Learning Combinations of Activation Functions

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Abstract—In the last decade, an active area of research has been devoted to design novel activation functions that are able to help deep neural networks to converge, obtaining better performance. The training procedure of these architectures usually involves optimization of the weights of their layers only, while non-linearities are generally pre-specified and their (possible) parameters are usually considered as hyper-parameters to be tuned manually. In this paper, we introduce two approaches to automatically learn different combinations of base activation functions (such as the identity function, ReLU, and tanh) during the training phase. We present a thorough comparison of our novel approaches with well-known architectures (such as LeNet-5, AlexNet, and ResNet-56) on three standard datasets (Fashion-MNIST, CIFAR-10, and ILSVRC-2012), showing substantial improvements in the overall performance, such as an increase in the top-1 accuracy for AlexNet on ILSVRC-2012 of 3.01 percentage points.

I. INTRODUCTION

In the last decade, deep learning has achieved remarkable results in computer vision, speech recognition and natural language processing, obtaining in some tasks human-like [1] or even super-human [2] performance. The roots of recent successes of deep learning can be found in: (i) the increase of the data available for training neural networks, (ii) the rising of commodity computational power needed to crunch the data, (iii) the development of new techniques, architectures, and activation functions that improve convergence during training of deeper networks, overcoming the obstacle of vanishing/exploding gradient [3], [4].

For many years, neural networks have usually employed logistic sigmoid activation functions. Unfortunately, this activation is affected by saturation issues. This problem reduces its effectiveness and, nowadays, its usage in feedforward networks is discouraged [5]. To overcome such weakness and improve accuracy results, an active area of research has been devoted to design novel activation functions.

The training procedure of these architectures usually involve optimization of the weights of their layers only, while non-linearities are generally pre-specified and their (possible) parameters are usually considered as hyper-parameters to be tuned manually. In this paper, we introduce two approaches able to automatically learn combinations of base activation functions (such as: the identity function, ReLU, and tanh) during training; our aim is to identify a search space for the activation functions by means of convex combination or affine combination of the base functions. To the best of our knowledge, this is one of the first attempts to automatically combine and optimize activation functions during the training phase.

We tested different well-known networks employing customary activation functions on three standard datasets and we compared their results with those obtained applying our novel approaches. The techniques proposed in this paper outperformed the baselines in all the experiments, even when using deep architectures: we found a 3.01 percentage points increase in the top-1 accuracy for AlexNet on ILSVRC-2012.

This paper is organized as follows: in Section II the related works are summarized; in Section III the proposed methods are described; in Section IV the experimental results are presented; finally, conclusions and future works are summarized in Section V.

II. RELATED WORK

Designing activation functions that enable fast training of accurate deep neural networks is an active area of research. The rectified linear activation function, introduced in [6] and argued in [7] to be a better biological model than logistic sigmoid activation function, has eased the training of deep neural networks by alleviating the problems related to weight initialization and vanishing gradient. Slight variations of ReLU have been proposed over the years, such as leaky ReLU (LReLU) [8], which addresses dead neuron issues in ReLU networks, thresholded ReLU [9], which tackles the problem of large negative biases in autoencoders, and parametric ReLU (PRelu) [10], which treats the leakage parameter of LReLU as a per-filter learnable weight.

A smooth version of ReLU, called softplus, has been proposed in [11]. Despite some theoretical advantages over ReLU (it is differentiable everywhere and it has less saturation issues), this activation function usually does not achieve better results [7] when compared to its basic version.

More recently, maxout has been introduced in [12] as an activation function aimed at enhancing dropouts abilities as a model averaging technique. Among its extensions, it is worth mentioning the probabilistic maxout [13], and the $L_p$ norm pooling activation [14] that is able to recover the maxout activation when $p \to \infty$.

Considering the last developments of activation functions in neural networks, it is important to mention the exponential linear unit function (ELU) [15] and the scaled exponential
linear unit function (SELU) \cite{15}. Like ReLU, LReLU, and PReLU, ELU reduces the vanishing gradient problem. Furthermore, ELU has negative values, allowing to push mean unit activations closer to zero like batch normalization, and speeding up the learning. SELU extends this property ensuring that activations close to zero mean and unit variance that are propagated through many network layers will converge towards zero mean and unit variance, even under the presence of noise and perturbations.

With the exception of PReLU, all previous activations are pre-specified (i.e. non-learnable). A first attempt to learn activations in a neural network can be found in \cite{17}, where the authors propose to randomly add or remove logistic or Gaussian activation functions using an evolutionary programming method. On the other hand, in \cite{18}–\cite{21} the authors proposed novel approaches to learn the best activation function per neuron among a pool of allowed activations by means of genetic and evolutionary algorithms.

A different method has been proposed in \cite{22}, which is able to learn the hardness parameter of a sigmoid function, similarly to the approach employed by PReLU to learn the leakage parameter.

However, all the previous learning techniques are limited by the fact that the family of functions over which the learning takes place is either finite or a simple parameterization of customary activation functions.

Recently, in \cite{23} the authors tackle the problem from a different angle using piecewise linear activation functions that are learned independently for each neuron using gradient descent. However, (i) the number of linear pieces is treated as a hyper-parameter of the optimization problem; (ii) the number of learned parameters increases proportionally to the amount of hidden units; (iii) all the learned piecewise linear activation functions are ReLU\(x)\) for \(x\) large enough (i.e. there exists \(u \in \mathbb{R}\) such that \(g(x) = \text{ReLU}(x)\) for \(x \geq u\)), thus reducing the expressivity of the learned activation functions by design.

It is worth mentioning that, in \cite{23} the authors propose the network in network approach where they replace activation functions in convolutional layers with small multi-layer perceptrons.

In this paper we try to overcome some of the limitations of the aforementioned approaches. Indeed, the two techniques explained in Section \ref{sec:methods} (i) increase the expressivity power of the learned activation functions by enlarging the hypothesis space explored during training with respect to \cite{17}–\cite{22}; (ii) restrict the hypothesis space with respect to \cite{23}, \cite{24}, in order to allow a faster training without the need of a careful initialization of network weights (see Proposition \ref{prop:conv} and the following lines).

### III. Methods

A neural network \(N_d\) made of \(d\) hidden layers can be seen as the functional composition of \(d\) functions \(L_i\) followed by a final mapping \(L\) that depends on the task at hand (e.g. classification, regression): \(N_d = L \circ L_d \circ \ldots \circ L_1\). In particular, each hidden layer function \(L_i\) can be written as the composition of two functions, \(g_i\) followed by \(\sigma_i\), the former being a suitable remapping of the layer input neurons, the latter being the activation function of the layer: \(L_i = \sigma_i \circ g_i\). In the most general case, both \(\sigma_i\) and \(g_i\) are parameterized and belongs to some hypothesis spaces \(\mathcal{H}_{\sigma_i}\) and \(\mathcal{H}_{g_i}\), respectively. Hence, the learning procedure of \(L_i\) amounts to an optimization problem over the layer hypothesis space \(\mathcal{H}_i = \mathcal{H}_{\sigma_i} \times \mathcal{H}_{g_i}\).

Usually, \(\sigma_i\) is taken as a non-learnable function; therefore, in the most common scenario \(\mathcal{H}_{\sigma_i}\) is a singleton: \(\mathcal{H}_i = \{ \{\sigma_i\} \times \mathcal{H}_{g_i}\}\). For example, for a fully-connected layer from \(\mathbb{R}^m\) to \(\mathbb{R}^m\), with ReLU activation we have that \(\mathcal{H}_{g_i}\) is the set of all affine transformations from \(\mathbb{R}^m\) to \(\mathbb{R}^m\), i.e. \(\mathcal{H}_i = \{\text{ReLU}\times \text{Lin}(\mathbb{R}^m, \mathbb{R}^m)\times \text{K}(\mathbb{R}^m)\}\), where \(\text{Lin}(A, B)\) and \(\text{K}(B)\) are the sets of linear maps between \(A\) and \(B\), and the set of translations of \(B\), respectively.

In this paper, we introduce two techniques to define learnable activation functions that could be plugged in all hidden layers of a neural network architecture. The two approaches differ in how they define the hypothesis space \(\mathcal{H}_{\sigma_i}\). Both of them are based on the following idea: (i) select a finite set of activation functions \(\mathcal{F} = \{f_1, \ldots, f_N\}\), whose elements will be used as base elements; (ii) define the learnable activation function \(\sigma_i\) as a linear combination of the elements of \(\mathcal{F}\); (iii) identify a suitable hypothesis space \(\mathcal{H}_{\sigma_i}\); (iv) optimize the whole network, where the hypothesis space of each hidden layer is \(\mathcal{H}_i = \mathcal{H}_{\sigma_i} \times \mathcal{H}_{g_i}\).

We will now give some basic definitions used throughout the paper. Note that, hereinafter, all activation functions from \(\mathbb{R}\) to \(\mathbb{R}\) will naturally extend to functions from \(\mathbb{R}^n\) to \(\mathbb{R}^m\) by means of entrywise application.

Given a vector space \(V\) and a finite subset \(A \subseteq V\), we can define the following two subsets of \(V\):

(i) the convex hull of \(A\), namely:

\[
\text{conv}(A) := \{ \sum_{i=1}^{n} c_i a_i \mid \sum_{i=1}^{n} c_i = 1, c_i \geq 0, a_i \in A\};
\]

(ii) the affine hull of \(A\), namely:

\[
\text{aff}(A) := \{ \sum_{i=1}^{n} c_i a_i \mid \sum_{i=1}^{n} c_i = 1, a_i \in A\}.
\]

We remark that, neither \(\text{conv}(A)\) nor \(\text{aff}(A)\) are vector subspaces of \(V\). Indeed, \(\text{conv}(A)\) is just a generic convex subset in \(V\) reducing to a \(|A| - 1\)-dimensional simplex whenever the elements of \(A\) are linearly independent. On the other hand, \(\text{aff}(A)\) is an affine subspace of \(V\) of dimension \(|A| - 1\), i.e. for an arbitrary \(a \in \text{aff}(A)\) the set \(\{a - a' \mid a' \in \text{aff}(A)\}\) is a linear subspace of \(V\) of dimension \(|A| - 1\). Clearly, \(\text{conv}(A) \subseteq \text{aff}(A)\).

Let \(F := \{f_0, f_1, \ldots, f_N\}\) be a finite collection of activation functions \(f_i\) from \(\mathbb{R}\) to \(\mathbb{R}\). We can define a vector space \(F\) from \(F\) by taking all linear combinations \(\sum_i c_i f_i\). Note that, despite \(F\) is (by definition) a spanning set of \(F\), it is not generally a basis; indeed \(|F| \geq \dim F\).

Since (almost everywhere) differentiability is a property preserved by finite linear combinations, and since \(\text{conv}(F) \subseteq \text{aff}(F) \subseteq F\), assuming that \(F\) contains (almost everywhere) differentiable activation functions, \(\text{conv}(F)\) and \(\text{aff}(F)\) are made of (almost everywhere) differentiable functions, i.e. valid
activation functions for a neural network that can be learned by means of gradient descent.

The activations used in real world scenarios are usually monotonic increasing functions. Unfortunately, the monotonicity is not ensured under arbitrary linear combination, meaning that even if all \( f_i \in \mathcal{F} \) are non-decreasing functions, an arbitrary element \( \hat{f} \in \mathcal{F} \) might be neither a non-decreasing nor a non-increasing function. As a matter of fact, considering only non-decreasing differentiable functions \( f'_i \geq 0 \ \forall f_i \in \mathcal{F} \), all non-decreasing differentiable functions in \( \mathcal{F} \) lie inside the convex cone \( \text{cone}(\mathcal{F}) \subset \mathcal{F} \), i.e.: 

\[
\text{cone}(\mathcal{F}) := \{ \sum_i c_i f_i \mid c_i \geq 0, f_i \in \mathcal{F} \}.
\]

Indeed, \( \forall g \in \text{cone}(\mathcal{F}) \) we have that \( g' \geq 0 \). Thanks to the definition of \( \text{aff}(\mathcal{F}) \), \( \text{cone}(\mathcal{F}) \), and \( \text{conv}(\mathcal{F}) \), we can conclude that \( \text{conv}(\mathcal{F}) = \text{cone}(\mathcal{F}) \cap \text{aff}(\mathcal{F}) \), which implies that monotonicity of the elements of \( \mathcal{F} \) is preserved by all the elements of \( \text{conv}(\mathcal{F}) \) but not by \( \text{aff}(\mathcal{F}) \) (see Figure 1). Nevertheless, in [25] it is shown that even non-monotonic activation functions can approximate arbitrarily complex functions for sufficiently large neural networks. Indeed, in [5] the authors trained a feedforward network using cosine activation functions not approximating the identity near origin (at least from one side).

(a) choose a finite set \( \mathcal{F} = \{ f_1, \ldots, f_N \} \), where each \( f_i \) is a (almost everywhere) differentiable activation function approximating the identity near origin (at least from one side);

(b) define a new activation function \( \hat{f} \) as a linear combination of all the \( f_i \in \mathcal{F} \);

(c) select as hypothesis space \( \mathcal{H}_f \) the sets \( \text{aff}(\mathcal{F}) \) or \( \text{conv}(\mathcal{F}) \).

In Section IV we present results using the following choices for \( \mathcal{F} \):

\[ \mathcal{F} := \{ \text{id}, \text{ReLU} \}, \quad \mathcal{F} := \{ \text{id}, \text{tanh} \}, \quad \mathcal{F} := \{ \text{ReLU}, \text{tanh} \}, \quad \mathcal{F} := \{ \text{id}, \text{ReLU}, \text{tanh} \}, \]

where \( \text{id} \) is the identity function. Clearly, other choices of \( \mathcal{F} \) may be available, provided the requirements in [1] are satisfied.

Since \( \text{conv}(\mathcal{F}) \subset \text{aff}(\mathcal{F}) \), the convex hull-based technique can be understood as a regularized version of the affine hull-based one, where the corresponding hypothesis space has been explicitly constrained to be compact. Such a regularization, in addition to restrict the complexity of the hypothesis space, guarantees that the final activation function is monotonic (provided all \( f_i \in \mathcal{F} \) are monotonic as well). Moreover, the convex hull-based technique together with \( \mathcal{F} := \{ \text{id}, \text{ReLU} \} \) recovers the learnable LReLU activation function, i.e. \( \text{LReLU}_\alpha(x) = x \) if \( x \geq 0 \), while \( \text{LReLU}_\alpha(x) = \alpha x \) otherwise, where \( 0 \leq \alpha \leq 1 \) (usually \( \alpha = 10^{-2} \)). Indeed, \( \text{conv}(\mathcal{F}) = \{ f := p \cdot \text{id} + (1 - p) \cdot \text{ReLU} \mid 0 \leq p \leq 1 \} \) and since \( \text{ReLU} = \text{id} \) for \( x \geq 0 \) and \( \text{ReLU} = 0 \) otherwise, we have that \( f = p \cdot \text{id} + (1 - p) \cdot \text{id} = \text{id} \) for \( x \geq 0 \) and \( f = p \cdot \text{id} + (1 - p) \cdot 0 = p \cdot \text{id} \) otherwise, i.e. \( \text{LReLU}_\alpha \).

It is worth mentioning that, as shown in Figure 2, layers using convex hull-based and affine hull-based activations with
n base functions can also be seen as the following two-stage pipeline: (i) n stacked fully connected layers, each of them featuring one of the base functions, and all of them sharing weights; (ii) a 1D-convolution with kernel of size 1 between the n separate channels), whose weights are constrained to sum to one (and to be positive in case of the affine hull-based technique).

IV. RESULTS

We tested the convex hull-based and the affine hull-based approaches by evaluating their effectiveness on three publicly available datasets used for image classification, greatly differing in the number of input features and examples. Moreover, each dataset was evaluated using different network architectures. These networks were trained and tested using as activation functions (for all their hidden layers) those learned during training phase by means of random horizontal flip and image shifting; the considered architectures are the following:

**LeNet-5**: a convolutional network made of two convolutional layers followed by two fully connected layers [32]. The convolutional layers have respectively 20 and 50 filters of size $5 \times 5$, while the hidden fully connected layer is made of 500 neurons. Max pooling with size $2 \times 2$ is used after each convolutional layer, without dropout. We assessed LeNet-5 on Fashion-MNIST and CIFAR-10 datasets, resulting in two networks with 431k and 657k parameters, respectively;

**KerasNet**: a convolutional neural network included in the Keras framework [33]. It is made of 4 convolutional layers and 2 fully connected layers, and it employs both max pooling and dropout. This architecture is presented in Table I. We tested KerasNet on both Fashion-MNIST and CIFAR-10 datasets, resulting in two networks with 890k and 1.2M parameters, respectively;

**ResNet-56**: a residual network made of 56 total layers, employing pooling and skip connection [34]. Its performance has been evaluated on CIFAR-10, corresponding to a network with 858k parameters;

**AlexNet**: a convolutional network made of 5 convolutional and 3 fully connected layers [31]. We tested it against ILSVRC-2012 dataset, resulting in a network with 61M parameters. Note that, as shown in [31], ReLU-based activation functions significantly outperform networks based on other activations. Therefore, in this context ReLU-based networks

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1 The 1D-convolution with kernel of size 1 can also be seen as a weighted average of the stacked fully connected layers (with possibly negative weights in case of the affine hull-based technique).
only have been implemented and tested for comparison.

All networks were implemented from scratch using the Keras framework [33] on top of TensorFlow [35]. Notice that, our AlexNet implementation is a porting to Keras of the Caffe architecture [34]. We trained LeNet-5, KerasNet, and ResNet-56 using RMSprop with learning rate $10^{-4}$ and learning rate decay over each mini-batch update of $10^{-6}$. For AlexNet we used SGD with starting learning rate $10^{-2}$, stepwise learning rate decay, weight-decay hyper-parameter $5 \cdot 10^{-4}$, and momentum 0.9.

Table III shows the top-1 accuracy for all the run experiments. The best configurations (shaded cells in the table) are always achieved using our techniques, where in 5 out of 6 experiments the affine hull-based approach outperformed convex hull-based ones. The uplift in top-1 accuracy using our techniques, where in 5 out of 6 experiments the affine hull-based approach outperformed convex hull-based ones.

Moreover, the proposed techniques usually achieve better results than their corresponding base activation functions (boldface in the table). Note that, the novel activations work well for very deep networks and also with various architectures involving different types of layers (e.g. residual unit, dropout, pooling, convolutional, and fully connected).

Furthermore, our experiments show how the learning of the leakage parameter achieved by the activation based on \text{conv}([[\text{id}, \text{ReLU}])] allows to outperform or to achieve the same results of LReLU.

In this paper we introduced two novel techniques able to learn new activations starting from a finite collection $F$ of base functions. Both our ideas are based on building an arbitrary linear combination of the elements of $F$ and on defining a suitable hypothesis space where the learning procedure of the linear combination takes place. The hypothesis spaces for the two techniques are \text{conv}($F$) and \text{aff}($F$). We showed that, provided all the elements of $F$ approximate the identity near origin, \text{aff}($F$) is the only set where it is possible to find combined activations that also approximate id near origin. Moreover, \text{aff}($F$) allows to explore non-monotonic activation functions, while \text{conv}([[\text{id}, \text{ReLU}]]) may be seen as a learnable LReLU activation function.

We tested the two techniques on various architectures (LeNet-5, KerasNet, ResNet-56, AlexNet) and datasets (Fashion-MNIST, CIFAR-10, ILSVR-C2012), comparing results with LReLU and single base activation functions.

In all our experiments, the techniques proposed in this paper achieved the best performance and the combined activation functions learned using our approaches usually outperform the corresponding base components. The effectiveness of the proposed techniques is further proved by the increase in performance achieved using networks with different depths and architectures.

In our opinion, an interesting extension of this work would be to analyze other sets ($F$) of base functions.

\section{Conclusion}

In this paper we introduced two novel techniques able to learn new activations starting from a finite collection $F$ of base functions. Both our ideas are based on building an arbitrary linear combination of the elements of $F$ and on defining a suitable hypothesis space where the learning procedure of the linear combination takes place. The hypothesis spaces for the two techniques are \text{conv}($F$) and \text{aff}($F$). We showed that, provided all the elements of $F$ approximate the identity near origin, \text{aff}($F$) is the only set where it is possible to find combined activations that also approximate id near origin. Moreover, \text{aff}($F$) allows to explore non-monotonic activation functions, while \text{conv}([[\text{id}, \text{ReLU}]]) may be seen as a learnable LReLU activation function.

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In all our experiments, the techniques proposed in this paper achieved the best performance and the combined activation functions learned using our approaches usually outperform the corresponding base components. The effectiveness of the proposed techniques is further proved by the increase in performance achieved using networks with different depths and architectures.

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TABLE II
EXPERIMENT RESULTS. THE TABLE SHOWS THE TOP-1 ACCURACY RESULTS FOR ALL THE ANALYZED NETWORKS ON FASHION-MNIST TEST SET, CIFAR-10 TEST SET, AND ILSVRC-2012 VALIDATION SET. CONVEX HULL-BASED AND AFFINE HULL-BASED ACTIVATIONS ACHIEVING TOP-1 ACCURACY RESULTS GREATER THAN THEIR CORRESPONDING BASE ACTIVATION FUNCTIONS ARE HIGHLIGHTED IN BOLDFACE. THE BEST RESULT FOR EACH NETWORK/DATASET IS SHADED.

| Activation | LeNet-5 KerasNet | LeNet-5 KerasNet ResNet-56 | AlexNet |
|------------|------------------|--------------------------|--------|
| id         | 90.50% 90.51%    | 75.27% 75.23%            | 42.60% |
| ReLU       | 91.06% 90.79%    | 80.37% 79.97%            | 89.18% |
| tanh       | 92.43% 93.43%    | 78.96% 82.86%            | 82.65% |
| LReLU      | 91.03% 91.13%    | 80.94% 80.07%            | 80.95% |
| conv(id, ReLU) | 91.87% 92.39% | 80.94% 84.74% | 90.51% |
| conv(tanh) | 92.36% 93.64%    | 79.45% 78.53%            | 86.46% |
| conv(tanh, ReLU) | 92.56% 92.04% | 80.21% 84.80% | 88.31% |
| conv(id,ReLU, tanh) | 92.21% 92.94% | 80.48% 85.21% | 89.60% |
| aff(id, ReLU) | 92.83% 93.37% | 82.52% 84.93% | 89.22% |
| aff(tanh)  | 92.65% 94.41%    | 78.97% 86.05%            | 82.97% |
| aff(tanh, ReLU) | 93.02% 93.48% | 81.23% 84.83% | 89.44% |
| aff(id, ReLU, tanh) | 92.80% 94.41% | 80.13% 87.45% | 88.62% |

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