Bayesian and L₁ Approaches for Sparse Unsupervised Learning

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Abstract

The use of L₁ regularisation for sparse learning has generated immense research interest, with many successful applications in diverse areas such as signal acquisition, image coding, genomics and collaborative filtering. While existing work highlights the many advantages of L₁ methods, in this paper we find that L₁ regularisation often dramatically under-performs in terms of predictive performance when compared to other methods for inferring sparsity. We focus on unsupervised latent variable models, and develop L₁ minimising factor models, Bayesian variants of "L₁", and Bayesian models with a stronger L₀-like sparsity induced through spike-and-slab distributions. These spike-and-slab Bayesian factor models encourage sparsity while accounting for uncertainty in a principled manner, and avoid unnecessary shrinkage of non-zero values. We demonstrate on a number of data sets that in practice spike-and-slab Bayesian methods outperform L₁ minimisation, even on a computational budget. We thus highlight the need to re-assess the wide use of L₁ methods in sparsity-reliant applications, particularly when we care about generalising to previously unseen data, and provide an alternative that, over many varying conditions, provides improved generalisation performance.

1. Introduction

Over the last decade, there has been tremendous excitement in learning parsimonious models using sparsity. Sparse learning is now a significant research topic – this significance being tied to the theoretical and practical advancement of sparse learning methods using the L₁ norm. The use of the L₁ norm in penalised regression problems such as the Lasso (Tibshirani, 1996), in natural scene understanding and image coding problems (Olshausen and Field, 1996), and more recently in compressed sensing (Candes, 2006), has served to cement the importance and efficacy of the L₁ norm as a means of inducing sparsity. Among its important properties, the L₁ norm is the closest convex norm to the L₀ norm, has a number of provable properties relating to the optimality of solutions and oracle properties (van de Geer and Bühlmann, 2009), and allows for the wide array of tools from convex optimisation to be used in computing sparse solutions. With the use of sparse methods in increasingly diverse application domains, it is timely to now contextualise the use of the L₁ norm and critically evaluate its behaviour in relation to other competing methods.

To achieve sparsity, the idealised but intractable sparsity criterion uses the L₀ norm to penalise the number of non-zero parameters. To more closely match the L₀ objective function, we develop here the use of discrete mixture priors for sparse learning, commonly referred to as spike-and-slab priors (Mitchell and Beauchamp, 1988; Ishwaran and Rao, 2005). A spike-and-slab is a discrete mixture of a point mass at zero (the spike) and any other continuous distribution (the slab). It is is similar to the L₀ norm in that it imposes a penalty on the number of non-zero parameters in a model. We show that spike-and-slab distributions provide improvements in learning, and that both Bayesian methods and the
use of the spike-and-slab distribution deserve more prominent attention in the vast literature for sparse modelling.

Our analysis focuses on unsupervised linear latent variable models (also known as matrix completion models), a class of models that are amongst the core tools in the machine learning practitioner’s toolbox. Factor analysis, the inspiration for this class of models, describes real-valued data by a set of underlying factors that are linearly combined to explain the observed data. This base model allows for many adaptations, such as generalisations to non-Gaussian data (Collins et al., 2002; Mohamed et al., 2008), or in learning sparse underlying factors (Dueck and Frey, 2004; Lee et al., 2009; Carvalho et al., 2008). In unsupervised learning, a sparse representation is desirable in situations where: 1) there are many underlying factors that could explain the data, 2) only a subset of which explain the data, and 3) the subset is different for each observation.

After introducing our framework for unsupervised models (section 2), we develop approaches for sparse Bayesian learning, culminating in a thorough comparative analysis. Our contributions include:

- We introduce new generalised latent variable models with strong sparsity, providing an important new class of sparse models that can readily handle non-Gaussian and heterogeneous data sets (sect. 4).
- We develop a spike-and-slab model for sparse unsupervised learning and derive a full MCMC algorithm for it. This MCMC method is applicable to other models based on discrete-continuous mixtures and is more efficient than naive samplers (sect. 3).
- We present the first comparison of approaches for sparse unsupervised learning based on optimisation methods, Bayesian methods using continuous sparsity-favouring priors, and Bayesian methods using the spike-and-slab. We bring these methods together and compare their performance in a controlled manner on both benchmark and real world data sets across a breadth of model types (sect. 6).
- Interestingly, our results show that strong sparsity in the form of spike-and-slab models can outperform the commonly used $L_1$ methods in unsupervised modelling tasks.

2. Unsupervised Latent Variable Models and Sparsity

We are concerned with models of the form:

$$X = V \Theta + E, \quad e_n \sim N(0, \Sigma),$$

which is the matrix factorisation problem in which we search for a set of underlying factors $V$ and weights $\Theta$ that are combined to explain the observed data $X$. We often consider Gaussian latent variables and Gaussian noise with diagonal or isotropic covariances, in which case this model recovers the familiar factor analysis and principal components analysis models, respectively. If $V$ is sparse then subsets of the underlying factors explain the data and different subsets explain each observed data point.

Increasingly we do not deal with real-data, which is well described by a Gaussian distribution, but data that may be binary, categorical, non-negative or a heterogeneous set of these. It is interesting to then consider generalisations of the basic model (1) in which the conditional probability of the observed data is defined using the exponential family of distributions, as:

$$x_n \mid v_n, \Theta \sim \text{Expon} \left( \sum_k \nu_{nk} \theta_k \right); \quad \theta_k \sim \text{Conj}(\lambda, \nu) \quad (2)$$

We use the shorthand $x_n \sim \text{Expon}(\psi)$ to represent the exponential family of distributions with natural parameters $\psi = v_n \Theta$. For this model, the natural parameters are a sum of the parameters $\theta_k$, weighted by $\nu_{nk}$, the points in the latent subspace corresponding to data point $x_n$. For the exponential family of distributions, the conditional probability of $x_n$ given parameter vector $\psi$ takes the form:

$$p(x_n \mid \psi) = h(x_n) \exp(s(x_n)\top \psi - A(\psi)),$$

where $s(x_n)$ are the sufficient statistics, $\psi$ is a vector of natural parameters and $A(\psi)$ is the log-partition function. Probability distributions that belong to the exponential family also have natural conjugate prior distributions, which we use to model the distribution of the parameters $\Theta$. We use the notation: Conj($\lambda, \nu$) as shorthand for the conjugate distribution, which has the form: $p(\theta_k) \propto \exp(\lambda \top \theta_k - \nu A(\theta_k))$, with hyperparameters $\lambda$ and $\nu$, and $A(\theta_k)$ is the same log-partition function from the likelihood function.

Figure 1 is a graphical representation of general unsupervised models; the shaded node $x_n$ represents the observed data item $n$. The plate notation represents replication of variables and the dashed node $\varphi$ represents any appropriate prior distribution for the latent variables $v_n$. The observed data forms an $N \times D$ matrix $X$, with rows $x_n$. $N$ is the number...
of data points and $D$ is the number of observed dimensions. $\Theta$ is a $K \times D$ matrix with rows $\theta_k$. $V$ is an $N \times K$ matrix with rows $v_n$, which are $K$-dimensional vectors, where $K$ is the number of latent factors.

The $K$ latent variables for each data point are generally assumed to be independent a priori: $v_n \sim \prod_{k=1}^K S(v_{nk}|\varphi)$, where $S$ is the prior on each variable with hyperparameters $\varphi$ (figure 1). The prior distribution $S(v_{nk})$ can be of any type. If the exponential family is Gaussian and we use Gaussian latent variables, we recover factor analysis; general exponential families corresponds to the well known exponential family PCA models (EPCA) (Collins et al., 2002; Mohamed et al., 2008). Considering non-Gaussian latent variables instantiates models such as ICA or the relevance vector machine (RVM) (Levin et al., 2009; Wipf and Nagarajan, 2008)

Unsupervised models with sparsity are obtained by employing sparsity-favouring distributions. A sparsity-favouring distribution can be any distribution with high excess kurtosis, indicating that it is highly peaked with heavy tails, or a distribution with a delta-mass at zero. The set of sparsity-favouring distributions includes the Normal-Gamma, Normal Inverse-Gaussian, Laplace (or double Exponential), Exponential, or generally the class of scale-mixtures of Gaussian distributions (Polson and Scott, 2010). Distributions that encourage sparsity fall into two classes: continuous sparsity-favouring or spike-and-slab distributions, which give rise to notions of weak and strong sparsity, respectively:

**Weak sparsity.** A parameter vector $\omega$ is considered to be ‘weakly sparse’ if none of its elements are exactly zero, but has most elements close to zero with a few large entries. This implies that a weakly sparse vector $\omega$ has a small $L_p$ norm for small $p$, or has entries that decay in absolute value according to some power law (Johnstone and Silverman, 2004).

**Strong sparsity.** A parameter vector $\omega$ is considered to be ‘strongly sparse’ if elements of $\omega$ are exactly zero. The spike-and-slab prior places mass explicitly on zero and is thus a prior suited to achieving this notion of sparsity in learning.

### 3. Strongly Sparse Bayesian Models

A Bayesian approach to learning averages model parameters and variables according to their posterior probability distribution given the data, rather than searching for a single best parameter setting as in an optimisation approach. To obtain Bayesian models with strong sparsity, we use a spike-and-slab prior (Mitchell and Beauchamp, 1988; Ishwaran and Rao, 2005): a discrete-continuous mixture of a point mass at zero referred to as the ‘spike’ and any other distribution known as the ‘slab’. This slab distribution is most often a uniform or Gaussian distribution, but may be any appropriate distribution. Since we have positive mass on zero, any samples produced include exact zeroes, thereby enforcing strong sparsity. The spike-and-slab can also be seen as placing a penalty on the number of non-zero parameters, and thus enforces sparsity in a manner similar to an $L_0$ norm penalisation. MCMC allows us to stochastically find suitable solutions in this setting, where this is not possible otherwise due to the combinatorial nature of the optimisation.

We construct a spike-and-slab prior using a binary indicator matrix $Z$ to indicate whether a latent dimension contributes to explaining the observed data or not. Each observed data point $x_n$ has a corresponding vector of Bernoulli indicator variables $z_n$. The spike components are combined with a Gaussian distribution, which forms the slab component:

$$p(z_n|\pi) = \prod_k B(z_{nk}|\pi_k) = \prod_k \pi_k^{z_{nk}} (1-\pi_k)^{1-z_{nk}}, \quad (3)$$

$$p(v_n|z_n, \mu, \Sigma) = \prod_k N(v_{nk}|\mu_k, z_{nk}\sigma_k^2), \quad (4)$$

where $N$ represents the Gaussian density with mean $\mu_k$ and variance $\sigma_k^2$. We place a Beta prior $\beta(\pi_k|\epsilon, f)$ on the Bernoulli parameters $\pi_k$. When $z_{nk} = 0$, $p(v_{nk})$ in equation 4 becomes a $\delta$-function at zero, indicating the spike being chosen instead of the slab. We complete the model specification by using a Gaussian-Gamma prior for the unknown mean $\mu_k$ and variance $\sigma_k^2$. We denote the set of unknown variables to be inferred as $\Omega = \{Z, V, \Theta, \pi, \mu, \Sigma\}$ and the set of hyperparameters $\Psi = \{\epsilon, f, \lambda, \nu\}$.

**MCMC Sampling Scheme**

Since the spike-and-slab is not differentiable, many popular MCMC techniques, such as Hybrid Monte Carlo, are not applicable. We proceed in the context of Metropolis-within-Gibbs sampling, where we sequentially sample each of the unknown variables using Metropolis-Hastings. Our sampling procedure iterates through the following steps: 1) Sample $Z$ and $V$ jointly; 2) Sample $\Theta$ by slice sampling (Neal, 2003); 3) Sample $\mu$, $\Sigma$ and $\pi$ by Gibbs sampling.

In sampling the latent factors $z_{nk}$ and $v_{nk}$ in step 1, we first decide whether a latent factor contributes to the data or not by sampling $z_{nk}$ having integrated out $v_{nk}$: $p(z_{nk} = 0|X, \pi, V_{-nk})$ and $p(z_{nk} = 1|X, \pi, V_{-nk})$, where $V_{-nk}$ are current values of $V$, with $v_{nk}$ excluded. Based on this decision, the
latent variable is sampled from the spike or the slab component. All variables \(v_{nk}\) associated with the slab components are sampled using slice sampling. Evaluating these probabilities involves computing the following integrals:

\[
p(z_{nk}=0|\mathbf{X}, \mathbf{V}, \nu_{nk}) = \int p(z_{nk}=0, v_{nk}=0|\mathbf{X}, \mathbf{V}, \nu_{nk}, \pi)dv_{nk} \\
= (1-\pi_k)p(\mathbf{X}|\mathbf{V}, \nu_{nk}, v_{nk}=0, \Theta) \\
\]

(5)

\[
p(z_{nk}=1|\mathbf{X}, \mathbf{V}, \nu_{nk}) = \int p(z_{nk}=1, v_{nk}|\mathbf{X}, \mathbf{V}, \nu_{nk}, \pi)dv_{nk} \\
= \pi_k \int p(\mathbf{X}|\mathbf{V}, \Theta)N(v_{nk}|\mu_k, \sigma_k^2)dv_{nk} \\
\]

(6)

While computing equation 5 is easy, the integral in equation 6 is not tractable in general. In the case of the Gaussian family, \(v_{nk}\) can be marginalised and we do exactly this. For other families the integral must be approximated. A number of approximation methods exist such as Monte Carlo integration, importance sampling and pseudo-marginal approaches, and the Laplace approximation, which we use here. The use of Laplace’s method introduces a bias due to the approximation of the target distribution. This problem has been studied by Guihenneuc-Jouyaux and Rousseau (2005) where the Laplace approximation is used in MCMC schemes with latent variables such as in our case, and show that such an approach can behave well. Guihenneuc-Jouyaux and Rousseau (2005) show that as the number of observations increases, the approximate distribution becomes close to the true distribution, and describe a number of assumptions for this to hold, such as requiring differentiability, a positive definite information matrix and conditions on the behaviour of the prior at boundaries of the parameter space.

At least three other approaches for sampling the latent variables can be considered: 1) A more naive sampling of alternating between \(\mathbf{V}\) and \(\mathbf{Z}\) without integrating out the slab. 2) Sampling \(\mathbf{V}\) after integrating \(\mathbf{Z}\). We found the collapsed scheme we describe in eq (5)–(6) quickly informs us of the state of the slab overall and resulted in faster mixing. 3) Reversible jump MCMC is also feasible and requires a different prior specification, also using a binary indicator vector but with a prior on the number of non-zero latent variables (e.g., using a Poisson).

We sample \(\mathbf{V}\) and \(\Theta\) in steps 1 and 2 by slice sampling (Neal, 2003), which can be thought of as a general version of the Gibbs sampler. Sampling proceeds by alternately sampling an auxiliary variable \(u\), the slice level, and then randomly drawing a value for the parameter from an interval along the slice. The variables \(\{\mu, \Sigma\}\) and \(\pi\) in step 3 have conjugate relationships with the latent variables \(\mathbf{V}\) and \(\mathbf{Z}\) respectively. Gibbs sampling is used since the full conditional distributions are easily derived. 1

4. Models with \(L_1\) norms and Sparsity-Favouring Priors

The \(L_1\) norm has become the established mechanism with which to encode sparsity into many problems, and has a strong connection to continuous densities that promote sparsity. The \(L_1\) norm has a number of appealing properties: it gives the closest convex optimisation problem to the \(L_0\) problem; there is an broad theoretical basis with provable properties \((L_0-L_1\text{ equivalence and exact recovery based on RIP}); and can be implemented efficiently based on the tools of convex optimisation (linear and semi-definite programming).

Sparsity Inducing Loss Functions

This leads us naturally to consider sparse latent variable models based on the \(L_1\) norm. If we assume that the latent distribution is a Laplace, \(S(v_n) \propto \exp(-\alpha \|v_n\|_1)\), the maximum a posteriori solution for \(\mathbf{V}\) is equivalent to \(L_1\) norm regularisation in this model. We define the following objective for sparse generalised latent variable modelling:

\[
\min_{\mathbf{V}, \Theta} \sum_n \ell(x_n, v_n; \Theta) + \alpha \|\mathbf{V}\|_1 + \beta R(\Theta), \\
\]

(7)

where the loss function \(\ell(x_n, v_n; \Theta) = -\ln p(x_n|v_n; \Theta)\), is the negative log likelihood obtained using equation 2. Equation 7 provides a unifying framework for sparse models with \(L_1\) regularisation. The regularisation parameters \(\alpha\) and \(\beta\), control the sparsity of the latent variables and the degree to which parameters will be penalised during learning. The function \(R(\Theta)\) is the regulariser for the model parameters \(\Theta\). This model is specified generally and applicable for a wide choice of regularisation functions \(R(\cdot)\), including the \(L_1\) norm. Such a loss function was described previously by Lee et al. (2009) – here we focus on unsupervised settings and specify the loss more generally, allowing for both sparse activations as well as basis functions. One configuration we consider is the use of the modified loss (7) with \(R(\Theta) = -\ln p(\Theta|\lambda, \nu)\). This loss allows sparsity in the latent variables and corresponds to finding the maximum a posteriori (MAP) solution. We shall refer to this model as the \(L_1\) model.

Optimisation is performed by alternating minimisation. Each step then reduces to established

\footnotetext{1}{Implementation notes online at: cs.ubc.ca/~shakirm}
problems for which, we can then rely on the extensive literature regarding $L_1$ norm minimisation. A number of methods exist to solve these problems: they can be recast as equivalent inequality constrained optimisation problems and solved using a modified LARS algorithm (Lee et al., 2006), recast as a second order cone program, or solved using a number of smooth approximations to the regularisation term (Schmidt et al., 2007), amongst others.

### Sparse Bayesian Learning

Continuous densities with high excess kurtosis such as the zero-mean Laplace distribution or Student’s-t distribution are often used in Bayesian models where sparsity is desired. For a model with priors that prefer sparsity, the Bayesian averaging process often results in non-sparse posteriors and give solutions that are nearly zero, resulting in weakly sparse models. We consider two models with sparsity in the latent variables $v_n$: 

**Laplace Model.** Using the Laplace distribution: $v_n \sim \prod_{k=1}^{K} \frac{1}{b_k} \exp (-b_k |v_{nk}|)$, a Bayesian version of the $L_1$ model described by equation 7 can be specified. The equivalence between this model and the $L_1$ model can be seen by comparing the log-joint probability using the Laplace distribution, to the $L_1$ loss of equation 7. We refer to Bayesian inference in this Laplace model as LXPCA, in contrast to the $L_1$ model, which is an optimisation-based method.

**Exponential Model.** If parameters or latent variables are to be positively constrained, the natural choice would be an exponential distribution peaked at zero: $v_n \sim \prod_{k=1}^{K} b_k \exp (-b_k v_{nk})$, which has similar shrinkage properties to the Laplace. We refer to this model as NXPCA.

These distributions are popular in sparse regression problems (Seeger. et al., 2007; Wipf and Nagarajan, 2008) and are natural candidates in the unsupervised models explored here. The hierarchical model specification is completed by placing a Gamma prior on the unknown rate parameters $b$, with shared shape and scale parameters $\alpha$ and $\beta$ respectively. We denote the set of unknown variables to be inferred as $\Omega = \{V, \Theta, b\}$ and the set of hyperparameters $\Psi = \{\alpha, \beta, \lambda, \nu\}$. The joint probability of the model is:

$$p(X, \Omega | \Psi) = p(X | V, \Theta) p(\Theta | \lambda, \nu) p(V | b) p(b | \alpha, \beta)$$  (8)

Inference in this model is accomplished using Markov Chain Monte Carlo (MCMC) methods, and the log of the joint probability (8) is central to this sampling. We use a sampling approach based on Hybrid Monte Carlo (HMC). This can be implemented easily, and we defer the algorithmic details to MacKay (2003).

### 5. Related Work

The body of related work is broad and the work described here is far from exhaustive, but attempts to capture many papers of relevance in contextualising approaches to, and applications of sparse learning. There is a wide body of literature for sparse learning in problems of feature selection, compressed sensing and regression using the $L_1$ norm, such as those by Tibshirani (1996); d’Aspremont et al. (2005); Candès (2006); Lee et al. (2006). Bayesian methods for sparse regression problems using continuous distributions have also been discussed by Seeger et al. (2007); Carvalho et al. (2010); O’Hara and Sillanpää (2009). Wipf and Nagarajan (2008) derive a relationship between automatic relevance determination (ARD), maximum likelihood and iterative $L_1$ optimization. Archambeau and Bach (2009) provide a nice exploration of ARD-related priors and variational EM for sparse PCA and sparse CCA.

Of relevance to unsupervised learning of real-valued data is sparse PCA and its variants (Zou et al., 2004; d’Aspremont et al., 2005; Rattray et al., 2009). The wide body of literature on matrix factorisation is also indirectly related (Airoldi et al., 2008). These methods do not deal with the exponential family generalisation and may yield sparse factors as a by-product, rather than by construction. There are also many other papers of relevance in bioinformatics, computer vision, ICA and blind deconvolution (Levin et al., 2009). The methods we develop here also have a strong bearing on the basis pursuit problem widely used in geophysics and other engineering fields and can allow not only for the solution of basis pursuit, but also in obtaining useful estimates of uncertainty.


### 6. Experimental Results

We consider the generalisation performance of unsupervised methods to unseen data, which appear as
missing data. To handle missing data, we divide the data into a set of observed and missing data, $X = \{X^{\text{obs}}, X^{\text{missing}}\}$, and condition on the set $X^{\text{obs}}$ in the inference. We create test sets by randomly selecting 10% of the elements of the data matrix. Test elements are set as missing values in the training data, and our learning algorithms have been designed in all cases to handle missing data. We calculate the predictive probability (negative log probability, NLP) and the root mean squared error (RMSE) using the testing data. We created 20 such data sets, each with a different set of missing data, and provide mean and one standard deviation error bars for each of our evaluation metrics. For fairness, the regularisation parameters $\alpha$ and $\beta$ in section 4 are chosen by cross-validation using a validation data set, which is chosen as 5% of the data elements. This set is independent of the data that has been set aside as training or testing data.

6.1. Benchmark Data

We use the block images data (Griffiths and Ghahramani, 2006) as a synthetic benchmark data set. The data consists of binary images, with each image $x_n$ represented as a 36-dimensional vector. The images were generated with four latent features, each being a type of block. The observed data is a combination of a number of these latent features. Noise is added by flipping bits in the images with probability 0.1. This data set consists of a number of latent factors, only a subset of which contributes to explaining any single data point. This data is synthetic, but not generated from any of the models tested.

Figure 2(a) shows the NLP and RMSE on this benchmark data set. The methods developed are compared to EPCA (Collins et al., 2002), BXPAC (Mohamed et al., 2008) and to binary ICA (Kaban and Bingham, 2006). A random predictor would have an NLP = $100 \times 36 \times 10\% = 360$ bits. The models tested here have performance significantly better than this. Both optimisation-based and Bayesian learning approaches do well, but the spike-and-slab model shows the best performance with smaller error bars.

6.2. Real Data

We summarise the real data sets we use in table 1 (which includes data in the $D > N$ regime). Natural images are the topic of much research based on $L_1$ regularisation. For the Olshausen and

<table>
<thead>
<tr>
<th>Data</th>
<th>$N$</th>
<th>$D$</th>
<th>Type</th>
</tr>
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<tbody>
<tr>
<td>Natural scenes</td>
<td>10,000</td>
<td>144</td>
<td>Real</td>
</tr>
<tr>
<td>Animal attributes</td>
<td>33</td>
<td>102</td>
<td>Binary</td>
</tr>
<tr>
<td>Newsgroups</td>
<td>100</td>
<td>200</td>
<td>Counts</td>
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<tr>
<td>Hapmap</td>
<td>100</td>
<td>200</td>
<td>Binary</td>
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Field (1996) image data set, we use $12 \times 12$ image patches extracted from a set of larger images. We use the Gaussian instantiation of the sparse generalised model (equation 2) and evaluate the performance of: $L_1$ optimisation; a Laplace-prior factor model; and the Bayesian spike-and-slab model. Our results are shown for both underdetermined and overcomplete bases ($K = 192$ as in Olshausen and Field (1996)) in figure 2(b). All methods perform similarly in the low-rank approximation cases, but as the model becomes overcomplete, Bayesian methods perform better with the spike-and-slab method much better than other methods, particularly in reconstructing held-out/missing data. The animal attributes data set of Kemp and Tenenbaum (2008) consists of animal species with ecological and biological properties as features. We use the binary unsupervised model and show results for various latent dimensions for NLP and RMSE in figure 3. For this data, the NLP of a random classifier is 336 bits and the models have NLP values much lower than this.

We also use a subset of the popular 20 news-groups data set, consisting of documents and counts of the words used in each document, with data sparsity of 93%. Figure 4(b) shows the performance of the Poisson unsupervised model using $L_1$ and spike-and-slab. Apart from the application of the model to count data, the results show that the spike-and-slab model is able to deal effectively with the sparse data and provides effective reconstructions and good predictive performance on held out data. We are also able to show the improved behaviour of the spike-and-slab model using the Hapmap data set. The comparative performance is shown in figure 4(c) showing the spike-and-slab has performance similar to $L_1$ in terms of RMSE at low $K$, but much better performance for large $K$.

7. Discussion and Conclusion

The common lore when using MCMC is that it is dramatically slower than optimisation methods. For

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2 Obtained from: https://mathgen.stats.ox.ac.uk/impute/
optimisation methods, the cross-validation procedure needed to set regularisation parameters $\alpha$ and $\beta$, is computationally demanding due to the need to execute the optimisation for many combinations of parameters. This approach is also wasteful of data, since a separate validation data set is needed to make sensible choices of these values and to avoid model overfitting. While individual optimisations may be quick, the overall procedure can take an extended time, which depends on the granularity of the grid over which regularisation values are searched for. These parameters can be learnt in the Bayesian setting and have the advantage that we obtain information about the distribution of our latent variables, rather than point estimates and can have significantly better performance.

Figure 4 demonstrates this tradeoff between running time and performance of the optimisation and the Bayesian approaches. $L_1$ was allowed to run to convergence and the spike-and-slab for 200 iterations. In this instance, the Bayesian method is seemingly slower, but produced significantly better reconstructions in both the human judgements and newsgroups data. We considered the setting where we have a fixed time budget and fixed the running time for the spike-and-slab to that used by the $L_1$ model (including time to search for hyperparameters). The result is shown (as S&S fixed) in figure 4, which shows that even with a fixed time budget, MCMC performs better in this setting. The table of figure 4(d) shows that the number of non-zeroes in the reconstructions for various $K$ for the newsgroups data, with the true number of non-zeroes being 1436. $L_1$ is poor in learning the structure of this sparse data set, whereas the spike-and-slab is robust to the data sparsity.

All our results showed the spike-and-slab approach to have better performance than other methods compared in the same model class. The models based on the $L_1$ norm or Bayesian models with continuous sparsity favouring priors enforce global shrinkage on parameters of the model. It is this property that induces the sparsity property, but which also results in the shrinkage of parameters of relevance to the data. This can be problematic in certain cases, such as the newsgroups data set which resulted in overly sparse data reconstructions. The spike-and-slab has the ability to give both global and local shrinkage, thus allowing sparsity in the model parameters while not restricting parameters that contribute to explaining the data.

Current approaches for sparse learning will have difficulty scaling to large data sets in this regime. We might think of EP as a potential solution, such as used by Hernández Lobato et al. (2010), but this is restricted to regression problems. For the standard Gaussian model, Rattray et al. (2009) discuss this issue and propose a hybrid VB-EP approach as one way of achieving fast inference, but such an approach is not ideal, leaving scope for future work.

We have demonstrated that improved performance can be obtained by considering sparse Bayesian approaches. In particular, Bayesian learning with spike-and-slab priors consistently showed the best perfor-
mance on held out data and produced accurate reconstructions, even in the ‘large p’ paradigm or with restricted running times. By considering the broad family of unsupervised latent variable models, we developed a sparse generalised model and provided new sampling methods for sparse Bayesian learning using the spike-and-slab distribution. Importantly, we have provided the first comparison of sparse unsupervised learning using three approaches: optimisation using the $L_1$ norm, Bayesian learning using continuous sparsity favouring priors, and Bayesian learning using the spike-and-slab prior. We have also demonstrated our methods in diverse applications including text modelling, image coding and psychology showing the flexibility of the sparse models developed. These results show that Bayesian sparsity and spike-and-slab methods warrant a more prominent role and wider use in sparse modelling applications.

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