Unsupervised Layer-Wise Model Selection in Deep Neural Networks

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Abstract. Deep Neural Networks (DNN) propose a new and efficient ML architecture based on the layer-wise building of several representation layers. A critical issue for DNNs remains model selection, e.g. selecting the number of neurons in each DNN layer. The hyper-parameter search space exponentially increases with the number of layers, making the popular grid search-based approach used for finding good hyper-parameter values intractable. The question investigated in this paper is whether the unsupervised, layer-wise methodology used to train a DNN can be extended to model selection as well. The proposed approach, considering an unsupervised criterion, empirically examines whether model selection is a modular optimization problem, and can be tackled in a layer-wise manner. Preliminary results on the MNIST dataset suggest the answer is positive. Further, some unexpected results regarding the optimal size of layers depending on the training process, are reported and discussed.

1 INTRODUCTION

The general question of model selection — including the selection of hyper-parameter values for a Machine Learning (ML) algorithm — remains at the core of the Statistics and Machine Learning studies [7]. From a practitioner viewpoint, the best practice relies on variants of the cross-validation procedure [6]: one should select the model and hyper-parameter setting yielding the best performance on average. From a theoretical viewpoint, although some intrinsic limitations of cross-validation have been pointed out in the ML literature, these appear to be negligible [2] comparatively to methodological errors [9]. From a computational viewpoint, one generally proceeds by exploring the hyper-parameter space using a grid search, possibly using racing-based approaches in order to decrease the overall computational cost [17]. Overall, theoreticians and practitioners would likely join and agree that the fewer hyper-parameters, the better.

A new ML field, Deep Networks [3, 11] however seems to go against such a parameter-light trend. The main claim behind Deep Networks can be schematized as: several levels of representations, stacked on top of each other, are required to represent complex concepts in a tractable way; a single-level representation, though in principle able to do the job, will not make it in practice. While the greater expressiveness and compactness obtained through the composition of representations had been known for decades, deep architectures were discarded as they could not be trained in an efficient way. The training bottleneck of deep architectures was overcome through an original, sequential approach [3, 11] aimed at the greedy optimization of a seemingly irrelevant criterion: while the goal of a Deep Network is to achieve supervised learning, each layer is pre-trained using unsupervised learning criteria (more in section 2).

The issue of DNN hyper-parameter selection however remains critical, as the number of hyper-parameters (e.g. for each layer: number of neurons and learning rate) linearly increases with the number of layers, exponentially increasing the cost of a grid search.

This paper investigates whether the “Unsupervised learning first!” principle at the root of DNNs can be applied to hyper-parameter selection too. Accordingly, an unsupervised criterion based on the Reconstruction Error is proposed. The underlying question is whether hyper-parameter selection is a modular optimization problem, meaning that the optimal overall parameter setting can be obtained by iteratively finding the optimal setting for layer \(i\); \(i\) iteratively finding the optimal setting for layer \(i + 1\), conditionally to the hyper-parameter values for layers \(1 \ldots i\).

The experimental validation of the approach for Restricted Boltzmann Machines (trained with Mean Field Contrastive Divergence [21] on the MNIST dataset) suggests a positive answer to the modularity question (section 4). Furthermore, some unexpected findings, concerning the optimal size of the representation with the number of gradient updates of the training procedure, are reported and discussed, raising new questions for further study.

2 DEEP NEURAL NETWORKS

For the sake of completeness, this section introduces Deep Neural Networks, focusing on the Restricted Boltzmann Machine and Auto-Associator approaches. The interested reader is referred to [11, 1, 13] for a comprehensive presentation.

2.1 Restricted Boltzmann Machine (RBM)

While a Boltzmann Machine is a completely connected network made of a bag of visible and hidden units, Restricted Boltzmann Machines (RBMs) only involve connections between visible units on the one hand and hidden units on the other hand (Fig. 1).

Let us denote \(v = v_1, \ldots, v_q\) (respectively \(h = h_1, \ldots, h_r\)) the set of visible (resp. hidden) units. For notational simplicity, it is assumed that both the visible and the hidden layers involve a bias unit always set to 1. An RBM is described from its set of weights \(W \in \mathbb{R}^{q \times r}\), where \(w_{ij}\) stands for the weight on the connection between \(v_i\) and \(h_j\). All units are boolean. Each visible unit \(v_i\) encodes the \(i\)-th domain attribute, while each hidden unit \(h_j\) encodes a hypothesis. Formally, the so-called energy of an RBM state \((v, h)\) is defined as:

\[
\text{Energy}(v, h) = -h^\top W v
\]
An RBM can be interpreted as a constraint satisfaction network, where the weight $w_{ij}$ reflects the correlation between $v_i$ and $h_j$ (if $w_{ij} > 0$ a lower energy is obtained for $v_i = h_j$ everything else being equal). As such, an RBM induces a joint probability measure on the space of RBM states:

$$P_W(v, h) = \frac{e^{-\text{Energy}(v, h)}}{Z} \quad (2)$$

where $Z$ denotes as usual the normalization factor. Simple calculations show that visible (resp. hidden) units are independent conditionally to the hidden (resp. visible) units, and the conditional probabilities can be expressed as follows, where $\text{sgm}(t) = \frac{1}{1 + e^{-t}}$:

$$P_W(h_i|v) = \text{sgm}(\sum_j w_{ij}v_j) \quad 1 \leq i \leq q \quad (3)$$

$$P_W(v_j|h) = \text{sgm}(\sum_i w_{ij}h_i) \quad 1 \leq j \leq r \quad (4)$$

An RBM thus defines a probabilistic generative model: Considering a uniform distribution on the visible units, a probability distribution on the hidden units is derived (Eq. (3)), which enables in turn to derive a probability distribution on the visible units (Eq. (4)) and this process can be iterated after the so-called Gibbs sampling (Monte-Carlo Markov Chain), converging toward $P_W$. Let us consider the probability distribution $P_D$, defined from the empirical data $D$ by taking a uniform sample $v$ in $D$, and sampling $h$ after $P_W(v|h)$. Intuitively, the RBM model best fitting the data is such that $P_W = P_D$ (a sample $v$ is equally likely generated after $P_W$ or by uniformly sampling $D$).

Accordingly, an RBM is trained by minimizing the Kullback Leibler divergence $KL(P_D||P_W)$, or a tractable approximation thereof, the Contrastive Divergence [10]. Contrastive divergence can itself be approximated using a Mean Field approach [21], yielding a deterministic and faster learning procedure, albeit with higher risk of overfitting.

### 2.2 Stacked RBMs

After the Deep Network principles [11], stacked RBMs (SRBMs) are built in a layer-wise manner (Fig. 2). The first layer $h$ is built from the training set and the visible units $v$ as explained above, and the $i$-th layer RBM is built using the same approach, with the hidden units $h_{i-1}$ in the $i-1$-th layer being used as the new RBM’s visible layer.

The rationale for iterating the RBM construction is that trained hidden units $h_i$ are not independent; rather, they are independent conditionally to $v$. A more refined generative model can thus in principle be defined by capturing the correlations between the hidden units $h_i$ via a second RBM layer. More generally, the $i$-th layer in a stacked RBM aims at modelling the correlations between the hidden units in the previous layer. Letting $W_1, \ldots, W_\ell$ denote the RBM parameters for layers $1 \ldots \ell$, with $h_0 = v$, comes the equation:

$$P(v) = \sum_{h^1 \cdots h^\ell} P_W(v|h^1)P_W(h^1|h^2)\cdots P_W(h^\ell) \quad (5)$$

Let $f_W$ (respectively $g_W$) denote the forward propagation of an input to the hidden layer according to $P_W(h|v)$ (resp. the backward propagation from the hidden layer from the input according to $P_W(v|h)$). Considering a $\ell$-layer RBM with weights $W_1, \ldots, W_\ell$ (Fig. 2), $F_\ell$ and $G_\ell$ are respectively defined as the forward propagation of the input to the $\ell$-th layer, and the backward propagation from the $\ell$-th layer to the input:

$$F_\ell = f_{W_\ell} \circ \cdots \circ f_{W_1} \quad G_\ell = g_{W_1} \circ \cdots \circ g_{W_\ell} \quad (6)$$

### 2.3 Stacked Auto-Associators

Deep Neural Networks can also be built by stacking auto-associators [13]. An auto-associator is a 1-hidden layer neural network aimed at reproducing its input; formally, it uses back propagation to minimize the Reconstruction error defined as

$$\sum_{x \in D} ||x - \Phi(x)||^2$$

where $\Phi$ is the function corresponding to forward propagation through the network.

Stacking Auto-Associators proceeds by setting the $i$-th DNN layer to the hidden layer of the Auto-Associator built from the $(i-1)$-th DNN layer.

### 3 UNSUPERVISED MODEL SELECTION

The model selection approach is inspired from the DNN unsupervised layer-wise methodology. After discussing the position of the problem, this section describes an unsupervised criterion supporting the model selection task. The presented approach focuses on finding the optimal number of neurons in each layer of a stacked RBM. The choice of a SRBM architecture is motivated by their good empirical results and their strong theoretical background [11, 13, 14].
3.1 Position of the problem
The goal is to optimize the size $n_i$ of the $i$-th layer $L_i$ conditionally to the size of the layers $1, \ldots, i - 1$; in this manner, model selection is brought to a sequence of $\ell$ scalar optimization problems, as opposed to a single optimization problem in $N^2$. A first question regards the type of optimization criterion to be used. After [3], it is better that supervised learning criteria only be considered at a late stage when learning a deep architecture. Extensive empirical evidence suggests that unsupervised learning actually regularizes the supervised optimization problem [8]. The proposed model selection approach will accordingly be based on an unsupervised criterion.

Another question raised by a layer-wise approach regards its consistency, aka the modularity of the model selection problem. Formally, the question is whether the global optimum of the considered criterion is found with a layer-wise sequential optimization approach. Lastly and most importantly, the question is whether the layer-wise unsupervised optimization approach eventually enforces good performances w.r.t. supervised learning.

3.2 Reconstruction Error
A most natural unsupervised criterion relevant to RBMs is the log-likelihood of test data under the model. This criterion would however require computing the normalization factor in Eq. (2), which is intractable in the general case. An alternative is to consider the Reconstruction Error inspired from the auto-associator approach (section 2.3). Formally, the Reconstruction Error of an RBM is computed by clamping $v$ to some data sample $x$, computing the (real-valued) hidden unit configuration after $P(h|v)$, backpropagating this hidden configuration onto the visible units, and computing the square Euclidean distance between $x$ and the visible unit configuration obtained. For the sake of computational tractability, the Mean Field approximation is used.

With some notations as in section 2.2, let $W^*_D$ denote the weights of the RBM trained from the dataset $D$ using the hyper-parameters $\theta$; then the associated Reconstruction Error is defined as:

$$L(D, \theta) = -\mathbb{E}_D [L(D, W^*_D)] = \sum_{x \in D} ||x - g_{W^*_D} \circ f_{W^*_D}(x)||^2$$

The Reconstruction Error corresponding to the whole $\ell$-layer RBM can be directly defined as:

$$L(D, W_1, \ldots, W_\ell) = \sum_{x \in D} ||x - G_\ell \circ F_\ell(x)||^2$$

For reference, examples of reconstructed digits from the MNIST dataset are given in Fig.3.

4 EXPERIMENTAL VALIDATION
This section reports on the experimental validation of the proposed unsupervised layer-wise approach for hyper-parameter selection in stacked RBMs and discusses possible extensions.

4.1 Goals of experiments
The goal of the experiments is to answer the following questions:

Feasibility: Does the considered unsupervised criterion clearly and steadily support some selection of the optimal number of neurons?

Stability: Does the proposed procedure offer some stability w.r.t. the experimental setting (number of samples, number of epochs)?

Efficiency: How does the model selected in an unsupervised layer-wise manner affect the supervised classification accuracy?

Consistency: Does the model globally optimized over $\ell$ layers coincide with the layer-wise optimization of the size of each layer?

Generality: A last question concerns the extension of the proposed approach to RBMs trained with plain Contrastive Divergence, and Auto-Associators.

4.2 Experimental setting
The experiments consider the MNIST dataset including 60,000 $28 \times 28$ images representing digits from 0 to 9 in grey level, intensively used in the DNN literature [11, 13]. The unsupervised learning stage considers 1,000, 10,000 or 60,000 examples. A disjoint test set including 1,000 examples is used to assess the (supervised or unsupervised) generalization performance.

For the sake of computational tractability, the experimental validation was limited to a 2-layer RBM setting. The above experiment goals were investigated in this restricted experimental setting, based on a grid-search systematic exploration of the first and second layer size. The overall computational effort is 300 days CPU time on a 1.8GHz Opteron processor. Using multiple cores, several models were trained in parallel making the total training time about 15 days.

4.3 Feasibility and stability
The Reconstruction Error on the MNIST test set for the first RBM is reported vs the number of neurons (in log scale) in Fig.4. As expected, the Reconstruction Error is a monotonically decreasing function of the number of neurons. The feasibility of the approach however is empirically established as the Reconstruction Error displays a plateau when the number of neurons increases. After these results,

4 Mean Field Contrastive Divergence is used in the unsupervised phase, and backpropagation with momentum is used in the supervised phase. The algorithm is available upon request.

5 Experiments conducted on the cifar-10 dataset likewise show a decreasing reconstruction error as the number of neurons increases. Due to the higher complexity of the cifar-10 dataset comparatively to MNIST, the plateau however was not reached for the largest considered networks (up to 12,000 neurons).
the Reconstruction Error criterion suggests a clear and stable hyper-
parameter selection of \( n_1 \geq n_1^* = 300 \) neurons.

Following the proposed layer-wise approach (section 3.1), we now
optimize the size of the second layer, conditionally to the selected
size of the first layer. Setting the size of the first layer to \( n_1 = 300 \),
the selection of the optimal number of neurons in the second layer
after the Reconstruction Error criterion (Fig. 5), was conducted in
the same way as above. Once again, the experimental results show a
plateau for the Reconstruction Error criterion after a certain threshold
value. The optimal number of neurons on the second layer condition-
ally to \( n_1 = 300 \) is obtained for \( n_2 \geq n_2^* = 200 \).

The stability of the criterion with respect to experimental settings
is illustrated on Fig. 4, which shows the Reconstruction Error for
three different sizes of the training set and same number of gradient
updates. This confirms that the optimal layer size does not depend on
the size of the training set, conditionally to the number of gradient
updates; this point will be discussed further in section 4.6.

4.4 Efficiency and consistency

The efficiency of the presented approach must eventually be assessed
with the classification accuracy. In the previous experiments, RBMs
with different hidden layer sizes were trained and evaluated against
the Reconstruction Error criterion only. In order to assess their clas-
sification performance, a last layer with 10 output neurons is built
on the top of the first RBM-layer (with weights uniformly initialized
in \([-1, 1]\)), and a backpropagation algorithm with early stopping is
applied on the whole network [3]. Fig. 6 depicts the classification ac-
curacy versus the number of neurons in the hidden layer. As shown
in Fig. 7, the classification accuracy increases as the Reconstruction
Error decreases. Overall, the unsupervised criterion used for model
selection yields same performance as that of the best 1-hidden layer
neural networks in the literature (1.9%) [3].

A last question regards the consistency of the modular approach,
that is, whether simultaneously optimizing several layers yields the
same result as sequentially optimizing each layer conditionally to
the optimal setting for the previous layers. The global Reconstruction
Error on a 2-layer RBM is depicted in Fig. 8, where the first (resp. sec-
ond) axis stands for the number of neurons in the first (resp. second)
layer.

As could have been expected, the Reconstruction Error decreases
as the number of units increases; and it is shown to plateau after a suf-
cient number of units in each layer. Furthermore, one cannot com-
pensate for the insufficient number of units in layer 1 by increasing
the number of neurons in layer 2. This behavior is unexpected in the
perspective of the mainstream statistical learning theory [20], where
the VC-dimension of the model space increases with the number of
weights in the network, whatever the distribution of the neurons on
the different layers, as confirmed empirically in the multi-layer per-
ceptron framework [18].

The Reconstruction Error isolines are approximately rectangular-
shaped: for a given number of units \( n_1 \) on the first layer, there exists a
best Reconstruction Error reached for $n_2^*$ greater to a minimal value, referred to as $n_2^*(n_1)$. Likewise, the best Reconstruction Error for a given number of units $n_2$ on the second layer, is reached whenever the number of hidden units on the first layer is sufficient. Overall, the optimal Reconstruction Error w.r.t. the simultaneous optimization of $n_1$ and $n_2$, given in Fig. 8, leads to the same optimal setting as the layer-wise procedure in Fig.4 and Fig.5 ($\text{argmin} (n_1, n_2^*) = (n_1^*, n_2^*|n_1^*)$), which empirically confirms the modularity of the optimization problem.

### 4.5 Generality

Complementary experiments show that the presented approach hardly applies when considering Contrastive Divergence (instead of Mean Field); on the one hand, the Reconstruction Error very slowly decreases as the number of neurons increases, making the plateau detection computationally expensive. Furthermore, the Reconstruction Error displays a high variance between runs due to the stochastic nature of the learning procedure.

In the meanwhile, the same approach was investigated in the Auto-Associator (AA) framework, which naturally considers the Reconstruction Error as training criterion. Fig. 9 shows the AA Reconstruction Error on the MNIST dataset vs the number of neurons. Interestingly, the optimal layer size is larger than in the RBM case; interpreting this fact is left for further work.

### 4.6 Model selection and training process

Intermediate experiments, aimed at reducing the computational cost and memory resources involved in the experiment campaign, led to consider datasets with sizes 60,000, 10,000 and 1,000 (Fig. 10) trained for 1 epoch. Quite interestingly, for the Reconstruction Error criterion, the optimal number of hidden units decreases with smaller datasets.

A second experiment was launched to see if the above results could be attributed to the increased diversity of the bigger datasets, or to the increased number of gradient updates (granted that 1 epoch was used for every dataset in Fig. 10).

For two datasets of 1,000 and 60,000 samples, the Reconstruction Error is evaluated at multiple points in the training process, each separated by 1,000 gradient updates. The results (Fig.11) show the Reconstruction Error w.r.t. the number of neurons for the two datasets as the training progresses from right to left. The Reconstruction Error isolines clearly show the decreasing number of neurons needed with the increasing number of gradient updates.

A tentative interpretation for this fact goes as follows. Firstly, the above results state that the optimal model size can decrease as training goes on. Secondly, a main claim at the core of DNNs [4, 12] is that they capture a more abstract and powerful description of the data domain than shallow networks. Therefore it is conjectured that the network gradually becomes more able to construct key features as training goes on. Further work will aim at investigating experimentally this conjecture, by examining the features generated in the deep layers depending on the input distribution, as done by [16].

### 5 Conclusion and Perspectives

The contribution of the paper is twofold. Firstly, an unsupervised layer-wise approach has been proposed to achieve model selection in Deep Networks. The selection of hidden layer sizes is a critical issue potentially hindering the large scale deployment of DNN. A frugal unsupervised criterion, the Reconstruction Error, has been proposed. A proof of principle for the feasibility and stability of Reconstruction Error-based Model Selection has been experimentally given on the MNIST dataset, based on an extensive experiment campaign. The merits of the approach have also been demonstrated from a supervised viewpoint, considering the predictive accuracy in classification for the supervised DNN learned after the unsupervised layer-wise parameter setting.
After these results, the model selection tasks related to the different layers can be tackled in a modular way, iteratively optimizing each layer conditionally to the previous ones. This result is unexpected in the perspective of standard Neural Nets, where the complexity of the model is dominated by the mere size of the weight vector. Quite the contrary, it seems that deep networks actually depend on the sequential acquisition of different "skills", or representational primitives. If some skills have not been acquired at some stage, these can hardly be compensated at a later stage.

Lastly, the dependency between the model selection task and the training effort has been investigated. Experimental results suggest that more intensive training efforts lead to a more parsimonious model. Further work will investigate in more depth these findings, specifically examining the properties of abstraction of the hidden layers in an Information Theoretical perspective and taking inspiration from [16]. Along the same lines, the choice of the examples (curriculum learning [5]) used to train the RBM, will be investigated w.r.t. the unsupervised quality of the hidden units: The challenge would be to define an intrinsic, unsupervised, measure to order the examples and construct a curriculum.

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REFERENCES