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A Bayesian Subset Specific Approach to
Joint Selection of Multiple Graphical Models

Peyman Jalali, Kshitij Khare and George Michailidis

Wells Fargo; and University of Florida

Abstract: The problem of joint estimation of multiple graphical models from high dimensional data has been studied in the statistics and machine learning literature, due to its importance in diverse fields including molecular biology, neuroscience and the social sciences. This work develops a Bayesian approach that decomposes the model parameters across the multiple graphical models into shared components across subsets of models and edges, and idiosyncratic ones. Further, it leverages a novel multivariate prior distribution, coupled with a jointly convex regression based pseudo-likelihood that enables fast computations through a robust and efficient Gibbs sampling scheme. We establish strong posterior consistency for model selection under high dimensional scaling, with the number of variables growing exponentially as a function of the sample size. The efficiency of the proposed approach in borrowing strength across models to identify jointly shared edges is illustrated on both synthetic and real data.

Key words and phrases: Pseudo-likelihood; Gibbs sampling; posterior consistency; Omics data.
1. Introduction

The problem of joint estimation of multiple related Gaussian graphical models has attracted a lot of interest in statistics and machine learning due to its wide application in biomedical studies involving Omics data - e.g. Pierson et al. (2015) and Kling et al. (2015) -, as well as text mining and roll call voting Guo et al. (2011). The key idea, which makes this approach preferable to separate network-wise estimation, is to “borrow strength” across the related models and thus enhance the “effective” sample size used for estimation of the model parameters. In high-dimensional settings, joint estimation is achieved primarily by the use of various penalty functions which induce sparsity/zeros in the group-specific inverse covariance (precision) matrices. Specifically, Guo et al. (2011), who first formulated the problem, modeled the elements of each inverse covariance matrix as a product of a common across all models component and an idiosyncratic (model specific) component and imposed an $\ell_1$ penalty on each one; thus, when the penalty sets the common component to zero, the corresponding edge is absent across all models, whereas if the common component is not zero, edges can be absent because the penalty sets the idiosyncratic one to zero for selected models. Another set of approaches aims to achieve a certain amount of “fusing” across all models under consideration, focusing both of the presence of common edges, as well as their absence across all models simultaneously. Ex-
amples of such approaches include Danaher et al. (2014) that employed a group lasso and/or a fused lasso penalty on each edge parameter across all models and Cai et al. (2016) that used a mixed $\ell_1/\ell_\infty$ norm for the same task.

However, in many application settings, shared connectivity patterns across models occurs only for a subset of edges, while the remaining ones exhibit different connectivity patterns in each model. In other settings, subsets of edges share common connectivity patterns across only a subset of models. In both instances, the previously mentioned approaches will exhibit a rather poor performance in discovering these more complex patterns. To address this issue, Ma and Michailidis (2016) proposed a supervised approach based on fusing through a group lasso penalty, wherein the various connectivity patterns across subsets of edges and subsets of models are a priori known. An alternative supervised approach Saegusa and Shojaie (2016) employed a similarity graph penalty for fusing across models, coupled with an $\ell_1$ penalty for obtaining sparse model estimates. The similarity graph is assumed to be a priori known.

The Bayesian paradigm comes with the advantage of natural uncertainty quantification through the posterior distribution, and also a natural structured mechanism for incorporating prior information. A Bayesian variant of the approach in Saegusa and Shojaie (2016) was introduced in Peterson et al. (2015), wherein a Markov random field prior distribution was used to capture model sim-
ilarity, followed by a spike-and-slab prior distribution on the edge model parameters. Another Bayesian approach was recently developed in [Tan et al. (2017)] which, similar to [Peterson et al. (2015)], uses $G$-Wishart prior distributions on the group-specific precision matrices given the sparsity patterns in each group, and then employs a multiplicative model based hierarchical prior on these networks to induce similarity/dependence. Recent papers including [Shaddox et al. (2018)], [Petersen et al. (2020)] extend the ideas in [Wang (2015)] to the joint estimation setting for improved computational scalability. However, this class of approaches still suffers from scalability issues beyond moderate dimensional settings with 150 or so variables. Two key computational roadblocks for posterior sampling are: (a) the fact that the precision matrices are restricted to be positive definite, and (b) the dependence structure between groups is induced through priors on large discrete spaces of sparsity patterns (graphs) for the precision matrices, and the conditional updates of relevant discrete/latent variables and respective hyper-parameters can be messy and significantly add to the computational burden.

For the single graphical model estimation problem based on $n$ i.i.d. observations from a distribution with inverse-covariance/precision matrix $\Omega$, the entries in the $i^{th}$ row of $\Omega$ can be interpreted as least squares coefficients when regressing the $i^{th}$ variable against the other variables (see (4.9)), an idea leveraged to develop quasi-likelihood/pseudo-likelihood based approaches, see [Meinshausen]
and Bühlmann (2006), Lin et al (2017), Atchade (2019). These approaches relax the positive definite (p.d.) constraint on $\Omega$, leading to significant improvement in computational speed. Note that the relaxation of the p.d. constraint does not create any issues/complications for model/sparsity selection in $\Omega$, which is often the key objectivel. If a p.d. estimate of $\Omega$ is needed for a downstream application, it can be obtained in a straightforward manner, for example, by computing the restricted MLE based on the estimated sparsity pattern.

Lin et al (2017) extend this idea to the joint graphical model estimation problem, by using the regression based neighborhood selection procedure of Meinshausen and Bühlmann (2006) in conjunction with an alternate version of the Markov random field priors in Peterson et al (2015) to induce dependence between groups. The relaxation of the p.d. constraint leads to significant improvement in the computational performance compared to the likelihood based approaches mentioned above. However, use of matrix inversions and latent variable updates etc. still leads to a steep increase in computational cost of the corresponding algorithm, labeled BNS, as the number of variables $p$ increases (see Section 5.2). “Maximization-based” Bayesian approaches for joint graphical estimation, which focus on obtaining posterior modes have been proposed in Li et al (2018) and Yang et al (2021). These approaches are computationally scalable, but unlike the other “sampling-based” approaches discussed above, these
methods do not generate samples from the posterior and are unable to provide detailed uncertainty quantification. For example, Yang et al. (2021) use relevant conditional posterior probabilities evaluated at the posterior mode to evaluate uncertainty for individual edges, but do not provide more nuanced uncertainty for joint inclusion/exclusion of multiple edges (see Table G.2 for example).

While uncertainty quantification through the posterior is an attractive feature of a Bayesian approach, in high-dimensional settings it is crucial to rigorously justify its validity. In the current context, this corresponds to establishing strong posterior selection consistency, i.e., proving that the posterior distribution of the combined sparsity pattern in the group-specific precision matrices asymptotically places all of its mass on the “true” sparsity pattern in the high-dimensional data-generating model. While Yang et al. (2021) establish selection consistency for the posterior mode, a high-dimensional strong selection consistency result for joint graphical model estimation has not been established for any of the Bayesian approaches discussed above.

With this background, the key objective of this paper is to develop a scalable sampling-based Bayesian approach with high-dimensional selection consistency guarantees for joint estimation and uncertainty quantification for multiple related Gaussian graphical models that exhibit complex edge connectivity patterns across models for different subsets of edges. We avoid the Markov random
field based approach for inducing group similarity, and instead take a more di-
rect and completely different approach based on a subset-specific decomposition
(see (2.2)) of the group-specific precision matrices. We then introduce a novel
Subset Specific ($S^2$) prior that for each edge aims to select the subset of mod-
els it is common to (see (3.6)-(3.8) and Remark 1). We couple these with the
jointly convex regression based pseudo-likelihood used in Khare et al. (2015) for
estimating a single Gaussian graphical model.

The above framework leads to an easy to implement and scalable Gibbs
sampling scheme for exploring the posterior distribution. This corresponding al-
gorithm, labeled BJNS (Bayesian Joint Network Selector), essentially involves
$O(p^2)$ univariate updates from relevant mixture distributions, and avoids the
need for matrix inversions or latent variables. As a result, the computational
performance of BJNS is an order of magnitude faster than the BNS algorithm of
Lin et al. (2017) (see Section 5.2). Our direct subset specific approach can lead
to significantly improved statistical selection performance compared to existing
methods (see Section 5). We also establish strong posterior model selection con-
sistency result (Theorem 1) for the proposed approach. Intuitively, the proposed
framework achieves the objectives set forth in the Ma and Michailidis (2016)
work, without requiring a priori specification of the shared edge connectivity
patterns; thus, the approach is fully unsupervised. Also, the availability of pos-
terior samples allows for uncertainty quantification in the form of subset-specific inclusion probabilities (see Figure D.1 and Table G.2).

Note that the main goal of the proposed working framework is model/sparsity selection. As described in detail in Section 2.1, a constraint imposing restrictions on the magnitude of relevant precision matrix entries is necessary for identifiability purposes. First, this constraint has no adverse effect on the main task of sparsity selection in the sense that the framework still produces a valid posterior distribution on the space of all possible sparsity patterns for the various group-specific precision matrices. As our simulation and consistency results show, BJNS performs very well in terms of sparsity selection compared to existing methods even when this constraint is not satisfied in the data-generating model. Second, as pointed out above, any regression based approach is not directly useful for magnitude estimation anyway, since the resulting estimates of the precision matrices are not guaranteed to be positive definite. If such estimates are needed for downstream applications, one can obtain them by using the respective MLEs restricted to the estimated sparsity pattern for each group-specific matrix (see (4.12)). Note that the resulting estimates are guaranteed to be positive definite and are free from the identifiability constraint in Section 2.1 (which is used only for sparsity selection and does not play any role in (4.12)).

We also provide a simulation study evaluating the (magnitude) estimation accu-
racy in Supplemental Section D.4.

The remainder of the paper is organized as follows. Section 2 formulates the problem and Section 3 introduces the $S^2$ prior. Section 4 shows how to obtain and sample from the posterior distribution. Section 5 presents extensive numerical results and comparisons, while Section 6 presents a metabolomics application using a case-control study on Inflammatory Bowel Disease. Section 7 establishes high-dimensional posterior consistency for BJNS.

2. Framework for Joint Sparsity Selection

Suppose we have data from $K$ a priori defined groups. For each group $k$ ($k = 1, 2, ..., K$), let $\mathcal{Y}_k := \{y_{ki}^{(k)}\}_{i=1}^{n_k}$ denote $p$-dimensional i.i.d observations from a multivariate normal distribution, with mean 0 and covariance matrix $(\Omega^k)^{-1}$, which is specific to group $k$. Based on the discussion in the introductory section, the $K$ group-specific precision matrices $\{\Omega^k\}_{k=1}^{K}$ can share common edge patterns across subsets of the $K$ models, as delineated next. Our goal is to jointly select the edge structures (or equivalently, the sparsity patterns) for all the $K$ precision matrices to account for these shared structures.

Let $\mathcal{P}(K)$ denote the power set of $\{1, \cdots, K\}$ and for $k = 1, ..., K$, define $\vartheta_k$ as follows:

$$\vartheta_k = \{r \in \mathcal{P}(K) \setminus \{0\} : k \in r\}, \quad k = 1, ..., K. \quad (2.1)$$
It is easy to check that each $\vartheta_k$ is the collection of subsets which contain $k$, and has $2^{K-1}$ members. Denote by $\Psi^r$ the matrix that contains common patterns amongst precision matrices $\{\Omega^j\}_{j \in r}$. For any singleton set $r = \{k\}$, the non-zero elements in the matrix $\Psi^r$ correspond to edges that are unique to group $k$, while for any other set $r$ containing more than a single element, the non-zero elements in the matrix $\Psi^r$ correspond to edges (and their magnitudes) that are common across all members in $r$ (and not present in other networks). For example, the non-zero elements in $\Psi^{123} := \Psi^{\{1,2,3\}}$ correspond to edges that are shared exclusively by networks 1, 2, and 3.

Therefore, each precision matrix $\Omega^k$ can be decomposed as

$$\Omega^k = \sum_{r \in \vartheta_k} \Psi^r, \quad k = 1, ..., K, \quad (2.2)$$

where $\sum_{r \in \vartheta_k} \Psi^r$ accounts for all the structures in $\Omega^k$ which are either unique to group $k$ (i.e. $\Psi^k$) or are shared exclusively between group $k$ and some combination of other groups (i.e. $\sum_{r \in \vartheta_k \setminus \{k\}} \Psi^r$). We further assume that $\Psi^k \in M_p^+$ for $k = 1, 2, ..., K$, where $M_p^+$ denotes the space of all $p \times p$ matrices with positive diagonal entries. Finally, the diagonal entries of every joint matrix $\Psi^r$, with $r \in \bigcup_{k=1}^K (\vartheta_k \setminus \{k\})$ are set to zero; in other words, the diagonals entries of $\Omega^k$ are contained in the corresponding $\Psi^k$.

To illustrate the notation, consider the case of $K = 3$ and following the notation in (2.1), define the sets: $\vartheta_1 = \{\{1\}, \{12\}, \{13\}, \{123\}\}$, $\vartheta_2 = \{\{2\}, \{12\}, \{23\}, \{123\}\}$,
and $\vartheta_3 = \{\{3\}, \{13\}, \{23\}, \{123\}\}$ and accordingly, decompose the precision matrices $\Omega^2$, $\Omega^1$, $\Omega^3$, as follows:

$$
\Omega^1 = \sum_{r \in \vartheta_1} \Psi^r = \Psi^1 + \Psi^{12} + \Psi^{13} + \Psi^{123}, \quad \Omega^2 = \sum_{r \in \vartheta_2} \Psi^r = \Psi^2 + \Psi^{12} + \Psi^{23} + \Psi^{123}, \quad \Omega^3 = \sum_{r \in \vartheta_3} \Psi^r = \Psi^3 + \Psi^{13} + \Psi^{23} + \Psi^{123},
$$

where the $\Psi^1$, $\Psi^2$, and $\Psi^3$ matrices contain group specific patterns, the $\Psi^{12}$, $\Psi^{13}$, $\Psi^{23}$ matrices contain patterns shared across pairs of models (for subsets of the edges) and the matrix $\Psi^{123}$ contains patterns shared across all models.

### 2.1 Identifiability Considerations

A moment of reflection shows that the model decomposition (2.2) is not unique. For example, for any arbitrary matrix $X$, the model (2.2) is equivalent to $\Omega^k = \sum_{r \in \vartheta_k} \Phi^r$ with $\Phi^r = \Psi^r + X$ and $\Phi^k = \Psi^k - \frac{1}{2^{k-1}} X$. Hence, without imposing appropriate identifiability constraints, meaningful inference is not feasible. To that end, rewrite the element-wise representation of model (2.2) as:

$$
\omega^k_{ij} = \sum_{r \in \vartheta_k} \psi^r_{ij}, \quad 1 \leq i < j \leq p, 1 \leq k \leq K,
$$

where $\omega^k_{ij}$ and $\psi^r_{ij}$ are the $ij^{th}$ coordinates of the matrices $\Omega^k$ and $\Psi^r$, respectively. We only consider the upper off-diagonal entries due to the symmetry of the precision matrix and thus define vectors $\theta_{ij}$ for every $1 \leq i < j \leq p$, as

$$
\theta_{ij} = \{\psi^r_{ij} \}_{r \in P(K) \setminus \{0\}},
$$

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where each $\theta_{ij}$ has $2^K - 1$ distinct parameters. For *identifiability purposes* we require that each vector $\theta_{ij}$ can have at most one non-zero element. Note that under this constraint, if an edge $(i, j)$ is shared amongst many groups, the non-zero element will be allocated to the maximal set $s \in \bigcup_{k=1}^{K} (\varnothing \setminus \{k\})$, while all subsets of $s$ will be allocated a zero value. As an example, consider again the case of $K = 3$ groups and an edge $(i, j)$ shared amongst all three groups. In this case, the edge will be allocated to the $\Psi_{123}$ component and not to any other components, such as $\Psi_{12}$ or $\Psi_{13}$. Hence $\Psi_{123}^{ij}$ will be non-zero, but $\Psi_{12}^{ij} = \Psi_{13}^{ij} = \Psi_{23}^{ij} = \Psi_{1}^{ij} = \Psi_{2}^{ij} = \Psi_{3}^{ij} = 0$. Next, we discuss the implications of this identifiability constraint.

The precision matrices $\{\Omega_k\}_{k=1}^{K}$ have a total of $Kp(p+1)/2$ parameters. We expand these precision matrices in terms of the subset specific matrices $\{\Psi_r\}_{r \in \mathcal{P}(K) \setminus \{0\}}$ (see (2.3)). This expanded set of parameter matrices has $Kp + (2^K - 1)p(p-1)/2$ parameters in all. *The identifiability constraint reduces the number of parameters to $Kp + p(p-1)/2$, and thereby helps significantly with computational scalability*. Further, sparsity constraints are introduced by the spike-and-slab priors described in Section 3.

Another consequence of the identifiability constraint is that if edge $(i, j)$ is shared by a subset $s$, then the magnitude of all $\{\omega_k^{ij}\}_{k \in s}$ is the same. Note that it makes sense in a variety of applications for shared edges to have similar
magnitudes and/or the same sign. In fact, existing approaches in Danaher et al. (2014), Li et al. (2018) based on group/fused lasso, encourage similarity or even exact equality of magnitudes across the shared edges. As demonstrated by the simulations results in Table 3, our working model still performs well in selecting the sparsity/skeleton as compared to existing methods when the shared edges have different magnitudes, but the same sign. This is also supported by the theoretical results in Section 7, where we allow the true precision matrices to have different magnitudes, but the same sign for shared edges.

Finally, since we employ a regression-based pseudo-likelihood (see (4.10)) which is well-defined as long as the relevant precision matrix has positive diagonal entries, we additionally relax the p.d. constraint on \( \{ \Omega^k \}_{k=1}^K \) for faster computation. Note that both the identifiability constraint and the p.d. relaxation do not in any way restrict the range of allowable sparsity patterns, and are only used in the working framework for sparsity selection. Once the sparsity patterns are selected, if needed, a simple refitting step is performed (see (4.12)) to obtain p.d. estimates of the \( K \) precision matrices which obey the selected sparsity patterns, and are completely free of the above identifiability constraint.
3. Subset Specific ($S^2$) Prior Distribution

Next, we construct a novel prior distribution that respects the introduced identifiability constraint, and encourages further sparsity in the parameters. For any generic symmetric $p \times p$ matrix $A$, define $\mathbf{a} = (a_{12}, a_{13}, \ldots, a_{p-1,p})$ and $\mathbf{d}_A = (a_{11}, a_{pp})$, where due to the symmetric nature of $A$, the vector $\mathbf{a}$ contains all the off-diagonal elements, while $\mathbf{d}_A$ the diagonal ones. In particular, $\mathbf{y}^r$ is the vectorized version of the off-diagonal elements of $\Psi^r$. Using the above notation, define $\Theta$ to be the vector obtained by combining the vectors $\{\mathbf{y}^r, r \in \mathcal{P}(K) \setminus \{0\}\}$. To illustrate, for $K = 3$ groups, $\Theta$ is given by

$$
\Theta = (\mathbf{y}^{123'}, \mathbf{y}^{23'}, \mathbf{y}^{13'}, \mathbf{y}^{12'}, \mathbf{y}^{3'}, \mathbf{y}^{2'}, \mathbf{y}^{1'})'.
$$

(3.5)

Note that $\Theta$ is a rearrangement of the vector $(\theta_{12}, \theta_{13}, \ldots, \theta_{p-1,p})'$. Thus, according to the location of the non-zero coordinates in $\theta_{ij}$ ($2^K$ possibilities), there are $2^{Kp(p-1)/2}$ possible sparsity patterns across the $K$ groups for $\Theta$. Let $\ell$ be a generic sparsity pattern for $\Theta$ and denote the set of all the $2^{Kp(p-1)/2}$ sparsity patterns by $\mathcal{L}$. To illustrate, consider $K = 2$ groups and $p = 3$ variables. In this case, each matrix has 3 off-diagonal edges ($\{ij : 1 \leq i < j \leq p\} = \{12, 13, 23\}$). Assume edge 12 is shared between the two groups, edge 13 is unique to group 2, and edge 23 is absent in both groups. In this case, $\Theta$ is given by $\Theta = (\psi_{12}^{12}, 0, 0, 0, \psi_{13}^{12}, 0, 0, 0, 0)^'$. and the sparsity pattern extracted
from $\Theta$ becomes: $\ell = ((1, 0, 0), (0, 1, 0), (0, 0, 0))'$. For every sparsity pattern $\ell$, let $d_\ell$ be the density (number of non-zero entries) of $\ell$, and $M_\ell$ be the space where $\Theta$ varies, when restricted to follow the sparsity pattern $\ell$. $\lambda \in \mathbb{R}^{2K-1}$, and $\Lambda = \text{diag}(\lambda, \lambda, \ldots, \lambda)$ is a diagonal matrix (with $p(p-1)/2$ diagonal blocks of $\lambda$) whose entries determine the amount of shrinkage imposed on the corresponding elements in $\Theta$. We specify the hierarchical prior distribution $S^2$ as follows:

$$
\pi(\Theta|\ell) = \frac{|\Lambda_{\ell\ell}|^{\frac{1}{2}}}{(2\pi)^{d_\ell/2}} \exp \left( -\frac{\Theta' \Lambda \Theta}{2} \right) I_{(\Theta \in M_\ell)}, \quad (3.6)
$$

$$
\pi(\ell) \propto \begin{cases} 
(2\pi)^{d_\ell/2} |\Lambda_{\ell\ell}|^{-\frac{1}{2}} q_1^{d_\ell} (1 - q_1)^{(p/2) - d_\ell} & d_\ell \leq \tau, \\
(2\pi)^{d_\ell/2} |\Lambda_{\ell\ell}|^{-\frac{1}{2}} q_2^{d_\ell} (1 - q_2)^{(p/2) - d_\ell} & d_\ell > \tau,
\end{cases} \quad (3.7)
$$

where $\Lambda_{\ell\ell}$ is a sub-matrix of $\Lambda$ obtained after removing the rows and columns corresponding to the zeros in $\Theta \in M_\ell$, and $q_1, q_2$ are edge inclusion probabilities, respectively, for the case of sparse ($d_\ell \leq \tau$) and dense ($d_\ell > \tau$) $\Theta$. An equivalent “spike-and-slab” representation of (3.6) is:

$$
\{\theta_{ij}\}_{1 \leq i < j \leq p} \text{ are conditionally independent given } \ell \text{ and }
\pi(\theta_{ij}|\ell) = 1_{\theta_{ij}=0}1_{\ell_{ij}=0} + \sum_{r \in \mathcal{P}(K) \setminus \{0\}} \sqrt{\lambda_r} e^{-\lambda_r \theta_{ij,r}^2 / 2} 1_{\theta_{ij,-r}=0}1_{\ell_{ij,r}=1} \quad (3.8)
$$

Note that the distribution of $\theta_{ij}$ is supported on the axes of $\mathbb{R}^{2K-1}$. The “spike” corresponds to the point mass at the origin, and the “slabs” correspond to a
normal distribution on an appropriate axis (when exactly one coordinate is non-zero). In fact, when \( q_1 = q_2 = q \), it follows by (5.7) that \( \{ \ell_{ij} \}_{1 \leq i < j \leq p} \) are a priori independent, and the prior distribution of \( \ell_{ij} \) is given by 

\[
P(\ell_{ij,k} = 1) = \sqrt{2\pi\lambda_k^{-1}} q / C \quad \text{for every } 1 \leq k \leq 2^K - 1, \text{ and}
\]

\[
P(\ell_{ij} = 0) = \frac{(1 - q)}{C}, \quad \text{where } C = 1 - q + q \left( \sum_{k=1}^{2^K-1} \sqrt{2\pi\lambda_k} \right).
\]

Hence, smaller values of \( q \) can be used to encourage sparser models in high-dimensional settings.

Note that the \( S^2 \) prior allows for at most one entry in each \( \theta_{ij} \) to be non-zero, and thus sets at least \( \frac{p(p-1)}{2}(2^K - 2) \) parameters to be exactly equal to zero. In particular, the \( S^2 \) prior considers the entire range of models allowable under the identifiability constraint discussed in Section 2.1: at one end we have the model with complete sparsity where \( \Theta = 0 \), and at the other end we have models with \( \binom{p}{2} \) parameters where each \( \theta_{ij} \) containing exactly one non-zero entry. In addition to forcing sparsity, the diagonal entries of \( \Lambda \) enforce shrinkage to the corresponding elements in \( \Theta \).

The vector \( \Theta \) only incorporates the off-diagonal entries of \( \Psi \) matrices. Regarding the diagonal entries, for every \( k \in \{1, ..., K\} \), we let \( \delta_{\Psi^k} \) be the vector comprising of the diagonal elements of the matrix \( \Psi^k \) and define \( \Delta \) to be the vector of all diagonal vectors \( \delta_{\Psi^k} \), i.e., \( \Delta = (\delta_{\Psi^1}, ..., \delta_{\Psi^K}) \). We assign an in-
dependent Exponential($\gamma$) prior on each coordinate of $\Delta$ (diagonal element of the matrices $\Psi_k$, $k = 1, \ldots, K$), i.e., $\pi(\Delta) \propto \exp(-\gamma 1' \Delta) I_{R^{Kp}}(\Delta)$. Selection of hyperparameters $\Lambda$, $\gamma$ is discussed in the next section. Since the diagonal entries of every joint matrix $\Psi^r$, with $r \in \bigcup_{k=1}^{K} (\vartheta_k \setminus \{k\})$ are set to zero, the specification of the prior is now complete.

**Remark 1.** Prior distributions with similarities to the subset specific one proposed here have been used in genetic association (eQTL) analysis with heterogeneous subgroups, see Wen and Stephens (2014); Flutre et al. (2013) and references therein. However, there are two crucial differences in these two settings. Firstly, in the eQTL setting, one has a single regression model with gene expression level as the response and a single predictor (genotype), whereas in the current joint graphical model setting, we employ a pseudo-likelihood consisting of $p$ different regressions (corresponding to the $p$ variables) and each of them has multiple ($p - 1$) predictors; see (4.9) and (4.10) below. Secondly, even though the p.d. constraint on the precision matrices is relaxed for sparsity selection, the symmetry constraint is not (we still need the sparsity patterns for each $\Omega^k$ to be symmetric). This symmetry couples the $p$ regressions forming the pseudo-likelihood. These complications lead to unique challenges in the methodological development and theoretical analysis for the joint graphical models setting.
4. The Bayesian Joint Network Selector (BJNS)

With a prior distribution in hand, sparsity/network selection is based on a pseudo-likelihood approach, leveraging the regression interpretation of the entries of $\Omega$. It can also be regarded as a weight function and as long as the product of the pseudo-likelihood and the prior density is integrable over the parameter space, one can construct a (pseudo) posterior distribution to carry out inference. The main advantage of using a pseudo-likelihood, as opposed to a full Gaussian likelihood, is that it allows for an easy to implement sampling scheme from the posterior distribution and in addition provides more robust results under deviations from the Gaussian assumption, as illustrated by work in the frequentist domain ([Khare et al., 2015; Peng et al., 2009]). Of course, the use of this pseudo-posterior distribution has to be justified by both theoretical consistency results and by assessing its performance in finite sample simulation settings. Both of these tasks will be undertaken in Section 5 and Section 6, respectively.

Note that if $Y \in \mathbb{R}^p$ with $\text{Cov}(Y) = \Omega^{-1}$, then

$$
\Omega_{j,-j} = (\Omega_{jk})_{1 \leq k \leq p, k \neq j} = \arg\min_{w \in \mathbb{R}^{p-1}} E (\Omega_{jj} Y_j + w' Y_{-j})^2.
$$

Hence, if $Y_1, Y_2, \ldots, Y_n$ are i.i.d. with covariance matrix $\Omega^{-1}$, then the above regression interpretation of $\Omega$, can be used to construct the CONCORD pseudo-
likelihood introduced in Khare et al. (2015), given by

\[
\exp \left\{ n \sum_{j=1}^{p} \log \omega_{jj} - \frac{n}{2} \text{tr} \left[ \Omega^2 S \right] \right\}, \tag{4.9}
\]

where \( S \) denotes the sample covariance matrix. The CONCORD pseudo-likelihood is jointly convex in the entries of \( \Omega \). This makes it both theoretically and computationally preferable over other variants of the regression based pseudo-likelihood (see discussion in Khare et al. (2015)). Further, since the regression interpretation does not depend on the Gaussian assumption, regression based approaches such as the one above generally provide more robust results under deviations from that assumption.

Let \( S^k \) denote the sample covariance matrix of the observations in the \( k^{th} \) group. Based on the above discussion, we employ the pseudo-likelihood in (4.9) and the model specification (2.2) to construct the joint pseudo-likelihood function for \( K \) precision matrices, as follows,

\[
\prod_{k=1}^{K} \exp \left\{ n \sum_{j=1}^{p} \log \psi_{jj}^k - \frac{n}{2} \text{tr} \left[ \left( \sum_{r \in \partial k} \Psi^r \right)^2 S^k \right] \right\}. \tag{4.10}
\]

Since we have parametrized the \( S^2 \) prior in terms of \( (\Theta, \Delta) \), we will rewrite the above pseudo-likelihood function in terms of \( (\Theta, \Delta) \), as well. Some straightforward algebra shows that

\[
\text{tr} \left[ \left( \sum_{r \in \partial k} \Psi^r \right)^2 S^k \right] = \begin{pmatrix} \Theta' & \Delta' \end{pmatrix} \begin{pmatrix} \Upsilon & A \\ A' & D \end{pmatrix} \begin{pmatrix} \Theta \\ \Delta \end{pmatrix}, \tag{4.11}
\]
where, \( \mathbf{Y} \) is a \( \frac{p(p-1)(2^K-1)}{2} \times \frac{p(p-1)(2^K-1)}{2} \) symmetric matrix whose entries are either zero or a linear combination of \( \{ s_{ij}^k \}_{1 \leq i < j \leq p} \); \( \mathbf{D} \) is a \( Kp \times Kp \) diagonal matrix with entries \( \{ s_{ii}^k \}_{1 \leq i \leq p} \); \( \mathbf{a} \) is a \( \frac{p(p-1)(2^K-1)}{2} \times 1 \) vector whose entries depend on \( \Delta \) and \( \{ s_{ij}^k \}_{1 \leq i < j \leq p} \); and finally \( \mathbf{A} \) is a \( \frac{p(p-1)(2^K-1)}{2} \times Kp \) matrix such that \( \mathbf{A} \Delta = \mathbf{a} \). The algebraic details of the equality in (4.11), structures of \( \mathbf{Y} \) and \( \mathbf{a} \), and the algebraic form of the joint and conditional posterior densities for \( \Theta \) and \( \Delta \) are provided in Supplemental Section A.1.

Gibbs Sampling Scheme for BJNS: Generating exact samples from the joint posterior of \( (\Theta, \Delta) \) is not feasible. We instead generate approximate samples by computing the full conditional distribution of the vectors \( \{ \theta_{ij} \}_{1 \leq i < j \leq p} \) and that of the diagonal entries \( \{ \psi_{ii}^k \}_{1 \leq i \leq p, 1 \leq k \leq K} \).

Each \( \theta_{ij} \) contains \( 2^K - 1 \) elements of which at most one is non-zero. For ease of exposition, let \( \theta_{l,ij} \) denote the \( l^{th} \) element of \( \theta_{ij} \) for \( l = 1, \ldots, 2^K - 1 \) (based on (2.4), every \( \theta_{l,ij} \) represents a \( \psi_{rr}^l \), for some \( r \in \mathcal{P}(K) \)). Using the same notation for the shrinkage parameters (diagonal elements of \( \Lambda \)), let \( \lambda_{l,ij} \) be the shrinkage parameter corresponding to \( \theta_{l,ij} \). Since there are \( 2^K \) possibilities for the location of the non-zeros in each \( \theta_{ij} \), one can see \( \theta_{ij} \) as an element in one of the disjoint spaces \( \mathbb{M}_0, \mathbb{M}_1, \ldots, \mathbb{M}_{2^K-1} \), where \( \mathbb{M}_0 \) is the singleton set consisting of the zero vector of length \( 2^K - 1 \) and \( \mathbb{M}_l \) \( (l = 1, \ldots, 2^K - 1) \) is the space spanned by the \( l^{th} \) unit vector of length \( 2^K - 1 \). Denote by \( \Theta_{-(ij)} \) the sub
vector $\Theta$ obtained by removing $\theta_{ij}$. It can be shown that (Supplemental Section B.1) the conditional density of $\theta_{ij}$ given $\Theta_{-ij}, \Delta$ is a mixture of univariate normal densities respectively supported on $\{M_i \}_{i=0}^{2K-1}$. Also, conditional on $\Theta$, the diagonal entries $\{\psi_{ii}^k\}_{1 \leq i \leq p}$ are a posteriori independent and an efficient algorithm to sample them is provided in Supplemental Section B.1.

Selection of Hyperparameters: Let $\theta$ and $\delta$ be generic elements of $\Theta$ and $\Delta$ and let $\lambda$ and $\gamma$ be their corresponding shrinkage parameters. Selecting appropriate values for the latter is an important task. In other Bayesian analysis of high dimensional models, shrinkage parameters are usually generated based on an appropriate prior distribution; see Park and Casella (2008); Kyung et al. (2010); Hans (2009) for regression analysis and Wang (2012) for graphical models. We assign independent gamma priors on each shrinkage parameter $\lambda$ or $\gamma$; specifically, $\lambda, \gamma \sim \text{Gamma}(r, s)$, for some hyper-parameters $r$ and $s$. The amount of shrinkage imposed on each element $\theta$ and $\delta$ is calculated by considering the posterior distribution of $\lambda$ and $\gamma$, given $(\Theta, \Delta)$. Straightforward algebra shows

$$
\lambda| (\Theta, \Delta) \sim \text{Gamma}(r + 0.5, 0.5\theta^2 + s), \quad \text{and} \quad \gamma| (\Theta, \Delta) \sim \text{Gamma}(r + 1, |\delta| + s).
$$

Note that $\mathbb{E}\{\lambda| (\Theta, \Delta)\} = \frac{r + 0.5}{0.5\theta^2 + s}$ and $\mathbb{E}\{\gamma| (\Theta, \Delta)\} = \frac{r + 1}{|\delta| + s}$. That is, our approach selects the shrinkage parameters $\lambda$ and $\gamma$ based on the current values of $\theta$ and $\delta$ in a way that larger (smaller) entries are regularized less (more).

The selection of the hyper-parameters $r$ and $s$ is also an important task and
in our experience can affect performance, especially for small sample sizes. As the sample size grows, the results are less sensitive to the choice of the hyperparameters (see Supplemental Section C for an illustration). In absence of any prior information, we recommend the non-informative choice \( r = 10^{-4} \) and \( s = 10^{-8} \), which corresponds to a flat prior for the \( \lambda \) and \( \gamma \) values, and is based on the suggestions made in Wang (2012) for hyperparameter selection. In general, we have found that this choice works well in the extensive simulations presented in Section 3. As far as the choice of \( q_1 \) and \( q_2 \) is concerned, our default choice again is the objective choice \( q_1 = q_2 = 1/2 \). To encourage sparser models in really high-dimensional situations, one can use the choice \( q_1 = 1/p, q_2 = q_1^p \) (essentially zero), and \( \tau = n/\log n \), based on the theoretical result in Section 7.

Finally, the construction of the Gibbs sampler proceeds as follows: matrices \( \{ \Psi^k \}_{k=1}^K \) are initialized as the identity matrix, while \( \{ \Psi^r \}_{r \in \mathcal{P}(K), \& |r| > 1} \) at zero. Then, in each iteration of the MCMC chain, we update the vectors \( \theta_{ij} \) and the diagonal entries \( \psi_{ii}^k \), one at a time, using their full conditional posterior densities given in (B.25) and (B.26), respectively. Procedure 1 in the Supplemental Document describes one iteration of the resulting Gibbs Sampler.

Procedure for Sparsity Selection and Uncertainty Quantification: Note that the conditional posterior probability density of the off-diagonal elements of \( \theta_{ij} \) is a mixture density that puts all of its mass on the events \( \{ \theta_{ij} : |\theta_{ij}|_0 \leq 1 \} \), where
$|\theta_{ij}|_0$ is the number of non-zero coordinates of $\theta_{ij}$. This property of BJNS allows for model selection, in the sense that for every (post burn-in) iteration of the Gibbs sampler one can check whether $\theta_{ij} = 0$ or which element of $\theta_{ij}$ (there could be at most one non-zero element) is non-zero. Note that each element of $\theta_{ij}$ corresponds to a subset of $\{1, \cdots, K\}$. The non-zero frequency during sampling for any given subset $S$, normalized by the total number of iterations, provides an estimate of the posterior probability that the edge $(i, j)$ is shared among elements of $S$. Hence, at the end of the procedure, we choose the element with the highest non-zero frequency during sampling. Denoting the chosen element (subset) by $\hat{S}_{ij}$, we set $\omega_{ij}^k = 0$ if $k \notin \hat{S}_{ij}$, and $\omega_{ij}^k \neq 0$ if $k \in \hat{S}_{ij}$.

Procedure for Positive Definite Estimation: Once sparsity selection in the form of subsets $\{\hat{S}_{ij}\}$ has been completed, these subsets can be used to obtain sparsity graphs $\{\hat{G}^k\}_{k=1}^K$ for the $K$ group-wise precision matrices (i.e., $(i, j)$ is in $\hat{G}^k$ if $k \in \hat{S}_{ij}$). Next, a p.d. estimate $\hat{\Omega}^k$ of $\Omega^k$ can be obtained as a solution to the following restricted optimization problem (implemented in R package glasso).

$$
\hat{\Omega}^k = \arg\min_{\Omega \in \mathbb{P}_{\hat{G}^k}} \left\{ \text{tr}(\Omega S^h_k) - \log \det \Omega \right\}.
$$

Here $\mathbb{P}_{\hat{G}^k}$ is the space of all positive definite matrices, wherein the $(i, j)^{th}$ entry is zero whenever $(i, j)$ is not an edge in $\hat{G}^k$. Hence, the resulting “refitted” estimates $\{\hat{\Omega}^k\}_{k=1}^K$ are positive definite, and are not constrained to obey any working sign/magnitude restrictions used in the sparsity selection process. Another
possibility is to obtain fully Bayesian refitted estimates by using $G$-Wishart priors on $P_{G^k}$ for each sub-group, and then computing the posterior mean for each group using one of the computationally efficient sampling methods developed in Mitsakakis et al. (2011); Lenkoski (2013); Khare et al. (2018).

5. Simulation Studies

In this section, we undertake a simulation-based evaluation of the statistical and computational properties of the BJNS algorithm. Most tables and figures, and several additional simulation results are provided in the Supplement.

5.1 Sparsity selection performance

We present two simulation studies to evaluate the sparsity selection performance of BJNS. In the first study (Section 5.1.1), we compare the performance of BJNS with other high-dimensional methodologies for joint sparsity selection: (1) separate estimation by Graphical lasso (Glasso) of each $\Omega^k$, joint estimation by Guo et al. (2011) (JEM-G), the two joint Graphical Lasso variants: GGL and FGL by Danaher et al. (2014), the Bayesian Neighbourhood Selection (BNS) by Lin et al. (2017). In the second study (provided in Supplemental Section D.1 due to space issues), we illustrate the strengths of BJNS in terms of sparsity selection and uncertainty quantification using two scenarios with four precision matrices
each. Also, a comparison with the Joint Structural Estimation Method (JSEM) by Ma and Michailidis (2016), which is a supervised approach that incorporates exact information regarding the shared sparsity structure, is provided in Supplemental Section D.3. Finally, the C++ code for BJNS is publicly available at https://www.github.com/PeymanJalali/BJNS.

5.1.1 Sparsity selection comparison with other high-dimensional methods

We consider two settings involving 6 networks. We first consider a scenario with \( K = 6 \) graphs each with \( p = 200 \) variables (see Figure G.4 in Supplemental Section G), where we generate the adjacency matrices corresponding to three distinct \( p \)-dimensional networks, so that the adjacency matrices in each column of the plot in Figure G.4 in Supplemental Section G are common. Next, we replace the connectivity structure of the bottom right diagonal block of size \( p/2 \) by \( p/2 \) in each adjacency matrix with that of another two distinct \( p/2 \)-dimensional networks, so that graphical models in each column exhibit the same connectivity pattern except in the bottom right diagonal block of their adjacency matrices. The resulting true decompositions are \( \Omega^i = \Psi^{i,i+1} + \Psi^{135} \) for odd \( i \), and \( \Omega^i = \Psi^{i-1,i} + \Psi^{246} \) for even \( i \).

The sparsity level for all six networks is set to 92% (equivalently, edge density set to 8%), while the proportion of common zeros (no edge present) across
all six networks is about 60%. Given the adjacency matrices, we then construct 
positive definite inverse covariance matrices with the nonzero off-diagonal en-
tries in each \( \Omega^k \) being uniformly generated from \([-0.6, -0.4] \cup [0.4, 0.6] \). Based 
on whether or not the corresponding shared edges have the same value, we con-
sider two settings: in the first the values for shared edges are set to be equal, 
while in the second one values for shared edges are only constrained to share the 
sign and hence their absolute values can differ across networks.

For each of these two settings, in addition to the scenario with edge density 
8 % described above, we consider another one, where 4% additional edges are 
added to each ‘true’ network to make the overall edge density 12%. For each 
of these scenarios, we generated \( n_k = 200, 300 \) independent samples for each 
\( k = 1, ..., K \) and examined the finite sample performance of different methods 
in identifying the true graphs/networks, using optimal choices of tuning param-
eters. To select the optimal value of the tuning parameters in all the penalized 
approaches, we search over a fine grid and chose the value which results in min-
imum BIC on the normalized data.

Once the off-diagonal elements are all chosen, we set the diagonal elements 
of each precision matrix to be bigger than the absolute value of its minimum 
eigenvalue. This enforcement will ensure that the resulting precision matrices 
are stable and invertable. Note that we need to compute the covariance matrices
to be able to generate synthetic data from multivariate normal distributions.

We assess the model/sparsity selection performance using the Matthews Correlation Coefficient (MCC), which is defined as

\[
MCC = \frac{TP \times TN - FP \times FN}{\sqrt{(TP + FP)(TP + FN)(TN + FP)(TN + FN)}},
\]

with TP, TN, FP and FN denoting the number of true positives, true negatives, false positives and false negatives, respectively. Larger values of MCC indicate better sparsity selection. Tables 1 and 2 show MCC values based on 50 replications for varying levels of sample size and true edge density.

Table 1: MCC values for various joint sparsity selection approaches across 6 networks, when the true sparsity patterns are random, and shared edges have exactly the same values. The MCC values are averaged over 50 replications.

<table>
<thead>
<tr>
<th>Edge density</th>
<th>Glasso</th>
<th>JEM-G</th>
<th>GGL</th>
<th>FGL</th>
<th>BNS</th>
<th>GemBag</th>
<th>BJNS</th>
</tr>
</thead>
<tbody>
<tr>
<td>n = 200</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8%</td>
<td>47 (0.009)</td>
<td>50 (0.010)</td>
<td>47 (0.009)</td>
<td>49 (0.009)</td>
<td>35 (0.010)</td>
<td>43 (0.010)</td>
<td><strong>57 (0.010)</strong></td>
</tr>
<tr>
<td>12%</td>
<td><strong>40 (0.008)</strong></td>
<td>35 (0.010)</td>
<td>35 (0.008)</td>
<td>39 (0.009)</td>
<td>30 (0.010)</td>
<td>34 (0.010)</td>
<td><strong>40 (0.010)</strong></td>
</tr>
<tr>
<td>n = 300</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8%</td>
<td>54 (0.009)</td>
<td>60 (0.010)</td>
<td>54 (0.008)</td>
<td>56 (0.008)</td>
<td>40 (0.010)</td>
<td>52 (0.010)</td>
<td><strong>70 (0.010)</strong></td>
</tr>
<tr>
<td>12%</td>
<td>47 (0.009)</td>
<td>43 (0.009)</td>
<td>42 (0.008)</td>
<td>47 (0.009)</td>
<td>35 (0.010)</td>
<td>42 (0.010)</td>
<td><strong>51 (0.010)</strong></td>
</tr>
</tbody>
</table>

It can be seen that BJNS significantly outperforms the competing methods across all settings considered in Tables 1 and 2. As expected, the overall performance of all methods improves with additional samples and worsens with
Table 2: MCC values for joint sparsity selection approaches across 6 networks, when the true sparsity patterns are random, and shared edges have the same sign but different values. The MCC values are averaged over 50 replications.

<table>
<thead>
<tr>
<th>Edge density</th>
<th>Glasso</th>
<th>JEM-G</th>
<th>GGL</th>
<th>FGL</th>
<th>BNS</th>
<th>GemBag</th>
<th>BINS</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
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<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n = 200</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8%</td>
<td>46 (0.009)</td>
<td>50 (0.011)</td>
<td>47 (0.010)</td>
<td>49 (0.009)</td>
<td>36 (0.011)</td>
<td>43 (0.010)</td>
<td><strong>54 (0.011)</strong></td>
</tr>
<tr>
<td>12%</td>
<td><strong>40 (0.008)</strong></td>
<td>36 (0.009)</td>
<td>36 (0.008)</td>
<td>39 (0.009)</td>
<td>29 (0.010)</td>
<td>35 (0.009)</td>
<td><strong>40 (0.009)</strong></td>
</tr>
<tr>
<td>n = 300</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8%</td>
<td>54 (0.010)</td>
<td>60 (0.010)</td>
<td>54 (0.009)</td>
<td>56 (0.008)</td>
<td>43 (0.010)</td>
<td>52 (0.011)</td>
<td><strong>67 (0.010)</strong></td>
</tr>
<tr>
<td>12%</td>
<td>48 (0.009)</td>
<td>44 (0.009)</td>
<td>43 (0.008)</td>
<td>48 (0.009)</td>
<td>37 (0.010)</td>
<td>43 (0.009)</td>
<td><strong>51 (0.009)</strong></td>
</tr>
</tbody>
</table>

higher edge density in the true precision matrices. Glasso is competitive in scenarios with smaller sample size (n = 200) and higher edge density (12%) due to the overall heterogeneity of the 6 networks. Note also that analogous performance patterns hold for the settings in Tables 1 and 2. Finally, to check whether these comparisons are impacted by the edge value range, we replicated Table 2, this time by generating the edges from \([-0.8, -0.3] \cup [0.3, 0.8]\), and obtain similar results. For more details please see Supplemental Section D.2.

5.2 Computational comparison with other sampling-based methods

Recent Gaussian likelihood based methods in Shaddox et al. (2018), Petersen et al. (2020) use the spike and slab prior based approach of Wang (2015) to im-
prove the computational scalability of the $G$-Wishart prior based approach of Peterson et al. (2015). These methods use a Gibbs sampler to generate samples from the resulting posterior. In addition to the precision matrix parameters, these models have latent variable parameters for edge inclusion, and other hyperparameters to encourage similarity between graphs. The overall updates for just the precision matrix parameters involve several matrix inversions, and have a computational complexity of $O(\min(np^3, p^4))$ per iteration. The updates for the other parameters including the latent variables for the sparsity patterns and other ‘relatedness’ hyperparameters require additional computation involving Metropolis-Hastings based moves. Hence, these methods are not typically scalable to settings beyond a hundred or so variables.

On the other hand, Lin et al. (2017) introduce a Bayesian analog of the regression based neighborhood selection approach of Meinshausen and Bühlmann (2006) for joint sparsity selection, called Bayesian Network Selector (BNS). They use an alternative Markov random field approach to encourage similarity among different groups. A Gibbs sampler is used to generate samples from the posterior distribution. Similar to Shaddox et al. (2018), BNS undertakes a column-wise update for the precision matrices which involves inversion of $(p - 1) \times (p - 1)$ matrices. While the computational complexity of the precision matrix parameter updates still remains $O(\min(np^3, p^3))$, the simpler
structure of the regression based approach results in several simplifications, including fewer number of matrix inversions per iteration etc. Also, the updates for the sparsity pattern based latent parameters and other hyperparameters are in general simpler than those in Shaddox et al. (2018). Hence, BNS provides significant computational improvement, and to the best of our knowledge, is the fastest sampling based Bayesian approach for joint graphical model selection.

Next, we derive the computational complexity for the BJNS Gibbs sampler described in Procedure 1 (see Supplement) which does not involve any matrix inversions. In particular, for each \((i, j)\), the full conditional posterior distribution of \(\theta_{ij}\) is a mixture of \(2^K\) univariate densities. The sparsity structure of \(\Upsilon_{(ij)(-ij)}\) and \(\Theta_{-(ij)}\) (see equations A.15 and A.16) and analysis similar to Khare et al. (2015, Lemma 6) imply that computation of each mean \(\mu_{l,ij}\) in (B.24) can be achieved in \(O(\min(n, p))\) operations. Hence, sampling from the mixture would require \(2^K O(\min(n, p))\) operations, implying that the overall worst case computational complexity per iteration is \(2^K O(\min(np^2, np^3))\).

Hence, the computational complexity of the vanilla BJNS algorithm is \(\frac{2^K}{Kp}\) times that of BNS. In many applications, such as those considered in Shaddox et al. (2018), Petersen et al. (2020) and the IBD data considered in Section 6, the number of groups \(K = 4\). The BJNS/BNS computational complexity ratio turns out to be 0.065 (OxPhos pathway for the COPD data in Shaddox et al. (2018)),
0.04 (Alzheimer’s MRI data in Petersen et al. (2020)) and 0.01 (IBD data in Section 6). In general, the computational complexity of BJNS is much smaller than that of BNS for typical genomics data sets, where $K$ is small/moderate and $p$ is much larger. To further illustrate the latter point, we provide a wall clock time comparison between BJNS and BNS in Table 3 for simulations with various number of variables $p$ and groups $K$. The true data generating process is identical to the one described in the previous subsection (we use $n = 3p/2$). In order to make a fair comparison, we compiled the MATLAB code of BNS provided by its authors, to machine language code. All computations were done on an intel CPU with 6Gb of memory. Overall, BJNS has a significantly lower wall clock time requirement compared to BNS across all settings.

### Table 3: Wall clock time for BJNS and BNS (4000 iterations).

<table>
<thead>
<tr>
<th></th>
<th>$p = 50$</th>
<th>$p = 200$</th>
<th>$p = 500$</th>
<th>$p = 50$</th>
<th>$p = 200$</th>
<th>$p = 500$</th>
</tr>
</thead>
<tbody>
<tr>
<td>BJNS</td>
<td>0.003h</td>
<td>0.092h</td>
<td>1.193h</td>
<td>0.016h</td>
<td>0.510h</td>
<td>5.738h</td>
</tr>
<tr>
<td>BNS $k = 2$</td>
<td>0.029h</td>
<td>0.282h</td>
<td>6.503h</td>
<td>$k = 4$</td>
<td>0.128h</td>
<td>0.672h</td>
</tr>
</tbody>
</table>

If $K$ is large, we develop a pre-processing step in Supplemental Section E to reduce the size of the mixture from $2^K$ to a much smaller user specified number $M_K$. Taking into account the pre-processing step, the overall per iteration computational complexity is further reduced to $(K^2 + M_K)O(\min(np^2, np^3))$. 

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6. An Application of BJNS to Metabolomics Data

In this section, we employ the proposed methodology to obtain networks across four groups of patients that participated in the Integrative Human Microbiome Project. The data were downloaded from the Metabolomics Workbench (Study ID ST000923) and correspond to measurements of 428 primary and secondary metabolites and lipids from stool samples of 542 subjects, partitioned in the following groups: inflammatory bowel disease (IBD) patients (males $n_1 = 202$ and females $n_2 = 208$) and non-IBD controls (males $n_3 = 72$ and females $n_4 = 70$), Groups 1-4, respectively. Since there are two factors in the study design, the following model was fitted to the data.

$$
\Omega^1 = \Psi^1 + \Psi^{12} + \Psi^{13} + \Psi^{1234}, \quad \Omega^3 = \Psi^3 + \Psi^{13} + \Psi^{34} + \Psi^{1234},
$$

$$
\Omega^2 = \Psi^2 + \Psi^{12} + \Psi^{24} + \Psi^{1234}, \quad \Omega^4 = \Psi^4 + \Psi^{24} + \Psi^{34} + \Psi^{1234}.
$$

The edge counts of the estimated precision matrices are shown in Table G.1 (set1: 289 lipids and set2: 139 proteins, set1.2: interaction edges between set1 and set2), together with the components in the proposed decomposition. We also applied JEM-G and the corresponding edge counts are also presented in Table G.1. Note that BJNS detects a large number of edges that are shared across all groups, indicating common patterns. Further, the component shared between male and female IBD patients has a fairly large number of edges, indicating that the disease status exhibits commonalities across genders. The graphs produced
by JEM-G are much more similar which is consistent with the fact that JEM-G differentiates the graphs only by a multiplicative factor. JEM-G also tends to provide significantly higher number of edges for each network.

Table G.2 illustrates the detailed uncertainty quantification provided by BJNS by providing posterior inclusion probabilities for all the 10 possible subsets for 10 chosen edges. Figure G.5 presents the common connectivity patterns shared across all four groups. The primary and secondary metabolites are depicted in red, while the lipids in blue. Not surprisingly, as shown in sub-figure G.5a, primary metabolites (those involved in cellular growth, development and reproduction) form a fairly strongly connected network. In addition, based on the sub-network in G.5b, different fairly strongly connected subnetworks amongst lipids are present including dicetylglycerols (DAG) with tristerylglycerols (TAG) that are main constituents of animal and vegetable fat (upper right corner of the plot) and various phospholipids (upper left corner of the plot). On the other hand, the connectivity between lipids (whose functions include storing energy, signaling and acting as structural components of cell membranes) to the metabolites is not particularly strong, see network in sub-figure G.5c. In general, the results reveal interesting patterns that can be used to understand progression of IBD disease.
7. High Dimensional Sparsity Selection Consistency

Let \( \{ \Omega^{k,0} \}_{k=1}^{K} \) denote the true precision matrices, and \( \mathbb{P}_0 \) denote the probability measure associated with the corresponding true data generating model. Note that the identifiability constraint in Section 2.1 assumes that whenever an edge \((i, j)\) is shared among a subset \( S \), the magnitudes of the \((i, j)^{th}\) entries in the corresponding precision matrices are exactly the same. **We allow \( \{ \Omega^{k,0} \}_{k=1}^{K} \) to deviate from this assumption, i.e., the entries in \( \{ \Omega^{k,0} \}_{k \in S} \) are allowed to have different magnitudes. We will show that as long as the deviation in the magnitudes is moderate, BJNS still leads to consistent high-dimensional model selection.**

Define matrices \( \{ \Omega^{k,0} \}_{k=1}^{K} \) as follows. For each \( 1 \leq i \neq j \leq p \) and \( 1 \leq k \leq K \), if \( \Omega^{k,0}_{ij} \neq 0 \), set

\[
\Omega^{k,0}_{ij} = \sum_{k' : \Omega^{k',0}_{ij} \neq 0} \Omega^{k',0}_{ij},
\]

and set \( \Omega^{k,0}_{ij} = \Omega^{k,0}_{ij} \) otherwise. The matrices \( \{ \Omega^{k,0} \}_{k=1}^{K} \) can be thought of as harmonized versions of the true precision matrices \( \{ \Omega^{k,0} \}_{k=1}^{K} \), which obey the identifiability constraint in Section 2.1. We define \( D_n = \max_{1 \leq i \neq j \leq p, 1 \leq k \leq K} \left| \Omega^{k,0}_{ij} - \Omega^{k,0}_{ij} \right| \) as a discrepancy measure between the true and harmonized precision matrices.

Let \( \left\{ \Psi^{r,0} , r \in \bigcup_{k=1}^{K} \vartheta_k \right\} \) be such that \( \Omega^{k,0} = \sum_{r \in \vartheta_k} \Psi^{r,0} \) corresponds to the decomposition of each harmonized precision matrix for \( k = 1, \ldots, K \). Let \( \Theta^0 \) be the vectorized version (see (3.5)) of the off-diagonal elements of the matrices \( \left\{ \Psi^{r,0} , r \in \bigcup_{k=1}^{K} \vartheta_k \right\} \). Let \( t \) denote the sparsity pattern in \( \Theta^0 \), \( \mathcal{M}_t \) denote the
corresponding parameter space, and $d_t$ denote the number of non-zeros in $\Theta^0$.

Note that our main objective is to accurately select the shared sparsity patterns in the off-diagonal entries of the group-specific precision matrices. Hence, following pseudo-likelihood based high-dimensional consistency proofs in Peng et al. (2009); Khare et al. (2015); Atchade (2019), we consider a setting where sufficiently accurate estimates $\left\{ \hat{\Omega}_{ii}^k \right\}_{1 \leq i \leq p}^{1 \leq k \leq K}$ of the diagonal entries are first obtained (see eq. (F.34) in the Supplemental Document, and the subsequent discussion for a quick way of obtaining such estimates using parallel lasso regressions). Denote the resulting estimates of the vectors $\Delta$ and $a$ by $\hat{\Delta}$ and $\hat{a}$, respectively. We now consider the accuracy of shared sparsity pattern selection for the off-diagonal entries after running the BJNS procedure with the diagonal entries fixed at $\hat{\Delta}$. For this section, we assume that the entries of $\lambda$ are fixed.

The following assumptions are needed to establish our consistency results.

**ASSUMPTION 1:** $d_t \sqrt{\frac{\log p}{n}} \to 0$, as $n \to \infty$.

This standard assumption essentially states that the number of variables $p$ has to grow slower than $e^{(\frac{n}{d_t^2})}$, see for example Banerjee and Ghosal (2014, 2015).

**ASSUMPTION 2:** (Sub-Gaussianity). There exists $c > 0$, independent of $n$ and $K$ such that $\mathbb{E} \left[ \exp \left( \alpha^\prime y_{k,i}^k \right) \right] \leq \exp (c\alpha^\prime \alpha)$. Theorem II below will show the BJNS procedure is robust (in terms of consistency) even under misspecification of the data generating distribution, as long as its tails are sub-Gaussian.
ASSUMPTION 3: (Bounded eigenvalues). There exists $\varepsilon_0 > 0$, independent of $n$ and $K$, such that the eigenvalues of $\Omega^{k,0}$ are uniformly bounded above and below by $\varepsilon_0$ and $1/\varepsilon_0$ respectively. This is a standard assumption in high dimensional consistency analysis, see for example Cao et al. (2019).

ASSUMPTION 4: (Signal Strength). Let $s_n$ be the smallest non-zero entry (in magnitude) in the vector $\Theta_0$. We assume $\frac{1}{n} \log s_n + nd_b \rightarrow 0$, where $b_n = \log p + nd_tD_n^2$. This is again a standard assumption. Similar assumptions on the signal size can be found in Khare et al. (2015); Peng et al. (2009).

ASSUMPTION 5: (Edge probability decay). Let $q_1 = e^{-a_2 d_b n}$, $q_2 = e^{-a_3 n b_n}$, and $\tau_n = \frac{\sqrt{n}}{\log p}$, for constants $a_1, a_2$ (not depending on $n$) specified in (F.39) in the Supplement. Assumption 5 a priori penalizes patterns with too many non-zeros (see Narisetty and He (2014); Cao et al. (2019) for similar assumptions). Next, we establish that the posterior mass assigned to the true model converges to one in probability (under the true model).

**Theorem 1.** (Strong Selection Consistency) Based on the joint posterior distribution given in (B.21), and under Assumptions 1-5, the following holds,

$$
\pi \left\{ \Theta \in \mathcal{M}_t \mid \hat{\Delta}, \mathcal{Y} \right\} \xrightarrow{P_0} 1, \quad \text{as} \quad n \rightarrow \infty.
$$

**Conclusion**

A comprehensive Bayesian methodology for joint estimation of multiple graphical models is introduced. Leveraging a novel multivariate prior distribution
along with a pseudo-likelihood, our model enables fast and provably accurate estimation. We illustrated how our methodology utilizes the information that is shared across the groups to provide higher accuracy. A computational strategy to deal with a large number of networks \( K \) is also developed and numerically investigated in Section E of the Supplement. Further, simulation studies illustrate the superior performance of BJNS vis-a-vis frequentist and Bayesian competitors. Finally, an application to IBD disease progression reveals interesting patterns.

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**References**


