Reweighted Wake-Sleep

Jörg Bornschein\textsuperscript{1} and Yoshua Bengio\textsuperscript{2}
\textsuperscript{1}CIFAR Global Scholar, \textsuperscript{2}CIFAR Senior Fellow
Department of Computer Science and Operations Research
University of Montreal
Montreal, Quebec, Canada

Abstract

Training deep directed graphical models with many hidden variables and performing inference remains a major challenge. Helmholtz machines and deep belief networks are such models, and the wake-sleep algorithm has been proposed to train them. The wake-sleep algorithm relies on training not just the directed generative model but also a conditional generative model (the inference network) that runs backward from visible to latent, estimating the posterior distribution of latent given visible. We propose a novel interpretation of the wake-sleep algorithm which suggests that better estimators of the gradient can be obtained by sampling latent variables multiple times from the inference network. This view is based on importance sampling as an estimator of the likelihood, with the approximate inference network as a proposal distribution. This interpretation is confirmed experimentally, showing that better likelihood can be achieved with this reweighted wake-sleep procedure. Based on this interpretation, we propose that a sigmoid belief network is not sufficiently powerful for the layers of the inference network, in order to recover a good estimator of the posterior distribution of latent variables. Our experiments show that using a more powerful layer model, such as NADE, yields substantially better generative models.

1 Introduction

Training directed graphical models – especially models with multiple layers of hidden variables – remains a major challenge. This is unfortunate because, as has been argued previously (Hinton et al., 2006; Bengio, 2009), a deeper generative model has the potential to capture high-level abstractions and thus generalize better. The exact log-likelihood gradient is intractable, be it for Helmholtz machines (Dayan et al., 1995), sigmoid belief networks (SBNs), or deep belief networks (DBNs) (Hinton et al., 2006), which are directed, or deep Boltzmann machines (DBMs), which are undirected. Even obtaining an unbiased estimator of the gradient of the DBN or Helmholtz machine likelihood is not something that has been achieved in the past. Here we show that it is possible to get an unbiased estimator of the likelihood (which unfortunately makes it a slightly biased estimator of the log-likelihood), using an importance sampling approach. Past proposals to train Helmholtz machines and DBNs rely on maximizing a variational bound as proxy for the log-likelihood (Hinton et al., 1995; Kingma and Welling, 2014; Rezende et al., 2014). The first of these is the wake-sleep algorithm (Hinton et al., 1995), which relies on combining a “recognition” network (which we call an approximate inference network, here, or simply inference network) with a “generative” network. In the wake-sleep algorithm, they basically provide targets for each other. We review these previous approaches and introduce a novel approach that we generalize the wake-sleep algorithm. Whereas the original justification of the wake-sleep algorithm has been questioned (because we are optimizing a KL-divergence in the wrong direction), a contribution of this paper is to shed a different light on the wake-sleep algorithm, viewing it as a special case of the proposed reweighted wake-sleep (RWS) algorithm, i.e., as reweighted wake-sleep with a single sample. This makes it clear that
wake-sleep corresponds to optimizing a somewhat biased estimator of the likelihood gradient, while using more samples (i.e., RWS) makes the estimator less biased (and asymptotically unbiased as more samples are considered). We empirically show that effect, with clearly better results obtained with \( K = 5 \) samples than with \( K = 1 \) (wake-sleep), and 5 or 10 being sufficient to achieve good results. Unlike in the case of DBMs, which rely on a Markov chain to get samples and estimate the gradient by a mean over those samples, here the samples are iid, avoiding the very serious problem of mixing between modes that can plague MCMC methods (Bengio et al., 2013) when training undirected graphical models.

Another contribution of this paper regards the architecture of the deep generative model and of the approximate inference network. We view the inference network as estimating the posterior distribution of latent variables given the observed input. With this view, it is plausible that the classical architecture of the inference network (an SBN, details below) is inappropriate and we test this hypothesis empirically. In the classical sigmoidal belief network (SBN) (e.g., in the DBN and Helmholtz machine), the conditional distribution of each layer of the inference network, given the previous layer, is a factorized Bernoulli (where the probability for each bit is computed as in a logistic regression with the previous layer bits as inputs). We find that more powerful parametrizations of each layer as a conditional probability model yields better results.

2 Reweighted Wake-Sleep

2.1 The Wake-Sleep Algorithm

The wake-sleep algorithm was proposed as a way to train Helmholtz machines, which are deep directed graphical models \( p(\mathbf{x}, \mathbf{h}) \) over visible variables \( \mathbf{x} \) and latent variables \( \mathbf{h} \), where the latent variables are organized in layers \( \mathbf{h}_k \), with the \( k \)-th layer taking as input the random vector generated by the previous layer in the generating sequence, \( \mathbf{h}_{k+1} \). In the Helmholtz machine (Hinton et al., 1995, Dayan et al., 1995), the top layer, \( \mathbf{h}_L \), has a factorized unconditional distribution, so that ancestral sampling can proceed from \( \mathbf{h}_L \) down to \( \mathbf{h}_1 \) and then the generated sample \( \mathbf{x} \) is generated by the bottom layer, given \( \mathbf{h}_1 \). In the deep belief network (DBN) (Hinton et al., 2006), the top layer is instead generated by a restricted Boltzmann machine (RBM), i.e., by a Markov chain, while simple ancestral sampling is used for the others. Each intermediate layer is specified by a conditional distribution parametrized as a stochastic sigmoidal layer (see Section 3 for details).

The wake-sleep algorithm is a training procedure for such generative models, which involves training an auxiliary network, called the inference network, that takes a visible vector \( \mathbf{x} \) as input and stochastically outputs samples \( \mathbf{h}_k \) for all layers \( k = 1 \) to \( L \). The inference network outputs samples are from a distribution that should estimate the conditional probability of the latent variables of the generative model (at all layers) given the input. Note that in these kinds of models (and this is generally the case with latent variables), exact inference, i.e., sampling from \( p(\mathbf{h}|\mathbf{x}) \) is intractable.

The wake-sleep algorithm proceeds in two phases. In the wake phase, an observation \( \mathbf{x} \) (from the real world) is sampled from the training distribution \( D \) and propagated stochastically up the inference network (one layer at a time), thus sampling latent values \( \mathbf{h} \) from \( q(\mathbf{h}|\mathbf{x}) \). Together with \( \mathbf{x} \), the sampled \( \mathbf{h} \) forms a target for training \( p \), i.e., one performs a step of gradient ascent update with respect to maximum likelihood over the generative model \( p(\mathbf{x}, \mathbf{h}) \), with the data \( \mathbf{x} \) and the inferred \( \mathbf{h} \). This is useful because whereas computing the gradient of the marginal likelihood \( p(\mathbf{x}) = \sum_\mathbf{h} p(\mathbf{x}, \mathbf{h}) \) is intractable, computing the gradient of the complete log-likelihood \( \log p(\mathbf{x}, \mathbf{h}) \) is easy. In addition, these updates decouple all the layers (because both the input and the target of each layer are considered observed). In the sleep phase, a “dream” sample is obtained from the generative network by ancestral sampling from \( p(\mathbf{x}, \mathbf{h}) \) and is used as a target for the maximum likelihood training of the inference network, i.e., \( q \) is trained to estimate \( p(\mathbf{h}|\mathbf{x}) \).

The justification for the wake-sleep algorithm that was originally proposed is based on the following variational bound,

\[
\log p(\mathbf{x}) \geq \sum_\mathbf{h} q(\mathbf{h}|\mathbf{x}) \log \frac{p(\mathbf{x}, \mathbf{h})}{q(\mathbf{h}|\mathbf{x})}
\]

that is true for any inference network \( q \), but the bound becomes tight as \( q(\mathbf{h}|\mathbf{x}) \) approaches \( p(\mathbf{h}|\mathbf{x}) \). Maximizing this bound with respect to \( p \) corresponds to the wake phase update. The update with
respect to \( q \) should minimize \( KL(q(h|x)||p(h|x)) \) (with \( q \) as the reference) but instead the sleep phase update minimizes the reversed KL divergence, \( KL(p(h|x)||q(h|x)) \) (with \( p \) as the reference).

### 2.2 An Importance Sampling View yields Reweighted Wake-Sleep

If we think of \( q(h|x) \) as estimating \( p(h|x) \) and train it accordingly (which is basically what the sleep phase of wake-sleep does), then we can reformulate the likelihood as an importance-weighted average:

\[
p(x) = \sum_h p(x, h) = \sum_h q(h | x) \frac{p(x, h)}{q(h | x)}
\]

\[
\approx \frac{1}{K} \sum_{h^{(k)} \sim q(h|x)} \frac{p(x, h^{(k)})}{q(h^{(k)} | x)}
\]

Eq. (1) is a consistent and unbiased estimator for the marginal likelihood \( p(x) \). The optimal \( q \) that results in a minimum variance estimator is \( q^*(h | x) = p(h | x) \). In fact we can show that this is a zero-variance estimator, i.e., the best possible one that will result in a perfect \( p(x) \) estimate even with a single arbitrary sample \( h \sim p(h | x) \):

\[
\mathbb{E}_{h \sim p(h|x)} \left[ \frac{p(h | x) p(x)}{p(h | x)} \right] = p(x) \mathbb{E}_{h \sim p(h|x)} [1] = p(x)
\]

(2)

Any mismatch between \( q \) and \( p(h|x) \) will increase the variance of this estimator, but it will not introduce any bias. In practice however, we are typically interested in an estimator for the log-likelihood. Taking the logarithm of (1) and averaging over multiple datapoints will result in a conservative biased estimate and will, on average, underestimate the true log-likelihood due to the concavity of the logarithm. Increasing the number of samples will decrease both the bias and the variance. Variants of this estimator have been used in e.g. (Kingma and Welling, 2014; Rezende et al., 2014) to evaluate trained models.

### 2.3 Training by Reweighted Wake-Sleep

We now consider the models \( p \) and \( q \) parameterized with parameters \( \theta \) and \( \phi \) respectively.

**Updating \( p_{\theta} \) for given \( q_{\phi} \):** We propose to use a importance sampling estimator based on eq. (1) to compute the gradient of the marginal log-likelihood \( \mathcal{L}(\theta, x) = \log p_{\theta}(x) \):

\[
\frac{\partial}{\partial \theta} \mathcal{L}_p(\theta, x \sim D) = \frac{1}{p(x)} \mathbb{E}_{h \sim q(h|x)} \left[ \frac{p(x, h)}{q(h | x)} \frac{\partial}{\partial \theta} \log p(x, h) \right]
\]

\[
\approx \sum_{k=1}^K \omega_k \frac{\partial}{\partial \theta} \log p(x, h^{(k)}) \text{ with } h^{(k)} \sim q(h|x)
\]

(3)

and the importance weights \( \omega_k \) are automatically normalized such that they sum up to one.

**Updating \( q_{\phi} \) for given \( p_{\theta} \):** In order to minimize the variance of the estimator (1) we would like \( q(h|x) \) to track \( p(h|x) \). We propose to train \( q \) using maximum likelihood learning with the loss \( \mathcal{L}_q(\phi, x, h) = \log q_{\phi}(x|h) \). There are at least two reasonable options how to obtain training data for \( \mathcal{L}_q \): (1) maximize \( \mathcal{L}_q \) under the empirical distribution of the data: \( x \sim D \), \( h \sim p(h | x) \), or (2) maximize \( \mathcal{L}_q \) under the generative model \( (x, h) \sim p_{\theta}(x, h) \). We will refer to the former as \textit{wake}-\textit{q}-learning and to the latter as \textit{sleep}-\textit{q}-learning. In the case of a DBN (where the top layer is generated
Algorithm 1 Reweighted Wake-Sleep training procedure and likelihood estimator. $K$ is the number of approximate inference samples and controls the trade-off between computation and accuracy of the estimators (both for the gradient and for the likelihood). We typically use a large value ($K=500$) for test set likelihood estimator but a small value ($K=5$) for estimating gradients. Both the wake-phase and sleep-phase update rules for $q$ are optionally included (either one or both can be used, and best results were obtained using both). The original wake-sleep algorithm has $K=1$ and only uses the sleep-phase update of $q$. To estimate the log-likelihood at test time, only the computations up to $L_L$ are required.

```
for number of training iterations do
    Sample example(s) $x$ from the data generating distribution
    for $k = 1$ to $K$ do
        Sample latent variables $h^{(k)}$ from $q(h|x)$ layerwise (first layer above $x$, second layer, etc. up to top hidden layer).
        Compute $q(h^{(k)}|x)$ and $p(x, h^{(k)})$
    end for
    Compute unnormalized weights $\omega_k = \frac{p(x, h^{(k)})}{q(h^{(k)}|x)}$
    Normalize the weights: $\tilde{\omega}_k = \frac{\omega_k}{\sum_{k'} \omega_{k'}}$
    Compute unbiased likelihood estimator $\hat{p}(x) = \text{average}_k \omega_k$, or asymptotically unbiased estimator of log-likelihood $\hat{L}_L(x) = \log \text{average}_k \omega_k$
    Wake-phase update of $p$. Compute asymptotically unbiased estimator of log-likelihood gradient w.r.t. $p$, $\sum_k \tilde{\omega}_k \frac{\partial \log p(x, h^{(k)})}{\partial \theta}$, and perform an update of $p$'s parameters using it
    Optionally, wake-phase update of $q$. Sample $(x', h')$ from $p$ and use gradient $\frac{\partial \log q(h'|x')}{\partial \theta}$
end for
```

by an RBM), there is an intermediate solution, which has been proposed in (Hinton et al. 2006) and called contrastive wake-sleep. In contrastive wake-sleep we sample $x$ from the data, propagate it stochastically into top layer and use that $h$ as starting point for a short Markov chain in the RBM, then sample the other layers in the generative network $p$ to generate the rest of $(x, h)$. The objective is to put the inference network’s capacity where it matters most, i.e., near the input configurations that are seen in the training set.

Analogous to eqn. (1) and (3) we can use importance sampling to derive gradients for wake-q-learning (the details of the derivation can be found in the supplement):

$$\frac{\partial}{\partial \phi} L_q(\phi, x \sim D) \approx \sum_{k=1}^{K} \tilde{\omega}_k \frac{\partial}{\partial \phi} \log q_\phi(h^{(k)}|x)$$

(4)

with the same importance weights $\tilde{\omega}_k$ as in (3). Note that this is equivalent to optimizing $q$ so as to minimize $KL(p(\cdot|x) \parallel q(\cdot|x))$. For sleep-q-learning we consider the model distribution $p(x, h)$ a fully observed system and can derive the gradients without further sampling:

$$\frac{\partial}{\partial \phi} L_q(\phi, (x, h) \sim p(x, h)) \approx \frac{\partial}{\partial \phi} \log q_\phi(h|x) \text{ with } x, h \sim p(x, h)$$

(5)

This gradient is equivalent to the $q$-update used in the classical wake-sleep algorithm.

2.4 Relation to Wake-Sleep and Variational Bayes

There has been a resurgence of interest in algorithms related to the wake-sleep algorithm for directed graphical models such as the sigmoidal belief networks (SBN) and the Helmholtz machine (which is a generative SBN that is paired with an approximate inference SBN).

In Neural variational inference and learning (NVIL, Mnih and Gregor (2014)) the authors propose to maximize the variational lower bound on the log-likelihood to get a joint objective for both $p_\theta$ and $q_\phi$. It was known that this approach results in a gradient estimate of very high variance for
the recognition network $q$ (Dayan and Hinton [1996]). In the NVIL paper the authors therefore use variance reductions techniques such as baselines to obtain a practical algorithm that enhances significantly over the original wake-sleep algorithm.

Recent examples for continuous latent variables include the auto-encoding variational Bayes (Kingma and Welling [2014]) and stochastic backpropagation papers (Rezende et al. [2014]). In both cases one maximizes a variational lower bound on the log-likelihood that is rewritten as two terms: one that is just log-likelihood reconstruction error through a stochastic encoder (approximate inference) - decoder (generative model) pair, and one that regularizes the output of the approximate inference stochastic encoder so that its marginal distribution matches the generative prior on the latent variables (and the latter is also trained, to match the marginal of the encoder output). Besides the fact that these variational auto-encoders are only for continuous latent variables, another difference with the reweighted wake-sleep algorithm proposed here is that in the former, a single sample from the approximate inference distribution is sufficient to get an unbiased estimator of the gradient of a proxy (the variational bound). Instead, with the reweighted wake-sleep, a single sample would correspond to regular wake-sleep, which gives a biased estimator of the likelihood gradient. On the other hand, as the number of samples increases, reweighted wake-sleep provides a less biased (asymptotically unbiased) estimator of the log-likelihood and of it’s gradient. Similar in spirit, but aimed at a structured output prediction task is the method proposed by Tang and Salakhutdinov (2013). The authors optimize the variational bound of the log-likelihood instead of the direct IS estimate but they also derive update equations for the proposal distribution that resembles many of the properties also found in reweighted wake-sleep.

3 Component Layers

Although the framework can be readily applied to continuous variables, we here restrict ourselves to distributions over binary visible and binary latent variables. We build our models by combining probabilistic components, each one associated with one of the layers of the generative network or of the inference network. The generative model can therefore be written as $p_{\theta}(x, h) = p_0(x|h_1) p_1(h_1|h_2) \cdots p_L(h_L)$, while the inference network has the form $q_{\phi}(h|x) = q_1(h_1|x) \cdots q_L(h_L|h_{L-1})$. For a distribution $P$ to be a suitable component we must have a way to efficiently compute $P(x^{(k)}, y^{(k)})$ given some samples $(x^{(k)}, y^{(k)})$, and we must have a method to efficiently draw iid samples $x^{(k)} \sim P(x|y)$ for a given $y$. In the following we will describe experiments containing three kinds of layers:

Sigmoid Belief Network (SBN) layer: A SBN layer (Saul et al. [1996]) is a directed graphical model with independent variables $x_i$ given the parents $y$. 

$$P_{\text{SBN}}(x_i = 1|y) = \sigma(W_i^y y + b_i).$$

(6)

Although a SBN is a very simple generative model given $y$, performing inference for $y$ given $x$ is in general intractable.

Autoregressive SBN layer (DARN):

An autoregressive sigmoid belief network layer (Frey [1998], Bengio and Bengio [2000], Gregor et al. [2014]) is similar to an SBN layer but with the output units $x_i$ not being independent of each other, given the layer’s input $y$. Instead their dependency is captured by a fully connected directed acyclic graph where the $x_i$ can be predicted like in a logistic regression in terms of its predecessors $x_{<i}$ and of the input of the layer, $y$:

$$P_{\text{DARN}}(x_i = 1|x_{<i}, y) = \sigma(W_i^{y|x} y + S_i^{<i} x_{<i} + b_i).$$

(7)

We use $x_{<i} = (x_1, x_2, \cdots, x_{i-1})$ to refer to the vector containing the first $i$-1 observed variables. The matrix $S$ is a lower triangular matrix that contains the autoregressive weights between the observed variables. With $S_i^{<j}$ we refer to the first $j$-1 elements of the $i$-th row of this matrix.

Conditional NADE layer: The neural autoregressive distribution estimator layer (NADE, Larochelle and Murray [2011]) is a model that uses an internal, accumulating hidden layer to predict observed variables $x$, given the vector containing all previously observed variables $x_j$. DARN is thus a special case of NADE without (deterministic) hidden layer to mediate the conditional dependence between $x_i$ and its predecessors. Instead of a logistic regression, that dependency is mediated
by an MLP (Bengio and Bengio, 2000).

\[
P(x_i = 1 \mid x_{<i}) = \sigma(W^i: x_{<i} + a) + b_i)
\]  

(8)

where \(W\) and \(V\) are the encoding and decoding matrices for the NADE hidden layer. For our purposes we need to condition this model on another layer of random variables \(y\):

\[
P^{\text{NADE}}(x_i = 1 \mid x_{<i}, y) = \sigma(V^i: \sigma(W^i: x_{<i} + U_{a\ y} + a) + U_{bi} y + b_i))
\]  

(9)

Such a conditional NADE has been used previously for modeling musical sequences (Boulanger-Lewandowski et al., 2012).

For each layer distribution we can construct an unconditioned distribution by removing the random variable \(y\) (i.e., setting \(y=0\)). We use such unconditioned distributions as top layer \(p(h)\) for the generative network \(p\). By removing \(y\) from an SBN layer we obtain a factorized Bernoulli distribution; by removing \(y\) from a DARN layer we obtain a fully-visible sigmoid belief network (FVSBN, (Frey, 1998)) and by removing \(y\) from a NADE layer we obtain a regular, unconditioned NADE.

4 Experiments

Here we present series of experiments on the MNIST and the CalTech-Silhouettes datasets. The supplement describes additional experiments on various smaller datasets from the UCI repository. With these experiments we (1) quantitatively analyze the influence of the number of samples \(K\); (2) demonstrate that using a more powerful layer-model for the inference network \(q\) can significantly enhance the results even when the generative model is a simple SBN; and (3) that we can reach (close-to) state-of-the-art performance, especially when using powerful layer models such as a conditional NADE. Our implementation is available at [https://github.com/jbornschein/reweighted-ws/](https://github.com/jbornschein/reweighted-ws/).

4.1 MNIST

We use the MNIST dataset that was binarized according to Murray and Salakhutdinov (2009) and downloaded in it’s binarized form from (Larochelle, 2011). We use the last 1000 datapoints as validation set and do early-stopping with a lookahead of 10 epochs. For training we use stochastic gradient decent with momentum (\(\beta=0.95\)) and set mini-batch size to 25. The experiments in this paragraph were run with learning rates of \(\{0.0003, 0.001, \text{and } 0.003\}\). From these three we always report the experiment with the highest validation log-likelihood. In the majority of our experiments a learning rate of 0.001 gave the best results, even across different layer models (SBN, DARN and NADE). If not noted otherwise we use \(K=5\) samples during training and \(K=500\) samples to estimate the final log-likelihood on the test set.

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1 Although we refer to them as log-likelihood estimates, we actually report the lower-bounds of the log-likelihood that can be arbitrarily tightened by increasing the number of test-samples (see Section 2.2).
Table 1: MNIST for different architectures and network depths. In the third column we cite the numbers reported by [Mnih and Gregor 2014]. Columns three and four report the variational NLL bounds; columns 5 to 8 report the NLL estimates (See Section 2.2).

| P-model | size   | NVIL (NLL bound) | wake-sleep (NLL bound) | wake-sleep (NLL est.) | reweighted WS Q-model: SBN | reweighted WS Q-model: NADE |
|---------|--------|------------------|------------------------|-----------------------|-----------------------------|-------------------------------|
| SBN     | 200    | 113.1            | 120.7                  | 116.3                 | 105.7                       | 96.7                          |
| SBN     | 200-200| 99.8             | 109.4                  | 106.9                 | 98.3                        | 92.3                          |
| SBN     | 10-200-200 | 96.7 | 104.4 | 101.3 | 97.8 | 94.7 | 91.8 | 88.9 |
| DARN    | 200    |                  |                        |                       |                             | 91.8                          |
| DARN    | 200-200|                  |                        |                       |                             | 95.2                          |
| NADE    | 200    |                  |                        |                       |                             | 86.8                          |
| NADE    | 200-200|                  |                        |                       |                             | 87.6                          |

Figure 2: A Final log-likelihood estimate wrt number of test samples used. Solid lines: estimated using the importance sampler; dashed-lines: variational bound estimator. B Samples from the SBN/SBN 10-200-200 generative model. C Samples from the NADE/NADE 250 generative model. (We show the probabilities from which each pixel is sampled)

To disentangle the influence of the different q-learning methods we setup p and q networks consisting of SBN layers with three hidden layers of 10, 200 and 200 units (SBN/SBN 10-200-200). After convergence, the model trained using sleep-q-learning reached a final estimated log-likelihood of $-93.4$, the model trained with wake-q-learning reached $-92.8$ and the model trained with both reached $-91.9$. As a control we trained a model that does not update $q_\phi$ at all. This model reached $-171.4$. We confirmed that combining wake-q-learning and sleep-q-learning generally gives the best results by repeating this experiment with various other architectures. For the remainder of this paper we therefore train all models with combined wake-q and sleep-q-learning.

Next we investigate the influence of the number of samples used during training. The results are visualized in Fig. 1 A. Although the results depend on the layer-distributions and on the depth and width of the architectures, we generally observe that the final estimated log-likelihood does not improve significantly when using more than 5 samples during training for NADE models, and using more than 25 samples for models with SBN layers. We can quantify the bias and the variance of the gradient estimator (1) using bootstrapping. While training a SBN/SBN 10-200-200 model with $K=100$ training samples, we use $K=5,000$ samples to get a high quality estimate of the gradient for a small but fixed set of 25 datapoints (the size of one mini-batch). By repeatedly resampling smaller sets of $\{1, 2, 5, \cdots, 5000\}$ samples with replacement and computing the gradient based on these, we get a measure for the bias and the variance of the small sample estimates relative the high quality estimate. These results are visualized in Fig. 1 B. In Fig. 2 A we finally investigate the quality of the log-likelihood estimator (equation 1) when applied to the MNIST test set.

Table 1 summarizes how different architectures compare to each other and how reweighted wake-sleep compares to related methods for training directed models. In Table 2 (left) we compare our best trained models to the state-of-the-art results published on MNIST: our model with the largest log-likelihood reaches 85.32 and is a shallow model composed of (conditional) NADEs with with
### Results on binarized MNIST

| Method                        | NLL bound | NLL est. |
|-------------------------------|-----------|----------|
| RWS (SBN/SBN 10-200-200)      | 90.79     |          |
| RWS (SBN/NADE 10-200-200)     | 88.47     |          |
| RWS (NADE/NADE 250)           | 85.23     |          |
| NADE (500 units, [1])         | 85.35     |          |
| EoNADE (2hl, 128 orderings, [2]) | 85.10   |          |
| DARN (500 units, [3])         | 84.13     |          |
| RBM (500 units, CD3, [4])     | 105.5     |          |
| RBM (500 units, CD25, [4])    | 86.34     |          |
| DBN (500-2000, [5])           | 86.22     | 84.55    |

### Results on CalTech 101 Silhouettes

| Method                        | NLL est. |
|-------------------------------|----------|
| RWS (SBN/SBN 10-50-100-300)   | 116.9    |
| RWS (SBN/NADE 10-50-100-300)  | 112.6    |
| RWS (NADE/NADE 150)           | 104.3    |
| NADE (500 hidden units)        | 110.6    |
| RBM (4000 hidden units, [6])  | 107.8    |

Table 2: Results for various reweighted wake-sleep trained directed models in relation to previously published methods: [1] Larochelle and Murray (2011), [2] Murray and Larochelle (2014), [3] Gregor et al. (2014), [4] Salakhutdinov and Murray (2008), [5] Murray and Salakhutdinov (2009), [6] Cho et al. (2013).

Figure 3: CalTech 101 Silhouettes: A Random selection of training data points. B Random samples from the SBN/SBN 10-50-100-300 generative network. C Random Samples from the NADE-150 generative network. (We show the probabilities from which each pixel is sampled)

250 hidden units. This model was trained using 50 epochs with a learning rate of 0.003 and $K=5$ samples and another 50 epochs with a learning rate of 0.001 and $K=25$ training samples.

#### 4.2 CalTech 101 Silhouettes

We applied reweighted wake-sleep to the 28 x 28 pixel CalTech 101 Silhouettes dataset. This dataset consists of 4100 examples in the training set, 2264 examples in the validation set and 2307 examples in the test set. We trained various architectures on this dataset using the same hyperparameter as for the MNIST experiments. Table 2 (right) summarizes our results. Note that our best SBN/SBN model is a relatively deep network with 4 hidden layers (300-100-50-10) and reaches a estimated LL of -116.9 on the test set. Our best network, a shallow NADE/NADE-150 network reaches -104.3 and improves over the previous state of the art (−107.8, an RBM with 4000 hidden units by Cho et al. (2013)).

#### 5 Conclusions

We have introduced a novel training procedure for deep generative models, which can have either discrete or continuous latent variables. It generalises and improves over the wake-sleep algorithm providing a lower bias and lower variance estimator of the log-likelihood gradient at the price of more samples from the inference network. We demonstrated that training directed models with reweighted wake-sleep results in competitive models that produce high quality samples and that are close to or improve upon state-of-the-art in terms of log-likelihood on several discrete datasets. In all cases we were able to train fairly deep networks with up to 4 layers without layerwise pretraining and without carefully tuning learning schedules.

We found that even if the generator network uses SBN layers, better results can be obtained with an inference network that has more powerful layers, such as DARN or NADE. However, while our best models with autoregressive layers in the generative network were always providing significantly
better results than the models using SBN layers only, these models where always shallow with only one hidden layer. At this point it is unclear if this is due to optimization problems.

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6 Supplement

6.1 Gradients for $p(x, h)$

$$\frac{\partial}{\partial \theta} L_p(\theta, x \sim D) = \frac{\partial}{\partial \theta} \log p_\theta(x) = \frac{1}{p(x)} \frac{\partial}{\partial \theta} \sum_h p(x, h)$$

$$= \frac{1}{p(x)} \sum_h p(x, h) \frac{\partial}{\partial \theta} \log p(x, h)$$

$$= \frac{1}{p(x)} \sum_h q(h|x) p(x, h) \frac{\partial}{\partial \theta} \log p(x, h)$$

$$= \frac{1}{p(x)} \sum_{h \sim q(h|x)} \left[ \frac{p(x, h)}{q(h|x)} \frac{\partial}{\partial \theta} \log p(x, h) \right]$$

$$\approx \frac{1}{\sum_k \omega_k} \frac{1}{K} \sum_{k=1}^K \omega_k \frac{\partial}{\partial \theta} \log p(x, h^{(k)})$$

(10)

with $\omega_k = \frac{p(x, h^{(k)})}{q(h^{(k)} | x)}$ and $h^{(k)} \sim q(h|x)$

6.2 Gradients for wake-q-learning

$$\frac{\partial}{\partial \phi} L_q(\phi, x \sim D) = \frac{\partial}{\partial \phi} \sum_h p(x, h) \log q_\phi(h|x)$$

$$= \frac{1}{p(x)} \sum_{h \sim q(h|x)} \left[ \frac{p(x, h)}{q(h|x)} \frac{\partial}{\partial \phi} \log q_\phi(h|x) \right]$$

$$\approx \frac{1}{\sum_k \omega_k} \frac{1}{K} \sum_{k=1}^K \omega_k \frac{\partial}{\partial \phi} \log q_\phi(h^{(k)} | x)$$

(11)

Note that we arrive at the same learning gradients when we set out to minimize the $KL(p \parallel q)$ for a given datapoint $x$:

$$\frac{\partial}{\partial \phi} KL(p_\theta(h|x) \parallel q_\phi(h|x)) = \frac{\partial}{\partial \phi} \sum_h p_\theta(h|x) \log \frac{p_\theta(h|x)}{q_\phi(h|x)}$$

$$= - \sum_h p_\theta(h|x) \frac{\partial}{\partial \phi} \log q_\phi(h|x)$$

$$\approx - \frac{1}{\sum_k \omega_k} \frac{1}{K} \sum_{k=1}^K \omega_k \frac{\partial}{\partial \phi} \log q_\phi(h^{(k)} | x)$$

(12)
6.3 Additional experimental results

6.3.1 Learning curves for MNIST experiments

![Learning curves for various MNIST experiments.](image)

6.3.2 Bootstrapping based $\log(p(x))$ bias/variance analysis

Here we show the bias/variance analysis from Fig. 1 B (main paper) applied to the estimated $\log(p(x))$ wrt the number of test samples.

![Bias and standard deviation of the low-sample estimated $\log(p(x))$.](image)
6.3.3 UCI binary datasets

We performed a series of experiments on 8 different binary datasets from the UCI database:

For each dataset we screened a limited hyperparameter space: The learning rate was set to a value in $0.001, 0.003, 0.01$. For SBNs we use $K=10$ training samples and we tried the following architectures: Two hidden layers with 10-50, 10-75, 10-100, 10-150 or 10-200 hidden units and three hidden layers with 5-20-100, 10-50-100, 10-50-150, 10-50-200 or 10-100-300 hidden units. We trained NADE/NADE models with $K=5$ training samples and one hidden layer with 30, 50, 75, 100 or 200 units in it.

| Model       | ADULT | CONNECT4 | DNA   | MUSHROOMS | NIPS-8-12 | OCR-LETTERS | RCV1 | WEB  |
|-------------|-------|----------|-------|-----------|-----------|-------------|------|------|
| FVSBN       | 13.17 | 12.39    | 83.64 | 10.27     | 276.88    | 39.30       | 49.84| 29.35|
| NADE$^*$    | 13.19 | 11.99    | 84.81 | 9.81      | 273.08    | 27.22       | 46.66| 28.39|
| EoNADE$^+$  | 13.19 | 12.58    | 82.31 | 9.68      | 272.38    | 27.31       | 46.12| 27.87|
| DARN$^3$    | 13.19 | 11.91    | 81.04 | 9.55      | 274.68    | 28.17       | 46.10| 28.83|
| RWS - SBN   | 13.65 | 12.68    | 90.63 | 9.90      | 272.54    | 29.99       | 46.16| 28.18|
| hidden units| 5-20-100, 10-50-150, 10-150, 10-50-150, 10-100-300, 10-50-200, 10-50-300|
| RWS - NADE  | 13.16 | 11.68    | 84.26 | 9.71      | 271.11    | 26.43       |      |      |
| hidden units| 30, 50, 100, 50, 75, 100 |

Table 3: Results on various binary datasets from the UCI repository. The top two rows quote the baseline results from Larochelle & Murray (2011); the third row shows the baseline results taken from Uria, Murray, Larochelle (2014). (NADE$^*$: 500 hidden units; EoNADE$^+$: 1hl, 16 ord)