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Bayesian Canonical Correlation Analysis

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Abstract

Canonical correlation analysis (CCA) is a classical method for seeking correlations between two multivariate data sets. During the last ten years, it has received more and more attention in the machine learning community in the form of novel computational formulations and a plethora of applications. We review recent developments in Bayesian models and inference methods for CCA which are attractive for their potential in hierarchical extensions and for coping with the combination of large dimensionalities and small sample sizes. The existing methods have not been particularly successful in fulfilling the promise yet; we introduce a novel efficient solution that imposes group-wise sparsity to estimate the posterior of an extended model which not only extracts the statistical dependencies (correlations) between data sets but also decomposes the data into shared and data set-specific components. In statistics literature the model is known as inter-battery factor analysis (IBFA), for which we now provide a Bayesian treatment.

Keywords: Bayesian modeling, canonical correlation analysis, group-wise sparsity, inter-battery factor analysis, variational Bayesian approximation

1. Introduction

Canonical correlation analysis (CCA), originally introduced by Hotelling (1936), extracts linear components that capture correlations between two multivariate random variables or data sets. During the last decade the model has received a renewed interest in the machine learning community as the standard model for unsupervised multi-view learning settings. In a sense, it is the analogue of principal component analysis (PCA) for two co-occurring observations, or views, retaining the positive properties of closed-form analytical solution and ease of interpretation of its more popular cousin.

A considerable proportion of the work has been on non-linear extensions of CCA, including neural network based solutions (Hsieh, 2000) and kernel-based variants (Bach and Jordan, 2002; Lai and Fyfe, 2000; Melzer et al., 2001). This line of research has been covered in a comprehensive overview by Hardoon et al. (2004), and hence will not be discussed in detail in this article. Instead, we review a more recent trend treating CCA as a generative model, initiated by the work of Bach and

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Jordan (2005). Most works in the generative approach retain the linear nature of CCA, but provide inference methods more robust than the classical linear algebraic solution and, more importantly, the approach leads to novel models through simple changes in the generative description or via the basic principles of hierarchical modeling.

The generative modelling interpretation of CCA is essentially equivalent to a special case of a probabilistic interpretation (Browne, 1979) of a model called inter-battery factor analysis (IBFA; Tucker, 1958). While the analysis part of Browne (1979) is limited to the special case of CCA, the generic IBFA model describes not only the correlations between the data sets but provides also components explaining the linear structure within each of the data sets. One way of thinking about IBFA is that it complements CCA by providing a PCA-like component description of all the variation not captured by the correlating components. If the analysis focuses on only the correlating components, or equivalently the latent variables shared by both data sets, the solution becomes equivalent to CCA. However, the extended model provides novel application opportunities not immediately apparent in the more restricted CCA model.

The IBFA model has recently been re-invented in the machine learning community by several authors (Klami and Kaski, 2006, 2008; Ek et al., 2008; Archambeau and Bach, 2009), resulting in probabilistic descriptions identical with that of Browne (1979). The inference has been primarily based on finding the maximum likelihood or maximum a posteriori solution of the model, with practical algorithms based on expectation maximization. Since the terminology of calling these models (probabilistic) CCA has already become established in the machine learning community, we will regard the names CCA and IBFA interchangeable. Using the term CCA emphasizes finding of the correlations and shared components, whereas IBFA emphasizes the decomposition into shared and data source-specific components.

In this paper we extend this IBFA/CCA work to a fully Bayesian treatment, extending our earlier conference paper (Virtanen et al., 2011), and in particular provide two efficient inference algorithms, a variational approximation and a Gibbs sampler, that automatically learn the structure of the model that is, in the general case, unidentifiable. The model is solved as a generic factor analysis (FA) model with a specific group-wise sparsity prior for the factor loadings or projections, and an additional constraint tying the residual variances within each group to be the same. We demonstrate how the model not only finds the IBFA solution, but also provides a CCA solution superior to the earlier Bayesian variants of Klami and Kaski (2007) and Wang (2007).

The technical description of the model and its connection to other models are complemented with demonstrations on practical application scenarios for the IBFA model. The main purpose of the experiments is to show that the tools find the intended solution, and to introduce prototypical application cases.

2. Canonical Correlation Analysis

Before explaining the Bayesian approach for canonical correlation analysis (CCA), we briefly introduce the classical CCA problem. Given two co-occurring random variables with N observations collected as matrices $\mathbf{X}_1 \in \mathbb{R}^{D_1 \times N}$ and $\mathbf{X}_2 \in \mathbb{R}^{D_2 \times N}$, the task is to find linear projections $\mathbf{U} \in \mathbb{R}^{D_1 \times K}$ and $\mathbf{V} \in \mathbb{R}^{D_2 \times K}$ so that the correlation between $\mathbf{u}_k^T \mathbf{X}^{(1)}$ and $\mathbf{v}_k^T \mathbf{X}^{(2)}$ is maximized for the components k , under the constraint that $\mathbf{u}_k^T \mathbf{X}^{(1)}$ and $\mathbf{u}_{k'}^T \mathbf{X}^{(1)}$ are uncorrelated for all $k \neq k'$ (and similarly for the

other view). The solution can be found analytically by solving the eigenvalue problems

$$\begin{aligned} \mathbf{C}_{11}^{-1}\mathbf{C}_{12}\mathbf{C}_{22}^{-1}\mathbf{C}_{21}\mathbf{u} &= \rho^2\mathbf{u}, \\ \mathbf{C}_{22}^{-1}\mathbf{C}_{21}\mathbf{C}_{11}^{-1}\mathbf{C}_{12}\mathbf{v} &= \rho^2\mathbf{v}, \end{aligned} \tag{1}$$

where

$$\mathbf{C} = \begin{bmatrix} \mathbf{C}_{11} & \mathbf{C}_{12} \\ \mathbf{C}_{21} & \mathbf{C}_{22} \end{bmatrix}$$

is the joint covariance matrix of $\mathbf{x}^{(1)}$ and $\mathbf{x}^{(2)}$ and ρ denotes the canonical correlation. In practice all components can be found by solving a single generalized eigenvalue problem. For more detailed discussion on classical CCA, see for instance the review of Hardoon et al. (2004).

3. Model

Our Bayesian approach to CCA is based on latent variable models and linear projections. At the core of the generative process is an unobserved latent variable $\mathbf{z} \in \mathbb{R}^{K \times 1}$, which is transformed via linear mappings to the observation spaces to represent the two multivariate random variables $\mathbf{x}^{(1)} \in \mathbb{R}^{D_1 \times 1}$ and $\mathbf{x}^{(2)} \in \mathbb{R}^{D_2 \times 1}$. The observed data samples are provided as matrices $\mathbf{X}^{(m)} = [\mathbf{x}_1^{(m)}, \dots, \mathbf{x}_N^{(m)}] \in \mathbb{R}^{D_m \times N}$ with N observations. To simplify the notation, we denote by $\mathbf{x} = [\mathbf{x}^{(1)}; \mathbf{x}^{(2)}]$ and $\mathbf{X} = [\mathbf{X}^{(1)}; \mathbf{X}^{(2)}]$ the feature-wise concatenation of the two random variables. Throughout the paper we use the superscript (m) , where m is 1 or 2, to denote the view or data set in question, though for scalar variables (such as D_m) we use the subscript without risk of confusion to streamline the notation. For matrices and vectors the subscripts are used to indicate the individual elements, with $\mathbf{X}_{:,n}$ denoting the whole n th column of \mathbf{X} (also denoted by \mathbf{x}_n to simplify the notation when appropriate) and $\mathbf{X}_{d,:}$ denoting the d th row treated as a column vector. Finally, we use $\mathbf{0}$ and \mathbf{I} to denote zero- and identity matrices of sizes which make sense in the context, without cluttering the notation.

3.1 Inter-battery Factor Analysis

In the latent variable model studied in this work,

$$\begin{aligned} \mathbf{z} &\sim \mathbf{N}(\mathbf{0}, \mathbf{I}), \\ \mathbf{z}^{(m)} &\sim \mathbf{N}(\mathbf{0}, \mathbf{I}), \\ \mathbf{x}^{(m)} &\sim \mathbf{N}(\mathbf{A}^{(m)}\mathbf{z} + \mathbf{B}^{(m)}\mathbf{z}^{(m)}, \Sigma^{(m)}), \end{aligned} \tag{2}$$

following the probabilistic interpretation of inter-battery factor analysis by Browne (1979).¹ The notation $\mathbf{N}(\boldsymbol{\mu}, \Sigma)$ corresponds to the normal distribution with mean $\boldsymbol{\mu}$ and covariance Σ . Here the $\Sigma^{(m)} \in \mathbb{R}^{D_m \times D_m}$ are diagonal matrices, indicating independence of the noise over the features. Our practical solutions will further simplify the model by assuming isotropic noise, but could easily be extended to generic diagonal noise covariances as well. A plate diagram of the model is given in Figure 1.

The conceptual meaning of the various terms in the model is as follows. The shared latent variables \mathbf{z} capture the variation common to both data sets, and they are transformed to the observation

1. The original definition by Browne (1979) allows for more relaxed definitions for the various covariance terms, but in practice he resorts to the choices made above in the actual analysis part of his work and makes the same independence assumptions.

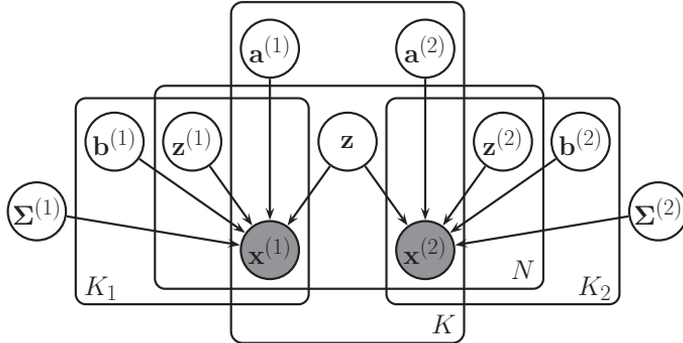


Figure 1: Graphical illustration of the inter-battery factor analysis (IBFA) model as a plate diagram. The shaded nodes $\mathbf{x}^{(m)}$ denote the two observed random variables, and the latent variables \mathbf{z} capture the correlations between them. The variation specific to each view is modeled with view-specific latent variables $\mathbf{z}^{(m)}$. The parameters of the model are the linear projections from the \mathbf{z} to the data ($\mathbf{A}^{(m)}$ with columns $\mathbf{a}^{(m)}$) and from the $\mathbf{z}^{(m)}$ to the data ($\mathbf{B}^{(m)}$ with columns $\mathbf{b}^{(m)}$), complemented by the residual noise covariance of each view denoted by $\Sigma^{(m)}$.

space by the linear mappings $\mathbf{A}^{(m)}\mathbf{z}$, where $\mathbf{A}^{(m)} \in \mathbb{R}^{D_m \times K}$. The remaining variation is modeled by the latent variables $\mathbf{z}^{(m)} \in \mathbb{R}^{K_m \times 1}$ specific to each data set, transformed to the observation space by another linear mapping $\mathbf{B}^{(m)}\mathbf{z}^{(m)}$, where $\mathbf{B}^{(m)} \in \mathbb{R}^{D_m \times K_m}$. The actual observations are then generated by adding up these two terms, followed by addition of noise that is independent over the dimensions. We assume zero-mean data without loss of generality; the model could include a separate mean parameter whose estimate would anyway converge close to the empirical mean which can equivalently be subtracted from the data prior to the analysis. We also assume fully observed data; techniques similar to what Ilin and Raiko (2010) propose for Bayesian PCA could be adopted to handle missing data.

In terms of classical models, the model can be interpreted as CCA complemented by two separate FA models (or PCA models if assuming isotropic noise) factorizing the residuals of the CCA within each data set. This connection will become more apparent in the following sections when the probabilistic interpretation of CCA is introduced. Typically both K and K_m are smaller than the corresponding data dimensionality, implying that the model provides low-rank approximations for the two data matrices.

3.2 Probabilistic Canonical Correlation Analysis

There exists a simple way of converting the IBFA model of (2) into a probabilistic version of CCA (Bach and Jordan, 2005; Browne, 1979; De Bie and De Moor, 2003). The process starts by integrating out the view-specific latent variables $\mathbf{z}^{(m)}$, to reach a model that has explicit components only for the shared variation similarly to how CCA only explains the correlations. Simple algebraic

manipulation gives the model

$$\begin{aligned} \mathbf{z} &\sim \mathcal{N}(\mathbf{0}, \mathbf{I}), \\ \mathbf{x}^{(m)} &\sim \mathcal{N}(\mathbf{A}^{(m)}\mathbf{z}, \mathbf{B}^{(m)}\mathbf{B}^{(m)T} + \Sigma^{(m)}). \end{aligned}$$

The latent representation of this model is simpler, only containing the \mathbf{z} instead of three separate sets of latent variables, but the diagonal covariance of the IBFA model is replaced with $\mathbf{B}^{(m)}\mathbf{B}^{(m)T} + \Sigma^{(m)}$. In effect, the view-specific variation is now modeled only implicitly, in form of correlating noise. If we further assume that the dimensionality of the $\mathbf{z}^{(m)}$ is sufficient for modeling all such variation, the model can be re-parameterized with $\Psi^{(m)} = \mathbf{B}^{(m)}\mathbf{B}^{(m)T} + \Sigma^{(m)}$ without loss of generality. This results in the model

$$\begin{aligned} \mathbf{z} &\sim \mathcal{N}(\mathbf{0}, \mathbf{I}), \\ \mathbf{x}^{(m)} &\sim \mathcal{N}(\mathbf{A}^{(m)}\mathbf{z}, \Psi^{(m)}), \end{aligned} \tag{3}$$

where $\Psi^{(m)}$ is a generic covariance matrix. This holds even if assuming isotropic noise in (2).

Browne (1979) proved that (3) is equivalent to classical CCA, by showing how the maximum likelihood solution finds the same canonical weights as regular CCA, up to a rotation. Bach and Jordan (2005) proved the same result through a slightly different derivation, whereas De Bie and De Moor (2003) provided a partial proof showing that the CCA solution is a stationary point of the likelihood. The fundamental result of these derivations is that the maximum likelihood estimates $\hat{\mathbf{A}}^{(m)}$ correspond to rank-preserving linear transformations of \mathbf{U} and \mathbf{V} , the solutions of (1). While the connection was shown for the case with a generic Ψ , it holds also for the IBFA model as long as the rank of $\mathbf{B}^{(m)}$ is sufficient for modeling all data set-specific variation. This is because the model itself is the same, it just explicitly includes the nuisance parameters $\mathbf{z}^{(m)}$ and $\mathbf{B}^{(m)}$.

Even though the generative formulation is equivalent to classical CCA in the sense that they both find the same subspace, one difference pointed out also by Browne (1979) is worth emphasizing: The generative formulation maintains a single latent variable \mathbf{z} that captures the shared variation, whereas CCA results in two separate but correlating variables obtained by projecting the observed variables into the correlating subspace. It is, however, possible to move between these representations; a single latent variable can be obtained by averaging the canonical scores ($\mathbf{u}_k^T \mathbf{X}^{(1)}$ and $\mathbf{v}_k^T \mathbf{X}^{(2)}$) of regular CCA; two separate latent variables can be produced with the generative formulation by estimating the distribution of \mathbf{z} conditional on having observed only one of the views ($p(\mathbf{z}|\mathbf{x}^{(1)})$ and $p(\mathbf{z}|\mathbf{x}^{(2)})$).

3.3 Identifiability

The model in (2) is in general unidentifiable in two respects. The first is shared with the marginalized version in (3): the models are invariant to rank-preserving linear transformations. For all invertible $\mathbf{R} \in \mathbb{R}^{K \times K}$ we have $\mathbf{A}^{(m)}\mathbf{z} = \mathbf{A}^{(m)}\mathbf{R}\mathbf{R}^{-1}\mathbf{z}$, and hence the solution is defined only up to such transformations. In other words, the model finds the same subspace as the classical CCA would, but extracting the specific components requires further constraints or postprocessing. Browne (1979) resorts to simple identifiability constraints borrowed from regular factor analysis, whereas Archambeau et al. (2006) provide a post-processing step that is close to applying regular CCA to the covariance matrices of the probabilistic solution.

The full IBFA model (2) has additional degrees of freedom in terms of component allocation. The model comes with three separate sets of latent variables with component numbers K , K_1 and K_2 . However, individual components can be moved between these sets without influencing the likelihood of the observed data; removal of a shared component can always be compensated by introducing two view-specific components, one for each data set, that have the same latent variables. In practice, all solutions for the full IBFA model hence need to carefully address the choice of model complexity. In the next section we will introduce one such solution, based on Bayesian inference.

3.4 The Role of View-specific Variation

The models (2) and (3) are both very closely related to probabilistic formulation of PCA, FA, and many other simple matrix factorizations. The crucial difference worth pointing out is the definition of the noise. Instead of assuming independent noise over the dimensions the CCA model allows for arbitrary correlations between them. This is done either by explicitly parameterizing the noise through a covariance matrix $\Psi^{(m)}$ as in (3) or by the separate view-specific components $\mathbf{B}^{(m)}\mathbf{z}^{(m)}$ as in (2).

Modeling the correlations in view-specific noise is crucial for extracting the true correlations between the views. This is easy to illustrate by constructing counter-examples where the correlating dimensions are of smaller scale than some strong view-specific variation. Any joint model assuming independent noise over the dimensions will find the view-specific variation as the most prominent components. It may be possible to identify these components as view-specific in a post-processing step to reach interpretation similar to CCA, but directly modeling the view-specific variation as separate components has obvious advantages.

The importance of modeling the variation within each view in addition to the shared effects is so subtle that even some authors claiming to work with CCA have ignored it. For example, Shon et al. (2006), Fujiwara et al. (2009), and Rai and Daumé III (2009) all describe their models as CCA, but eventually resort to assuming independent noise on top of the shared components. This is a reasonable assumption that simplifies computation dramatically, but it also means that the models do not correspond to CCA but are instead variants of collective matrix factorization (CMF; Singh and Gordon 2008). They are useful tools for multi-view data, but it is important to realize that the simplifying assumption dramatically changes the nature of the model. In particular, such models are likely to misinterpret strong view-specific variation as a shared effect, since they have no means of explaining it otherwise. Our choice of modeling the view-specific variation as a low-rank process results in similar computational performance as ignoring the view-specific variation, but retains the capability of modeling also view-specific variation.

4. Inference

For learning the IBFA model we need to infer both the latent signals \mathbf{z} and $\mathbf{z}^{(m)}$ as well as the linear projections $\mathbf{A}^{(m)}$ and $\mathbf{B}^{(m)}$ from data. For this purpose, we need to estimate the posterior distribution $p(\mathbf{z}, \mathbf{z}^{(1)}, \mathbf{z}^{(2)}, \mathbf{A}^{(1)}, \mathbf{A}^{(2)}, \mathbf{B}^{(1)}, \mathbf{B}^{(2)}, \Sigma^{(1)}, \Sigma^{(2)} | \mathbf{X}^{(1)}, \mathbf{X}^{(2)})$, and marginalize over the possibly uninteresting variables. In this section we first review the earlier inference solutions for the Bayesian CCA model without presenting the technical details, and then proceed to explaining our solution for the full Bayesian IBFA model.

Before explaining the Bayesian inference solutions, we mention earlier maximum likelihood solutions for completeness. Bach and Jordan (2005) gave an expectation maximization algorithm

for the pure CCA model, whereas Klami and Kaski (2008) extended it for the IBFA model. Both approaches generalize immediately to seeking maximum a posteriori (MAP) estimates, which provides a justified way for adding regularization in the solution. Here, however, we are interested in analysis of the full posterior distribution.

4.1 Bayesian Inference

For Bayesian analysis, the model needs to be complemented with priors for the model parameters. Bayesian treatments of CCA were independently proposed by Klami and Kaski (2007) and Wang (2007). Both formulations use the inverse-Wishart distribution as a prior for the covariance matrices $\Psi^{(m)}$ in (3) and apply the automatic relevance determination (ARD; Neal, 1996) prior for the linear mappings $\mathbf{A}^{(m)}$.

The ARD is a Normal-Gamma prior for the projection weights. For each component (column) $\mathbf{a}_k^{(m)}$ the prior specifies a precision $\alpha_k^{(m)}$ that controls the scale of the values for that component:²

$$\begin{aligned} \text{ARD}(\mathbf{A}^{(m)}|\alpha_0, \beta_0) &= \prod_{k=1}^K p(\mathbf{a}_k^{(m)}|\alpha_k^{(m)})p(\alpha_k^{(m)}|\alpha_0, \beta_0), \\ \alpha_k^{(m)} &\sim \text{Gamma}(\alpha_0, \beta_0), \\ \mathbf{a}_k^{(m)} &\sim \text{N}(\mathbf{0}, (\alpha_k^{(m)})^{-1}\mathbf{I}). \end{aligned}$$

The hyperpriors α_0, β_0 are set to small values (in our experiments to $\alpha_0 = \beta_0 = 10^{-14}$) to obtain a relatively noninformative prior with wide support.³ The posterior of the model then becomes one where the number of components is automatically selected by pushing $\alpha_k^{(m)}$ of unnecessary components towards infinity. A justification for this observation is obtained by integrating $\alpha_k^{(m)}$ out in the above prior; we then get a heavy-tailed prior for the elements of $\mathbf{A}^{(m)}$ with considerable posterior mass around zero. The component choice can be made more robust by further assuming $\alpha^{(1)} = \alpha^{(2)}$, which corresponds to placing the ARD prior for $\mathbf{A} = [\mathbf{A}^{(1)}; \mathbf{A}^{(2)}]$ (Klami and Kaski, 2007). More data will then be used for determining the activity of each component, but data sets with comparable scale are required. Further insights into the ARD prior are provided by Wipf and Nagarajan (2008).

For the covariance matrices $\Psi^{(m)}$ a natural choice is to use a conjugate inverse-Wishart prior

$$\Psi^{(m)} \sim \text{IW}(\mathbf{S}_0, \nu_0)$$

with ν_0 degrees of freedom and scale matrix \mathbf{S}_0 , which results in positive definite draws as long as the degrees of freedom (which for N samples becomes $N + \nu_0$ in the posterior) is at least equal the data dimensionality. Both Klami and Kaski (2007) and Wang (2007) adopted this choice.

Given the above priors, several inference techniques for the posterior are feasible. Wang (2007) provided a variational mean-field algorithm, whereas Klami and Kaski (2007) used Gibbs sampling. Both of these algorithms are fairly straightforward and easy to derive, since all conditional distributions are conjugate. The former is more efficient in determining the correct model complexity due

2. Note that ARD generates both the matrix $\mathbf{A}^{(m)}$ as well as the scales $\alpha^{(m)}$, and hence notation $\text{ARD}(\mathbf{A}^{(m)}, \alpha^{(m)}|\alpha_0, \beta_0)$ would be more accurate. However, since $\alpha^{(m)}$ is irrelevant for the rest of the model, we adopt the more compact notation.

3. The distribution is flat over the positive real line, but slightly favors values near zero.

to the ARD prior updates being more efficient in the variational framework, whereas the latter is easier to extend, as demonstrated by Klami and Kaski (2007) by using the Bayesian CCA as part of a non-parametric hierarchical model. Further extensions of the CCA model, described in Section 6, have used both approaches.

Despite the apparent simplicity of the derivation, it is worth pointing out that inference of the Bayesian CCA model is difficult for large dimensionalities. This is because we need to estimate the posterior distribution over the $D_m \times D_m$ covariance matrices $\Psi^{(m)}$. The inference algorithms generally need to invert those matrices in every step, resulting in $O(D_m^3)$ complexity. More importantly, providing accurate estimates would require extremely large sample sizes; the covariance matrix has $O(D_m^2)$ parameters, which is often well above the data set size. Hence the direct Bayesian treatment of CCA needs to resort to either using very strong priors (for example, favoring diagonal covariance matrices and hence regularizing the model towards Bayesian PCA), or it will end up doing inference over a very wide posterior. Consequently, all the practical applications of Bayesian CCA in the earlier works have been for relatively low-dimensional data; the original works by Klami and Kaski (2007) and Wang (2007) had at most 8 dimensions in any of their experiments. Later applications have typically used some alternative dimensionality reduction techniques to make Bayesian CCA feasible for otherwise too high-dimensional data (Huopaniemi et al., 2009, 2010).

4.2 Group-wise Sparsity for IBFA

While the above solutions are sufficient for the CCA model, barring the difficulties with high dimensionality, the full IBFA model requires more advanced inference methods. Next we will introduce a novel inference solution extending our earlier conference paper (Virtanen et al., 2011). Besides providing a Bayesian inference technique for the IBFA model, the algorithm is applicable also to the regular CCA case and, as will be shown later, actually is superior to the earlier solutions also for that scenario.

A main challenge in learning the Bayesian IBFA (BIBFA) model, as discussed in Section 3.3, is that it requires learning three separate sets of components and the solution is unidentifiable with respect to allocating components to the three groups. A central element in our solution is to replace these three sets with just one set, and solve the allocation by requiring the projections to be sparse in a specific structured way. This is done in a way that does not change the model itself, but allows automatic complexity selection.

We start with a straightforward re-formatting of the model. We define $\mathbf{y} = [\mathbf{z}; \mathbf{z}^{(1)}; \mathbf{z}^{(2)}] \in \mathbb{R}^{K_c \times 1}$, where $K_c = K + K_1 + K_2$, as the concatenation of the three latent variables and set

$$\mathbf{W} = \begin{bmatrix} \mathbf{A}^{(1)} & \mathbf{B}^{(1)} & \mathbf{0} \\ \mathbf{A}^{(2)} & \mathbf{0} & \mathbf{B}^{(2)} \end{bmatrix} \tag{4}$$

and

$$\Sigma = \begin{bmatrix} \Sigma^{(1)} & \mathbf{0} \\ \mathbf{0} & \Sigma^{(2)} \end{bmatrix}.$$

Now we can write (2) equivalently as

$$\begin{aligned} \mathbf{y} &\sim \mathcal{N}(\mathbf{0}, \mathbf{I}), \\ \mathbf{x} &\sim \mathcal{N}(\mathbf{W}\mathbf{y}, \Sigma). \end{aligned} \tag{5}$$

In other words, we are now analyzing the feature-wise concatenation of the data sources with a single latent variable model with diagonal noise covariance $\Sigma \in \mathbb{R}^{D \times D}$, where $D = D_1 + D_2$, and a projection matrix $\mathbf{W} \in \mathbb{R}^{D \times K_c}$ with the specific structure shown in (4).

Ignoring the structure in \mathbf{W} , the model is actually a Bayesian factor analysis model (Ghahramani and Beal, 2000). If Σ is further assumed to be spherical ($\Sigma = \sigma^2 \mathbf{I}$), the model equals the Bayesian PCA (Bishop, 1999) with the specific structure in \mathbf{W} . We use $\Sigma^{(m)} = \sigma_m^2 \mathbf{I}$ with a Gamma prior for the noise precisions $\tau_m = \sigma_m^{-2}$. That is, we make the PCA assumption separately for both data sets. However, it is important to remember that the model still allows for dependencies between the features of both views, by modeling them with $\mathbf{B}^{(m)} \mathbf{z}^{(m)}$. Hence, the spherical noise covariance does not restrict the flexibility of the model but decreases the number of parameters. Alternatively, we could allow each dimension to have their own variance parameter as in factor analysis models; this could be useful when the scales of the variables are very different.

Since efficient inference solutions are available for regular factor analysis, the only challenge in learning the BIBFA model is in obtaining the right kind of structure for \mathbf{W} . We solve the BIBFA model by doing inference directly for (5) and learn the structure of \mathbf{W} by imposing group-wise sparsity for the components (columns of \mathbf{W}), which results in the model automatically converging to a solution that matches (4) (up to an arbitrary re-ordering of the columns). In other words, we do not directly specify the matrices $\mathbf{A}^{(m)}$ and $\mathbf{B}^{(m)}$, but instead learn a single \mathbf{W} matrix. To implement the group-wise sparsity, we divide the variables in \mathbf{x} into two groups corresponding to the two data sets, and construct a prior that encourages sparsity over these groups. For each component \mathbf{w}_k the elements corresponding to one group are either pushed all towards zero, or are all allowed to be active. Recently Jia et al. (2010) introduced a similar sparsity constraint for learning factorized latent spaces; our approach can be seen as a Bayesian realization of the same idea, applied to canonical correlation analysis.

It turns out that the correct form of sparsity can easily be obtained by a simple extension of the ARD prior used for component selection in many Bayesian component models, including the Bayesian CCA described in the previous section. We define the group-wise ARD as

$$p(\mathbf{W}) = \prod_{m=1}^2 \text{ARD}(\mathbf{W}^{(m)} | \alpha_0, \beta_0),$$

with separate ARD prior for each $\mathbf{W}^{(m)}$. Here $\mathbf{W}^{(1)}$ denotes the first D_1 rows of \mathbf{W} and $\mathbf{W}^{(2)}$ refers to the remaining D_2 rows. Similarly to how ARD has earlier been used to choose the number of components, the group-wise ARD makes unnecessary components $\mathbf{w}_k^{(m)}$ inactive for each of the views separately. The components needed for modeling the shared response will have small $\alpha_k^{(m)}$ (that is, large variance) for both views, whereas the view-specific components will have small $\alpha_k^{(m)}$ for the active view and a large one for the inactive one. Finally, the model still selects automatically the total number of components by making both views inactive for unnecessary components. To our knowledge, Virtanen et al. (2011) is the first to consider this simple extension of ARD into multiple groups. Later Virtanen et al. (2012a) and Damianou et al. (2012) discussed the prior in more detail, presenting also extensions to more than two groups.

In practice, the elements of the inactive $\mathbf{w}_k^{(m)}$ will not become pushed exactly to zero, but instead to very small values. For most applications of BIBFA this is not a problem, since we need not identify the components. For example, the demonstration in Section 7.2 that uses CCA for predicting one view from the other automatically ignores the view-specific components even when $\mathbf{w}_k^{(m)}$ is not

exactly zero. Similarly, the explorative data analysis experiment illustrated in Figures 8 and 9 is invariant to the actual components and only relies on the total amount of contribution each feature has on the shared variation. However, in case the individual components are needed, the structure of (4) can be obtained by thresholding small values to zero, for example based on the amount of relative variance explained, and re-ordering the components. The problem is essentially identical to choosing the threshold for PCA models, and hence the techniques suggested for Bayesian PCA apply directly. The ARD prior efficiently pushes the variance of inactive components towards zero, and hence selecting the threshold is often easy in practice.

We apply variational approximation for inference, using the factorized distribution

$$q(\mathbf{W}, \boldsymbol{\tau}_m, \boldsymbol{\alpha}^{(m)}, \mathbf{Y}) = \prod_{n=1}^N q(\mathbf{y}_n) \prod_{m=1}^2 \left(q(\boldsymbol{\tau}_m) q(\boldsymbol{\alpha}^{(m)}) \right) \prod_{d=1}^{D_1+D_2} q(\mathbf{W}_{d,:}).$$

Here $\mathbf{W}_{d,:}$ corresponds to the d th row of \mathbf{W} , a vector spanning over the K different components. The different terms $q(\cdot)$ in the approximation are updated alternately to minimize the Kullback-Leibler divergence $D_{KL}(q, p)$ between $q(\mathbf{W}, \boldsymbol{\tau}_m, \boldsymbol{\alpha}^{(m)}, \mathbf{Y})$ and $p(\mathbf{W}, \boldsymbol{\tau}_m, \boldsymbol{\alpha}^{(m)}, \mathbf{Y}|\mathbf{X})$ to obtain an approximation best matching the true posterior. Equivalently, the task is to maximize the lower bound

$$\mathcal{L}(q) = \log p(\mathbf{X}) - D_{KL}(q, p) = \int q(\mathbf{W}, \boldsymbol{\tau}_m, \boldsymbol{\alpha}^{(m)}, \mathbf{Y}) \log \frac{p(\mathbf{W}, \boldsymbol{\tau}_m, \boldsymbol{\alpha}^{(m)}, \mathbf{Y}, \mathbf{X})}{q(\mathbf{W}, \boldsymbol{\tau}_m, \boldsymbol{\alpha}^{(m)}, \mathbf{Y})}$$

for the marginal likelihood, where the integral is over all of the variables in $q(\mathbf{W}, \boldsymbol{\tau}_m, \boldsymbol{\alpha}^{(m)}, \mathbf{Y})$. Since all priors are conjugate, variational optimization over $q(\cdot)$, constrained to be probability densities, automatically specifies the functional form of all of the terms. Furthermore, we get closed-form updates for each of them, conditional on the choices for all other terms, resulting in straightforward update rules for an EM-style algorithm. The details are given in Appendix A.

As mentioned in Section 3.3, the model is unidentifiable with respect to linear transformations of \mathbf{y} , and only the prior $p(\mathbf{y})$ is influenced by the allocation of the components into shared and view-specific ones. The above procedure for learning the component complexities via group-wise ARD tremendously helps with the latter issue; even though the likelihood part would be equal for a model that splits a true shared component into two separate ones, the variational lower bound will be considerably better for a choice that does not need to replicate the latent variables. In particular, being able to completely drop a component means that the Kullback-Leibler divergence between $q(\mathbf{y}_k)$ and $p(\mathbf{y}_k)$ becomes essentially zero; this advantage would be lost if a shared component was replicated as two view-specific ones.

Interestingly, the variational approximation solves implicitly also the rotational invariance. Even though the likelihood is invariant with respect to right-multiplication of \mathbf{W} with any invertible matrix \mathbf{R} , the variational lower bound is maximal for a specific rotation. Since the \mathbf{R} does not influence the likelihood, it can only improve the lower bound by transforming the approximation into one that best matches the prior distribution. The prior, in turn, assumes independent latent variables, implying that the optimal solution will result in an \mathbf{R} that makes the latent variables of the posterior approximation also maximally independent.⁴ The model is hence identified in the same sense as

4. For the current model, the independence corresponds to orthogonality of the latent variables. We prefer to use the phrase independence as it better fits the notion of assuming independent latent variables and may become more precise in extensions; for other priors and inference algorithms independence need not equal orthogonality.

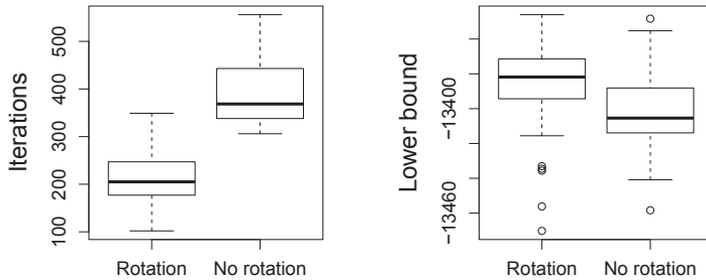


Figure 2: Illustration of the effect of parameter-expanded VB, where the variational lower bound is explicitly optimized with respect to a linear transformation \mathbf{R} to make the updates less correlated. The number of iterations until convergence is reduced clearly when optimizing for the rotation (left), while the lower bound is still on average slightly better (right). However, since \mathbf{R} is optimized numerically the total computation time might still be smaller without the rotation optimization, depending on the data and parameter values (such as K_c). These plots were drawn for the data analyzed in Section 7.1.1 and Figure 9; the boxplots show the results of 50 runs with random initialization. The overall picture is similar for other data sets, but the actual differences vary depending on the complexity of the data. In practice, we recommend trying both with and without the rotation for each data type, and to choose the solution resulting in a better lower bound.

the classical CCA solution is; the latent variables are assumed orthogonal, instead of assuming orthogonal projections (like in PCA).

That property also allows deriving a more efficient algorithm for optimizing the variational approximation, following the idea of parameter-expanded variational Bayes (Qi and Jaakkola, 2007; Luttinen and Ilin, 2010). We introduce explicit parameter $\mathbf{R} \in \mathbb{R}^{K_c \times K_c}$ in the approximation and optimize the lower bound also with respect to it. Transforming the parameters with \mathbf{R} improves the convergence speed dramatically, due to lower correlations between the EM updates for \mathbf{y} and \mathbf{W} , and often also results in slightly better lower bound. Both of the properties are illustrated in Figure 2, and the details for how the rotation can be optimized are given in Appendix B.

In summary, the above model formulation with the associated variational approximation provides a fully Bayesian treatment for the IBFA model. It can also be used for solving the CCA problem with a low-rank assumption for the view-specific noise. The model automatically selects the complexity of the three separate component types through a group-wise ARD prior applied for a joint FA model (that additionally shares the noise variances for all variables within a view), and disambiguates between different rotations by maximizing the orthogonality of the latent variables for improved interpretability and computational efficiency. Open-source implementation of the model, written in the R language, is available in CRAN: <http://cran.r-project.org/package=CCAGFA>.

4.3 On the Choice of Group-sparsity Prior

The above derivation uses group-wise ARD for inferring the component activities. This particular choice is, however, not the only possibility. In fact, any reasonable prior that results in group-wise sparse projection matrix \mathbf{W} could be adopted. Here we briefly discuss possible alternatives, and derive one practical implementation that uses sampling-based inference instead of the variational approximation described above.

BIBFA is essentially a linear model for the concatenation of the two sources, made interpretable by the group-wise sparsity. Hence, a sufficient requirement for a model to implement the BIBFA concept is that it can make $\mathbf{W}_{:,k}$ sparse in the specific sense of favouring solutions where all of the elements corresponding to the first D_1 or the last D_2 dimensions (or both) are driven to zero. This can be achieved in two qualitatively different ways, called weak and strong sparsity by Mohamed et al. (2012). The ARD prior is an example of the former, a continuous sparsity-inducing prior that results in elements that are close to zero but not exactly so. Other priors that induce weak sparsity could also be considered, such as the group-wise extensions of the Laplace and scale-mixture priors Archambeau and Bach (2009) and Guan and Dy (2009) used for sparse PCA, but as we will demonstrate in the empirical experiments, already the ARD prior works well. Hence, we use it as a representative of weak sparsity priors. As general properties, such priors allow continuous inference procedures that are often efficient, but it is not always trivial to separate low-activity components from inactive ones for interpretative purposes. This is because the elements are not made exactly zero even for the components deemed inactive, but instead the values are pushed to very small values.

In our applications in Section 7, we do not need to accurately identify the active components, since already near-zero effects become irrelevant for the predictive measures used. In case more precise determination of the active components is needed, it may be better to switch to strong sparsity, using priors that provide exact zeroes in \mathbf{W} . For this purpose, we here extend the element-wise sparse factor analysis model of Knowles and Ghahramani (2011) for the BIBFA setup. The original model is based on the spike-and-slab prior, where each element of \mathbf{W} is drawn from a two-component prior. One of the components, the spike, is a delta distribution centered at zero, whereas the other, the slab, is a Gaussian distribution. Hence, each element can either become exactly zero or is drawn from a relatively noninformative distribution. To create a BIBFA method based on this idea, we introduce the group-wise spike-and-slab prior with the prior

$$\begin{aligned}
 p(\mathbf{W}, \mathbf{H}, \alpha_b, \boldsymbol{\pi} | \alpha_0, \beta_0) &= p(\boldsymbol{\pi}) p(\mathbf{H} | \boldsymbol{\pi}) \prod_{m=1}^2 p(\mathbf{W}^{(m)} | \mathbf{H}_{m,:}, \boldsymbol{\alpha}^{(m)}) p(\boldsymbol{\alpha}^{(m)} | \alpha_0, \beta_0), \quad (6) \\
 \mathbf{W}_{d,k}^{(m)} | \mathbf{H}_{m,k}, \boldsymbol{\alpha}_k^{(m)} &\sim \mathbf{H}_{m,k} \mathcal{N}(0, (\boldsymbol{\alpha}_k^{(m)})^{-1}) + (1 - \mathbf{H}_{m,k}) \delta_0, \\
 \mathbf{H}_{m,k} | \boldsymbol{\pi}_m &\sim \text{Bernoulli}(\boldsymbol{\pi}_m), \\
 \boldsymbol{\pi}_m &\sim \text{Beta}(1, 1), \\
 \boldsymbol{\alpha}_k^{(m)} | \alpha_0, \beta_0 &\sim \text{Gamma}(\alpha_0, \beta_0),
 \end{aligned}$$

where δ_0 denotes a point-density at zero. That is, the view-specific $\boldsymbol{\pi}_m$ tells the probability for a component to be active, binary $\mathbf{H}_{m,k}$ drawn from the Bernoulli distribution tells whether component k is active in view m , and finally $\mathbf{W}_{:,k}$ is either exactly zero or its elements are all drawn independently from a Gaussian distribution with precision $\boldsymbol{\alpha}_k^{(m)}$ depending on whether $\mathbf{H}_{m,k}$ is zero or one, respectively.

For inference, we use Gibbs sampler by Knowles and Ghahramani (2011) with small modifications. In particular, the elements of \mathbf{H} now depend on D_m features instead of just a single one. This, however, does not make the inference more complicated; the dimensions are independent and hence we get the conditional density by multiplying element-wise terms that still integrate $\mathbf{W}_{d,k}^{(m)}$ out. Another change is motivated by the fact that we only need to estimate a $2 \times K$ matrix \mathbf{H} , instead of a $D \times K$ matrix needed for element-wise sparsity. Since we only have two choices for each component, it does not make sense to use the Indian Buffet Process (IBP) prior for \mathbf{H} ; there cannot be any interesting structure in \mathbf{H} . Hence, we simplify the model to merely draw each entry of \mathbf{H} independently. The details of the resulting sampler are presented in Appendix D.

4.4 Model Summary

We will next briefly summarize the Bayesian CCA model and lay out the two alternative inference strategies. These methods will be empirically demonstrated and compared in the following sections.

4.4.1 BAYESIAN CCA WITH LOW-RANK COVARIANCE, OR BAYESIAN IBFA (BIBFA)

The assumption of low-rank covariance results in the IBFA model of (5). Efficient inference is done in the factor analysis model for $\mathbf{x} = [\mathbf{x}^{(1)}; \mathbf{x}^{(2)}]$ with group-wise sparsity prior for the projection matrix:

$$\begin{aligned} \mathbf{y} &\sim \mathbf{N}(\mathbf{0}, \mathbf{I}), \\ \mathbf{x} &\sim \mathbf{N}(\mathbf{W}\mathbf{y}, \mathbf{\Sigma}), \\ \mathbf{W}^{(m)} &\sim \text{ARD}(\alpha_0, \beta_0). \end{aligned} \tag{7}$$

Here $\mathbf{W}^{(m)}$ denotes the dimensions (rows) of \mathbf{W} corresponding to the m th view, and $\mathbf{\Sigma}$ is a diagonal matrix with D_1 copies of τ_1^{-1} and D_2 copies of τ_2^{-1} on its diagonal. The noise precision parameters are given Gamma priors $\tau_m \sim \text{Gamma}(\alpha_0^\tau, \beta_0^\tau)$. Inference for the model is done according to the updates provided in Appendices A and B.

An alternative inference scheme replaces the above ARD prior with the group-wise spike-and-slab prior of (6) and draws samples from the posterior using Gibbs sampling.

4.4.2 BAYESIAN CCA WITH FULL COVARIANCE (BCCA)

The Bayesian CCA as presented by Wang (2007) and Klami and Kaski (2007) models the view-specific variation with a free covariance parameter. The full model is specified as

$$\begin{aligned} \mathbf{z} &\sim \mathbf{N}(\mathbf{0}, \mathbf{I}), \\ \mathbf{x}^{(m)} &\sim \mathbf{N}(\mathbf{A}^{(m)}\mathbf{z}, \mathbf{\Psi}^{(m)}), \\ \mathbf{A}^{(m)} &\sim \text{ARD}(\alpha_0, \beta_0), \\ \mathbf{\Psi}^{(m)} &\sim \text{IW}(\mathbf{S}_0, \nu_0), \end{aligned} \tag{8}$$

and inference follows the variational updates provided by Wang (2007). When $\mathbf{A} = [\mathbf{A}^{(1)}; \mathbf{A}^{(2)}]$ is drawn from a single ARD prior, the lower bound can analytically be optimized with respect to a rotation \mathbf{R} (see Appendix C), resulting in considerable speedup.

The variational approximations for both BCCA and BIBFA are deterministic and will converge to a local optimum that depends on the initialization. We initialize the model by sampling the latent variables from the prior, and recommend running the algorithm multiple times and choosing the solution with the best variational lower bound. In the experiments we used 10 initializations.

All of the above require pre-specifying the number of components K . However, since the ARD prior (or the spike-and-slab prior for the Gibbs sampler variant) automatically shuts down components that are not needed, the parameter can safely be set large enough; the only drawback of using too large K is in increased computation time. In practice, one can follow a strategy where the model is first run with some reasonable guess for K . In case all components remain active, try again with a larger K .

5. Illustration

In this section we demonstrate the BIBFA model on artificial data, in order to illustrate the factorization into shared and data set-specific components, as well as to show that the inference proceduree converge to the correct solution. Furthermore, we provide empirical experiments demonstrating the importance of making the low-rank assumption for the view-specific noise, in terms of both accuracy and computational speed, by comparing BIBFA (7) with BCCA (8).

The results are illustrated primarily from the point-of-view of the variational inference solution; the variational approximation is easier to visualize and compare with alternative methods. The Gibbs sampler produced virtually identical results for these examples, as demonstrated in Figures 4 and 6.

5.1 Artificial Example

First, we validate the model on artificial data drawn from a model from the same model family, with parameters set up so that it contains all types of components (view-specific and shared components). The latent signals \mathbf{y} were manually constructed to produce components that can be visually matched with the true ones for intuitive assessment of the results. Also the $\alpha^{(m)}$ parameters, controlling the activity of each latent component in both views, were manually specified. The projections \mathbf{W} were then drawn from the prior, and noise with fixed variance was added to the observations.

The left column of Figure 3 illustrates the data generation, showing the four latent components, two of which are shared between the two views. We generated $N = 100$ samples with $D_1 = 50$ and $D_2 = 40$ dimensions, and applied the BIBFA model with $K = 6$ components to show that it learns the correct latent components and automatically discards the excess ones. The results of the variational inference are shown in the middle column of Figure 3; the Gibbs sampler produces virtually indistinguishable results. The learned matrix of α -values (and the corresponding elements in \mathbf{W}) reveals that the model extracted exactly four components, correctly identifying two of them as shared components and two as view-specific ones (one for each data set). The actual latent components also correspond to the ones used for generating the data. The components are presented in the order returned by the model, which is invariant to the order. We also see how the model is invariant to the sign of \mathbf{y} , but that it gives the actual components instead of a linear combination of those, demonstrating that the variational approximation indeed solves the rotational disambiguity that would remain for instance in the maximum likelihood solution.

The BIBFA results are further illustrated in Figure 4. The plot shows the approximate posterior for two of the model parameters, namely the residual noise levels τ_m , demonstrating that the model

has found the true generating parameters. We see that the true parameter values fall nicely within the posterior and both the variational approximation and the Gibbs sampler provide almost the same posterior. We performed similar comparison for all parameters, and the results are also similar, indicating that both variants model this simple data correctly. Furthermore, the results do not here depend notably on the initialization; the model always converges to the right solution. Finally, we also studied that the model is robust with respect to the number of components K . We reran the experiment with multiple values of K upto 30, always getting the same result where only 4 components remain active. This demonstrates that we can safely overestimate the number of components, the only negative side being increased computation time.

5.2 Quantitative Comparison

Next we proceed to quantitatively illustrating that the solution obtained with the BIBFA model is superior to both classical CCA and earlier Bayesian CCA variants not making the low-rank assumption for view-specific noise. We use the same data generation process as above, but explore the two main dimensions of potential applications by varying the number of samples N and the data dimensionality D_m . For the easy case of large N and small D all methods work well since there is enough information for determining the correlations accurately. Classical CCA has an edge in computational efficiency, due to the analytic solution, but also the Bayesian variants are easy to compute since the complexity is only linear in N . Below we will study in more detail the more interesting cases where either D is large or N is small, or both.

We generated data with 4 true correlating components drawn from the prior, drawing N independent samples of $D_1 = D_2$ dimensions, and measure the performance by comparing the average of the four largest correlations ρ_k , normalized by the ground truth correlations. For the Bayesian variants we estimate the correlation between the expectations $\langle \mathbf{Y}_{k,:} | \mathbf{X}^{(1)} \rangle$ and $\langle \mathbf{Y}_{k,:} | \mathbf{X}^{(2)} \rangle$ that are easy to compute by a slight modification of the inference updates. Here $\mathbf{Y}_{k,:}$ contains the k th latent variable for all N samples. Note that $\mathbf{Y}_{k,:} | \mathbf{X}^{(m)}$ follows the prior for the components k that are switched off for the m th view.

We compare the two variants of the Bayesian CCA, denoting by BCCA a model parameterized with full covariance matrices (8) and by BIBFA the fully factorized model (7), with both classical CCA and a regularized CCA (RCCA). For BCCA we used $\nu_0 = D_m$ degrees of freedom and the scale matrix $\mathbf{S}_0 = 0.01 * \mathbf{I}$ to give a reasonably flat prior over the covariances, and for BIBFA we gave a flat prior $\alpha_0^c = \beta_0^c = 10^{-14}$ for the precisions; the other parameters for BCCA and BIBFA were identical. We followed Gonzales et al. (2008) as the reference implementation of a regularized CCA, but replaced the leave-one-out validation for the two regularization parameters with 20-fold cross-validation instead, after verifying that it does not result in statistically significant differences in accuracy compared to the proposed scheme. With leave-one-out validation the computational complexity of RCCA would be quadratic in N , which would have made it severely too slow. To keep the computational load manageable we further devised a two-level grid for choosing the two regularization parameter values: We first try values in a loose two-dimensional grid of 7×7 values and then search for the optimal value in a dynamically created 7×7 grid around the best values.

Figure 5 illustrates the accuracy of the four methods for various scenarios, showing the relative correlations for both training and test data. The main observation is that BIBFA and RCCA are consistently the best methods, with BIBFA having a slightly better accuracy. Classical CCA without regularization breaks down completely for large D/N ratios, as does BCCA with full covariance

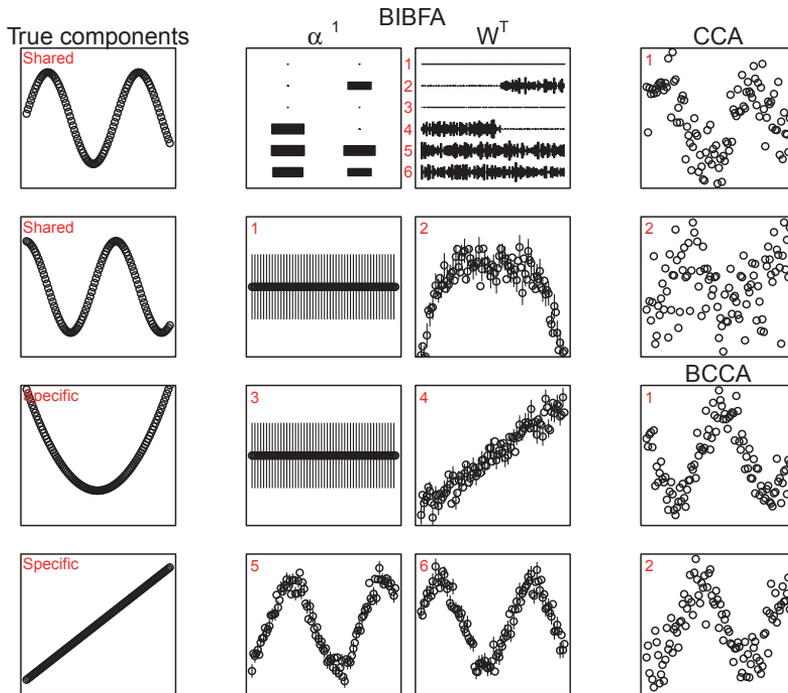


Figure 3: Illustration that Bayesian IBFA (BIBFA) finds the correct underlying latent components. The left column shows four components in the generated data, the first two being shared between the two views and the last two being specific to just one view. BIBFA finds all four components while ignoring the excess ones. The top row of the BIBFA block shows the Hinton-plot (the area of each block indicating the value) of the component variances and the elements of $|\mathbf{W}|$, and the remaining six plots show the estimated latent variables, the small red numbers indicating the link between the latent components and the rows of \mathbf{W}^T . Components 5 and 6 are shared, revealed by non-zero variance for both views, components 2 and 4 are the two view-specific components, and the unnecessary components 1 and 3 have been suppressed to the prior in the sense that their mean and variance match those of the prior. The small lines depict one standard deviation, revealing that the model is more confident on its predictions for the shared components, due to more data ($D_1 + D_2$ features compared to just D_1 or D_2) available for inferring them. The classical CCA (top two plots in the right column), which is only applicable for extracting the shared components, finds much noisier versions of the components, and for slightly higher dimensionality would return only noise. Bayesian CCA with full covariance matrices (bottom two plots in the right column) does better than classical CCA, but does not capture the components as well as BIBFA. For all methods the latent variables have here been estimated for held-out test data.

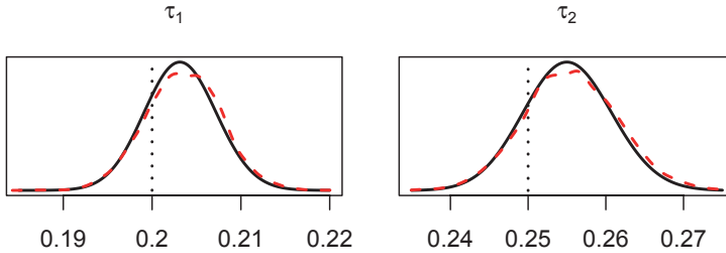


Figure 4: Illustration that Bayesian IBFA (BIBFA) finds also the correct posterior distributions for the model parameters. The plot shows the approximate distributions $q(\tau_m)$ for the variational approximation (solid black line) as well as the posterior obtained with the Gibbs sampling algorithm of the spike-and-slab variant (dashed red line), revealing how both capture the true generating values denoted by dotted black lines. The modes are not exactly at the true value due to the small sample size, but both inference strategies provide the same result.

matrices. Both of these are understandable observations since the methods estimate $D_m \times D_m$ covariance matrices with few or no constraints. To further illustrate the behavior of BIBFA, we plot in Figure 6 the estimated number of shared and view-specific components for both the variational and Gibbs sampling variants. For the purpose of this illustration, we considered a component of the variational inference solution to be active if $\alpha_k^{(m)}$ was below 50 (the true value for active components was 1) and shared if the relative variance of $\alpha_k^{(1)}$ and $\alpha_k^{(2)}$ was below 10, whereas for the Gibbs sampler $\mathbf{H}_{m,k}$ directly reveals the activities. We see that both inference algorithms are conservative in the sense that for very small sample sizes they miss some of the components, using the residual noise term to model the variation that cannot be reliably explained with so limited data. Starting from $N = 64$ (which is still smaller than the data dimensionality for two of the plots) the ranks are estimated correctly.

Another important dimension is the computational time. CCA, RCCA and BCCA all require inverting $D_m \times D_m$ covariance matrices, which results in $O(D_m^3)$ complexity, whereas BIBFA is linear in N and D_m and cubic only with respect to K . The computational times are illustrated in Figure 7, revealing clearly how the lower complexity of BIBFA realizes as faster computation. For very small D the regularized CCA solution is slightly faster than BIBFA, but for large D it becomes impractically slow, even with our faster cross-validation scheme. The overall trend hence is that despite its iterative inference algorithm BIBFA is a much faster solution for high-dimensional CCA problems than regularized CCA solutions that require matrix inversion and cross-validation for tuning the regularization.

The overall summary of these illustrations is that the BIBFA model solves the CCA problem well, even in cases (large dimensionality and/or small sample size) where regular CCA and Bayesian CCA with full covariance matrices do not work at all. Carefully regularized CCA finds the correlations roughly as well as BIBFA, but it is considerably slower for large dimensionalities and lacks interpretable view-specific components, and cannot be extended as easily to directions discussed in the next section. While K was here small, making the gap between BIBFA and the rest of the models

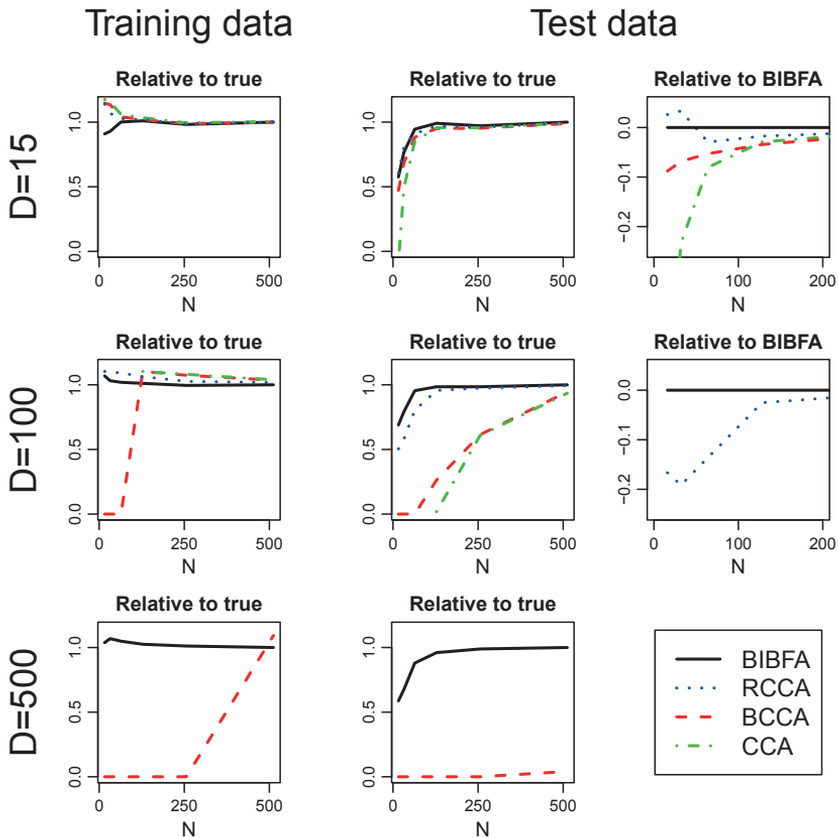


Figure 5: Illustration of the relative performance of the BIBFA model, Bayesian CCA with inverse-Wishart priors (BCCA), regularized CCA (RCCA) and classical CCA for various sample sizes (N ; x-axis) and dimensionalities D (row). RCCA is missing for the last row due to too high computational cost, and CCA could not be computed for $D > N$. The first column shows the sum of the first four correlations (the data has four non-zero correlations) on the training data, normalized so that 1 matches the true value (y-axis). All methods but BIBFA overfit for small N and D , whereas BCCA severely underfits for small N and large D , not finding any reliable correlations. The second column shows the same measure for test data, revealing how BIBFA outperforms the other methods for all cases, except RCCA for very small N and D . The third column shows a zoomed inset for the most interesting region, this time normalized so that the result of BIBFA is used as the baseline, revealing more clearly the advantage BIBFA has over RCCA for all but the smallest samples sizes for $D = 15$.

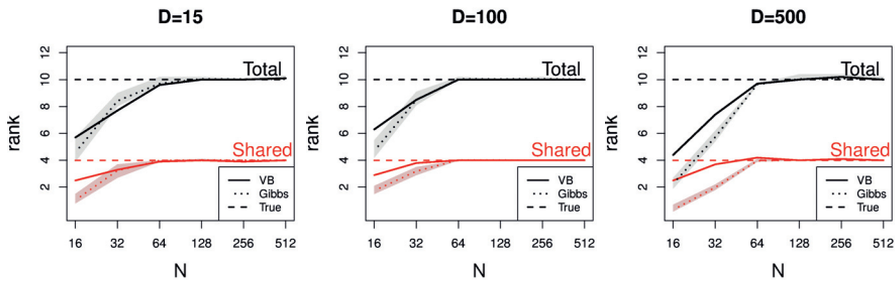


Figure 6: Learning the rank of the data. For reasonable number of samples both the variational approximation (VB) and the spike-and-slab sampler (Gibbs) learn the correct number of both shared components (red lines) and total components (black lines) for all three dimensionalities (subplots). The difference between these two curves corresponds to the sum of residual noise ranks of the two views, which are not shown to avoid cluttering the image. The solid lines correspond to the results of the variational approximation, averaged over 10 random initializations, whereas the dashed line shows the mean of the posterior samples for the Gibbs sampler and the shaded region covers the values between the 5% and 95% quantiles. The two inference algorithms perform roughly as well, and a notable observation is that both methods underestimate the number of components for very small sample sizes, especially for the higher dimensionalities. This is the correct behavior when there is not enough evidence to support the findings.

bigger than in most real applications, the empirical experiments with real-world data in Section 7 reveal that for plenty of practical applications with thousands of dimensions it is sufficient to use values of K in the range of tens. Hence, the computational advantage will hold in real applications as well, making BIBFA a feasible model for scenarios where D would be clearly too large for direct inversion of the covariance matrices.

6. Variants and Extensions

The key advantage of the Bayesian treatment, besides robustness for small sample sizes, is that it enables easy modifications and extensions. In this section we will review a number of extensions presented for the Bayesian CCA model, to provide an overview of the possibilities opened up by the probabilistic treatment of the classical model.

6.1 Modifying the Generative Model

Since the latent variable model is described through a generative process, it is straightforward to change the distributional assumptions in the model to arrive at alternatives designed for specific purposes. Typically these modifications will need to be accompanied by changes in the inference process that are not necessarily trivial, but without the probabilistic formulation extensions like these would be more difficult to keep consistent and justify.

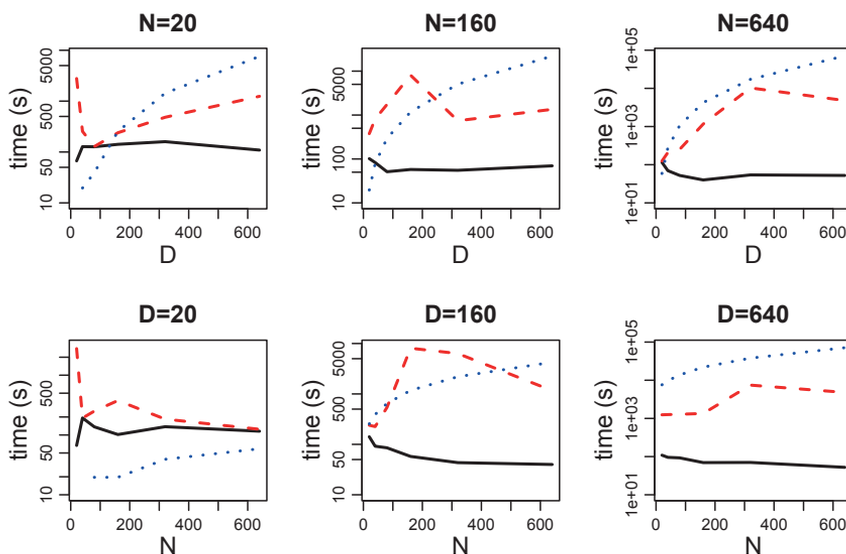


Figure 7: Illustration of the computational time (in seconds) for the BIBFA model (solid black line), Bayesian CCA with inverse-Wishart prior for covariances (BCCA; dashed red line), and regularized CCA (RCCA; dotted blue line). The Bayesian variants assume 10 random restarts, whereas the regularized CCA uses 20-fold cross-validation over a two-stage grid of the two regularization parameters, requiring a total of $20 \cdot (49+49)$ runs. The top row shows how regularized CCA becomes very slow for large dimensionality D , irrespective of the number of samples N . The bottom row shows the linear growth as a function of N for regularized CCA, effectively constant time complexity for BIBFA, and illustrates an interplay of N and D for the BCCA model (it needs more iterations for convergence when D is roughly N). For BIBFA the theoretical complexity is linear in both N and D , but the number of iterations needed for convergence depends on the underlying data in a complex manner and hence the trend is not visible here. Instead, for this data the computational time is almost constant.

The first improvement over the classical CCA brought by the probabilistic interpretation was to replace the Gaussian noise in (3) with the multivariate Student’s t distribution (Archambeau et al., 2006). This makes the model more robust for outlier observations, since observations not fitting the general pattern will be better modeled by the noise term. The maximum likelihood solution provided for the robust CCA by Archambeau et al. (2006) was later extended to the Bayesian formulation with a variational approximation by Viinikanoja et al. (2010).

Klami et al. (2010) extended Bayesian CCA by generalizing from the Gaussian noise assumption to noise with any distribution in the exponential family. Using the natural parameter formulation of exponential family distributions, a generic formulation applicable for any choice was de-

rived. The solution was built on top of the Gibbs-sampling scheme of Klami and Kaski (2007), with considerable technical extensions to cope with the fact that conjugate priors are no longer justified.

Recently, Virtanen et al. (2012b) extended the IBFA-type modelling to count data, introducing a multi-view topic model that generates the observed counts similarly to how IBFA generates continuous data. That is, the model automatically learns topics that are shared between the views as well as topics specific to each view, using a hierarchical Dirichlet process (HDP; Teh et al., 2006) formulation.

Another line of extensions changes the prior for the projections $\mathbf{A}^{(m)}$. Archambeau and Bach (2009) presented a range of sparse models based on various prior distributions. They introduced sparsity priors and associated variational approximations for Bayesian PCA and the full IBFA model, but did not provide empirical experiments with the latter. Another sparse variant was provided by Fujiwara et al. (2009), using an element-wise ARD prior to obtain sparsity, though the method is actually not a proper CCA model since it does not model view-specific variation at all. Rai and Daumé III (2009) built similarly motivated sparse CCA models via a non-parametric formulation where an Indian Buffet Process prior (Ghahramani et al., 2007) is used to switch projection weights on and off. The same non-parametric prior also controls the overall complexity of the model. The inference is based on a combination of Gibbs and more general Metropolis-Hastings steps, but again the model lacks the crucial CCA property of separately modeling view-specific variation.

Leen and Fyfe (2006) and Ek et al. (2008) extended the probabilistic formulation to create Gaussian process latent variable models (GP-LVM) for modeling dependencies between two data sets. They integrate out the projections $\mathbf{A}^{(m)}$, giving a representation that enables replacing the outer product with a kernel matrix, resulting in non-linear extensions. Leen and Fyfe (2006) formulated the model as direct generalization of probabilistic CCA, whereas Ek et al. (2008) modeled explicitly also the view-specific variation. Recently, Damianou et al. (2012) extended the approach to a Bayesian multi-view model that uses group-wise sparsity to identify shared and view-specific latent manifolds for a GP-LVM model, using an ARD prior very similar to the one used by Virtanen et al. (2011) and here for BIBFA.

The conceptual idea of CCA has also been extended beyond linear transformation and continuous latent variables. As a practical example, multinomial latent variables provide clustering models. Both Klami and Kaski (2006) and Rogers et al. (2010) presented clustering models that capture the dependencies between two views with the cluster structure while modeling view-specific variation with another set of clusters. Recently, Rey and Roth (2012) followed the same idea, modeling arbitrary view-specific structure within the clusters with copulas. The Bayesian CCA approach has also been extended beyond vectorial data representations; van der Linde (2011) provided a Bayesian CCA model for functional data, building on the variational approximation.

Haghighi et al. (2008) and Tripathi et al. (2011) extended probabilistic CCA beyond the underlying setup of co-occurring data samples. They complement regular CCA learning by a module that infers the relationship between the samples in the two views, by finding close neighbors in the CCA subspace. This enables both computing CCA for setups where the pairing of (some of) the samples is not known but also applications where learning the pairing is the primary task. Recently, Klami (2012) presented a variational Bayesian solution to the same problem, extending BIBFA to include a permutation parameter re-ordering the samples.

Some related methods not described in the terminology of Bayesian CCA are also worth mentioning, due to the close relationship between both the task and the models. Singh and Gordon

(2008) introduced collective matrix factorization (CMF), where the task is to learn simultaneous matrix factorizations of the form $\mathbf{X}^{(m)} = \mathbf{V}^{(m)}\mathbf{Z}$ for multiple (in their application three) views, which is equivalent to the Bayesian CCA formulation. However, the exact definition of the noise additive to the factorization is crucial; BIBFA includes explicit components for modeling view-specific variation (or they are modeled with full covariance matrices as in the earlier Bayesian CCA solutions). CMF, in turn, assumes that all variation is shared, by factorizing the noise over the dimensions. Hence, CMF is more closely related to learning PCA for the concatenated data sources, and is incapable of separating the shared variation from the view-specific one. Recently, Agarwal et al. (2011) extended CMFs to localized factor model (LMF) that allows separate latent variables $\mathbf{z}^{(m)}$ for the views and models them as a linear combination of global latent profiles \mathbf{u} . This extended model is capable of implementing the CCA idea by selectively using only some of the global latent profiles for each of the views, though it is not explicitly encouraged and the authors do not discuss the connection. The residual component analysis by Kalaitzis and Lawrence (2012) is also closely related; it is a framework that includes probabilistic CCA as a special case. They assume a model where the data is already partly explained by some components and the rest is explained by a set of factors. By iteratively treating the view-specific and shared components as the explanatory factors they can learn the maximum likelihood solution of IBFA (and hence CCA) through eigen-decompositions, but their general formulation also applies to other data analysis scenarios.

Finally, a number of papers have discussed extensions of probabilistic CCA into more than two views. Already Archambeau and Bach (2009) mention that the generative model directly generalizes to more than two views, but they do not show that their inference solution would provide meaningful results for multiple views. Recently, Virtanen et al. (2012a) presented the first practical multi-view generalization of Bayesian CCA, coining the method group factor analysis (GFA), and Damianou et al. (2012) described a GP-LVM -based solution for multiple views. We do not discuss the multi-view generalizations further in this article, since the extended model cannot be directly interpreted as CCA; the concept of correlation does not directly generalize to multiple views.

6.2 Building Block in Hierarchical Models

The generative formulation of probabilistic models extends naturally to complex hierarchical representations. The Bayesian CCA model itself is already a hierarchical model, but can also be used as a building block in more complex hierarchical models. In essence, most Bayesian models operating on individual data sets can be generalized to work for paired data by incorporating a CCA-type latent variable formulation as a part of the model.

The first practical examples considered the simplest hierarchical constructs. Klami and Kaski (2007) introduced an infinite mixture of Bayesian CCA models, accompanied with a Gibbs sampling scheme. Later Viinikanoja et al. (2010) provided a variational approximation for mixtures of robust CCA models, resulting in a computationally more efficient algorithm for the same problem. These kinds of mixture models can be thought of as locally linear models that partition the data space into clusters and fit a separate CCA model within each. The clustering step is, however, integrated in the solution and is also influenced by the CCA models themselves.

Recently, some authors have used Bayesian CCA as an integral part in more complex hierarchical models. Huopaniemi et al. (2009) integrate a dimensionality reduction step into Bayesian CCA by clustering the original features and applying Bayesian CCA to the latent variables that aggregate features within a cluster, to make BCCA feasible for high-dimensional metabolomics data with

very limited sample size. Huopaniemi et al. (2010) addresses the same application domain, this time combining BCCA with multivariate analysis of variance (ANOVA). Nakano et al. (2011), in turn, created a hierarchical topic trajectory model (HTTM) by using CCA as the observation model in a hidden Markov model (HMM).

7. Applications

In this section we will discuss some of the applications of CCA, covering both general application fields and concrete problem setups. Some of the examples are from fields where the probabilistic variants have not been widely applied yet, but where the need for CCA-type modeling is apparent and the properties of the data suit well the strengths of the Bayesian approach.

We have divided the applications into two broad categories. The first category considers CCA as a tool for exploratory data analysis, seeking to evaluate the amount of correlation or dependency between various information sources or to illustrate which of the dimensions correlate with the other view. The other category uses CCA as a predictive model, building on the observation that CCA is a good predictor for multiple outputs, correctly separating the information useful for prediction from the noise.

7.1 Data Analysis

One of the key strengths of the Bayesian approach is that it enables justified analysis of small samples, providing estimates of the reliability of the results. For the application fields with plenty of data also the classical and kernel-based CCA solutions work well, as has been demonstrated for example in analysis of relationships between text documents and image content (Vinokourov et al., 2003). Hence, we focus here on applications where the amount of data is typically limited.

Life sciences are a prototypical example of a field with limited sample sizes. In many analysis scenarios the samples correspond to individuals, and high cost of measurements prevents collecting large data sets. There are also several application scenarios where the number of samples is restricted for biological reasons, for example when studying rare diseases or effects specific to an individual instead of a population.

CCA has received a lot of attention in analysis of omics data, including genomics measured with microarrays as well as proteomics and metabolomics measured by mass spectrometry. Huopaniemi et al. (2009, 2010) applied extensions of Bayesian CCA to find correlations between concentrations of biomolecules in different tissues and species to build “translational” models. In their studies, the samples correspond to individual mice and humans with a sample size in the order of tens, whereas the features correspond to concentrations of hundreds of lipids. Similar setups but still much more extreme ratios of D_m/N are encountered frequently in microarray analysis, where the dimensions correspond to tens of thousands of genes. Due to the limitations of the earlier models the Bayesian solutions have not yet been used with full strength in such applications.

Another typical application scenario is in brain activity analysis, where the samples typically correspond to time-slices of an experiment and the features span the brain activity measured either through BOLD (blood-oxygen-level-dependent) signal activity in small brain volumes called voxels (in functional magnetic resonance imaging fMRI) or through magnetometers on the scalp (magnetoencephalography MEG). Fujiwara et al. (2009) used sparse Bayesian CCA to predict the visual stimuli from fMRI data, and Koskinen et al. (2012) inferred the identity of short speech segments using mixture of robust Bayesian CCAs applied to MEG. Several authors have also applied classical

CCA or its multiset extensions for fMRI data; Ylipaavalniemi et al. (2009) studied the relationships between brain activity and naturalistic stimuli features, Deleus and Hulle (2011) explored functional connectivity between multiple brain areas, and Rustandi et al. (2009) integrated fMRI data of multiple subjects. Similar tasks could also be solved with the Bayesian variants, in particular with the BIBFA model.

7.1.1 ILLUSTRATION

To demonstrate the use of CCA in an exploratory data analysis scenario, we apply it to the problem of cancer gene prioritization based on co-occurring gene expression and copy number data (Lahti et al., 2012). DNA alterations frequent in cancers, measured by the copy number data, are known to induce changes in the expression levels, and hence cancer-associated genes can be mined by searching for such interactions.

One approach is to proceed over the whole genome in a gene-by-gene fashion, searching for correlations between the gene expression and copy number modification. Lahti et al. (2009) adapted CCA for this task, using it to estimate the amount of correlation inside short continuous windows of the genome. A collection of cancer and control patients are treated as samples, and the features are the genes within a window. For each window they computed so-called similarity-constrained CCA and labeled the genes within that window with the resulting correlation. That is, a gene is assumed to be cancer-related if CCA finds strong correlation within a small chromosomal window around it. This approach is one of the leading solutions for finding cancer-associated genes from integrated copy number and gene expression data, as shown in the recent comparison by Lahti et al. (2012).

For computing the association scores for N genes the above process requires running N separate CCA models, one for each neighborhood. Within each window, the CCA is ran for N' samples (the patients, on the order of 30-50 for typical data sets) and $D_x = D_y$ features (the genes within the window). The authors used window sizes of roughly 10-20, to guarantee that the number of samples exceeds the number of features, satisfying the usual requirement for CCA-style models.

The BIBFA model (7) has been specifically designed to tackle the issue of high dimensionality, and hence it allows a much more direct approach. Instead of measuring the amount of correlation for several small windows, we simply run CCA considering the patients as samples and all of the genes in the whole genome as features. Direct inspection of the weights in the shared components then reveals the cancer-associated genes; a high weight implies an association between the copy number and gene expression, relating the gene to the cancer under study.

We applied BIBFA, using the ARD prior and variational inference, on the two publicly available data sets used in the recent comparison of various integrative cancer-gene mining tools by Lahti et al. (2012), the Pollack and Hyman data sets. We repeated their experimental setup to obtain results directly comparable with their study, and measured the performance by the same measure, the area under curve (AUC) for retrieving known cancer genes (37 out of 4247 genes in Pollack, and 47 out of 7363 genes in Hyman). We ran the BIBFA model for K_c between 5 and 60 components and chose the model with the best variational lower bound, resulting in $K_c = 15$ for Hyman and $K_c = 40$ for Pollack. The full results of BIBFA and the comparison methods are reported in Figure 8, revealing that our method outperforms all of the alternatives for both data sets. The results would be similar for a wide range of values of K_c ; for all values we beat the alternative methods. We also applied the Gibbs-sampler variant, which produced very similar results.

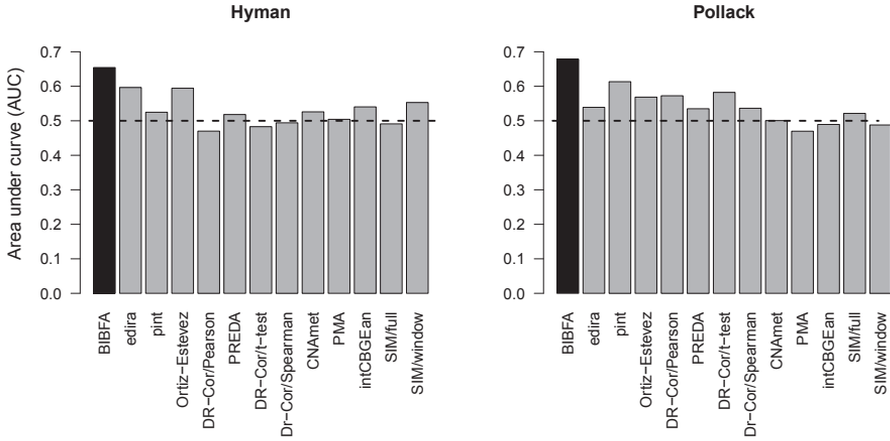


Figure 8: Comparison of AUC scores for the various methods in detecting cancer-related genes in genome-wide data. The BIBFA model ranks the genes based on the weight of that gene in both $\mathbf{W}^{(1)}$ and $\mathbf{W}^{(2)}$, and finds the cancer genes with better accuracy than any of the methods studied in the recent comparison by Lahti et al. (2012). The accuracies for all the other methods are taken from the publicly available results provided by the authors; see their article for the names and details of the methods. Of these methods, pint is the most closely related to ours. It screens the genome by computing CCA for narrow windows and ranks the genes according to the strength of dependency found within each window.

For ranking the genes we used the measure $s_g = \sum_{k=1}^{K_c} |\langle \mathbf{W}_{g,k}^{(1)} \rangle \langle \mathbf{W}_{g,k}^{(2)} \rangle|$. That is, for each component we multiply the expected projection vectors corresponding to gene expression and copy number change, to emphasize effects seen in both views. We then simply sum the absolute values of these quantities over all components, to reach the measure s_g for each gene g . Note that the view-specific components have no effect on the score, since either $\langle \mathbf{W}_{:,k}^{(1)} \rangle$ or $\langle \mathbf{W}_{:,k}^{(2)} \rangle$ will be zero for all genes. To further illustrate the approach, Figure 9 plots the quantity over one chromosome in the Pollack data (chromosome 17, the one most strongly associated with the breast cancer studied in that data) and compares the result with the activity profile provided by the similarity-constrained CCA model (also called pint, after the name of the public software implementation) of Lahti et al. (2009).

7.2 Multi-label Prediction

Another interesting application for CCA is in prediction. Even though the model is symmetric with respect to $\mathbf{x}^{(1)}$ and $\mathbf{x}^{(2)}$, it is surprisingly efficient as a predictive model. In a sense, the model can be seen as a combination of purely unsupervised and supervised learning; both of the views can be considered as supervising the other view, yet the model is (here) defined as a generative description of the whole data collection.

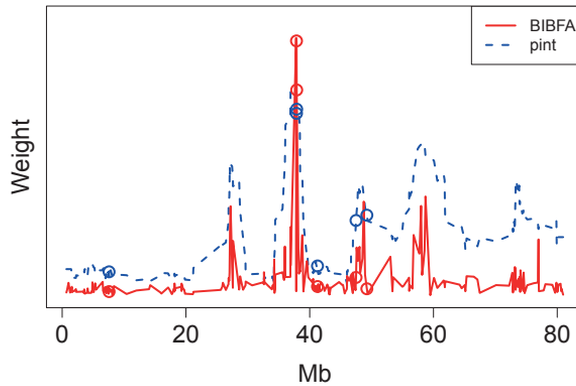


Figure 9: Illustration of the weights s_g (y-axis) learned for the 272 genes in Chromosome 17, depicted in the chromosomal order (x-axis), with BIBFA (red) and the window-based comparison method of pint (blue). Since the gene detection is based on the ranks of the genes, the two weight vectors have here been re-scaled into comparable scales for visual inspection. Both methods reveal a similar overall trend over the chromosome, capturing especially the known cancer-related region around 40Mb, but BIBFA does it in much more direct fashion: The BIBFA profile shown here is the outcome of a single run, whereas 272 applications of similarity-constrained CCA were needed to create the profile for pint. Note that BIBFA gets this result without making any assumptions about mutations in a region causing gene expression changes in the same region, showing the power of the algorithm but resulting in some noise in the result. Pint, in contrast, analyzes local neighborhoods and hence necessarily results in similar values for nearby chromosomal regions. The six known cancer genes are marked with circles. Neither method captures all of them, but BIBFA finds the top two cancer genes at higher ranks. On the other hand, BIBFA seems to here miss some cancer-related genes around the 47Mb region; there is no guarantee that the method would always be more accurate, but the numerical comparisons in Figure 8 show that it on average outperforms pint.

The CCA model is closely related to multiple regression. Breiman and Friedman (1997) showed that CCA is particularly efficient in multiple regression when the response variables are correlated. In recent years, for example Ji et al. (2008), Rai and Daumé III (2009), and Sun et al. (2011) have all demonstrated good predictive performance for CCA-type models in multi-output prediction or multiple regression tasks.

The Bayesian formulation through the BIBFA model helps in understanding why CCA works so well for multiple regression tasks. The predictive distribution of interest is $p(\mathbf{x}^{(1)}|\mathbf{x}^{(2)})$, which cannot be computed in close form, but for which it is easy to obtain expectations from the variational

approximation. The mean prediction is given by

$$\langle \mathbf{x}^{(1)} | \mathbf{x}^{(2)} \rangle = \langle \mathbf{A}^{(1)} \mathbf{z} \rangle_{q(\mathbf{A}^{(1)})q(\mathbf{z}|\mathbf{x}^{(2)})} = \langle \mathbf{A}^{(1)} \rangle \langle \sigma_2^{-2} \rangle \Sigma \langle \mathbf{A}^{(2)T} \rangle \mathbf{x}^{(2)},$$

$$\text{where } \Sigma = \left(\mathbf{I} + \langle \sigma_2^{-2} \rangle \langle \mathbf{A}^{(2)T} \mathbf{A}^{(2)} \rangle \right)^{-1}.$$

The notable observation is that neither $\mathbf{B}^{(1)}$ nor $\mathbf{B}^{(2)}$ (or the corresponding latent variables $\mathbf{z}^{(1)}$ or $\mathbf{z}^{(2)}$) appear in the formula. This implies the model correctly neglects variation specific to $\mathbf{X}^{(2)}$ when making the prediction, and additionally does not attempt to predict the variation in $\mathbf{X}^{(1)}$ that cannot be predicted. In other words, the model squeezes the prediction through the shared latent variables \mathbf{z} , the lower-dimensional representation capturing all the information flow from one data set to another. Note that in practice the prediction can be directly written in terms of \mathbf{W} and \mathbf{y} , without needing to explicitly extract $\mathbf{A}^{(m)}$ and $\mathbf{B}^{(m)}$, since the zeroes induced in \mathbf{W} due to the group-wise sparsity will cancel the unnecessary components out automatically.

Similar observation holds also for the Gibbs sampler variant using the spike-and-slab prior (6); again the mean prediction only depends on the shared components. However, making predictions with that model is considerably more time-consuming, since the predictions need to be averaged over the posterior samples. When making predictions for new samples, we need to store all of the posterior samples and then run the sampler again for each of those estimate the latent variables and the predicted $\mathbf{X}^{(1)}$. Due to this extra computational overhead for the sampler, we will next demonstrate the BIBFA in multi-label prediction tasks using only the variational inference variant.

7.2.1 ILLUSTRATION

To measure the multi-label classification accuracy of the BIBFA model we applied it on 10 benchmark data sets from the Mulan library (Tsoumakas et al., 2010), using the split to train and test samples given in the library. Each of these data sets includes several binary labels that are not mutually exclusive, and we encode them into $\mathbf{X}^{(1)}$ so that each column represents one label. Then the task becomes predicting $\mathbf{X}^{(1)}$ from $\mathbf{X}^{(2)}$. Since the labels are discrete, we feed the predicted values through a simple threshold filter, with the threshold for each class chosen to maximize the accuracy on the training data. The Bayesian formulation would enable integrating also more advanced ways of handling with the binary data (Klami et al., 2010), but here the primary purpose is to demonstrate the application of the basic principle instead of developing a fully-fledged multi-label prediction model.

We compare the BIBFA model (7) with both classical CCA and the standard Bayesian CCA with full covariance matrices (8), using the variational approximation of Wang (2007). For BCCA we set the maximal number of components to $K = \min(D_1, D_2, 50)$, and for classical CCA we chose the number of components by 10-fold cross-validation within the training set. For BIBFA we set K_c to the minimum of 100 and the number of components extracted by Bayesian PCA ran on the concatenation of the two data views. Overall, these choices constitute a fair way of selecting the model complexity for each of the methods. For BIBFA and BCCA we started the optimization from 10 different random initializations and chose the solution that resulted in the best lower bound for the training data, whereas CCA is a deterministic algorithm and always provides the global optimum. The regularized CCA model studied earlier in Section 5.2 was left out due to its immensely high computational cost for most of the data sets, but preliminary studies showed that it did not outperform even classical CCA on the ones with sufficiently few dimensions. Besides showing the

Data Set	D_1	D_2	N_{train}	BIBFA	CCA	BCCA	RML	RAKEL	MLKNN
emotions	6	72	391	0.223	0.232	0.329	0.225	0.223	0.209
scene	6	294	1211	0.105	0.332	0.162	0.127	0.115	0.0953
yeast	14	103	1500	0.202	0.205	0.211	-	0.233	0.198
genbase	27	1186	463	9.3e-4	-	9.3e-4	-	0.0011	0.0052
medical	45	1449	333	0.0124	-	0.0276	-	0.0113	0.0188
enron	53	1001	1123	0.0465	-	0.0607	-	0.0509	0.0514
mediamill	101	120	30933	0.0309	0.161	0.0305	-	0.0335	0.0314
bibtex	159	1836	4880	0.0131	0.0138	-	-	0.0144	0.0140
Corel5-k	374	499	4500	0.0094	0.0099	0.0098	-	0.0096	0.0093
delicious	983	500	12920	0.0182	0.0183	-	-	0.0185	0.0183

Table 1: Prediction errors (Hamming loss) for 10 benchmark data sets sorted by the increasing number of labels D_1 . For each data set the error for the best method has been boldfaced. The proposed Bayesian inter-battery factor analysis model (BIBFA) outperforms the classical CCA and Bayesian CCA with full covariance matrices (BCCA) for almost all data sets. For cases with a large number of labels BIBFA outperforms also designated multi-label prediction models RAKEL and MLKNN, showing that modeling the dependencies between the labels helps more when the number of labels is high. The figures for the reverse multi-label prediction model (RML) were taken from Petterson and Caetano (2010), N_{train} is the number of training samples, and D_2 is the input dimensionality. The values missing for BCCA were excluded due to too long computation time (more than 5 hours per run), and classical CCA was not ran for data sets where the dimensionality of either view is higher than the number of samples (for the enron data, $D_2 > N$ for the cross-validation runs needed for setting the threshold).

relationships between the various CCA-based methods, we also compared BIBFA with three multi-label prediction models with publicly available code or results, RAKEL (Tsoumakas and Vlahavas, 2007) and MLKNN (Zhang and Zhou, 2007) as implemented in the Mulan library, and reverse multi-label prediction model by Petterson and Caetano (2010). For measuring the performance we use the Hamming distance between the predictions and the true labels, penalizing equally much for both false negatives and false positives.

Table 1 shows that BIBFA is the best of the CCA variants on all but one of the data sets. Furthermore, Table 2 demonstrates how it is again considerably faster than BCCA model, even though we analytically optimized for the rotation \mathbf{R} in the Bayesian CCA model. BIBFA also outperforms the other comparison models systematically for the cases with very large number of labels (D_1), with the exception of the *Corel5-k* data set. This demonstrates that CCA-type models are particularly useful for multi-label prediction tasks with an extreme number of labels, most likely because more information can then be extracted from the dependencies between the labels. The improvements presented in this paper are needed especially for that domain, since BIBFA is most useful for analysis of high-dimensional data.

For cases with a low number of labels (below 20 for the first three data sets), MLKNN outperforms IBFA. This is understandable as it is a model specifically designed for multi-label prediction and it explicitly maximizes the prediction accuracy, in contrast to BIBFA that is a generative model

Method	emotions	scene	yeast	genbase	medical
BIBFA	3	13	2	11	14
BCCA	1	8	2	46	79
Method	enron	mediamill	bibtex	Corel5-k	delicious
BIBFA	13	33	14	4	26
BCCA	289	95	>300	104	>300

Table 2: Average computation times for BIBFA and BCCA (in minutes) until convergence (relative change of the lower bound below 10^{-6}). For small dimensionalities the computational demands of the methods are comparable, but for high dimensionality the BCCA model becomes infeasible.

for both data sources. Nevertheless, BIBFA outperforms the two other comparison methods even for data sets with few labels.

8. Discussion

In this paper we have reviewed the works on probabilistic and Bayesian canonical correlation analysis, with particular focus on the extensions made possible by the probabilistic interpretation. While the solutions presented here are linear, as opposed to the possibly nonlinear kernel-based CCA models (Haroon et al., 2004), the extensions and the ease of including CCA as a sub-model in larger hierarchical models clearly showcase the importance of probabilistic treatment of the problem. Works by Fujiwara et al. (2009) and Huopaniemi et al. (2010) have recently demonstrated how the Bayesian solution has enabled analysis of life science data sets with very low sample sizes.

Besides reviewing the earlier work, we introduced a novel solution that that results in considerably more efficient inference for the Bayesian CCA model, especially for high-dimensional data. The key is to make a low-rank assumption for the noise specific to each data set, which results in re-formulation of CCA as a more complex latent variable model called inter-battery factor analysis (IBFA) in the statistics literature (Tucker, 1958). While the extended model seems more complex due to having more unknown latent variables, it has the advantage of diagonal noise that reduces the risk of overfitting and simplifies the computation to the extent that it is actually much more efficient to learn the Bayesian IBFA (BIBFA) model than it is to directly learn the Bayesian CCA solution.

The computational difficulties stemming from introducing the extra latent variables are solved by clever usage of group-wise sparsity assumption. Instead of explicitly instantiating several latent variables, we re-cast the IBFA model as a straightforward joint factor analysis model with a specific prior driving the component group-wise sparse, showing how the resulting model is equivalent to IBFA. For inference we proposed two alternative solutions with alternative sparsity-inducing priors. One uses parameter expanded variational approximation with automatic relevance determination (ARD) prior, whereas the other uses Gibbs sampling with spike-and-slab prior. Both variants work well in practice, and usually seem to produce very similar results.

Given the efficient inference solution we believe the necessary tools for real-world application of Bayesian CCA are now available, making the approach feasible for scenarios that were previ-

ously not possible to solve. In this work we demonstrated how earlier serial computation of several low-dimensional CCA models could be replaced by single use of BIBFA for the original high-dimensional data in the task of extracting cancer-related genes. Similar conceptual shifts should be possible for other domains as well, in particular in life sciences where the sample sizes are typically in the order of tens while the dimensionality may be thousands or even larger.

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Appendix A. Variational Updates for the BIBFA Model

The joint likelihood of the BIBFA model is

$$p(\mathbf{X}, \mathbf{W}^{(1)}, \mathbf{W}^{(2)}, \mathbf{Y}, \boldsymbol{\alpha}^{(1)}, \boldsymbol{\alpha}^{(2)}, \boldsymbol{\tau}^{(1)}, \boldsymbol{\tau}^{(2)}) = \prod_{m=1}^2 p(\mathbf{W}^{(m)} | \boldsymbol{\alpha}^{(m)}) p(\boldsymbol{\alpha}^{(m)}) p(\boldsymbol{\tau}_m) \prod_{n=1}^N p(\mathbf{y}_n) p(\mathbf{x}_n^{(m)} | \mathbf{W}^{(m)}, \mathbf{y}_n, \boldsymbol{\tau}_m),$$

where

$$\begin{aligned} \mathbf{x}_n^{(m)} &\sim \mathcal{N}(\mathbf{W}^{(m)} \mathbf{y}_n, \boldsymbol{\tau}_m^{-1} \mathbf{I}), \\ \mathbf{W}_{:,k}^{(m)} &\sim \mathcal{N}(\mathbf{0}, (\boldsymbol{\alpha}_k^{(m)})^{-1} \mathbf{I}), \\ \boldsymbol{\alpha}_k^{(m)} &\sim \text{Gamma}(\alpha_0, \beta_0), \\ \boldsymbol{\tau}_m &\sim \text{Gamma}(\alpha_0, \beta_0), \\ \mathbf{y}_n &\sim \mathcal{N}(\mathbf{0}, \mathbf{I}). \end{aligned}$$

We use mean field variational approximation to approximate the posterior, with the factorization

$$Q(\Theta) = q(\mathbf{Y}) \prod_{m=1}^2 q(\mathbf{W}^{(m)}) q(\boldsymbol{\alpha}^{(m)}) q(\boldsymbol{\tau}_m),$$

where Θ denotes all of the parameters and latent variables. For the latent variables we further assume column-wise independence (that is, the latent variables of observations are independent) and for projections row-wise independence (that is, each component is independent). The distributions are found by maximizing the lower bound of the marginal log-likelihood

$$\mathcal{L}(Q) = \int q(\Theta) \log \frac{p(\mathbf{X}, \Theta)}{q(\Theta)} d\Theta \leq \log p(\mathbf{X}),$$

and free-form optimization of the factored variables in $Q(\Theta)$ results in analytically tractable distributions due to conjugate priors. The forms of these distributions and the matrix-form updates

rules for efficient computations are given below, after introducing the necessary notation for the expectations.

For normal distribution $\mathbf{x} \sim \mathcal{N}(\boldsymbol{\mu}, \boldsymbol{\Sigma})$ the first moment is given as $\langle \mathbf{x} \rangle = \boldsymbol{\mu}$ and the second moment as $\langle \mathbf{x}\mathbf{x}^T \rangle = \langle \mathbf{x} \rangle \langle \mathbf{x} \rangle^T + \boldsymbol{\Sigma}$. We also introduce the notation $\langle \mathbf{X} \rangle = [\boldsymbol{\mu}_1, \dots, \boldsymbol{\mu}_N]$ and $\langle \mathbf{X}\mathbf{X}^T \rangle = \langle \mathbf{X} \rangle \langle \mathbf{X} \rangle^T + N\boldsymbol{\Sigma}$ to indicate the moments of N independent draws of \mathbf{x} with different $\boldsymbol{\mu}_n$ but the same covariance matrix. For gamma distribution $\alpha \sim \text{Gamma}(\alpha_0, \beta_0)$ the expectation is given by $\langle \alpha \rangle = \frac{\alpha_0}{\beta_0}$.

For the projections we get row-wise independent factorial approximation

$$\begin{aligned} q(\mathbf{W}^{(m)}) &= \prod_{d=1}^{D_m} \mathcal{N}(\mathbf{W}_{d,:}^{(m)} | \boldsymbol{\mu}_{\mathbf{W}_{d,:}^{(m)}}, \boldsymbol{\Sigma}_{\mathbf{W}^{(m)}}), \\ \boldsymbol{\Sigma}_{\mathbf{W}^{(m)}} &= (\langle \boldsymbol{\alpha}^{(m)} \rangle^{-1} + \langle \boldsymbol{\tau}^{(m)} \rangle \langle \mathbf{Y}\mathbf{Y}^T \rangle)^{-1}, \\ \langle \mathbf{W}^{(m)} \rangle &= \mathbf{X}^{(m)} \langle \mathbf{Y} \rangle^T \boldsymbol{\Sigma}_{\mathbf{W}^{(m)}} \langle \boldsymbol{\tau}^{(m)} \rangle, \end{aligned}$$

where $\langle \mathbf{W}^{(m)} \rangle^T = [\boldsymbol{\mu}_{\mathbf{W}_{1,:}^{(m)}}, \dots, \boldsymbol{\mu}_{\mathbf{W}_{D_m,:}^{(m)}}]$, and for the latent variables the update is given by

$$\begin{aligned} q(\mathbf{Y}) &= \prod_{n=1}^N \mathcal{N}(\mathbf{y}_n | \boldsymbol{\mu}_{\mathbf{y}_n}, \boldsymbol{\Sigma}_{\mathbf{Y}}), \\ \boldsymbol{\Sigma}_{\mathbf{Y}} &= (\mathbf{I} + \sum_{m=1}^2 \langle \boldsymbol{\tau}^{(m)} \rangle \langle \mathbf{W}^{(m)T} \mathbf{W}^{(m)} \rangle)^{-1}, \\ \langle \mathbf{Y} \rangle &= \sum_{m=1}^2 \langle \boldsymbol{\tau}^{(m)} \rangle \boldsymbol{\Sigma}_{\mathbf{Y}} \langle \mathbf{W}^{(m)} \rangle^T \mathbf{X}^{(m)}. \end{aligned}$$

For the ARD parameters the updates are

$$\begin{aligned} q(\boldsymbol{\alpha}^{(m)}) &= \prod_{k=1}^K \text{Gamma}(\alpha_k^{(m)} | a_{\alpha_k}^{(m)}, b_{\alpha_k}^{(m)}), \\ a_{\alpha_k}^{(m)} &= \alpha_0 + D_m/2, \\ b_{\alpha_k}^{(m)} &= \beta_0 + \langle \mathbf{W}^{(m)T} \mathbf{W}^{(m)} \rangle_{k,k}/2, \end{aligned}$$

where $\langle \alpha_k^{(m)} \rangle = \frac{a_{\alpha_k}^{(m)}}{b_{\alpha_k}^{(m)}}$. Finally, for the noise precision parameters we have

$$\begin{aligned} q(\boldsymbol{\tau}_m) &= \text{Gamma}(a_{\boldsymbol{\tau}}^{(m)}, b_{\boldsymbol{\tau}}^{(m)}), \\ a_{\boldsymbol{\tau}}^{(m)} &= \alpha_0 + ND_m/2, \\ b_{\boldsymbol{\tau}}^{(m)} &= \beta_0 + (\sum_{dn} \mathbf{X}_{dn}^{(m)2} + \text{Trace}[\langle \mathbf{W}^{(m)T} \mathbf{W}^{(m)} \rangle \langle \mathbf{Y}\mathbf{Y}^T \rangle] \\ &\quad - 2\text{Trace}[\langle \mathbf{W}^{(m)} \rangle \langle \mathbf{Y} \rangle \mathbf{X}^{(m)T}]) / 2. \end{aligned}$$

The algorithm proceeds by updating the parameters of the above factors sequentially until convergence. In the experiments included in this paper we determined convergence as relative change of $\mathcal{L}(\mathcal{Q})$ falling below 10^{-6} .

Appendix B. Parameter-expanded Variational Bayes for BIBFA

To make the correlations between the mean-field updates of BIBFA smaller, we can optimize $\mathcal{L}(Q)$ also with respect to a likelihood-invariant rotation \mathbf{R}

$$\mathbf{x}^{(m)} = \mathbf{W}^{(m)}\mathbf{y} = \mathbf{W}^{(m)}\mathbf{R}\mathbf{R}^{-1}\mathbf{y} = \mathbf{W}^{(m)*}\mathbf{y}^*,$$

where the asterisk is used to denote the transformed variables. We perform this optimization after updating \mathbf{W} and \mathbf{Y} according to the equations in Appendix A, but before learning the ARD parameters.

Given the new \mathbf{R} , the transformed factorial distributions are given as

$$\begin{aligned} q^*(\mathbf{W}^{(m)}) &= \prod_{d=1}^{D_m} \mathcal{N}(\mathbf{W}_{d,:}^{(m)} | \mathbf{R}^T \boldsymbol{\mu}_{\mathbf{W}_{d,:}^{(m)}}, \mathbf{R}^T \boldsymbol{\Sigma}_{\mathbf{W}^{(m)}} \mathbf{R}), \\ q^*(\mathbf{Y}) &= \prod_{n=1}^N \mathcal{N}(\mathbf{y}_n | \mathbf{R}^{-1} \boldsymbol{\mu}_{\mathbf{y}_n}, \mathbf{R}^{-1} \boldsymbol{\Sigma}_{\mathbf{Y}} \mathbf{R}^{-T}), \\ q^*(\boldsymbol{\alpha}^{(m)}) &= \prod_{k=1}^K \text{Gamma}(\alpha_0 + D_m/2, \beta_0 + \mathbf{r}_k^T \langle \mathbf{W}^{(m)T} \mathbf{W}^{(m)} \rangle_{\mathbf{r}_k} / 2), \end{aligned}$$

where \mathbf{r}_k is the k th column of \mathbf{R} . The cost function for optimizing \mathbf{R} is

$$\begin{aligned} \mathcal{L}_{\mathbf{R}} &= \langle \log p(\mathbf{Y}) \rangle^* - \langle \log q^*(\mathbf{Y}) \rangle^* + \\ &\quad \sum_{m=1}^2 \langle \log p(\mathbf{W}^{(m)} | \boldsymbol{\alpha}^{(m)}) \rangle^* - \langle \log q^*(\mathbf{W}^{(m)}) \rangle^* + \\ &\quad \langle \log p(\boldsymbol{\alpha}^{(m)}) \rangle^* - \langle \log q^*(\boldsymbol{\alpha}^{(m)}) \rangle^*, \end{aligned}$$

where $\langle \cdot \rangle^*$ denotes the expectation with respect to the transformed distribution. The first four individual terms can be written, omitting constants independent of \mathbf{R} , as

$$\begin{aligned} \langle \log p(\mathbf{Y}) \rangle^* &= -\text{Trace}[\mathbf{R}^{-1} \langle \mathbf{Y}\mathbf{Y}^T \rangle \mathbf{R}^{-T}] / 2, \\ -\langle \log q(\mathbf{Y}) \rangle^* &= -N \log |\mathbf{R}|, \\ \langle p(\mathbf{W}^{(m)} | \boldsymbol{\alpha}^{(m)}) \rangle^* &\approx -D_m/2 \log \prod_{k=1}^K \mathbf{r}_k^T \langle \mathbf{W}^{(m)T} \mathbf{W}^{(m)} \rangle_{\mathbf{r}_k}, \\ -\langle \log q(\mathbf{W}^{(m)}) \rangle^* &= D_m \log |\mathbf{R}|. \end{aligned}$$

The last two terms can be accurately approximated as constants, since the prior is effectively non-informative ($\alpha_0 = \beta_0 \approx 0$). The same argument has been used to approximate the third term; a part that is effectively constant has been left out. Finally, the cost function to be maximized as a function of \mathbf{R} is

$$\begin{aligned} \mathcal{L}_{\mathbf{R}} &= -\text{Trace}[\mathbf{R}^{-1} \langle \mathbf{Y}\mathbf{Y}^T \rangle \mathbf{R}^{-T}] / 2 + \left(\sum_{m=1}^2 D_m - N \right) \log |\mathbf{R}| \\ &\quad - \sum_{m=1}^2 D_m/2 \log \prod_{k=1}^K \mathbf{r}_k^T \langle \mathbf{W}^{(m)T} \mathbf{W}^{(m)} \rangle_{\mathbf{r}_k}. \end{aligned} \tag{9}$$

For finding the optimum we calculate the gradient of the cost and use standard L-BFGS algorithm for the optimization, with the initial solution of $\mathbf{R} = \mathbf{I}$.

After the optimization converges the expectations can be transformed as

$$\begin{aligned}\langle \mathbf{Y} \rangle &\leftarrow \mathbf{R}^{-1} \langle \mathbf{Y} \rangle, \\ \Sigma_{\mathbf{Y}} &\leftarrow \mathbf{R}^{-1} \Sigma_{\mathbf{Y}} \mathbf{R}^{-T}, \\ \langle \mathbf{Y} \mathbf{Y}^T \rangle &\leftarrow \langle \mathbf{Y} \rangle \langle \mathbf{Y} \rangle^T + N \Sigma_{\mathbf{Y}}, \\ \langle \mathbf{W}^{(m)} \rangle &\leftarrow \langle \mathbf{W}^{(m)} \rangle \mathbf{R}, \\ \Sigma_{\mathbf{W}} &\leftarrow \mathbf{R}^T \Sigma_{\mathbf{W}} \mathbf{R}, \\ \langle \mathbf{W}^{(m)T} \mathbf{W}^{(m)} \rangle &\leftarrow \langle \mathbf{W}^{(m)} \rangle^T \langle \mathbf{W}^{(m)} \rangle + D_m \Sigma_{\mathbf{W}}.\end{aligned}$$

Appendix C. Parameter-expanded Variational Bayes for BCCA

Appendix B explains how to optimize the BIBFA lower bound with respect to the linear transformation. It is also possible to do the same for the BCCA model with full covariance matrices (8). In particular, for the choice of $\alpha^{(1)} = \alpha^{(2)}$ we can solve for optimal \mathbf{R} analytically. In the experiments conducted in this paper we always used this optimization step when computing BCCA.

The derivation follows closely the derivation provided for parameter-expanded factor analysis by Luttinen and Ilin (2010), and hence we only summarize here the main steps. We start with the expression in (9), and note that the last term,

$$\langle p(\mathbf{W}|\alpha) \rangle^* \approx -D/2 \log \prod_{k=1}^K \mathbf{r}_k^T \langle \mathbf{W}^T \mathbf{W} \rangle \mathbf{r}_k,$$

is maximized if $\mathbf{R}^T \langle \mathbf{W}^T \mathbf{W} \rangle \mathbf{R}$ is a diagonal matrix. Hence, we can add that as a constraint in the cost function. For a diagonal matrix we can easily compute the trace, and the term simplifies to

$$\langle p(\mathbf{W}|\alpha) \rangle^* \approx -D/2 \log \prod_{k=1}^K \mathbf{r}_k^T \langle \mathbf{W}^T \mathbf{W} \rangle \mathbf{r}_k = -D \log |\mathbf{R}|,$$

canceling $\langle \log q(\mathbf{W}) \rangle^* = D \log |\mathbf{R}|$ out. By reparameterizing \mathbf{R} through its singular value decomposition we can find the optimum for the remaining terms in (9) by computing the left singular vectors by eigendecomposition of $\langle \mathbf{Z} \mathbf{Z}^T \rangle / N$. The right singular vectors are then chosen to make $\langle \mathbf{W}^T \mathbf{W} \rangle$ diagonal.

Appendix D. BIBFA with Spike-and-slab Prior

For inference with the spike-and-slab prior (6) we use Gibbs sampling, following closely the updates given for element-wise sparse FA model by Knowles and Ghahramani (2011). Here we summarize the necessary changes for adopting their model for group-wise sparsity in BIBFA.

D.1 Sampling \mathbf{H} and \mathbf{W}

We sample each entry $\mathbf{H}_{m,k}$ independently, based on the relative likelihoods of the two possible values. For $\mathbf{H}_{m,k} = 1$ we integrate out $\mathbf{W}_{:,k}^{(m)}$, which can be done independently for each element.

This results in the relative probability

$$\begin{aligned} \frac{p(\mathbf{H}_{m,k} = 1 | \mathbf{X}^{(m)})}{p(\mathbf{H}_{m,k} = 0 | \mathbf{X}^{(m)})} &= \frac{\pi_m \prod_{d=1}^{D_m} \int p(\mathbf{X}^{(m)} | \mathbf{W}_{d,k}^{(m)}) p(\mathbf{W}_{d,k}^{(m)} | 0, (\boldsymbol{\alpha}_k^{(m)})^{-1}) d\mathbf{W}_{d,k}^{(m)}}{(1 - \pi_m) \prod_{d=1}^{D_m} p(\mathbf{X}^{(m)} | \mathbf{W}_{d,k}^{(m)} = 0)} \\ &= \frac{\pi_m}{(1 - \pi_m)} \left(\frac{(\boldsymbol{\alpha}_k^{(m)})^{-1}}{\lambda} \right)^{D_m/2} \exp\left(\frac{1}{2} \lambda \boldsymbol{\mu}^T \boldsymbol{\mu}\right), \end{aligned}$$

where conditioning on the rest of the variables has been dropped for clarity. Here $\lambda = \tau_m \mathbf{Y}_{k,:}^T \mathbf{Y}_{k,:} + \boldsymbol{\alpha}_k^{(m)}$ and $\boldsymbol{\mu} = \frac{\tau_m}{\lambda} (\mathbf{X}^{(m)} - \sum_{j \neq k} \mathbf{W}_{:,j}^{(m)} \mathbf{Y}_{j,:}^T) \mathbf{Y}_{k,:}$. Compared to Knowles and Ghahramani (2011), we need to multiply D_m separate terms to reach the final ratio. On the other hand, we need not consider new components since we have replaced the Indian Buffet Process (IBP) prior with a simple Bernoulli prior; IBP would not be useful since we only have two realizations for each component.

While the above step integrates over $\mathbf{W}^{(m)}$, we will still need the projections for sampling other parameters of the model. Hence, we instantiate them by drawing $\mathbf{W}_{:,k}^{(m)} \sim \mathcal{N}(\boldsymbol{\mu}, \lambda^{-1} \mathbf{I})$ if $\mathbf{H}_{m,k} = 1$. Otherwise, we set the whole vector to $\mathbf{0}$ as dictated by the prior.

D.2 Sampling the Rest of the Parameters

The sampling for the rest of the parameters does not depend on the prior used for \mathbf{W} , since they depend directly on the current values of \mathbf{W} . The conditional distributions are very close to the updates used for the variational approximation, only now they are conditional on the current values instead of the expectations. Below we show the sampling equations for \mathbf{y}_n as an example; the updates for $\boldsymbol{\alpha}^{(m)}$ and $\tau^{(m)}$ can be easily modified from the variational updates given in Appendix A.

$$\begin{aligned} \mathbf{y}_n &\sim \mathcal{N}(\boldsymbol{\mu}_{\mathbf{y}_n}, \boldsymbol{\Sigma}_{\mathbf{Y}}), \\ \boldsymbol{\Sigma}_{\mathbf{Y}} &= (\mathbf{I} + \sum_{m=1}^2 \tau^{(m)} \mathbf{W}^{(m)T} \mathbf{W}^{(m)})^{-1}, \\ \boldsymbol{\mu}_{\mathbf{y}_n} &= \sum_{m=1}^2 \tau^{(m)} \boldsymbol{\Sigma}_{\mathbf{Y}} \mathbf{W}^{(m)T} \mathbf{x}_n^{(m)}. \end{aligned}$$

Finally, we need to update the variables π_m , drawing them from their Beta-posterior

$$\pi_m \sim \text{Beta}\left(1 + \sum_k \mathbf{H}_{m,k}, 1 + K - \sum_k \mathbf{H}_{m,k}\right).$$

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