SCATTERING CONVOLUTIONAL HIDDEN MARKOV TREE

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ABSTRACT

A Scattering convolutional hidden Markov tree proposes a new inference mechanism for high-dimensional signals by combining the interesting signal representation created by the scattering transform to a powerful probabilistic graphical model. A wavelet scattering network computes a translation invariant and stable to deformations representation of a signal while still preserving its informative content. Such properties are acquired by cascading wavelet transform convolutions with non-linear modulus and averaging operators. The network's structure and its distributions are described using a Hidden Markov Tree. This yield a generative model for highdimensional inference. It offers a mean for performing several inference tasks among which are predictions. The scattering convolutional hidden Markov tree displays promising results on classification tasks of complex images in the difficult case were the number of training examples is extremely limited.

Index Terms— Scattering network, Hidden Markov Model, Classification, Deep network

1 Introduction

The standard approach to classify high dimensional signals can be expressed as a two steps procedure. First the data are projected in a feature space where the task at hand is simplified. Then prediction is done using a simple predictor in this new representational space. The mapping can either be hand-built —e.g. Fourier transform, wavelet transform— or learned. In the last decade methods for learning the projection have drastically improved under the impulsion of the so called deep learning. Deep neural networks (sometime enriched by convolutional architecture) have been able to learn very effective representations for a given dataset and a given task [1–3]. Such method have achieved state of the art on many standard problems [4, 5] as well as real world applications [6].

However deep learning methods are only efficient when we have access to a vast quantity of training examples [7]. But in some cases, such as in medical or defence applications for example, datapoints are rare or using an expert for handlabelling them is either time-consuming, costly or subjective. Hence in situations where training examples are expensive to collect, learning has to be performed on smaller datasets. In that case using a fix hand crafted set of filters seems to be one of the best solution [8]. Recently Mallat described the scattering transform [9] a fixed bank of wavelet filters used to generate data representation in a convolutional neural networks like architecture. This representational method associated to a support vector machine classifier achieved close to state of the art performances of some standard datasets [10]. Furthermore this method can be accurately applied to relatively smaller datasets [11].

When an extremely low number of training examples are available —one-shot learning [12]— generative classification methods provide better performances than their discriminative counterparts [13]. Model a signal representation using generative probabilistic graphical models have been successfully done for various wavelet transform [14,15]. In those work Hidden Markov trees are used to model the wavelet coefficients distribution.

In a similar fashion, we propose to model Mallat's scattering convolutional network [10] using hidden Markov trees. This combines a recently proposed deterministic analytically tractable transformation inspired by deep convolutional to a probabilistic graphical model in order to create a powerful probabilistic tool to handle high dimensional prediction problems. As opposed to previous works modeling wavelet trees, the use of scattering transforms allow us to leverage their interesting representational properties but also force us to adapt the HMT model to non-homogeneous, non regular trees. In contrast to passing the raw scattering coefficients into a classifier, our proposed method captures dependencies between different layers in a generative probabilistic model. Moreover as opposed to the commonly used simple classification method, once trained our model can tackle prediction problems but also other inference tasks —e.g. generation, sensitivity analysis... and can also outperform SVMs when only a very low number of training examples are available.

The remainder of this paper introduces the scattering convolutional hidden Markov tree and is organised as follow. In section 2 we review the scattering transform and some of its properties. Section 3 introduces the proposed Scattering Hidden Markov Tree (SCHMT). Section 4 we perform classification on a selection of standard datasets restricted to only a few training samples. We draw conclusions in Section 5.

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2 Scattering networks

Scattering convolutional networks (SCNs) [16] are Convolutional Neural Networks (CNNs) [3] using a fixed filter bank of wavelets. Those filters can be hand-crafted to yield descriptors with the desired invariances [9]. For image classification tasks, one is interested in descriptors that are —at least—stable to deformations and invariant to translations. Note that SCNs producing more complexes set of invariances exist but on the remainder of this paper we consider only on descriptors with the previously mentioned properties.

2.1 Scattering transform

Wavelets are localized functions stable to deformations. They are thus well adapted to construct descriptor that would also be translation invariant. A two-dimensional spatial wavelet transform W is obtained by scaling by 2^j and rotating by r_θ a mother wavelet ψ ,

$$\psi_{\lambda}(u) = \psi_{j,\theta}(u) = 2^{-2j}\psi(2^{-j}r_{\theta}p) \tag{1}$$

In the remainder of this paper we restrict to Morlet wavelet transforms defined on $\Lambda = G \times [\![0, J]\!]$ where G is a finite group of rotations of cardinal L and where the wavelet is taken at scale J,

$$W_J \mathbf{x} = \{ \mathbf{x} * \phi_J(u); \mathbf{x} * \psi_{\lambda}(u) \}_{n \in \mathbb{R}^2, \lambda \in \Lambda}$$
 (2)

While the averaging part ϕ_J of the wavelet transform is invariant to translations, the high frequency part ψ_λ is covariant to them [9]. Invariance within a limited range inferior 2^J can be achieved by averaging the positive envelope with a smooth window,

$$S_J[\lambda]\mathbf{x}(u) = |\mathbf{x} * \psi_{\lambda}| * \phi_J(u)$$
(3)

Such non-linearised averaged wavelet coefficients are used in various form in computer vision (SIFT [17], DAISY [18]), but the scattering transform proposes a new non-linearity as well as a layer based architecture.

2.2 Scattering convolutional network

While providing local translation invariance, the averaging convolution introduced in 3 also removes the spatial variability of the wavelet transform. SCNs cascade this wavelet modulus operator to recover the lost information and compute progressively more invariant descriptors. Let combine the wavelet transform and modulus operations into a single wavelet modulus operator,

$$\mathcal{U}_{J}\mathbf{x} = \{S[\emptyset]\mathbf{x}; U[\lambda]\mathbf{x}\}_{\lambda \in \Lambda_{J}} = \{\mathbf{x} * \phi_{J}; |\mathbf{x} * \psi_{\lambda}|\}_{\lambda \in \Lambda_{J}}, (4)$$

A scattering transform can be interpreted as a CNN [19] illustrated in Figure 1 which propagates a signal $\mathbf x$ across multiple layers of the network and which outputs at each layer m the scattering invariant coefficients $S[p_m]\mathbf x$ where $p_m=(\lambda_1\ldots\lambda_m)$ is a path of m orientations and scales.

The scattering energy is mainly concentrated along frequency decreasing paths, i.e. for which $|\lambda_{k+1}| \leq |\lambda_k|$ [9].

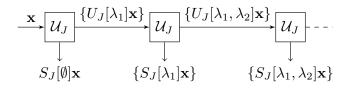


Fig. 1. Scattering networks can be seen as neural networks iterating over wavelet modulus operators \mathcal{U}_j . Each layer m outputs the averaged invariants $S[p_m]\mathbf{x}$ and covariant coefficients $U[p_{m+1}]\mathbf{x}$.

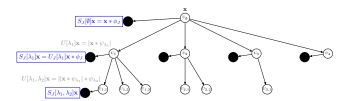


Fig. 2. Frequency decreasing scattering convolution network with J=4, L=1 and M=2. A node i at scale j_i generates $(j_i-1)\times L$ nodes.

The energy contained in the other paths is negligible and thus for applications only frequency decreasing paths are considered. Moreover there exist a path length M>0 after which all longer paths can be neglected. For signal processing applications, this decay appears to be exponential. And for classification applications, paths of length M=3, i.e. two convolutions, provides the most interesting results [10, 20].

This restrictions yield an easier parametrization of a scattering network. Indeed its now completely defined by the mother wavelet ϕ , the maximum path length considered M, the finest scale level considered J and the number of orientation considered L.

Hence for a given set of parameter (ψ, M, J, L) , let $ST_{(\psi, M, J, L)}(\mathbf{x})$ denotes the unique frequency decreasing windowed scattering convolutional network with those parameters evaluated for signal \mathbf{x} . Each node i of this network generates a -possibly empty- set of of nodes of size $(j_i-1)\times L$ where j_i is the scale of node i and i is the number of orientations considered and it has the architecture displayed by Figure 2.

2.3 Scattering convolutional classifier

In the original framework [16], the scattering network $ST_{(\psi,M,J,L)}(.)$ is used for classification task using a SVM classifier on the outputs of the network. Performance can be slightly improved by adding a feature selection step performing PCA on the scattering coefficients and keeping only the most informative ones. This classification framework provides results comparable with the state of the art on several datasets [10].

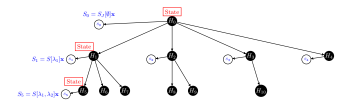


Fig. 3. Scattering convolutional hidden Markov tree.

3 The Scattering hidden Markov tree

State of the art performance can be achieved using SCNs associated to SVMs. However this approach is not adapted to very small training sets or to deliver a probabilistic output. To overcome those limitations we propose an adaptation of Crouse [14] and Durand [21] wavelet hidden Markov trees to the non-regular non-homogeneous tree structure of SCNs.

3.1 Hidden Markov tree model

The HMT models the marginal distribution of each real ST coefficient S_i as a Gaussian mixture. To each S_i , we associate a discrete hidden state H_i that takes on values in $[\![1,K]\!]$ with probability mass function (pmf) $P(H_i)$. Conditioned on $H_i=k, S_i$ is Gaussian with mean $\mu_{i,k}$ and variance $\sigma_{i,k}$. Thus, its overall marginal PDF is given by $P(w_i) = \sum_{k=1}^K P(H_i=m)P(S_i|H_i=k)$ with $P(S_i|H_i=k) \sim \mathcal{N}(\mu_{i,k},\sigma_{i,k})$. While each scattering coefficient S_i is conditionally gaussian given its state variable H_i , overall it has a non-Gaussian density. Finally the probability for the hidden node H_i to be in a state k given its father's state g is characterized by a transition probability such that $\epsilon_i^{(gk)} = P(H_i=k|H_{\rho(i)}=g)$. This yields $P(H_i=k) = \sum_{g=1}^K \epsilon_i^{(gk)} P(H_{\rho(i)}=g)$.

Such a model is pictured in Figure 3 and for a given scattering architecture —i.e. fixed M, J and L— the SCHMT model is fully parametrized by,

$$\Theta = (\pi_0, \{\epsilon_i, \{\theta_{k,i}\}_{k \in \llbracket 1, K \rrbracket}\}_{i \in \mathcal{T}}). \tag{5}$$

This model implies two assumptions on the scattering transform. First — K-populations— that a signal's scattering coefficients can be described by K clusters. This is a common assumptions for standard wavelets [15] and hence it can be extended to the scattering transform. The SCHMT also assumed —persistence— that the informative character of a coefficients is propagated across layers. This assumption is sound since scattering coefficients are highly correlated [19].

3.2 Learning the tree parameters

SCHMT is trained using the smoothed version of the Expectation-Maximization (EM) algorithm for hidden Markov trees proposed by [22] and adapted to non-homogeneous and non-binary trees.

Let $\bar{S}_i = \bar{s}_i$ be the observed sub-tree rooted at node i. By convention \bar{S}_0 denotes the entire observed tree. The smoothed

version of the E-step requires the computation of the conditional probability distributions $\xi_i(k) = P(H_i = k | \bar{\mathcal{S}}_i = \bar{s}_i)$ (smoothed probability) and $P(H_i = k, H_{\rho(i)} = g | \bar{\mathcal{S}}_i = \bar{s}_i)$ for each node $i \in \mathcal{T}$ and states k and g. This can be achieved through an upward-downward recursion displayed in Algorithm 1 and 2. The output from the downward step are used M-step as showed in Algorithm 3.

```
| // Initialization: | for All the nodes i of the tree \mathcal{T} do | P_{\theta_{k,i}}(s_i) = \mathcal{N}(s_i|\mu_{k,i},\sigma_{k,i}) end | for All the leaves i of the tree \mathcal{T} do | \beta_i(k) = \frac{P_{\theta_{k,i}}(s_i)P(H_i=k)}{\sum_{g=1}^K P_{\theta_{g,i}}(s_i)P(H_i=g)} | \beta_{i,\rho(i)}(k) = \sum_{g=1}^K \frac{\beta_i(g)\epsilon_i^{(kg)}}{P(H_i=g)}.P(H_{\rho(i)}=k) | l_i=0 end | // Induction: | for All non-leaf nodes i of the tree \mathcal{T} (Bottom-up) do | M_i = \sum_{k=1}^K P_{\theta_{k,i}}(s_i)\prod_{j\in c(i)}\frac{\beta_{j,i}(k)}{P(H_i=k)^{n_i-1}} | l_i=\log(M_i)+\sum_{j\in c(i)}l_j | \beta_i(k)=\frac{P_{\theta_{k,i}(s_i)}\prod_{j\in c(i)}(\beta_{j,i}(k))}{P(H_i=k)^{n_i-1}M_i} | for All the children nodes j of node i do | \beta_{i, \setminus c(i)}(k)=\frac{\beta_i(k)}{\beta_{i,j}(k)} | end | \beta_{i, \rho(i)}(k)=\sum_{g=1}^K \frac{\beta_i(g)\epsilon_i^{(kg)}}{P(H_i=g)}.P(H_{\rho(i)}=k) | end
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Algorithm 1: Smoothed upward algorithm.

Algorithm 2: Smoothed downward algorithm.

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\begin{split} & \text{// Initialization:} \\ & \pi_0(k) = \frac{1}{N} \sum_{n=1}^N P(H_0^n = m | s_0^n, \Theta^l) \\ & \text{// Induction:} \\ & \textbf{for All nodes } i \text{ of the tree } \mathcal{T} \backslash \{0\} \, \textbf{do} \\ & P(H_i = k) = \frac{1}{N} \sum_{n=1}^N P(H_i^n = k | \bar{s}_0^n, \Theta^l), \\ & \epsilon_i^{gk} = \frac{\sum_{n=1}^N P(H_i^n = k, H_{\rho(i)}^n = g | \bar{s}_0^n, \Theta^l)}{NP(H_{\rho(i)} = k)}, \\ & \mu_{k,i} = \frac{\sum_{n=1}^N s_i^n P(H_i^n = k | \bar{s}_0^n, \Theta^l)}{NP(H_i = k)}, \\ & \sigma_{k,i}^2 = \frac{\sum_{n=1}^N (s_i^n - \mu_{k,i})^2 P(H_i^n = k | \bar{s}_0^n, \Theta^l)}{NP(H_i = k)}. \\ & \textbf{end} \end{split}
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Algorithm 3: M-step of the EM algorithm.

3.3 MAP classification

Let Θ_c now be a set of parameters for an SCHMT \mathcal{T} learned on a training set $\{\bar{S}_{0,c}^n\}_{n\in \llbracket 1,N\rrbracket}=\{ST_{(\psi,J,M,L)}(\mathbf{x}_c^n)\}_{n\in \llbracket 1,N\rrbracket}$ composed of the scattering representations of N realizations of a signal of class c. Let also \mathbf{x}^{new} be another realization of this signal, not used for training and \mathcal{T}^{new} be the instance of the SCHMT generated by this realization.

| Training | | MNIST | | KTH-TIPS | |
|----------|--------|------------------|----------------------------|-------------------------|---|
| samples | | SCHMT | SCN+SVM | SCHMT | SCN+SVM |
| 2 | (20) | 29.7% | 32.0% | 28.9% | 32.2% |
| | (Full) | $ar{30}.ar{1}\%$ | $\bar{26}.\bar{1}\%$ | $ar{29}.ar{1}\%$ | $\overline{}2\overline{5}.\overline{6}\%$ |
| 5 | (20) | 45.0% | 56.5 % | 35.1% | 46.5 % |
| | (Full) | $ar{45}ar{5}\%$ | $\bar{4}\bar{3}.\bar{2}\%$ | $ar{35}ar{5}ar{\%}^{-}$ | $\bar{3}\bar{3}.\bar{2}\%$ |
| 10 | (20) | 51.3% | 64.8% | 49.1% | 57.7 % |
| | (Full) | $\bar{5}1.2\%$ | $ar{6}ar{1}.ar{4}\%$ | 49.4% | $oxed{ar{55.3}\%}$ |
| 20 | (20) | 50.0% | 82.3 % | 48.3% | 78.0% |
| | (Full) | 50.0% | $ar{79}.ar{8}\%$ | 48.7% | $ar{76.1}\%$ |

Fig. 4. Classification score on 200 samples (top row) and on the full testset (bottom row) of MNIST and KTH-TIPS trained using only a limited number of training points per class

In this context the MAP algorithm [22] aims at finding the optimal hidden tree $\hat{h}_0^{new}=(\hat{h}_0^{new}\dots\hat{h}_{I-1}^{new})$ maximizing the probability of this sequence given the model's parameters $P(\bar{\mathcal{H}}_0=\hat{h}_0^{new}|\mathcal{T}^{new},\Theta_c)$. The MAP framework also provides \hat{P} the value of this maximum.

The MAP algorithm can be used in a multi-class classification problem by training an SCHMT model per class and then, when presented with a new realization \mathbf{x}^{new} , comparing the probability of the MAP hidden tree provided by each model.

4 Classification results

We compare the performance of SCHMT to those of a SCN combined to an SVM (SCN+SVM) on restrictions to small number of training examples of two standard datasets. For both datasets and classification methods we use a scattering transform with M=3 orders, J=5 scales, L=3 orientations and a Morlet mother wavelet. The hidden Markov tree has K=2 states and is using a mixture of Gaussian to describe the relationship between the scattering coefficients and the hidden states. For the SVM, the best parameters are selected by cross-validation.

4.1 MNIST

We first test SCHMTs on the digit classification dataset MNIST [23]. SCHMT and SCN+SVM are both trained on a limited number of training examples per class and tested on 200 test samples (20 per class) and on the full testset (10000 images). The best results for each models are displayed in Figure 4.

When the number of training points is extremely limited—i.e. 2 or 5— SVM can reach acceptable score on a small testset. However the model learned has very poor generalization's properties and highly decreases in performance when the testset is scaled up. As opposed to this, SCHMT displays better generalization as its score remains almost constant while increasing the number of test samples. Finally it outperforms SVMs on large testset. As expected, when the number of training samples grow large enough—i.e. 10 and onward— for

the SVMs, SCN+SVMs reach better maximum classification score on both the small and the large testset.

We can also notice that EM algorithm performances are undermined by convergence to local minima issues [24]. This sometime yields poor learning quality for the SCHMT. However when convergence occurs correctly SCHMTs has better generalization performances than the best SVMs.

While confirming the superiority of a generative model in terms of generalization capacities for limited number of training points, this experiment also highly some weakness of the SCHMTs as sometime convergence is problematic. However when convergence occurs correctly SCHMT provides good classification score for such a low number of training examples.

4.2 KTH Texture

Further experiments are run on the KTH-TIPS texture dataset [25]. Similarly to what we have done for MNIST we perform the classification using limited training examples per class and perform the testing on both 20 and 155—i.e. rest of the dataset— samples per class. SVM's parameters are optimized using cross-validation. Results are displayed in Figure 4.

This texture dataset is more challenging than MNIST since the intra-class variability is higher. However the consequences are limited thanks to the scattering transform invariance properties. Again SCHMT perfoms well when given few training examples. The generalization capacity of SCHMT are also good since we see no change in performace when testing it on the 1550 remaining examples.

5 Conclusion

SCHMTs propose a new framework to process a highdimensional signal from the raw data to the prediction task. The scattering transform projects the data into a representational space of even higher dimensionality but of reduced volume along the invariants in the data. Then a probabilistic graphical model —hidden Markov tree— is used to fit a generative model to the distribution of the representation of the data. This yield a powerful tool combining the interesting properties of the scattering transform for signal representation and the representational power of the hidden Markov models. The modeled distribution can be used to perform efficient classification tasks even given low information. Even though this document considers only classification, a generative model is much more versatile than a simple -yet efficientdiscriminative counterpart. Because they model the full distribution of the data they can express more complex relationships between the observed and the unknown variables than simple discrimination.

To enhance SCHMT and especially the chance of converging toward a good minima during the EM learning, we plan on developing a variational to learn the model parameters [26].

6 References

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