model raises a fundamental question:

Question 1: What is the effective sample size for the statistical inference of μ_n and ρ_n ?

This question would be straightforward if μ_n and ρ_n were fixed, as it would fall under standard maximum likelihood estimation. However, when μ_n and ρ_n depend on *n*-the regime where the network is sparse-the inference of these parameters has been only partially explored in Krivitsky and Kolaczyk (2015). That work assumes that the effective sample sizes for μ_n and ρ_n are of the same order. Extending the analysis to allow different sparsity levels for μ_n and ρ_n provides a more comprehensive solution to Question 1, offering deeper insights into the effective sample sizes required for a broader range of network structures. Related, Chen et al. (2021) examines the effective sample size in the context of the Erdős–Rényi model under arbitrary sparsity, focusing on a single density parameter similar in spirit to μ_n . The examination of the interplay between the two parameters, μ_n and ρ_n , under differing sparsity regimes represents a new and more nuanced perspective, offering insights beyond those provided by models with a single density parameter.

More importantly, a complete answer to this question will pave the way for developing new models. As an example, we extend the BR model to the following:

$$p_{1.5} \text{ model}: \quad p_{ij}(0,0) \propto 1, \quad p_{ij}(1,0) \propto \exp\left(\mu_n + X_i^T \gamma_1 + Y_j^T \gamma_2\right), \\ p_{ij}(0,1) \propto \exp\left(\mu_n + X_j^T \gamma_1 + Y_i^T \gamma_2\right), \\ p_{ij}(1,1) \propto \exp\left(2\mu_n + \left(X_i^T + X_j^T\right)\gamma_1 + \left(Y_i^T + Y_j^T\right)\gamma_2 + \rho_n + V_{ij}^T\delta\right), \end{cases}$$

with additional parameters γ_1 , γ_2 , and δ , where $X_i \in \mathbb{R}^{d_1}$ represents covariates related to node *i*'s outgoingness, $Y_i \in \mathbb{R}^{d_2}$ relates to its incomingness, and $V_{ij} \in \mathbb{R}^{d_3}$ governs the reciprocity between nodes *i* and *j*. The model in (0.2) allows for node-specific heterogeneity via $X_i^T \gamma_1$ for outgoingness and $Y_j^T \gamma_2$ for incomingness, and $V_{ij}^T \delta$ to model heterogeneity in reciprocal relationships. Assuming that the parameters associated with the covariates are fixed, we further pose the following question:

Question 2: What are the effective sample sizes for the statistical inference of γ_1 , γ_2 , and δ ?

The model in (0.2) has a close relationship with the p_1 model introduced by Holland and Leinhardt (1981), where the p_1 model employs node-specific fixed effects without explicitly accounting for link-specific reciprocity. Our model in (0.2) parametrizes these fixed effects through covariates, achieving a more parsimonious structure. Although it may lack some of the flexibility of the p_1 model, this approach offers certain advantages, such as enabling link prediction for new nodes not used in model fitting. Additionally, a key advantage of the model in (0.2) lies in its suitability for sparser networks. We show that inference is feasible as long as the number of links diverges. In contrast, the p_1 model, with its large number of parameters, typically requires much denser networks to ensure the existence and asymptotic normality of its estimators, though no formal inference procedures are currently available for these estimators (see literature review below). Additionally, the model in (0.2) shares features with the p_2 model (Van Duijn et al., 2004), which also includes random effects for outgoingness and incomingness. As our model conceptually bridges the p_1 and p_2 models, we refer to it as the $p_{1.5}$ model.

1. The BR Model

We begin by examining the effective sample sizes for the BR model as specified in (0.1). For the sake of theoretical analysis and notational convenience, it is beneficial to work with the parameters (μ_n, τ_n) , where $\tau_n = 2\mu_n + \rho_n$. The negative log-likelihood function with respect to (μ_n, τ_n) can be expressed as:

$$\ell_n^{(1)}(\mu_n, \tau_n) = \sum_{i < j} \log(k_{n,ij}) - \mu_n \sum_{i < j} \left(A_{ij}(1 - A_{ji}) + A_{ji}(1 - A_{ij}) \right) - \tau_n \sum_{i < j} A_{ij} A_{ji},$$

where $k_{n,ij} = 1 + 2 \exp(\mu_n) + \exp(\tau_n)$ serves as the normalizing constant. It is important to note that the likelihood functions defined in terms of (μ_n, ρ_n) and (μ_n, τ_n) are equivalent, as are their corresponding maximum likelihood estimators. This leads us to the following lemma:

Lemma 1.1. Suppose $(\hat{\mu}_n, \hat{\tau}_n) = \operatorname{argmin}_{(\mu_n, \tau_n) \in \mathbb{R}^2} \ell_n^{(1)}(\mu_n, \tau_n)$. Then, it follows that $(\hat{\mu}_n, \hat{\tau}_n - 2\hat{\mu}_n) = \operatorname{argmin}_{(\mu_n, \rho_n) \in \mathbb{R}^2} \ell_n^{(2)}(\mu_n, \rho_n)$, where $\ell_n^{(2)}(\mu_n, \rho_n)$ denotes the negative log-likelihood function parametrized by μ_n and ρ_n . The reverse direction also holds.

Given this equivalence, we focus on estimating μ_n and τ_n . Inspired by the role of $-\log n$ in the Erdős–Rényi model for sparse networks, we define

$$\mu_n = -a\log n + \mu, \quad \tau_n = -b\log n + \tau_n$$

where $\mu \in [-M_{\mu}, M_{\mu}]$, $\tau \in [-M_{\tau}, M_{\tau}]$, and a, b > 0. The constant *a* preceding $\log n$ directly reflects network sparsity, though similar asymptotic normality results may arise from other scaling factors beyond $\log n$. From $\ell_n^{(1)}(\mu_n, \tau_n)$, we interpret *a* as the sparsity index for non-reciprocal links and *b* for reciprocal links.

This transformation clarifies the dependence of sparsity on n while allowing for intuitive statistical inference on the fixed parameters μ and τ . For further discussions on this topic, we refer to Krivitsky and Kolaczyk (2015) and Chen et al. (2021). It is important to note that the constants a, μ , b, and τ are not identifiable or estimable. To address these challenges, we will later develop a straightforward inference procedure for μ_n and τ_n .

Under the given scaling, we find that the expected number of non-reciprocal links is $E\left(\sum_{i,j=1}^{n} A_{ij} - \sum_{i < j} A_{ij} A_{ji}\right) \approx n^{2-a}$, while the expected number of reciprocated links is $E\left(\sum_{i < j} A_{ij} A_{ji}\right) \approx n^{2-b}$. Consequently, the total expected number of links is of order n^{2-a} if $a \leq b$, or n^{2-b} if a > b. This scaling choice highlights that the two quantities can indeed differ in magnitude. Notably, Krivitsky and Kolaczyk (2015) examined a special case of our framework when a = b = 1, leading to comparable expected numbers of non-reciprocal and reciprocated links. For sparse networks, the sufficient statistics $\left(\sum_{i < j} A_{ij} + A_{ji}, \sum_{i < j} A_{ij} A_{ji}\right)$ in the BR model can be efficiently computed using a sparse adjacency matrix. As a result, the time complexity for computing the maximum likelihood estimator is $O(n^{2-\min\{a,b\}})$, which is lower than $O(n^2)$ when $\min\{a,b\} > 0$.

We now derive the effective sample sizes for μ and τ , assuming that a and b are known. We begin by expressing the negative log-likelihood function as follows:

(1.1)
$$\ell_n(\mu,\tau) = \sum_{i < j} \log(k_{ij}) - \mu \sum_{i < j} \left(A_{ij}(1 - A_{ji}) + A_{ji}(1 - A_{ij}) \right) - \tau \sum_{i < j} A_{ij} A_{ji},$$

where $k_{ij} = 1 + 2n^{-a} \exp(\mu) + n^{-b} \exp(\tau)$ serves as the normalizing constant. Our maximum likelihood estimator is defined as

$$(\hat{\mu}, \hat{\tau}) = \operatorname{argmin}_{(\mu, \tau) \in \Omega_1} \frac{1}{\binom{n}{2}} \ell_n(\mu, \tau),$$

with $\Omega_1 = [-M_{\mu}, M_{\mu}] \times [-M_{\tau}, M_{\tau}]$. To derive the asymptotic results, we make the following assumptions:

Assumption 1.2. (Sparse network) Assume 0 < a, b < 2. The true values (μ_0, τ_0) lie within the interior of Ω_1 .

The conditions a > 0 and b > 0 ensure that the resulting graph is sparse, while a < 2 and b < 2 are necessary to guarantee that the total numbers of reciprocal and non-reciprocal links approach infinity. Without these conditions, consistent estimation would not be achievable. We now present the following result regarding the maximum likelihood estimator (MLE). All our results hold under Assumption 1.2, meaning they apply to arbitrarily sparse networks.

Proposition 1.3. (Asymptotic normality of the MLE in BR model) Under Assumption 1.2, as *n* approaches infinity, the MLE $(\hat{\mu}, \hat{\tau})$ is consistent and asymptotically normal, specifically:

$$\left(\sqrt{n^{2-a}(\hat{\mu}-\mu_0)}, \sqrt{n^{2-b}(\hat{\tau}-\tau_0)}\right)^T \rightsquigarrow N(0, \Sigma^{-1}),$$

where

$$\Sigma = \begin{pmatrix} \exp(\mu_0) & 0\\ 0 & \exp(\tau_0)/2 \end{pmatrix}.$$

Following the reasoning in Krivitsky and Kolaczyk (2015) and Chen et al. (2021), we can interpret n^{2-a} and n^{-b} as the effective sample sizes for μ and τ , respectively. This

interpretation is intuitive, as from equation (1.1), μ can be seen as the density parameter for the configuration (1,0) and (0,1), while τ represents the density parameter for the configuration (1,1).

REFERENCES

- Chen, M., Kato, K., and Leng, C. (2021). Analysis of networks via the sparse β -model. *Journal of the Royal Statistical Society Series B: Statistical Methodology*, 83(5):887–910.
- Erdős, P. and Rényi, A. (1959). On random graph. *Publicationes Mathematicate*, 6:290–297.
- Erdős, P. and Rényi, A. (1960). On the evolution of random graphs. *Publ. math. inst. hung. acad. sci*, 5(1):17–60.
- Holland, P. W. and Leinhardt, S. (1981). An exponential family of probability distributions for directed graphs. *Journal of the American Statistical association*, 76(373):33–50.
- Krivitsky, P. N. and Kolaczyk, E. D. (2015). On the question of effective sample size in network modeling: an asymptotic inquiry. *Statistical Science*, 30(2):184–198.
- Van Duijn, M. A., Snijders, T. A., and Zijlstra, B. J. (2004). p2: a random effects model with covariates for directed graphs. *Statistica Neerlandica*, 58(2):234–254.

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A REGRESSION TREE APPROACH TO MISSING DATA

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Keywords: imputation, incomplete data, prediction, recursive partitioning

The standard method to fitting a prediction model to incomplete data that have missing values in the predictor variables is to first complete the data by imputing (i.e., estimating) the missing values. This approach may not be logical if the "missing" values are non-existent instead of missing due to non-response. One example is the variable "age of spouse" for people who are single. Another common example occurs in so-called "skip questions", where variable $x_1 = 1$ if a person has a credit card and $x_1 = 0$ otherwise, and x_2 is the credit card balance. Here, x_2 would be reported as missing for people who do not have credit cards.

This talk introduces a new approach to missing values that makes missing-value imputation unnecessary. It accomplishes this by means of the GUIDE regression tree algorithm [4, 5], which fits a binary decision tree model to the incomplete data. A major strength of GUIDE is that it treats missing values as observed qualitative information and sends them to the left or right subnode at each split according to the values of the outcome (y) variable relative to those with non-missing values. In particular, it allows for splits that send missing values and only missing values to one subnode [6]. Other regression tree algorithms either impute the missing values before splitting the node [2], or send observations with missing values randomly to the left or right subnode [3]. The method is demonstrated on a dataset to predict death or intubation in patients hospitalized for Covid-19 [1].

REFERENCES

- Baker, T. B., Loh, W.-Y., Piasecki, T. M., Bolt, D. M., Smith, S. S., Slutske, W. S., Conner, K. L., Bernstein, S. L., and Fiore, M. C. (2023). A machine learning analysis of correlates of mortality among patients hospitalized with COVID-19. *Scientific Reports*, 13(4080).
- [2] Breiman, L., Friedman, J. H., Olshen, R. A., and Stone, C. J. (1984). *Classification and Regression Trees*. Chapman & Hall/CRC.
- [3] Hothorn, T., Hornik, K., and Zeileis, A. (2006). Unbiased recursive partitioning: a conditional inference framework. *Journal of Computational and Graphical Statistics*, 15:651–674.
- [4] Loh, W.-Y. (2002). Regression trees with unbiased variable selection and interaction detection. *Statistica Sinica*, 12:361–386.
- [5] Loh, W.-Y. (2009). Improving the precision of classification trees. Annals of Applied Statistics, 3:1710– 1737.
- [6] Loh, W.-Y., Eltinge, J., Cho, M. J., and Li, Y. (2019). Classification and regression trees and forests for incomplete data from sample surveys. *Statistica Sinica*, 29:431–453.

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ASYMPTOTIC THEORY OF EIGENVECTORS FOR LATENT EMBEDDINGS WITH GENERALIZED LAPLACIAN MATRICES

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Classification AMS 2020: 60F05, 62E20, 94A15

Keywords: Graph and manifold embeddings; Asymptotic distributions; Eigenvectors and eigenvalues; Local laws; RMT under dependency; High dimensionality

Laplacian matrices are commonly employed in many real applications, encoding the underlying latent structural information such as graphs and manifolds. The use of the normalization terms naturally gives rise to random matrices with dependency. It is well-known that dependency is a major bottleneck of new random matrix theory (RMT) developments. To this end, in this paper, we formally introduce a class of generalized (and regularized) Laplacian matrices, which contains the Laplacian matrix and the random adjacency matrix as a specific case, and suggest the new framework of the asymptotic theory of eigenvectors for latent embeddings with generalized Laplacian matrices (ATE-GL). Our new theory is empowered by the tool of generalized quadratic vector equation for dealing with RMT under dependency, and delicate high-order asymptotic expansions of the empirical spiked eigenvectors and eigenvalues based on local laws. The asymptotic normalities established for both spiked eigenvectors and eigenvalues will enable us to conduct precise inference and uncertainty quantification for applications involving the generalized Laplacian matrices with flexibility. We discuss some applications of the suggested ATE-GL framework and showcase its validity through some numerical examples. This is a joint work with Jianging Fan, Yingying Fan, Fan Yang and Diwen Yu.

Graphs and manifolds are commonly associated with sequence data such as texts. To enable text modeling and token generation, one may first construct Word2Vec embeddings of individual words and then build a graph of short sequences, where each short sequence can be viewed as a node of the graph and also be viewed as a point in a latent low-dimensional manifold. The link strengths between each pair of nodes can be calculated using a certain similarity measure of the embedding vectors, giving rise to a high-dimensional random matrix representing the graph data. For network applications, an important question is how to uncover the latent structural information underlying the graphs via low-dimensional manifold representations, often much lower than the ambient embedding dimensionality of each node. The Laplacian matrices for network data have been widely used to construct latent embeddings of graphs, where the nodes of the graph are represented in a latent subspace spanned by the corresponding leading eigenvectors of the Laplacian matrix. A natural question is how to characterize the asymptotic distributions of the leading eigenvectors and eigenvalues of the Laplacian matrix. The existing results in random matrix theory (RMT) have focused almost always on the setting of independent entries modulo symmetry, which is a major bottleneck of new RMT developments. Due to the use of the normalization terms, the Laplacian matrix is an example of a random matrix with dependency. To enable more flexible latent embeddings of graphs, we will extend the concept of the Laplacian matrix to that of the generalized (regularized) Laplacian matrix with index $\alpha \in [0, \infty)$. A key question we aim to address in this paper is how to characterize the asymptotic distributions of the leading eigenvectors and eigenvalues of the generalized (regularized) Laplacian matrices, a *new* class of high-dimensional random matrices with *dependency* representing the network data.

The primary objective of this paper is to investigate the asymptotic behaviors of the empirical spiked eigenvalues and eigenvectors of the generalized (regularized) Laplacian matrix (with some commonly used regularization terms) for the signal-plus-noise model when the signals are above a certain threshold. In particular, we will derive both the law of large numbers (LLN) and central limit theorems (CLTs) for the spiked sample eigenvalues and eigenvector components. Our results extend significantly previous the works [Fan, Fan, Han, and LvFan et al.2022a, Fan, Fan, Lv, and YangFan et al.2024] to the context of the generalized Laplacian matrix framework. These prior studies established the LLN and CLTs for spiked sample eigenvalues and eigenvector components of the adjacency matrices of large networks, which can be viewed as a special case of our results when $\alpha = 0$. Our results also compensate for the results of a recent work [Ke and WangKe and Wang2024], where entrywise large-deviation bounds for the eigenvectors associated with the largest eigenvalues of the Laplacian matrix for the DCMM model were established through the leave-one-out strategy. Additionally, in [Tang and PriebeTang and Priebe2018], the CLTs for the components of eigenvectors pertaining to the adjacency matrix and the Laplacian matrix of a random dot product graph were established, under the assumption of a prior distribution on the mean adjacency matrix.

Our results can be of independent theoretical interest due to the important role played by Laplacian matrices in the spectral graph theory. On the other hand, they can also serve as crucial ingredients for statistical inference concerning large networks and more general models. For example, they may enhance the characterization of the community membership probability matrix Π through spectral clustering methods for community detection, a widely used and scalable tool in the literature, as demonstrated in [Von LuxburgVon Luxburg2007, AbbeAbbe2017, JinJin2015, Le, Levina, and VershyninLe et al.2016, Lei and RinaldoLei and Rinaldo2015, Rohe, Chatterjee, and YuRohe et al.2011], or may enable hypothesis testing with network data, a prevalent technique utilized in various contexts such as [Arias-Castro and VerzelenArias-Castro and Verzelen2014,

Verzelen and Arias-CastroVerzelen and Arias-Castro2015,

Bickel and SarkarBickel and Sarkar2016,

LeiLei2016,

Wang and BickelWang and Bickel2017, Fan, Fan, Han, and LvFan et al.2022b, Fan, Fan, Lv, and YangFan et al.2024]. Due to the length constraint, we leave the investigation of various important applications of our theoretical results obtained in this paper to future work.

References

- [AbbeAbbe2017] Abbe, E. (2017). Community detection and stochastic block models: recent developments. *Journal of Machine Learning Research 18*(1), 6446–6531.
- [Arias-Castro and VerzelenArias-Castro and Verzelen2014] Arias-Castro, E. and N. Verzelen (2014). Community detection in dense random networks. *The Annals of Statistics* 42(3), 940–969.
- [Bickel and SarkarBickel and Sarkar2016] Bickel, P. J. and P. Sarkar (2016). Hypothesis testing for automated community detection in networks. *Journal of the Royal Statistical Society Series B* 78(1), 253–273.
- [Fan, Fan, Han, and LvFan et al.2022a] Fan, J., Y. Fan, X. Han, and J. Lv (2022a). Asymptotic theory of eigenvectors for random matrices with diverging spikes. *Journal of the American Statistical Association 117*, 996–1009.
- [Fan, Fan, Han, and LvFan et al.2022b] Fan, J., Y. Fan, X. Han, and J. Lv (2022b). SIMPLE: statistical inference on membership profiles in large networks. *Journal of the Royal Statistical Society Series B* 84, 630–653.
- [Fan, Fan, Lv, and YangFan et al.2024] Fan, J., Y. Fan, J. Lv, and F. Yang (2024). SIMPLE-RC: group network inference with non-sharp nulls and weak signals. *arXiv preprint arXiv:2211.00128*.
- [JinJin2015] Jin, J. (2015). Fast community detection by SCORE. *The Annals of Statistics* 43(1), 57–89.
- [Ke and WangKe and Wang2024] Ke, Z. T. and J. Wang (2024). Optimal network membership estimation under severe degree heterogeneity. *Journal of the American Statistical Association* (just-accepted), 1–28.
- [Le, Levina, and VershyninLe et al.2016] Le, C. M., E. Levina, and R. Vershynin (2016). Optimization via low-rank approximation for community detection in networks. *The Annals of Statistics* 44(1), 373–400.
- [LeiLei2016] Lei, J. (2016). A goodness-of-fit test for stochastic block models. *The Annals of Statistics* 44(1), 401–424.
- [Lei and RinaldoLei and Rinaldo2015] Lei, J. and A. Rinaldo (2015). Consistency of spectral clustering in stochastic block models. *The Annals of Statistics* 43(1), 215–237.
- [Rohe, Chatterjee, and YuRohe et al.2011] Rohe, K., S. Chatterjee, and B. Yu (2011). Spectral clustering and the high-dimensional stochastic blockmodel. *The Annals of Statistics* 39(4), 1878–1915.
- [Tang and PriebeTang and Priebe2018] Tang, M. and C. E. Priebe (2018). Limit theorems for eigenvectors of the normalized Laplacian for random graphs. *The Annals of Statistics* 46(5), 2360–2415.
- [Verzelen and Arias-CastroVerzelen and Arias-Castro2015] Verzelen, N. and E. Arias-Castro (2015). Community detection in sparse random networks. *The Annals of Applied Probability* 25(6), 3465–3510.
- [Von LuxburgVon Luxburg2007] Von Luxburg, U. (2007). A tutorial on spectral clustering. *Statistics and Computing* 17, 395–416.
- [Wang and BickelWang and Bickel2017] Wang, Y. R. and P. J. Bickel (2017). Likelihood-based model selection for stochastic block models. *The Annals of Statistics* 45(2), 500–528.

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GRAPH MATRICES AND TENSOR NETWORKS

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Keywords: graph matrices, tensor networks, tensors

Graph matrices are a type of matrix which is a powerful tool for analyzing problems on random inputs. Graph matrices have been used extensively for sum of squares lower bounds on average case problems [5, 11, 10, 19, 15, 18, 16, 24, 17, 21] and have also recently been used to analyze power-sum decompositions of polynomials [3], to analyze the ellipsoid fitting conjecture [20], [13], and to analyze a class of first-order iterative algorithms including belief propagation and approximate message passing [14]. That said, we only have a partial understanding of graph matrices. We currently know the following about graph matrices:

- (1) We have general norm bounds for graph matrices [2, 15, 22, 4, 17, 23, 24].
- (2) The limiting distribution of the singular values as $n \to \infty$ has been determined for a family of graph matrices called multi-Z-shaped graph matrices [7, 8].
- (3) A certain family of graph matrices behaves like Hermite polynomials of Gassian random variables [14].

When the random input is $G(n, \frac{1}{2})$, graph matrices are defined as follows:

Definition 0.1 (Fourier characters over $G(n, \frac{1}{2})$). Given a set of potential edges E, we define $\chi_E(G) = (-1)^{|E \setminus E(G)|} = \prod_{e \in E} \chi_{\{e\}}(G)$ where $\chi_{\{e\}}(G) = 1$ if $e \in E(G)$ and -1 if $e \notin E(G)$.

Proposition 0.2. $E_{G \sim G(n, \frac{1}{\alpha})}[\chi_E(G)\chi_{E'}(G)] = 1$ if E' = E and 0 if $E' \neq E$.

Definition 0.3 (Shapes). A shape α consists of a graph with vertices $V(\alpha)$ and edges $E(\alpha)$ together with two distinguished tuples of vertices U_{α} and V_{α} which are subsets of $V(\alpha)$.

Definition 0.4 (Graph matrices). Given a shape α , we define the graph matrix M_{α} to be the $\frac{n!}{(n-|U_{\alpha}|)!} \times \frac{n!}{(n-|V_{\alpha}|)!}$ matrix whose rows and columns are indexed by tuples of size $|U_{\alpha}|$ and $|V_{\alpha}|$ with entries

$$M_{\alpha}(A,B) = \sum_{\pi: V(\alpha) \to V(G): \pi \text{ is injective}, \pi(U_{\alpha}) = A, \pi(V_{\alpha}) = B} \chi_{\pi(E(\alpha))(G)}$$

Definition 0.5. A vertex separator of a shape α is a set of vertices $S \subseteq V(\alpha)$ such that every path from U_{α} to V_{α} (including paths of length 0) must contain a vertex in S.

Theorem 0.6 (AMP20). For all shapes α which have no isolated vertices outside of U_{α} and V_{α} , with high probability, $||M_{\alpha}||$ is $\tilde{O}(n^{\frac{|V(\alpha)|-s_{\alpha}}{2}})$ where s_{α} is the minimum size of a vertex separator of α and the \tilde{O} contains factors depending on the size of α and logarithmic factors.

In my talk, I started by describing tensor networks (using the paper "Hand-waving and Interpretive Dance: An Introductory Course on Tensor Networks" [6] as a guide). I then described graph matrices, norm bounds on graph matrices, and the close connection between tensor networks and graph matrices. In particular, tensor networks which are flattened into matrices can be transformed into graph matrices by replacing the indices with vertices and replacing the matrix/tensor entries with edges/hyperedges. Finally, I illustrated the power of graph matrices by showing how they can be used to easily rederive part of the analysis for tensor PCA, the faster tensor PCA algorithm in [12], and the tensor decomposition algorithm in [9].

REFERENCES

- [1] Kwangjun Ahn, Dhruv Medarametla, and Aaron Potechin. Graph Matrices: Norm Bounds and Applications. arXiv 1604.03423, 2020.
- [2] Mitali Bafna, Jun-Ting Hsieh, Pravesh Kothari, and Jeff Xu. Polynomial-Time Power-Sum Decomposition of Polynomials. FOCS 2022.
- [3] Afonso Bandeira, Kevin Lucca, Petar Nizić-Nikolac, and Ramon van Handel. Matrix Chaos Inequalities and Chaos of Combinatorial Type. STOC 2025.
- [4] Boaz Barak, Samuel Hopkins, Jonathan Kelner, Pravesh Kothari, Ankur Moitra, and Aaron Potechin. A Nearly Tight Sum-of-Squares Lower Bound for the Planted Clique Problem. SIAM Journal on Computing Vol. 48, Issue 2, p.687-735, 2019.
- [5] Jacob Bridgeman and Christopher Chubb. Hand-waving and Interpretive Dance: An Introductory Course on Tensor Networks. arXiv:1603.03039, 2017.
- [6] Wenjun Cai and Aaaron Potechin. The Spectrum of the Singular Values of Z-Shaped Graph Matrices. arXiv 2006.14144, 2020.
- [7] Wenjun Cai and Aaaron Potechin. On Mixing Distributions Via Random Orthogonal Matrices and the Spectrum of the Singular Values of Multi-Z Shaped Graph Matrices. arXiv 2206.02224, 2022.
- [8] Rong Ge and Tengyu Ma. Decomposing overcomplete 3rd order tensors using sum-of-squares algorithms. APPROX/RANDOM 2015.
- [9] Mrinalkanti Ghosh, Fernando Granha Jeronimo, Chris Jones, Aaron Potechin, and Goutham Rajendran. Sum-of-Squares Lower Bounds for Sherrington-Kirkpatrick via Planted Affine Planes. FOCS 2020.
- [10] Samuel Hopkins, Pravesh Kothari, Aaron Potechin. Prasad Raghavendra, Tselil Schramm, and David Steurer. The Power of Sum-of-Squares for Detecting Hidden Structures. FOCS 2017.
- [11] Samuel Hopkins, Tselil Schramm, Jonathan Shi, and David Steurer. Fast spectral algorithms from sum-of-squares proofs: tensor decomposition and planted sparse vectors. STOC 2016.
- [12] Jun-Ting Hsieh, Pravesh Kothari, Aaron Potechin, and Jeff Xu. Ellipsoid Fitting Up to a Constant. ICALP 2023.
- [13] Chris Jones and L. Pesenti. Fourier Analysis of Iterative Algorithms. arXiv 2404.07881, 2024.
- [14] Chris Jones, Aaron Potechin, Goutham Rajendran, Madhur Tulsiani, and Jeff Xu. Sum-of-Squares Lower Bounds for Sparse Independent Set. FOCS 2021.
- [15] Chris Jones, Aaron Potechin, Goutham Rajendran, and Jeff Xu. Sum-of-Squares Lower Bounds for Densest k-Subgraph. STOC 2023.
- [16] Pravesh Kothari, Aaron Potechin, and Jeff Xu. Sum-of-Squares Lower Bounds for Independent Set on Ultra-Sparse Random Graphs. STOC 2024.
- [17] Shuo Pang. SOS lower bound for exact planted clique. CCC 2021.
- [18] Aaron Potechin and Goutham Rajendran. Machinery for Proving Sum-of-Squares Lower Bounds on Certification Problems. arXiv 2011.04253, 2020.
- [19] Aaron Potechin, Paxton Turner, Prayaag Venkat, and Alex Wein. Near-optimal fitting of ellipsoids to random points. COLT 2023.
- [20] Aaron Potechin and Jeff Xu. Sum-of-Squares Lower Bounds for Coloring Random Graphs. STOC 2025.

- [21] Goutham Rajendran and Madhur Tulsiani. Concentration of polynomial random matrices via Efron-Stein inequalities. SODA 2023.
- [22] Madhur Tulsiani and June Wu. Simple Norm Bounds for Polynomial Random Matrices via Decoupling. ITCS 2025.
- [23] Jeff Xu. Switching Graph Matrix Norm Bounds: from i.i.d. to Random Regular Graphs. CCC 2025.

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REGRESSION UNDER NETWORK INTERFERENCE

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Keywords: Dependent Data, Generalized Linear Models, Minorization-Maximization, Pseudo-Likelihood

This extended abstract is based on Fritz, Schweinberger, Bhadra, and Hunter (2024) and Stewart and Schweinberger (2025).

1. Network interference

In connected populations, the treatments and outcomes of units can affect the outcomes of other units, which implies that the outcomes of units are interdependent. To study causal and non-causal relationships among attributes under network interference, a comprehensive regression framework for dependent predictors X, outcomes Y, and connections Z is needed.

2. Regression under network interference

We introduce a comprehensive regression framework for dependent predictors X, outcomes Y, and connections Z (Fritz et al., 2024). The regression framework can be used for studying non-causal and causal relationships among attributes (X, Y) of connected units and captures attribute-attribute, attribute-connection, and connection-connection dependencies, while retaining the advantages of linear regression, logistic regression, and other regression models by being interpretable and widely applicable. Scalable statistical computing is based on convex optimization of pseudo-likelihoods using minorization-maximization algorithms. An application to hate speech on social media demonstrates the advantages of the regression framework.

3. THEORETICAL GUARANTEES

Theoretical guarantees for regression under network interference are non-trivial, because the outcomes and connections $(\mathbf{Y}, \mathbf{Z}) \mid \mathbf{X} = \mathbf{x}$ are dependent. We provide theoretical guarantees by generalizing results of Stewart and Schweinberger (2025) for dependent connections \mathbf{Z} to dependent outcomes and connections $(\mathbf{Y}, \mathbf{Z}) \mid \mathbf{X} = \mathbf{x}$.

Lemma 1 of Stewart and Schweinberger (2025). Let $g : \mathbb{R}^p \mapsto \mathbb{R}^p$ $(p \ge 1)$ be a homeomorphism and $\|\cdot\|$ be a vector norm with induced matrix norm $\|\cdot\|$. Consider any $\theta^* \in \mathbb{R}^p$ and any $\epsilon > 0$, and define

$$\delta(\epsilon) \ \coloneqq \ \inf_{\boldsymbol{\theta} \,\in \, \mathrm{bd}\, \mathscr{B}(\boldsymbol{\theta}^\star, \epsilon)} \, \|g(\boldsymbol{\theta}) - g(\boldsymbol{\theta}^\star)\|,$$

where $\mathscr{B}(\boldsymbol{c}, \rho) \coloneqq \{\boldsymbol{a} \in \mathbb{R}^p : \|\boldsymbol{a} - \boldsymbol{c}\| < \rho\}$ is a ball with center $\boldsymbol{c} \in \mathbb{R}^p$ and radius $\rho > 0$ and $\operatorname{bd} \mathscr{B}(\boldsymbol{c}, \rho)$ is the boundary of $\mathscr{B}(\boldsymbol{c}, \rho)$. If $g(\boldsymbol{\theta})$ is continuously differentiable and

 $\mathcal{I}(\boldsymbol{\theta}) \coloneqq \nabla_{\boldsymbol{\theta}} g(\boldsymbol{\theta})$ is invertible for all $\boldsymbol{\theta} \in \mathscr{B}(\boldsymbol{\theta}^{\star}, \epsilon)$, then

$$\frac{\epsilon}{\sup_{\boldsymbol{\theta} \in \mathscr{B}(\boldsymbol{\theta}^{\star},\epsilon)} \|\!\!|\!| \mathcal{I}(\boldsymbol{\theta})^{-1} \|\!\!|} \leq \delta(\epsilon). \quad \Box$$

Lemma 1 helps "transport" concentration-of-measure between homeomorphic spaces, facilitating rates of convergence. To demonstrate, consider regression models with exponential-family densities of the form $f_{\theta^*}(t) \propto e^{\langle \theta^*, t \rangle}$, where $\theta^* \in \mathbb{R}^p$ and $\mu(\theta^*) \coloneqq \mathbb{E}_{\theta^*} T \in \mathbb{R}^p$ are the data-generating natural and mean-value parameters of the exponential family, and $\hat{\theta}$ and $\mu(\hat{\theta}) = T$ are the maximum likelihood estimators of θ^* and $\mu(\theta^*) = \mathbb{E}_{\theta^*} T$, respectively. Since the natural and mean-value parameter spaces of exponential families are homeomorphic, Lemma 1 implies that

$$\begin{split} \mathbb{P}(\boldsymbol{\theta} \in \mathscr{B}(\boldsymbol{\theta}^{\star}, \epsilon)) &= \mathbb{P}(\boldsymbol{T} \in \boldsymbol{\mu}(\mathscr{B}(\boldsymbol{\theta}^{\star}, \epsilon))) & \text{because } \boldsymbol{\mu} \text{ is a homeomorphism} \\ &\geq \mathbb{P}(\boldsymbol{T} \in \mathscr{B}(\boldsymbol{\mu}(\boldsymbol{\theta}^{\star}), \delta(\epsilon))) & \text{by definition of } \delta(\epsilon) \\ &\geq 1 - \alpha(\delta(\epsilon)) & \text{by concentration of } \boldsymbol{T} \\ &\geq 1 - \alpha\left(\frac{\epsilon}{\sup_{\boldsymbol{\theta} \in \mathscr{B}(\boldsymbol{\theta}^{\star}, \epsilon)} \||\mathcal{I}(\boldsymbol{\theta})^{-1}\||}\right) & \text{by Lemma 1 applied to } \boldsymbol{\mu}, \end{split}$$

where $\alpha(.)$ is a non-increasing function that quantifies the strength of concentration of Taround $\mu(\theta^*) = \mathbb{E}_{\theta^*} T$. In other words: If the probability mass of $\mu(\hat{\theta}) = T$ concentrates around $\mu(\theta^*) = \mathbb{E}_{\theta^*} T$, then the probability mass of $\hat{\theta}$ concentrates around θ^* , paving the way for convergence rates for $\hat{\theta}$ based on $\mu(\hat{\theta}) = T$ (compare Theorems 1 and 2 of Stewart and Schweinberger, 2025). While specific convergence rates depend on additional properties of the data-generating model, the above argument suggests that the convergence rate of maximum likelihood estimators $\hat{\theta}$ depends on

- the precision in a neighborhood of θ^* as quantified by $\sup_{\theta \in \mathscr{B}(\theta^*, \epsilon)} \| \mathcal{I}(\theta)^{-1} \|$;
- the strength of concentration of *T* as quantified by α(.), which depends on the tails of the distribution of *T* and the dependence induced by the model.

The above argument applies to all exponential families (e.g., generalized linear models, graphical models, and Gaussian and non-Gaussian Markov random fields), and helps establish theoretical guarantees for regression based on independent or dependent observations, including regression under network interference (Fritz et al., 2024).

REFERENCES

- Fritz, C., M. Schweinberger, S. Bhadra, and D. R. Hunter (2024). A regression framework for studying relationships among attributes under network interference. *Available from: arXiv:2410.07555*.
- Stewart, J. R. and M. Schweinberger (2025). Pseudo-likelihood-based *M*-estimators for random graphs with dependent edges and parameter vectors of increasing dimension. *The Annals of Statistics*. To appear.

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A MULTILAYER PROBIT NETWORK MODEL FOR COMMUNITY DETECTION WITH DEPENDENT LAYERS

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Classification AMS 2020: 62A09, 91D30

Keywords: inter-layer dependence; sparse networks; multivariate probit model; stochastic block model.

Multilayer networks often exhibit various dependence structures between network layers. Various inter-layer dependence modeling highlights the importance of incorporating such dependencies for more accurate and efficient network analysis. For example, [3] introduced the autoregressive stochastic block model (SBM) to capture inter-layer dependence with a time series structure; [7] proposed the multilayer Ising model to capture the full inter-layer dependence. However, it remains unclear how to extend [3] to accommodate more general dependence structures, whereas the method in [7] appears to have difficulty in estimating connection probabilities due to the intractable computation cost of the partition function [5]. Moreover, very little has been done in the literature to theoretically investigate the impact of dependence structures on the community detection accuracy.

In this work, we introduce a novel multilayer probit network model that integrates the classical multilayer SBM [4, 2] with the multivariate probit model [1]. It incorporates diverse inter-layer dependence structures between layers into network modeling so as to achieve better estimation of the homogeneous community structure.

Let \mathcal{G} denote a multilayer network comprising M network layers on N common nodes, where each network layer can be represented via its adjacency matrix $\mathbf{A}^{(b)} = (A_{ij}^{(b)})_{N \times N} \in \{0,1\}^{N \times N}$ for $b \in [M]$. Here, $A_{ij}^{(b)} = A_{ji}^{(b)} = 1$ if an edge exists between nodes i and j in the b-th layer, and $A_{ij}^{(b)} = A_{ji}^{(b)} = 0$ otherwise. We consider the following multilayer probit network model,

$$\begin{aligned} A_{ij}^{(b)} &= \mathbb{I} \big\{ \mu_{e_i e_j}^{(b)} + \varepsilon_{ij}^{(b)} > 0 \big\}, \text{ for any } b \in [M], \\ \big(\varepsilon_{ij}^{(1)}, \cdots, \varepsilon_{ij}^{(M)} \big)^\top &\sim N \big(0, \mathbf{\Sigma}_{e_i e_j} \big), \text{ for any } i \neq j, \end{aligned}$$

where $\mathbb{I}(\cdot)$ is the indicator function, $e_i \in [K]$ denotes the homogeneous community membership of node *i* across *M* layers, $\boldsymbol{\mu}^{(b)} \in \mathbb{R}^{K \times K}$ denotes the mean matrix for each network layer, and $\boldsymbol{\Sigma}_{kl} \in \mathbb{R}^{M \times M}$ is positive definite for any $k, l \in [K]$. Note that $P(A_{ij}^{(b)} = 1) = P(\varepsilon_{ij}^{(b)} > -\mu_{e_ie_j}^{(b)}) = \Phi(\mu_{e_ie_j}^{(b)})$, where $\Phi(\cdot)$ is the cumulative distribution function of N(0, 1).

Let $\boldsymbol{\mu} = (\mu_{kl}^{(b)})_{k,l \in [K], b \in [M]}$ and $\boldsymbol{\mu}^{(b)} = (\mu_{kl}^{(b)})_{k,l \in [K]}$ for each $b \in [M]$. Further, let $\boldsymbol{\Sigma} = (\Sigma_{kl}^{(bd)})_{k,l \in [K], b, d \in [M]}$ and $\boldsymbol{\Sigma}_{kl} = (\Sigma_{kl}^{(bd)})_{b,d \in [M]}$ for any $k, l \in [K]$. Define $\boldsymbol{\Theta} = (\boldsymbol{\mu}, \boldsymbol{\Sigma})$ with $\boldsymbol{\Theta}_{kl}^{(bd)} = (\mu_{kl}^{(b)}, \mu_{kl}^{(d)}, \Sigma_{kl}^{(bd)})$. Denote $\boldsymbol{Z} = (Z_{ik})_{i \in [N]; k \in [K]}$ as the homogeneous community membership matrix, where $Z_{ik} = 1$ if $e_i = k$, and $Z_{ik} = 0$ otherwise. Since the full likelihood of the multilayer network is computationally inefficient, we consider

a pairwise likelihood function as an alternative, which largely facilitates the computation without compromising estimation accuracy [6]. Specifically, we replace the full likelihood $\mathbb{P}(\mathbf{A}_{ij}; \{\mu_{kl}^{(b)}\}_{b=1}^{M}, \boldsymbol{\Sigma}_{kl})$ with

$$\prod_{1 \le b < d \le M} \mathbb{P}\left(A_{ij}^{(b)}, A_{ij}^{(d)}; \boldsymbol{\Theta}_{kl}^{(bd)}\right),$$

where

$$\mathbb{P}(A_{ij}^{(b)}, A_{ij}^{(d)}; \Theta_{kl}^{(bd)}) = \alpha_1 \left(\Theta_{kl}^{(bd)}\right)^{A_{ij}^{(b)}A_{ij}^{(d)}} \times \alpha_2 \left(\Theta_{kl}^{(bd)}\right)^{A_{ij}^{(b)}(1-A_{ij}^{(d)})} \times \alpha_3 \left(\Theta_{kl}^{(bd)}\right)^{(1-A_{ij}^{(b)})A_{ij}^{(d)}} \times \alpha_4 \left(\Theta_{kl}^{(bd)}\right)^{(1-A_{ij}^{(b)})(1-A_{ij}^{(d)})}$$

The terms $\alpha_1, \alpha_2, \alpha_3$ and α_4 are defined as (0.1)

$$\begin{aligned} \alpha_1(\boldsymbol{\Theta}_{kl}^{(bd)}) &= \mathbb{P}(A_{ij}^{(b)} = 1, A_{ij}^{(d)} = 1; \boldsymbol{\Theta}_{kl}^{(bd)}) = \Phi_2(\mu_{kl}^{(b)}, \mu_{kl}^{(d)}, \Sigma_{kl}^{(bd)}), \\ \alpha_2(\boldsymbol{\Theta}_{kl}^{(bd)}) &= \mathbb{P}(A_{ij}^{(b)} = 1, A_{ij}^{(d)} = 0; \boldsymbol{\Theta}_{kl}^{(bd)}) = \Phi(\mu_{kl}^{(b)}) - \Phi_2(\mu_{kl}^{(b)}, \mu_{kl}^{(d)}, \Sigma_{kl}^{(bd)}), \\ \alpha_3(\boldsymbol{\Theta}_{kl}^{(bd)}) &= \mathbb{P}(A_{ij}^{(b)} = 0, A_{ij}^{(d)} = 1; \boldsymbol{\Theta}_{kl}^{(bd)}) = \Phi(\mu_{kl}^{(d)}) - \Phi_2(\mu_{kl}^{(b)}, \mu_{kl}^{(d)}, \Sigma_{kl}^{(bd)}), \\ \alpha_4(\boldsymbol{\Theta}_{kl}^{(bd)}) &= \mathbb{P}(A_{ij}^{(b)} = 0, A_{ij}^{(d)} = 0; \boldsymbol{\Theta}_{kl}^{(bd)}) = 1 - \Phi(\mu_{kl}^{(b)}) - \Phi(\mu_{kl}^{(b)}) + \Phi_2(\mu_{kl}^{(b)}, \mu_{kl}^{(d)}, \Sigma_{kl}^{(bd)}), \end{aligned}$$

where $\Phi(\cdot)$ is the cumulative distribution function of N(0,1), and $\Phi_2(\cdot, \cdot, \sigma)$ is the cumulative distribution function of $N_2(\begin{pmatrix} 0 \\ 0 \end{pmatrix}, \begin{pmatrix} 1 & \sigma \\ \sigma & 1 \end{pmatrix})$. The pairwise log-likelihood then becomes

$$\mathcal{L}(\Theta, \mathbf{Z}) = \sum_{k,l} \sum_{i,j} \sum_{b < d} Z_{ik} Z_{jl} \Big\{ A_{ij}^{(b)} A_{ij}^{(d)} \log \alpha_1(\Theta_{kl}^{(bd)}) + A_{ij}^{(b)} (1 - A_{ij}^{(d)}) \log \alpha_2(\Theta_{kl}^{(bd)}) \\ + (1 - A_{ij}^{(b)}) A_{ij}^{(d)} \log \alpha_3 \left(\Theta_{kl}^{(bd)}\right) + (1 - A_{ij}^{(b)}) (1 - A_{ij}^{(d)}) \log \alpha_4(\Theta_{kl}^{(bd)}) \Big\} \\ = : \sum_{k,l} \mathcal{L}_{kl}(\Theta, \mathbf{Z}).$$

Denote S_{kl} as the pre-specified, shape-constrained set for Σ_{kl} . Specifically, we focus on two scenarios, the sparse covariance matrix scenario with

$$\mathcal{S}_{kl} = \left\{ \boldsymbol{X} \in \mathbb{R}^{M \times M} \mid \boldsymbol{X} \succ 0, \operatorname{diag}(\boldsymbol{X}) = \mathbf{1}_M, \operatorname{Supp}(\boldsymbol{X}) = T_{kl} \right\},$$

and the sparse precision matrix scenario with

$$S_{kl} = \left\{ \boldsymbol{X} \in \mathbb{R}^{M \times M} \mid \boldsymbol{X} \succ 0, \operatorname{diag}(\boldsymbol{X}) = \mathbf{1}_M, \operatorname{Supp}(\boldsymbol{X}^{-1}) = T_{kl} \right\}.$$

In both cases, $T_{kl} \subseteq [M] \times [M]$ represents the set of positions, known a priori, with $|T_{kl}| = s_{kl}^*$. Two examples for each scenario are the multilayer Ising model [7] and the autoregressive SBM [3]. Define the parameter space as

$$\boldsymbol{\Omega} = \left\{ \boldsymbol{\omega} = (\boldsymbol{\mu}, \boldsymbol{\Sigma}, \boldsymbol{Z}) \mid \boldsymbol{Z} \in \{0, 1\}^{N \times K}, \ \boldsymbol{Z} \boldsymbol{1}_{K} = \boldsymbol{1}_{N}, \ c_{l} \rho_{N,M} \leq \Phi(\mu_{kl}^{(b)}) \leq c_{u} \rho_{N,M}, \\ \boldsymbol{\Sigma}_{kl} \in \mathcal{S}_{kl}, \ \text{and} \ \sup_{k,l} \| \operatorname{ndiag}(\boldsymbol{\Sigma}_{kl}) \|_{\max} \leq D_{N,M} \right\},$$

where $c_l < 1 < c_u$ are two constants and $\rho_{N,M}$ controls the network sparsity level. Note that the magnitudes of s_{kl}^* and $D_{N,M}$ specify the inter-layer dependence structures and the strength of dependence across different layers, respectively. Denote the true

parameters as $\omega^* = (\Theta^*, Z^*) = (\mu^*, \Sigma^*, Z^*)$ and assume $\omega^* \in \Omega$. Lemma 0.1 shows that the pairwise likelihood function in (0.2) is Fisher consistent in Ω .

Lemma 0.1. Let $e(\boldsymbol{\omega}^*, \boldsymbol{\omega}) = \frac{1}{N^2 M^2} \sum_{k,l} \mathbb{E} (\mathcal{L}_{kl}(\boldsymbol{\Theta}^*, \boldsymbol{Z}^*) - \mathcal{L}_{kl}(\boldsymbol{\Theta}, \boldsymbol{Z}))$, then it holds true that $e(\boldsymbol{\omega}^*, \boldsymbol{\omega}) \geq 0$ for any $\boldsymbol{\omega} \in \boldsymbol{\Omega}$.

Lemma 0.1 shows that ω^* is a maximizer of $\mathbb{E}(\mathcal{L}(\Theta, \mathbb{Z}))$, and thus justifies the use of the pairwise likelihood function in estimating ω^* . Therefore, we estimate (Θ^*, \mathbb{Z}^*) via the constrained maximum pairwise log-likelihood estimate,

(0.2)
$$(\widehat{\Theta}, \widehat{Z}) = \operatorname*{argmax}_{(\Theta, Z) \in \Omega} \mathcal{L}(\Theta, Z).$$

We also adopt an alternative updating algorithm to solve the constrained optimization problem. Theoretically, we establish the asymptotic consistency of the proposed method for both parameter estimation and community detection under mild conditions.

We demonstrate how the inter-layer dependence structures and strength affect the accuracy of community detection in theory. In the autoregressive SBM, the proposed method exhibits a smaller misclassification rate than [3] when $\rho_{N,M} \gtrsim \frac{1}{\log(NM)}$ and $M \leq N$. In the multilayer Ising model [7] with $K \leq \log(NM), s_{kl}^* \asymp M^2, M \asymp N$, the required sparsity condition there is that $\rho_{N,M} \gg (\frac{1}{N})^{\frac{1}{1+c}}$ for some constant c > 0, up to some logarithmic terms. In contrast, the proposed method can achieve $\rho_{N,M} \gg \frac{1}{N}$, up to some logarithmic terms, which achieves a better sparsity condition. Moreover, through extensive simulations and a real-world multilayer international trade network, we demonstrate the superior numerical performance of the proposed method compared to several popular competitors.

REFERENCES

- [1] Patrick J Heagerty and Subhash R Lele. A composite likelihood approach to binary spatial data. *Journal of the American Statistical Association*, 93(443):1099–1111, 1998.
- [2] Paul W Holland, Kathryn Blackmond Laskey, and Samuel Leinhardt. Stochastic blockmodels: First steps. *Social networks*, 5(2):109–137, 1983.
- [3] Binyan Jiang, Jialiang Li, and Qiwei Yao. Autoregressive networks. *Journal of Machine Learning Research*, 24(227):1–69, 2023.
- [4] Jing Lei, Kehui Chen, and Brian Lynch. Consistent community detection in multilayer network data. *Biometrika*, 107(1):61–73, 2020.
- [5] Pradeep Ravikumar, Martin J Wainwright, and John D Lafferty. High-dimensional ising model selection using l1-regularized logistic regression. *The Annals of Statistics*, 38(3):1287–1319, 2010.
- [6] Cristiano Varin, Nancy Reid, and David Firth. An overview of composite likelihood methods. *Statistica Sinica*, 21(1):5–42, 2011.
- [7] Jingnan Zhang, Junhui Wang, and Xueqin Wang. Consistent community detection in inter-layer dependent multi-layer networks. *Journal of the American Statistical Association*, 119(548):3141–3151, 2024.

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ONLINE INFERENCE FOR LOW-RANK REINFORCEMENT LEARNING

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Keywords: online inference, online decision-making, low-rank models, reinforcement learning, stochastic gradient descent.

Reinforcement learning (RL) deals with how intelligent agents leverage contextual information and historical data to take actions in an uncertain environment in order to maximize the cumulative reward [1]. It has achieved phenomenal success in diverse fields, such as video games, robotics, autonomous driving, precision medicine, and recommendation systems. In modern applications, such context format can be rich and can often be formulated as a matrix or higher order tensor. This is evident in scenarios such as monitoring brain activity in real-time during clinical research, tracking dynamic user preferences in online recommender systems, and analyzing the evolving relationships in social network analysis. Consider neuroscience, where dynamic treatments may be tailored to a patient based on their neuroimaging. Here, the neuroimaging data forms a tensor state, while the treatment, such as dynamic sleep intervention, represents an action in the RL framework. Such high-dimensional higher-order tensor contexts necessitate the incorporation of low-rank structures in RL models.

Why inference in RL? While existing RL algorithms mainly focus on minimizing regret or choosing the best action with respect to some oracle policy, less attention has been paid to the statistical inference for RL models where the data are adaptively collected. In real-world applications of RL, we are often not just interested in obtaining the point estimate of the value function, but also a measure of the statistical uncertainty associated with the estimate. This is especially relevant to fields such as personalized medicine, mobile health and autonomous driving, where it is often risky to run a policy without a statistically sound estimate of its quality. For example, online A/B testing has been widely conducted by technological/pharmaceutical companies to compare a new product with an old one. Recent studies [2] have used various bandit or RL methods to form sequential online A/B testing procedures. In these online evaluation tasks, it is important to quantify the uncertainty of the point estimate for constructing a valid Moreover, the information obtained by conducting statistical hypothesis testing. inference of parameters or value functions, can eventually help experimenters to yield a better understanding in the used RL reward model, and this increase of knowledge can potentially improve the design of the experiments [3].

Why are new tools needed? When data is collected in an adaptive manner, even simple ordinary least squares can exhibit non-normal asymptotic behavior [3]. In this case, the confidence intervals constructed from traditional estimators induce bias and lead to wrong coverage. In extensive numerical studies, [4] empirically illustrate that common statistical hypothesis tests lead to as much as double the false positive rate and

false negative rate using adaptive data collected in the bandit setting. While the use of adaptively collected data for inferential purposes has gained popularity in recent years, existing inferential methods are primarily developed under simple settings. These include adaptive linear regression [5], linear bandit [3], batch Markov decision process [6], and linear stochastic approximation under Markov noise [7]. While these contributions serve as crucial initial steps in statistical inference for adaptive data, practical RL applications involving higher-order tensor contexts call for more sophisticated inference tools.

In this talk, we discuss provable online inferential tools tailored for low-rank reinforcement learning. We first introduce an efficient online low-rank stochastic gradient descent (SGD) method and establishes its non-asymptotic rate of convergence. Building upon this foundation, we propose a simple yet powerful online debiasing approach for the sequential statistical inference of low-rank tensor learning. The entire online procedure studied in this context, encompassing both estimation and inference, eliminates the need for data splitting or storing historical data, making it suitable for on-the-fly hypothesis testing. We then progress to low-rank contextual bandit by incorporating online decision-making policies, where sequential decisions rely on higher-order contextual information. By conducting hypothesis testing on entries of the parameter tensor, one can assess the impact of a specific region of the tensor context on the reward. The challenges of this inference arise from two sources of bias: the first due to the low-rank structure of the parameter, and the second originating from the decision-making policy, as the chosen action depends on all historical data. We discuss an online double debiasing procedure for statistical inference within the low-rank contextual bandit framework, and establish the validity of the resulting confidence interval. Additionally, we identify an intriguing tradeoff between parameter inference and regret minimization, prompting a formulation of this trade-off as a minimax multi-objective optimization problem.

References

- [1] R. S. Sutton and A. G. Barto. Reinforcement Learning: An Introduction. MIT Press, 2018.
- [2] C. Shi, S. Luo, H. Zhu, and R. Song. An online sequential test for qualitative treatment effects. *Journal of Machine Learning Research*, 22(286):1–51, 2021.
- [3] K. Zhang, L. Janson, and S. Murphy. Statistical inference with m-estimators on adaptively collected data. *Advances in Neural Information Processing Systems*, 34:7460–7471, 2021.
- [4] J. J. Williams, J. Nogas, N. Deliu, H. Shaikh, S. S. Villar, A. Durand, and A. Rafferty. Challenges in statistical analysis of data collected by a bandit algorithm: An empirical exploration in applications to adaptively randomized experiments. *arXiv preprint arXiv:2103.12198*, 2021.
- [5] K. Khamaru, Y. Deshpande, L. Mackey, and M. J. Wainwright. Near-optimal inference in adaptive linear regression. *arXiv preprint arXiv:2107.02266*, 2021.
- [6] C. Shi, S. Zhang, W. Lu, and R. Song. Statistical inference of the value function for reinforcement learning in infinite-horizon settings. *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, 2021.
- [7] P. Ramprasad, Y. Li, Z. Yang, Z. Wang, W. W. Sun, and G. Cheng. Online bootstrap inference for policy evaluation in reinforcement learning. *Journal of the American Statistical Association*, 2022 (just-accepted).

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THEORY AND APPLICATION OF HIGH-DIMENSIONAL SMOOTH TENSORS

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Keywords: Smooth tensors, Nonparametric Methods, Latent Permutation

Abstract. We consider the problem of structured tensor denoising in the presence of Such data problems arise commonly in recommendation unknown permutations. system, neuroimaging, community detection, and multiway comparison applications. Here, we develop a general family of smooth tensor models up to arbitrary index permutations; the model incorporates the popular tensor block models and Lipschitz hypergraphon models as special cases. We show that a constrained least-squares estimator in the block-wise polynomial family achieves the minimax error bound. A phase transition phenomenon is revealed with respect to the smoothness threshold needed for optimal recovery. In particular, we find that a polynomial of degree up to (m-2)(m+1)/2 is sufficient for accurate recovery of order-m tensors, whereas higher degree exhibits no further benefits. This phenomenon reveals the intrinsic distinction for smooth tensor estimation problems with and without unknown permutations. Furthermore, we provide an efficient polynomial-time Borda count algorithm that provably achieves optimal rate under monotonicity assumptions. The efficacy of our procedure is demonstrated through both simulations and Chicago crime data analysis.

Model. Let $\Theta \in \mathbb{R}^{d \times \cdots \times d}$ be an order-*m d*-dimensional tensor, $\pi : [d] \to [d]$ be an index permutation, and $\Theta(i_1, \ldots, i_m)$ the tensor entry indexed by (i_1, \ldots, i_m) . We sometimes also use shorthand notation $\Theta(\omega)$ for tensor entries with indices $\omega = (i_1, \ldots, i_m)$. Suppose we observe an order-*m d*-dimensional data tensor from the following model,

$$\mathcal{Y} = \Theta \circ \pi + \mathcal{E},$$

where \circ represents the function composition, $\pi: [d] \to [d]$ is an unknown latent permutation, $\Theta \in \mathbb{R}^{d \times \cdots \times d}$ is an unknown signal tensor under certain smoothness (to be specified in next paragraph), and \mathcal{E} is a noise tensor consisting of zero-mean, independent sub-Gaussian entries with variance bounded by σ^2 . The general model allows continuous- and binary-valued tensors. For instance, in binary tensor problems, the entries in \mathcal{Y} are $\{0, 1\}$ -labels from Bernoulli distribution, and the entrywise variance of \mathcal{E} depends on the mean. For ease of presentation, we assume $\sigma = 1$ throughout the paper. We call (0.1) the Gaussian model if the \mathcal{E} consists of i.i.d. $\mathcal{N}(0, 1)$ entries, and call (0.1) the sub-Gaussian model if \mathcal{E} consists of independently (but not necessarily identically) distributed sub-Gaussian entries.

We now describe the smooth model on the signal Θ . Suppose that there exists a multivariate function $f: [0,1]^m \to \mathbb{R}$ underlying the signal tensor, such that

(0.2)
$$\Theta(i_1,\ldots,i_m) = f\left(\frac{i_1}{d},\ldots,\frac{i_m}{d}\right), \text{ for all } (i_1,\ldots,i_m) \in [d]^m.$$

For a multi-index $\kappa = (\kappa_1, \ldots, \kappa_m) \in \mathbb{N}^m$ and a vector $\boldsymbol{x} = (x_1, \ldots, x_m)^T$, we denote $|\kappa| = \sum_{i \in [m]} \kappa_i$, $\kappa! = \prod_{i \in [m]} \kappa_i!$, $\boldsymbol{x}^{\kappa} = \prod_{i \in [m]} x_i^{\kappa_i}$, and the derivative operator $\nabla_{\kappa} = \frac{\partial^{|\kappa|}}{\partial x_1^{\kappa_1} \cdots \partial x_m^{\kappa_m}}$. The generative function f in (0.2) is assumed to be in the α -Hölder smooth family [5].

Definition 0.1 (α -Hölder smooth). Let $\alpha > 0$ and L > 0 be two positive constants. A function $f: [0,1]^m \to \mathbb{R}$ is called α -Hölder smooth, denoted as $f \in \mathcal{F}(\alpha, L)$, if

(0.3)
$$\sum_{\kappa:|\kappa|=\lceil\alpha-1\rceil} \frac{1}{\kappa!} |\nabla_{\kappa} f(\boldsymbol{x}) - \nabla_{\kappa} f(\boldsymbol{x}_0)| \le L \|\boldsymbol{x} - \boldsymbol{x}_0\|_{\infty}^{\alpha-\lceil\alpha-1\rceil}$$

holds for every $\boldsymbol{x}, \boldsymbol{x}_0 \in [0, 1]^m$.

The Hölder smooth function class is one of the most popular function classes considered in the nonparametric regression literature [3, 2]. In addition to the function class $\mathcal{F}(\alpha, L)$, we also define the smooth tensor class based on discretization (0.2),

(0.4) $\mathcal{P}(\alpha, L) = \{ \Theta \in \mathbb{R}^{d \times \dots \times d} : \Theta \text{ is generated from (0.2) and } f \in \mathcal{F}(\alpha, L) \}.$

Combining (0.1) and (0.2) yields our proposed *permuted smooth tensor model*. The unknown parameters are the smooth tensor $\Theta \in \mathcal{P}(\alpha, L)$ and latent permutation $\pi \in \Pi(d, d)$. The model is visualized in Figure 1(a) for the case m = 2 (matrices).



FIGURE 1. (a): Illustration of order-m d-dimensional permuted smooth tensor models with m = 2. (b): Phase transition of mean squared error (MSE) (on $-\log_d$ scale) as a function of smoothness α and tensor order m. Bold dots correspond to the critical smoothness level above which higher smoothness exhibits no further benefits to tensor estimation.

Results Summary. We develop a suite of statistical theory, efficient algorithms, and related applications for permuted smooth tensor models. Our contributions are summarized below. First, we develop a general permuted α -smooth tensor model of arbitrary smoothness level $\alpha > 0$. We establish the statistically optimal error rate and its dependence on model complexity. Specifically, we express the optimal rate as a function of tensor order *m*, tensor dimension *d*, and the smoothness level α , given by

(0.5)
$$\operatorname{Rate}(d) := d^{-\frac{2m\alpha}{m+2\alpha}} \vee d^{-(m-1)} \log d.$$

Our framework substantially generalizes earlier works which focus on only matrices with m = 2 [2, 3] or Lipschitz function with $\alpha = 1$ [1, 4]. The generalization enables us to obtain results previously impossible: i) As tensor order m increases, we demonstrate

the failure of pervious clustering-based algorithms [1, 2], and we develop a new block-wise polynomial algorithm for tensors of order $m \geq 3$; ii) As smoothness α increases, we demonstrate that the error rate converges to a fast rate $\mathcal{O}(d^{-(m-1)})$, thereby disproving the conjectured lower bound $\mathcal{O}(d^{-2m/(m+2)})$ posed by earlier work [1]. The results showcase the accuracy gain of our new approach, as well as the intrinsic distinction between matrices and higher-order tensors.

Second, we discover a phase transition phenomenon with respect to the smoothness needed for optimal recovery in the model (0.1) and (0.2). Figure 1(b) plots the dependence of estimation error in terms of smoothness level α for tensors of order m. We characterize two distinct error behaviors determined by a critical smoothness threshold. Specifically, the accuracy improves with α in the regime $\alpha \leq m(m-1)/2$, whereas the accuracy becomes a constant of α in the regime $\alpha > m(m-1)/2$. The results imply a polynomial of degree (m-2)(m+1)/2 = [m(m-1)/2 - 1] is sufficient for accurate recovery of order-m tensors of arbitrary smoothness in the model (0.1) and (0.2)., whereas higher degree brings no further benefits. The phenomenon is distinctive from matrix problems [3, 2] and classical *non-permuted* smooth function estimation [5], thereby highlighting the fundamental challenges in our new setting. These statistical contributions, to our best knowledge, are new to the literature of permuted smooth tensor problems.

Third, we propose two estimation algorithms with accuracy guarantees: the least-squares estimation and Borda count estimation. The least-squares estimation, although being computationally hard, reveals the fundamental model complexity in the problem. The result serves as the benchmark and a useful guide to the algorithm design. Furthermore, we develop an efficient polynomial-time Borda count algorithm that provably achieves a minimax optimal rate under an extra Lipschitz monotonic assumption. The algorithm handles a broad range of data types, including continuous and binary observations.

Lastly, we illustrate the efficacy of our method through both simulations and data applications. A range of practical settings are investigated in simulations, and we show the outperformance of our method compared to alternative approaches. Application to Chicago crime data is presented to showcase the usefulness of our method. We identify the key global pattern and pinpoint local smooth structure in the denoised tensor. Our method will help practitioners efficiently analyze tensor datasets in various areas. Toward this end, the package and all data used are released at CRAN link https://cloud.r-project.org/web/packages/SmoothTensor/index.html.

REFERENCES

[1] Balasubramanian, K. (2021). Nonparametric modeling of higher-order interactions via hypergraphons. *Journal of Machine Learning Research 22*, 1–25.

- [2] Gao, C., Y. Lu, and H. H. Zhou (2015). Rate-optimal graphon estimation. *The Annals of Statistics* 43(6), 2624–2652.
- [3] Klopp, O., A. B. Tsybakov, and N. Verzelen (2017). Oracle inequalities for network models and sparse graphon estimation. *The Annals of Statistics 45*(1), 316–354.
- [4] Li, Y., D. Shah, D. Song, and C. L. Yu (2019). Nearest neighbors for matrix estimation interpreted as blind regression for latent variable model. *IEEE Transactions on Information Theory* 66(3), 1760–1784.

[5] Tsybakov, A. B. (2009). *Introduction to nonparametric estimation*. Springer Science & Business Media.

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REGRET MINIMIZATION AND STATISTICAL INFERENCE IN ONLINE DECISION MAKING WITH HIGH-DIMENSIONAL COVARIATES

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Classification AMS 2020:

Keywords: linear contextual bandit, sparsity, ε -greedy, inference, regret

We Investigate regret minimization, statistical inference, and their interplay in high-dimensional online decision-making based on the sparse linear contextual bandit model. We integrate the ε -greedy bandit algorithm for decision-making with a hard thresholding algorithm for estimating sparse bandit parameters and introduce an inference framework based on a debiasing method using inverse propensity weighting. Under a margin condition, our method achieves either $O(\sqrt{T})$ regret or classical $O(\sqrt{T})$ -consistent inference, indicating an unavoidable trade-off between exploration and exploitation. If a diverse covariate condition holds, we demonstrate that a pure-greedy bandit algorithm-i.e., exploration-free-combined with a debiased estimator based on average weighting can simultaneously achieve optimal $O(\log T)$ regret and $O(\sqrt{T})$ -consistent inference. We also show that a simple sample mean estimator can provide valid inference for the optimal policy's value. Numerical simulations and experiments on Warfarin dosing data validate the effectiveness of our methods.

To the best of our knowledge, this work is the first to investigate regret minimization, statistical inference, and their interplay in high-dimensional online decision-making based on the sparse-LCB model. Our contributions are summarized as follows:

General Inference Framework and Tradeoff with Regret. We propose a novel statistical inference framework for adaptively collected high-dimensional data. Our approach integrates the ε -greedy bandit algorithm with hard-thresholding (HT), resulting in a biased estimator due to the adaptive data collection and implicit regularization introduced by the HT algorithm. To mitigate this bias, we introduce an online debiasing technique based on IPW that maintains low computational and storage complexity. Under a margin condition with parameter ν , the debiased estimator is asymptotically normal, enabling the construction of confidence intervals and hypothesis tests for both individual arm parameters and their differences. Additionally, we identify a trade-off between regret performance and the estimator's asymptotic variance, which affects inference efficiency by determining the width of confidence intervals and the p-values of Specifically, when the algorithm achieves a regret upper bound of test statistics. $O(T^{1-\gamma} + T^{(\gamma-1)(1+\nu)/2})$ with margin parameter ν , and some user-specified $\gamma \in [0, 1)$ —which characterizes the exploration probability, the estimator's asymptotic variance is $O(T^{-(1-\gamma)})$. For example, when $\nu = 1$, setting $\gamma = \frac{1}{2} + o(1)$ yields a regret bound of $O(T^{1/2})$ and an estimator variance of $O(T^{-1/2})$, which does not attain the classic \sqrt{T} -consistency; setting $\gamma = 0$ yields a trivial regret bound of O(T) and an

asymptotically normal estimator which is \sqrt{T} -consistent. While IPW is effective for debiasing, it unfortunately inflates the variance of the final estimator.

Simultaneous Optimal Inference and Regret. We demonstrate that optimal inference efficiency and regret performance can be simultaneously achieved under an additional covariate diversity assumption, commonly employed in high-dimensional bandit literature ([Bastani et al.(2021)Bastani, Bayati, and Khosravi, Ren and Zhou(2024)] and references therein). This assumption is motivated by the observation that when covariates are sufficiently diverse, an exploration-free algorithm (i.e., setting the exploration probability $\varepsilon = 0$ in the ε -greedy algorithm) can still adequately explore each arm. This automatic exploration facilitates debiasing through a simple average weighting (AW) approach, bypassing IPW and thereby avoiding variance inflation. Specifically, our approach achieves an optimal $O(\log T)$ regret upper bound, and the resulting estimators of arm parameters are asymptotically normal with a variance of $O(T^{-1})$, thereby attaining the classic \sqrt{T} -consistency and optimal inference efficiency. Additionally, we introduce an inference procedure for the optimal policy's value, often referred to as the Q-value, within this framework. We provide a straightforward method to assess the maximum total reward achievable by the optimal policy.

Empirical Result. We evaluate the empirical performance of our algorithm and inference framework through numerical simulations and a real-world data experiment. Specifically, we apply this framework to the aforementioned Warfarin dosing problem. Our approach identifies several significant variables that determine the appropriate dosage, with findings that are consistent with existing medical literature while also offering novel insights.

References

- [Bastani et al.(2021)Bastani, Bayati, and Khosravi] Hamsa Bastani, Mohsen Bayati, and Khashayar Khosravi. Mostly exploration-free algorithms for contextual bandits. *Management Science*, 67(3):1329–1349, 2021.
- [Ren and Zhou(2024)] Zhimei Ren and Zhengyuan Zhou. Dynamic batch learning in high-dimensional sparse linear contextual bandits. *Management Science*, 70(2):1315–1342, 2024.

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REVISIT TENSOR DECOMPOSITION: STATISTICAL OPTIMALITY AND COMPUTATIONAL GUARANTEES

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Keywords: Tucker decomposition, CP decomposition, High-dimensional statistics, Signal-to-noise ratio, Statistical inference, Minimax optimality, Computational complexity, Alternating least squares, Low-rank tensors, Tensor PCA, Statistical-computational trade-off, Robust initialization

Tensor decomposition is a foundational tool in modern data analysis, enabling the extraction of structured, low-dimensional representations from high-dimensional, multiway data. In this talk, we revisit two of the most widely used tensor decomposition frameworks, Tucker decomposition and Canonical Polyadic (CP) decomposition, through the lens of statistical optimality and computational guarantees. Our focus is on both the fundamental limits and practical algorithms for reliable use of tensor methods in noisy, high-dimensional settings.

We begin with Tucker decomposition, which models a low-rank tensor through multilinear projections along each mode [1]. This approach is particularly suited for applications in computational imaging and social sciences, where data are high-order and often corrupted by noise. We analyze the Tucker model in the presence of additive Gaussian noise, where the underlying signal tensor exhibits low multilinear rank. Our results characterize the three-phase behavior of statistical estimation under varying signal-to-noise ratios (SNR): (i) in the strong SNR regime, the Higher-Order Orthogonal Iteration (HOOI) algorithm achieves minimax-optimal rates for estimating the singular subspaces and the tensor itself; (ii) in the weak SNR regime, no consistent estimator exists; and (iii) in the moderate SNR regime, a statistical-computational gap emerges—consistent estimation is possible in theory but computationally intractable under standard complexity assumptions.

We further explore inference procedures in Tucker decomposition [2]. Building on recent developments, we establish asymptotic distributions for singular subspace estimators derived from alternating minimization, allowing for the construction of confidence regions. Importantly, unlike matrix-based settings where debiasing is often necessary, our results show that no debiasing is required for valid inference in tensor models—underlining a key distinction introduced by the multilinear structure and the tensor-specific computational landscape.

Next, we turn to CP decomposition, where a tensor is represented as a sum of rank-one components [3]. Despite its wide empirical use, the theoretical understanding of CP decomposition, especially under noise, non-orthogonality, and higher-rank scenarios, has remained limited. We address this gap by analyzing the Alternating Least Squares (ALS) algorithm in a signal-plus-noise model. We show that ALS, when properly initialized, achieves non-asymptotic, minimax-optimal error bounds for tensors

of arbitrary order, dimension, and rank. We propose a robust initialization method—Tucker-based Approximation with Simultaneous Diagonalization (TASD)—which compresses the tensor and stabilizes subsequent optimization. When used with ALS, the resulting estimator (TASD-ALS) is both statistically consistent and computationally efficient, achieving optimal estimation rates in practice.

Additionally, we provide a rigorous convergence analysis of ALS. We prove that in the rank-one setting, ALS achieves optimal error bounds in just one or two iterations. For general rank, we uncover a two-phase convergence pattern: an initial quadratic phase followed by a linear refinement, with rates determined by coherence properties of the underlying components. These findings give the first formal justification of the fast empirical convergence observed for ALS in structured tensor settings.

In summary, this talk bridges a significant gap between statistical theory and algorithmic practice in tensor decomposition. Our results provide sharp insights into the limits of estimation and inference, while offering provably effective algorithms that scale to modern high-dimensional, multi-modal data.

REFERENCES

- [1] Zhang, A., & Xia, D. (2018). Tensor SVD: Statistical and computational limits. *IEEE Transactions on Information Theory*, 64(11), 7311-7338.
- [2] Xia, D., Zhang, A. R., & Zhou, Y. (2022). Inference for low-rank tensors—no need to debias. *The Annals of Statistics*, 50(2), 1220-1245.
- [3] Tang, R., Chhor, J., Klopp, O., & Zhang, A. R. (2025). Revisit CP Tensor Decomposition: Statistical Optimality and Fast Convergence. *arXiv preprint arXiv:2505.23046*.

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A DYNAMIC NETWORK AUTOREGRESSIVE MODEL FOR TIME-VARYING NETWORK-LINK DATA

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Keywords: Global homogeneity, group structure, network-linked data, dynamic network.

Network-linked data, which refers to a group of units that are observed connected by a network and have a set of available attributes, has attracted much attention in the past few decades (Michell and West, 1996; Lee et al., 2010; Li et al., 2019, 2023; Huang et al., 2021). Yet its extension, time-varying network-link data, has received less investigation.

Existing methods for time-varying network-link data usually assume that units' attributes evolve over time, whereas the network remains unchanged as time increases. (Zhu et al., 2017; Wu, 2019; Zhu and Pan, 2020; Zhu et al., 2022; Chen et al., 2023; Zhu et al., 2023; Li et al., 2023). Zhu et al. (2017) firstly proposed a network vector autoregressive model (NAR) to incorporate network structure. Specifically, they assume that

(0.1)
$$Y_{it} = \mu + \mathbf{X}_i^{\top} \mathbf{\gamma} + \eta_0 n_i^{-1} \sum_{j=1}^N a_{ij} Y_{j(t-1)} + \eta_1 Y_{i(t-1)} + \epsilon_{it},$$

where $Y_t = (Y_{1t}, \ldots, Y_{Nt})^{\top}$ is the high-dimensional response vector with N being the number of nodes in network \mathcal{G} , the node-specific covariate vector X_i is independent and identically distributed random, $A = (a_{ij})_{i,j=1}^N \in \{0,1\}^{N \times N}$ is the adjacent matrix of \mathcal{G} with $a_{ij} = 1$ if there exists an edge between nodes i and j and $a_{ij} = 0$ otherwise, $n_i = \sum_{j=1}^n a_{ij}$ is the degree of node i, and $(\mu, \gamma, \eta_0, \eta_1)$ are parameters to be estimated. Since Zhu et al. (2017), many extensions of NAR model have been studied. For example, Wu (2019) extends model (0.1) to a time-varying setting by allowing $(\mu, \gamma, \eta_0, \eta_1)$ to change with t. Zhu et al. (2022) further extends to the functional varying coefficient setting. Another extension route is to assume that there exists some group structure among N nodes to capture the heterogeneity of nodes (Zhu and Pan, 2020; Chen et al., 2023; Zhu et al., 2023), which means that parameters are the same within each group but different across different groups. Li et al. (2023) studied a grouped time-varying NAR model by assuming the time-varying functional coefficients share some group structure.

However, all aforementioned methods have the following drawbacks. The first one is that network \mathcal{G} does not change over time. In real world, edges among nodes usually change frequently and drastically as time increases (Matias and Miele, 2017; Liu et al., 2018). The second one is that they assume that A is deterministic. In practice, it is well known that network data are collected with errors (Le and Li, 2022). It is common to assume that network data are generated by some parametric model, such as the Erdős Rényi model (Erdős et al., 1960) and the stochastic block model (Holland et al., 1983).

The third one is that the heterogeneity captured by group structure is not sufficient. Nodes within the same group should still behave differently, which corresponds to the degree-corrected stochastic block model (Karrer and Newman, 2011). Besides, as argued by Li et al. (2019) and Le and Li (2022), the parametric form of autoregressive neighborhood average in model (0.1) may be inappropriate to model the network effect to network-linked data, and thus leads to unsatisfactory performance. The last one is the assumption that X_{it} 's are identically and independently distributed across $1 \le i \le N$ and $1 \le t \le T$ may be inappropriate for real data.

In this paper, we propose a novel dynamic network autoregressive model to tackle the above problems for time-varying network-linked data. Specifically, we consider that networks are also evolving as time changes. Then, the dynamic networks are modeled with a tensor CANDECOMP/PARAFAC(CP) decomposition method (Kolda and Bader, 2009), where node and time features of networks are captured by some embedding vectors in low-dimensional Euclidean space. By assuming node-embedding vectors concentrate around some centers, we allow heterogeneity for nodes within the same group. Next, we reformulate the NAR model (0.1) with the help of node and time-embedding vectors. Nodes with similar embedding vectors will have similar contributions to the response variable Y_{it} . Moreover, we consider a flexible framework for the effect of covariate vector X_{it} , where both within-group and global homogeneities are allowed. We also allow non-random covariate vector X_{it} .

The main contribution of the proposed model is the development of a novel framework to model time-varying network-linked data, which mainly integrates a tensor decomposition method and the NAR model (0.1). Instead of considering a deterministic network without changing over time, we model dynamic networks via tensor decomposition. To the best of our knowledge, this is the first attempt to consider dynamic networks for network-linked data. More importantly, we propose a new dynamic network autoregressive model, which incorporates node-embedding and time-embedding vectors as dynamic network impact factors. It is more natural than the neighborhood average effect in literature. Node-embedding and time-embedding vectors and the group structure are estimated using the tensor power update algorithm (Zhang et al., 2023). To solve the resultant optimization task for the dynamic network autoregressive model, we employ a group lasso-type penalty and develop an efficient alternative update algorithm. Further, we establish the asymptotic consistencies for the proposed method whether the global effect of covariate vector exists or not. The superior numerical performance of the proposed method is supported by extensive simulated examples and a real application on time-varying network-linked fund data.

References

- Chen, E. Y., Fan, J., and Zhu, X. (2023). Community network auto-regression for highdimensional time series. *Journal of Econometrics*, 235(2):1239–1256.
- Erdős, P., Rényi, A., et al. (1960). On the evolution of random graphs. *Publ. Math. Inst. Hung. Acad. Sci*, 5(1):17–60.
- Holland, P. W., Laskey, K. B., and Leinhardt, S. (1983). Stochastic blockmodels: First steps. *Social networks*, 5(2):109–137.
- Huang, D., Zhu, X., Li, R., and Wang, H. (2021). Feature screening for network autoregression model. *Statistica Sinica*, 31:1239.

- Karrer, B. and Newman, M. E. (2011). Stochastic blockmodels and community structure in networks. *Physical review E*, 83(1):016107.
- Kolda, T. G. and Bader, B. W. (2009). Tensor decompositions and applications. *SIAM review*, 51(3):455–500.
- Le, C. M. and Li, T. (2022). Linear regression and its inference on noisy networklinked data. *Journal of the Royal Statistical Society Series B: Statistical Methodology*, 84(5):1851–1885.
- Lee, L.-f., Liu, X., and Lin, X. (2010). Specification and estimation of social interaction models with network structures. *The Econometrics Journal*, 13(2):145–176.
- Li, D., Peng, B., Tang, S., and Wu, W. (2023). Inference of grouped time-varying network vector autoregression models. *arXiv preprint arXiv:2303.10117*.
- Li, T., Levina, E., and Zhu, J. (2019). Prediction models for network-linked data. *The Annals of Applied Statistics*, 13(1):132–164.
- Liu, F., Choi, D., Xie, L., and Roeder, K. (2018). Global spectral clustering in dynamic networks. *Proceedings of the National Academy of Sciences*, 115(5):927–932.
- Matias, C. and Miele, V. (2017). Statistical clustering of temporal networks through a dynamic stochastic block model. *Journal of the Royal Statistical Society Series B: Statistical Methodology*, 79(4):1119–1141.
- Michell, L. and West, P. (1996). Peer pressure to smoke: the meaning depends on the method. *Health education research*, 11(1):39–49.
- Wu, B. (2019). Time-varying network vector autoregression model.
- Zhang, Y., Zhang, J., Sun, Y., and Wang, J. (2023). Change point detection in dynamic networks via regularized tensor decomposition. *Journal of Computational and Graphical Statistics*, (just-accepted):1–22.
- Zhu, X., Cai, Z., and Ma, Y. (2022). Network functional varying coefficient model. *Journal of the American Statistical Association*, 117(540):2074–2085.
- Zhu, X. and Pan, R. (2020). Grouped network vector autoregression. *Statistica Sinica*, 30(3):1437–1462.
- Zhu, X., Pan, R., Li, G., Liu, Y., and Wang, H. (2017). Necwork vector autoregression. *The Annals of Statistics*, 45(3):1096–1123.
- Zhu, X., Xu, G., and Fan, J. (2023). Simultaneous estimation and group identification for network vector autoregressive model with heterogeneous nodes. *Journal of Econometrics*, page 105564.

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