Learning Exponential Families

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Abstract

Recently much attention has been paid to implicit probabilistic models – models defined by mapping a simple random variable through a complex transformation, often a deep neural network. These models have been used to great success for variational inference, generation of complex data types, and more. In most all of these settings, the goal has been to find a particular member of that model family: optimized parameters index a distribution that is close (via a divergence or classification metric) to a target distribution (such as a posterior or data distribution). Much less attention, however, has been paid to the problem of *learning a model* itself. Here we define implicit probabilistic models with specific deep network architecture and optimization procedures in order to learn intractable exponential family models (not a single distribution from those models). These exponential families, which are central to some of the most fundamental problems in probabilistic inference, are learned accurately, allowing operations like posterior inference to be executed directly and generically by an input choice of natural parameters, rather than performing inference via optimization for each particular realization of a distribution within that model. We demonstrate this ability across a number of non-conjugate exponential families that appear often in the machine learning literature.

1 Introduction

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Probability models, the fundamental object of Bayesian machine learning, have long challenged researchers with the tradeoff between tractability and expressivity. Though well understood that a model should be chosen to instantiate a set of assumptions and capture existing domain knowledge [1, 2, 3], for many years too-simple models were chosen for their practical advantanges (such as conditional conjugacy), which left much to be desired in terms of expressive performance and scalability of these models.

More recently the pendulum has swung, via a resurgence in models which map a latent random 26 variable $w \sim q_0$ through a member of a highly expressive function family $\mathcal{G} = \{q_\theta : \theta \in \Theta\}$, the 27 composition resulting in an implicit probability model $\mathcal{M} = \{q(g_{\theta} \circ w) : \theta \in \Theta\}$ (where $q(\cdot)$ is 28 the pushforward density, i.e. the density induced on the image of the random variable w under 29 the function q_{θ}). Choosing \mathcal{G} to be a parameter-indexed family of neural networks has both a rich 30 history [4, 5], and has recently been used to produce exciting results for density estimation [6, 7, 8], 31 generation of complex data [9], variational inference [10, 11, 12], and more. A noted advantage of 33 these implicit density network models is that in many cases they make minimal assumptions about the data generative (or posterior inference) process. On the other hand, since these models have 34 been chosen to be generic and flexible, they can lack the classic stipulation that a model instantiates 35 existing domain knowledge. The downsides of a too-flexible model with finite data (albeit large) - and the corresponding bias-variance benefit of a restricted model - are textbook knowledge [13, \$7.3], and work on generalization and compressibility in deep networks suggests that this broad class of function families are indeed quite large, perhaps problematically so [14].

Is all the flexibility of an implicit density network model \mathcal{M} always necessary? Consider the case of variational inference, where a generative model $p(z)p_{\beta}(X|z)$ (latent z, observed data X) is stipulated in the classic sense to embody modeling assumptions (hierarchical model, topic model, Bayesian logistic regression, etc.). When such a model is intractable, it is increasingly common to deploy an implicit "recognition network" model for variational inference [10], which finds a $q_{\theta^*}(z) \in \mathcal{M}$ such that an evidence bound is optimized with respect to the true posterior p(z|X). However, note the widely recognized fact [15] that many such true posteriors p(z|X) belong to models that can be written as exponential families (albeit intractable, due to the choice of sufficient statistics t(z)), of the form: $\mathcal{P} = \left\{\frac{h(z)}{A(\eta)} \exp\left\{\eta^{\top} t(z)\right\} : \eta \in H\right\}$. Some effort has been made to learn single members of exponential families from their mean parameters [16], but here we are focused on the natural parameterization and the model itself (not simply members thereof).

Should we be able to learn a tractable approximation to this exponential family model, we would in the very least get the bias-variance benefits of an intelligently restricted model space, and at best would get inference "for free" in the sense that we could evaluate approximate posteriors directly without separate optimization for each dataset encountered (a different form of amortized inference [17, 10, 11, 18]). In this paper we aim to learn a restricted model $Q = \{q(z; \eta : \eta \in H)\}$ that will be a strict subset of the deep implicit model \mathcal{M} and will closely approximate a target exponential family \mathcal{P} . Note the critical difference between this aim and much of the literature that seeks to learn a density $q_{\theta}^* \in \mathcal{M}$ (we explore this distinction in depth both algorithmically and empirically).

To proceed, we must first specify a set of models $\mathbb{Q} = \{\mathcal{Q}_{\phi} : \phi \in \Phi\}$, from which we can learn a single model \mathcal{Q}_{ϕ^*} , and we must second define a sensible parameter space H of each model. To the first, we restrict Θ , the parameter space of \mathcal{M} , to be itself the image of a second deep *parameter network* family $\mathcal{F} = \{f_{\phi} : \phi \in \Phi\}$, such that $\{f_{\phi}(\eta) : \eta \in H\} \subset \Theta$. The second part is answered immediately by our choice of target \mathcal{P} , an exponential family which by definition has *natural* parameterization $\eta \in H$. Thus, appealingly, we know that H is precisely the correct parameter space for \mathcal{Q} (as it defines \mathcal{P}), and that the image of H under f_{ϕ} will be of the correct dimensionality within the codomain Θ ; approximation error between \mathcal{Q} and \mathcal{P} will be caused by the flexibility and learnability of the parameter network f_{ϕ} and the density network $g_{f_{\phi}(\eta)}$.

We define this two-network architecture, which we term an *exponential family network* (EFN), and we specify a stochastic optimization procedure over a variant of the typical Kullback-Leibler divergence. We then demonstrate the ability of EFNs to approximately learn exponential families, both known tractable families and well-used intractable families, including hierarchical Dirichlet and truncated normal Poisson families. Finally we demonstrate the utility of this approach in an example inferring the posterior distribution of the latent intensity of a point-process, given neural spike train data. In all, our contributions include:

- a novel implicit model: a two-network deep architecture to learn a probability model along with a doubly stochastic optimization that samples over both natural parameters (the family member to be learned) and data points (observations of the target density);
- analysis of the connections between approximately learning a model and approximate variational inference, and an empirical study that gives insight to possible improvements to variational inference;
- empirical results confirming performance against ground truth in known tractable exponential families and in common intractable exponential families.

2 Exponential family networks

To define exponential family networks (EFN), we begin with relevant context for our modeling choice of exponential families (§2.1). We then describe the primary network architectural constraint and the background we leverage to satisfy that constraint (§2.2). We then introduce EFN in detail, including the optimization algorithm used for learning (§2.3). The similarities with variational inference are then explored in depth in §2.4.

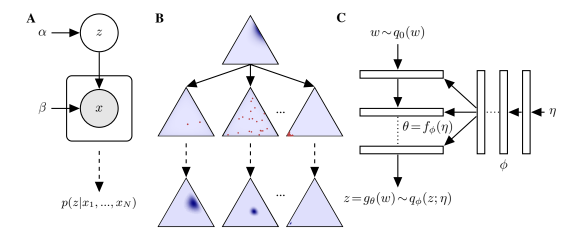


Figure 1: (A) Graphical model for conditionally iid sampling from an exponential family likelihood. (B) Hierarchical Dirichlets – prior $p_0(z)$ (top), three sample conditional Dirichlet datasets X of N=2, N=20, N=100 (middle), and three corresponding posteriors that themselves form an exponential family \mathcal{P} (bottom). (C) Architecture for exponential family network (EFN) – density network running top to bottom; parameter network running right to left.

2.1 Exponential families as target model \mathcal{P}

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91 92 93 We will focus on a fundamental problem setup in probabilistic inference, that of a latent variable $z \in \mathcal{Z}$ with prior belief $p_0(z)$, and where we observe a dataset $X = \{x_1, ..., x_N\} \subset \mathcal{X}$ as conditionally independent draws given z. Updating our belief with data produces the posterior $p(z|X) \propto p_0(z) \prod_{i=1}^N p(x_i|z)$. This setup is shown as a graphical model in Figure 1A.

In rare cases these posterior distributions are tractable due to either known conjugacy or to careful historical work (often an inversion, transformation-rejection, or similar custom numerical strategy) that has made these distributions computationally indistinguishable from tractable [19]. It is intriguing then to reflect upon the success that deep networks have offered to function approximation, and ask to what extent we can automate this numerical process, widening the class of effectively tractable distributions.

If we restrict our attention to priors and likelihoods that belong to exponential families $\mathcal{P} = \begin{cases} \frac{h(\cdot)}{A(\eta)} \exp\left\{\eta^\top t(\cdot)\right\} : \eta \in H \end{cases}$, the posterior can be also viewed as an exponential family, albeit intractable [15]. For simplicity we will hereafter suppress the base measure $h(\cdot)$. Consider:

$$p_0(z) = \frac{1}{A_0(\alpha)} \exp \left\{ \alpha^{\top} t_0(z) \right\} , \quad p(x_i|z) = \frac{1}{A(z)} \exp \left\{ \nu(z)^{\top} t(x_i) \right\},$$

where $t(\cdot)$ is the sufficient statistic vector, and $\nu(z)$ is the natural parameter of the likelihood in natural form [20]. The posterior then has the form:

$$p(z|x_1, ..., x_N) \propto \exp\left\{\begin{bmatrix} \alpha \\ \sum_i t(x_i) \end{bmatrix}^\top \begin{bmatrix} t_0(z) \\ \nu(z) \\ \log A(z) \end{bmatrix}\right\},$$
 (1)

which again is an exponential family, albeit intractable.

To give a concrete example, consider the hierarchical Dirichlet – a Dirichlet prior $z \sim Dir(\alpha)$ (of dimension $|\mathcal{Z}|$) with conditionally iid Dirichlet draws $x_i|z \sim Dir(\beta z)$, which has been considered historically [21], and is perhaps most notable for its nonparametric extension [22] (and has relevance for multi-corpus extensions of topic models [23, 24]). Figure 1B shows the prior for

a given α (top), and three examples of datasets that could arise via this generative model (middle). A set of basic manipulations shows the hierarchical Dirichlet posterior p(z|X) to be itself an exponential family with natural parameter $\eta = [\alpha - 1, \sum_i \log(x_i), -N]^{\top}$ and sufficient statistic $t(z) = [\log(z), \beta z, \log(B(\beta z))]^{\top}$. The corresponding posteriors are shown in Figure 1B (bottom). 112 113 Note importantly that, because the likelihood was chosen to be an exponential family (which is closed 114 under sampling), this form will not change for any choice of |Z|-dimensional hiearchical Dirichlet - any draw from the prior, any N, or any particular realization of observed data X (technically the 116 prior need not be exponential family, but we leave it as such for simplicity). The exponential family 117 is clearly sufficient for this property, and the Pitman-Koopman Lemma further clarifies that it is also 118 necessary (under reasonable conditions) [20, §3.3.3]. 119

The critical observation here is that, if we can approximately learn an intractable exponential family (the model itself), then it becomes trivial to perform posterior inference: we simply use the dataset to index into the natural parameter η of the intractable family, and the posterior distribution is produced. This is the goal of EFN.

124 2.2 Density networks as generic approximating family \mathcal{M}

Implicit probability models, which we will use for our approximating model family \mathcal{M} , can be defined by any base random variable $w \sim p_0$ mapped through any measurable, parameter-indexed function family $\mathcal{G} = \{g_\theta : \theta \in \Theta\}$; we denote the induced density on $z = g_\theta(w)$ as $q_\theta(z)$. Though trivial to sample from $q_\theta(z)$ for any choice of family \mathcal{G} , we here additionally require that we be able to explicitly calculate $q_\theta(z)$. This goal can be readily achieved by designing \mathcal{G} to contain only bijective functions, ideally with a Jacobian form that is convenient to compute. Designing that bijective \mathcal{G} as a deep neural network family, as we do here, is a well-established idea that has recently seen many variants and applications [5, 25, 26, 7, 6, 27, 28, 8, 29]. Specifically, let $z = g_\theta(w) = g_L \circ ... \circ g_1(w)$ for bijective vector-valued functions g_ℓ (where for clarity we have surpressed the dependence of each on θ), and denote $J_\theta^\ell(z)$ as the Jacobian of the function g_ℓ at the layer activation corresponding to z. Then we have:

$$q_{\theta}(z) = q_0 \left(g_1^{-1} \circ \dots \circ g_L^{-1}(z) \right) \prod_{\ell=1}^L \frac{1}{|J_{\theta}^{\ell}(z)|}.$$

The specific form of the layers g_{ℓ} can be chosen based on empirical considerations; we clarify our choice in §3. For the remainder (and to avoid confusion when we introduce a second network) we call this deep bijective neural architecture the *density network*; this network is shown vertically oriented (flowing from w down to z) in Figure 1C.

This density network induces the model $\mathcal{M} = \{q(g_{\theta} \circ w) : \theta \in \Theta\}$, which previous work has searched to find a single optimized distribution (such as a posterior or data generative density), on the

searched to find a single optimized distribution (such as a posterior or data generative density), on the assumption and subsequent empirical evidence that the target exponential family member is close to (or approximately belongs to) \mathcal{M} . We make the same assumption for the exponential family itself and seek to intelligently restrict \mathcal{M} in order to learn the exponential family.

2.3 Exponential family networks as approximating model Q

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Having introduced our target model \mathcal{P} , an exponential family with natural parameters $\eta \in H$, and the density network family \mathcal{M} , we now seek to learn $\mathcal{Q} \approx \mathcal{P}$, where $\mathcal{Q} \subset \mathcal{M}$. To do so we will parameterize θ , the parameters of the density network, as the image of a second *parameter network* family $\mathcal{F} = \{f_{\phi} : H \to \Theta, \phi \in \Phi\}$. This network is shown flowing from right to left in Figure 1C. Using a second meta-network to aid or restrict network learning has been used in a variety of settings; a few examples include parameterizing the optimization algorithm in the so-called "learning to learn" setting [30], and a more closely related work that used a second network to condition on observations for local latent variational inference [27], a connection which we explore closely in the following section.

Any choice of parameter network parameters ϕ induces a |H|-dimensional submanifold (the image $f_{\phi}(H)$) of the density network parameter space Θ , and as such defines a restricted model $\mathcal{Q}_{\phi}=$

¹To be clear this model is an exponential family if β is fixed or treated as a latent variable; this fact however will not be important for the development of this paper.

 $\{q_{f_{\phi}}(z;\eta):\eta\in H\}\subset\mathcal{M};$ by our choice of H as the natural parameter space of the exponential family target $\mathcal{P},$ this model restriction is at least of the correct dimensionality. Our goal then is to search over the implied set of models $\mathbb{Q}=\{\mathcal{Q}_{\phi}:\phi\in\Phi\}$ to find an optimal ϕ^* such that $\mathcal{Q}_{\phi^*}\approx\mathcal{P}.$

Given the connections between the exponential family and Shannon entropy, we will measure the error between Q_{ϕ} and \mathcal{P} with Kullback-Leibler divergence. Consider for the moment a fixed choice of natural parameter η ; we seek to minimize, over ϕ :

$$D\left(q_{\phi}(z;\eta)||p(z;\eta)\right) \propto \mathbb{E}_{q_{\phi}}\left(\log q_{\phi}(z;\eta) - \eta^{\top}t(z)\right) = \mathbb{E}_{q_{\phi}}\left(q_{0}\left(g_{\theta}^{-1}(z)\right) + \sum_{\ell=1}^{L}\log|J_{\theta}^{\ell}(z)| - \eta^{\top}t(z)\right),$$

where again we note that $\theta = f_{\phi}(\eta)$, and thus for a fixed eta, this objective depends only on ϕ . Indeed, the target $\eta^{\top}t(z)$ is linear in η (an obvious restatement of the log-linear exponential family form), giving us some hope that we may be able to learn this model. As a side note, this objective can also produce approximations of the log partition (as the intercept term implied by this linear target), which we have found to be reasonably accurate, though nuanced schemes are likely appropriate [31]; we do not explore that further here.

Of course we seek to approximate not just a single target exponential family member $(p(z;\eta))$ for a fixed η), but rather the entire model $\mathcal{P}=\{p(z;\eta):\eta\in H\}$. For optimization we thus need to introduce a distribution $p(\eta)$ (for sampling), leading to the objective:

$$\mathop{\rm argmin}_{\phi} \mathbb{E}_{p(\eta)} \left(D \left(q_{\phi}(z;\eta) || p(z;\eta) \right) \right) = \mathop{\rm argmin}_{\phi} D \left(q_{\phi}(z;\eta) p(\eta) || p(z;\eta) p(\eta) \right).$$

Unbiased estimates of this objective are immediate: $q_{\phi}(z;\eta)$ is sampled by computing calculating the density network parameters $\theta=f_{\phi}(\eta)$ (using the parameter network), sampling the latent $w\sim p_0(w)$, and running that w through the density network; $p(\eta)$ is user defined and thus trivial to sample. Stochastic optimization can then be carried out on the estimator:

$$\mathbb{L}(\phi) = \frac{1}{K} \frac{1}{M} \sum_{k=1}^{K} \sum_{m=1}^{M} \left(q_0 \left(g_{\theta^k}^{-1} \left(z^m \right) \right) + \sum_{\ell=1}^{L} \log |J_{\theta^k}^{\ell} \left(z^m \right)| - \eta_k^{\top} t \left(z^m \right) \right), \tag{2}$$

where $\theta^k = f_\phi(\eta_k)$. Successful optimization over ϕ should thus result in $\mathcal{Q}_{\phi^*} \in \mathbb{Q}$ that accurately approximates the target exponential family; that is, $\mathcal{Q} \approx \mathcal{P}$. We call this two-network architecture and optimization an exponential family network (EFN). What remains for empirical implementation is to make particular choices of hyperparameters, network layers, and optimization algorithm, which we specify in §3 below.

2.4 Relation to variational inference

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A tremendous amount of work in recent years has gone into variational inference (VI), and its 169 similarity to EFN warrants careful attention. In the following, we aim to carefully (and somewhat 170 pedantically) dissect this question. As such, though EFN can address any target exponential familiy, 171 to bring us closest to VI let us here restrict the EFN target model \mathcal{P} to be a family of posterior 172 distributions. 173 The typical role of variational inference is to infer an approximate posterior $q_{\phi}(z) \approx p(z|X)$. In this 174 setting, the difference with EFN is stark, in so much as VI learns this single posterior approximation, 175 whereas the main goal of the EFN is to approximate the model $\mathcal{P}=p_{\eta}(z|X):\eta inH$: to learn the family of distributions. More recently, much focus has gone into the particular instance of 177 VI for local variables z_i , for example $\prod_{i=1}^{N} p(z_i) p(x_i|z_i)$ (such as a variational autoencoder [10]) 178 or $p(u) \prod_{i=1}^{N} p(z_i|u) p(x_i|z_i)$ (latent Dirichlet allocation being a canonical example [23, 32]), the 179 result of which is often an amortized inference/recognition network that produces a local variational distribution $q_{\phi^*}(z_i|x_i)$. This local variational distribution is typically parameterized explicitly: the inference network $\mu_{\phi}(x_i)$ induces a local parametric distribution, often a Gaussian $q(z_i|x_i) \sim$ $\mathcal{N}(z_i; \mu_{\phi}(x_i))$ [10, for example]. Viewed this way, local-latent-variable VI methods induce a model $\{q_{\phi^*}(z_i|x_i): x_i \in X\}$ for a finite dataset X. In that sense, EFN and VI are similar 'model learning'

approaches. Even more closely, as part of a long-standing desire to add structure to VI beyond meanfield (classically [33, 34]; more recently [35, 36], to name but a few), in several cases a inference network has been used to parameterize a deep implicit model (in a two-network inference architecture, to say nothing of whether or not the generative model itself is a deep implicit model); closest to the EFN architecture is [27] (cf. Figure 2 of [27] with Figure 1C here). Thus EFN (when used for posterior families) can be seen as a close generalization of VI.

However, even accepting this VI-as-a-model view, the difference between the finite dataset X and 191 the natural parameter space H persists when viewed at a mechanical level; well-known are the 192 overfitting/generalization issues associated with a finite dataset compared with access to a distribution 193 $p(\eta)$. Thus one goal of EFN is to allow the model $Q_{\phi^*} \approx \mathcal{P}$ to be learned in the absence of a finite 194 dataset, such that inference on that dataset can then be executed without concerns of overfitting to 195 that set (and of course without having to run a VI optimization for every new dataset). Perhaps more 196 importantly, the "model" implied by VI is parameterized by x_i , and indeed the inference network 197 takes x_i as input. The EFN on the other hand is considerably more general: as Equation 1 shows, the posterior includes the natural parameters of the prior, allowing the EFN architecture to learn across a 199 more general setting that VI can not (since any VI inference network is only parameterized by data). 200 One final difference made clear by Equation 1 is that the observations are given to the EFN in natural 201 form (that is, $t(x_i)$, not x_i) [20]. This choice is a novel insight: by exploiting the known sufficiency 202 of $t(x_i)$ in the target model \mathcal{P} , some difference in performance for VI may be observed. We explore 203 this empirically in the following section. 204

Accordingly, while EFN and VI do at a high level bear multiple similarities, the differences are both material and provoke interesting speculation about means to improve both VI and EFN.

3 Results

We perform a number of experiments to investigate the performance of EFN. First, we test the ability 208 of EFN to approximate the target model \mathcal{P} when this model is a known, tractable exponential family: 209 this choice provides a simple ground truth and calibrates us to expected performance vs alternatives. 210 The main advantage of learning an EFN is to make tractable a previously intractable exponential 211 family (at least approximately). This confers major benefits in terms of test-time: for example, rather 212 than optimization needing to be run for variational inference with each particular dataset realized from a model class, EFN will allow immediate lookup. This benefit is orders of magnitude and is not 214 instructive to view, so here we focus our analyses on the costs of doing so: what approximation loss 215 is suffered when learning a whole family vs a single distribution. 216

To make this comparison, we use two alternatives. First, we restrict our algorithm to a single η ; that 217 is, K=1 in Equation 2, and further that choice of η is fixed throughout the course of optimization 218 (not stochastically sampled at every time). This is then a direct comparison that asks, given the same exact implicit model architecture, what cost is paid to learn a full model vs a single distribution. We call this alternative EFN1, which optimizes over ϕ as in the EFN. Second, it seems unnecessary to 221 carry around an entire parameter network $f_{\phi}(\eta)$ if that η will not change; thus our second alternative 222 (which is in some ways mechanically closest to traditional VI) is to dispose of the parameter network 223 and train the density network directly over θ (again with a deterministic choice of a single η); we call 224 this alternative NF1. 225

We also must make some particular architectural choices for these experiments. We considered a variety of density network architectures; in all the results we use the planar flow layer introduced in [27]. The parameter network need not be a density network, so we chose that more generically to be XXXXXXX.

In many of the results below we will analyze EFNs across a range of problem dimensionality D (that is, $z \in \mathcal{Z} \subseteq \mathbb{R}^D$). In all cases then we have also D layers in the density network, with units per layer **XXXXXXX**. The number of layers in the parameter network is **XXXXXXX**.

All code was implemented in tensorflow, and will be available at www.github.com/<anonymous>.

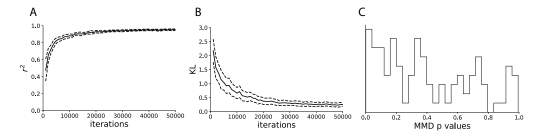


Figure 2: 25-dimensional Dirichlet exopnential family network. (A) Distribution of r^2 between log density of EFN samples and ground truth across choices of η throughout optimization. (B) Distribution of KL divergence throughout optimization. (C) Distribution of maximum mean discrepancy p-values between EFN samples and ground truth after optimization.

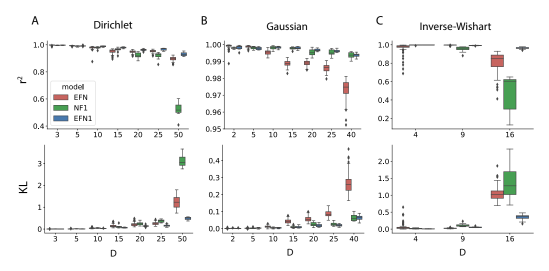


Figure 3: Scaling exponential family networks: D denotes the dimensionality of the family being learned, and comparisons are between EFN and its K=1 alternatives NF1 and EFN1 (see text). (A) Dirichlet family (B) Gaussian family (C) Inverse-Wishart family.

3.1 Tractable exponential families

Here we study the Dirichlet, Gaussian, and inverse-Wishart families, which offer a known ground truth and intuition about the range of performance that EFN – learning a model – can see with respect to its single-distribution counterparts (NF1 and EFN1).

First, to validate the basic EFN approach, we train the D=25-dimensional Dirichlet family. We chose $p(\eta)$, the prior on the α parameter vector of the Dirichlet, as **XXXXXX**. The number of η samples K at each iteration was XXX, and the minibatch size in z was M=XXXX. Figure 2 shows a high accuracy fit to this Dirichlet model: Figures 2A and 2B shows rapid convergence to high r^2 and low Kullback-Leibler divergence. r^2 is a convenient metric in so much as we are here doing distribution regression, so we calculate the coefficient of determination between the model predictions $q_{\phi}(z_i;\eta_k)$ and their known targets $\eta_k^{\top}t(z_i)$. We can then perform a standard MMD-based kernel two-sample test [37] between distributions chosen from \mathcal{P} and \mathcal{Q}_{ϕ^*} : the unstructured distribution of p values clarifies that the EFN model \mathcal{Q}_{ϕ^*} is not statistically significantly different than the true target Dirichlet family \mathcal{P} (using a test with **XXXX** samples).

Second, in Figure 3 we consider how this performance scales across dimensionality. Consider EFN vs EFN1, where again the only difference is that EFN attempts to learn the entire model (as in $\eta \in H$), whereas EFN1 chooses a single η and thus learns a single distribution. In both the Dirichlet and the Gaussian (Figure 3A and 3B), there is very minor (but statistically significant) loss from the EFN1 to EFN (but note the zoomed axis in Figure 3B; this difference is less than it may appear). This is quite encouraging: though training an entire model as opposed to a single distribution, performance holds

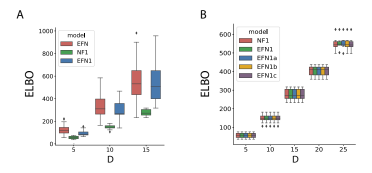


Figure 4: Scaling Dir-Dir

up adequately. If this performance level is adequate, using such a model is immediate; of course, failing that, the EFN could be used on a case by case basis to initialize the parameters $\theta_0 = f_\phi(\eta)$ for further optimization in θ . Performance in the inverse-Wishart is considerably less impressive when comparing the EFN to the EFN1, though we have found no satisfactory explanation for the shortcoming. It is also important to note that the distribution $p(\eta)$ can have material consequence on performance: the less entropic that distribution, the closer EFN gets to EFN1 by definition. The Dirichlet family has in our experience been robust to that choice, though perhaps surprisingly the Gaussian family has been less so (we swept the degrees of freedom of a Wishart prior on the covariance of the Gaussian $\nu=1,5,100XXXXXX$; the middle choice is shown here, the other two having very strong and very poor performance). Quite surprising is the performance of NF1. As a reminder the NF1 trains the density network directly over θ . One would think that, in so much as θ is typically of lower dimension than ϕ , that the NF1 would fit more easily; this expectation was only found in Figure 3B, though in Figure 3A and 3C EFN1 and EFN tended to outperform and scale better than NF1.

3.2 Intractable exponential families

Hierarchical Dirichlets Hierarchical dirichlets are useful and have some history; most notable is with the Hierarchical Dirichlet Process [22], but historically this was done in the finite case also [21]. Here is some math. Note that this matters for multi-corpus LDA generally as well [23, 24].

272 Truncated- and log-normal Poisson used a lot [38][39][40, 41]

273 Figure 4:

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- EFN in intractable exp fams (connecting to above, but with hard distribs and the ELBO)
- Panel A: Dir-Dir ELBO by dimensionality for NF1 and EFN and EFN1
- Panel B: Dir-Dir ELBO by dimensionality for EFN1 vs EFN1a vs 1b vs 1c vs NF1 (with N=1 data point)

3.3 Neural spike train analysis

- 280 Figure 5
- Panel A TNP picture example of prior and posterior with a few samples, just for feel good
- PANEL B: ELBO on held out data as a function of R, for a middle choice of training dataset size N and D.
- PANEL C: ELBO on held out data as a function of N, for a middle choice of number of samples in the posterior R.
- the posterior R.

PANEL D (optional): (ELBO EFN - ELBO NF1) as a surface plot as a function of R, N. That is, positive places is where EFN outperforms, negative NF1.

The key point with these is that, while you have the *same exact* flow network architecture, now you have to optimize over ϕ with a limited single dataset. Learning a restricted model space is good for the bias-variance tradeoff! Do this many times so that variance will become clear.

—other thoughts—Real data analysis and posterior inference. Key real data result on TNP.

Get some data from CRCNS that has many spike trains x_i for i = 1, ..., N (ask Gabriel, as he has done some poking around recently; or look at some of the above TNP/LNP refs).

Those spike trains should be conditionally independent draws from the same underlying intensity function z. (for example, trials under the same stimulus)

Bin the length of time T into $\approx 20-30$ equally spaced time bins. Thus z is now a vector in \mathbb{R}^20 .

Now each spike train x_i is a conditionally independent Poisson vector observation, with rate vector z.

Learn the 20 dimensional TNP exp fam, without any regard to this dataset X.

No: Panel No: TNP ELBO by dimensionality for NF1 and EFN and EFN1

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Panel A TNP picture example of prior and posterior with a few samples, just for feel good

Now we want to learn the posterior p(z) some fixed number R of data points).

To do this for an EFN, just plug in those R points $x_{i_1},...,x_{i_R}$ and the prior as a natural parameter, and job done.

To do this for an NF1, train a VI model by taking the log joint with R data points, then go through and resample R points every time from your training dataset with N data points.

PANEL A: ELBO on held out data as a function of R, for a middle choice of training dataset size N.

PANEL B: ELBO on held out data as a function of N, for a middle choice of number of samples in the posterior R.

PANEL C: (ELBO EFN - ELBO NF1) as a surface plot as a function of R, N. That is, positive places is where EFN outperforms, negative NF1.

The key point with these is that, while you have the *same exact* flow network architecture, now you have to optimize over ϕ with a limited single dataset. Learning a restricted model space is good for the bias-variance tradeoff! Do this many times so that variance will become clear. **Panel C v2:**

Possibly want to explicitly plot variance of EFN and NF1 to focus on the variance tradeoff

Panel C v3: change time bin granularity from 10 to 50 to show how this story changes in D.

My thought is that all will be exhausted by dimensionality sweeps by this point, so no.

also Notice one pain here is that these panels requires training a new EFN1 at every choice of N and R (but only one EFN). Sorry.

We hope and expect this will show that when the dataset gets small, this "traditional VI" will get arbitrarily bad (can't learn a network); eventually, there will be so much data that the VI will match or outperform the EFN... outperform because VI can focus specifically on this distribution rather than over the whole family, so the EFN has less effective data for this η (but not because it has a broader range of models, since we believe the EFN contains the closest member). Performance metric should be ELBO on some held out data or something like that (it's a posterior, so log likelihood doesn't really make sense). Test data anyway. Check VI papers for usual metrics. A key point to make here is that one great virtue of EFNs is is learning a restricted model, which should demonstrate the usual bias-variance tradeoff (see for example Hastie and Tibshirani book, Fig 7.2). Or Figure 4 is bias-variance and some sample posteriors in 2-d (showing how nicely it works), and then Fig 5 is the above performance, with both train and test.

This will be for one real example X. As such, to get error bars, just take a big dataset and randomly subsample the test set. Then the posterior performance is really for that very dataset, so the sem is coherent and the right thing to calculate/show. Important to clarify that doing so *does not* test how well this does across the entire exp fam, but just this one posterior. ((To test that, we would do it in simulation: generate *many datasets* X, then do the above for every one of them. Same computation for EFN (since its just plugging in a dataset), but VI alternatives 1 and 2 now need to be rerun for every dataset. And it's still simulated data, not really offering something fundamentally more than Fig 3 (well ok it's an intractable model, but I'm not sure that offers so much)...let's skip that altogether)).

342 4 Conclusion

Snappy closing remarks!

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